Direct Estimation of Tracer-Kinetic Parameter Maps From Highly Undersampled Brain Dynamic Contrast Enhanced MRI

Yi Guo,¹* Sajan Goud Lingala,¹ Yinghua Zhu,¹ R. Marc Lebel,² and Krishna S. Nayak¹

Purpose: The purpose of this work was to develop and evaluate a T₁-weighted dynamic contrast enhanced (DCE) MRI methodology where tracer-kinetic (TK) parameter maps are directly estimated from undersampled (k,t)-space data. Theory and Methods: The proposed reconstruction involves solving a nonlinear least squares optimization problem that includes explicit use of a full forward model to convert parameter maps to (k,t)-space, utilizing the Patlak TK model. The proposed scheme is compared against an indirect method that creates intermediate images by parallel imaging and compressed sensing before to TK modeling. Thirteen fully sampled brain tumor DCE-MRI scans with 5-second temporal resolution are retrospectively undersampled at rates R = 20, 40, 60, 80, and 100 for each dynamic frame. TK maps are quantitatively compared based on root mean squared-error (rMSE) and Bland-Altman analysis. The approach is also applied to four prospectively R = 30 undersampled whole-brain DCE-MRI data sets. Results: In the retrospective study, the proposed method performed statistically better than indirect method at R ≥ 80 for all 13 cases. This approach provided restoration of TK parameter values with less errors in tumor regions of interest, an improvement compared to a state-of-the-art indirect method. Applied prospectively, the proposed method provided whole-brain, high-resolution TK maps with good image quality. Conclusion: Model-based direct estimation of TK maps from k,t-space DCE-MRI data is feasible and is compatible up to 100-fold undersampling. Magn Reson Med 000:000–000, 2016. © 2016 International Society for Magnetic Resonance in Medicine.

Key words: dynamic contrast enhanced (DCE) MRI; kinetic parameter mapping; constrained reconstruction; model-based reconstruction

INTRODUCTION

Brain dynamic contrast enhanced (DCE) MRI is used to measure neurovascular parameters, such as blood–brain barrier permeability, in a variety of conditions, including brain tumor (1,2), multiple sclerosis (3,4), and Alzheimer’s disease (5). DCE-MRI involves collecting a series of T₁-weighted images during administration of a T₁-shortening contrast agent (6,7). tracer-kinetic (TK) modeling is then performed on the dynamic images to estimate physiological parameters, such as vascular permeability (Ktrans), fractional plasma volume (v_p), and extravascular-extracellular volume fraction (v_e) (8,9). The anatomical dynamic images are primarily used to derive the TK maps (2,10–12).

For many applications, current DCE-MRI with Nyquist sampling is unable to simultaneously provide high spatiotemporal resolution and adequate volume coverage. Compressed sensing- (13) and parallel imaging-based (14,15) schemes have been proposed to accelerate acquisition process, primarily to achieve better spatial resolution and coverage while maintaining the same temporal resolution. Notably, Lebel et al (16) used a temporal high-pass filter and multiple spatial sparsity constraints to achieve an undersampling rate (R) of 36 × and showed excellent quality of anatomical images in brain tumor cases. A recent pilot study in brain tumor patients indicated that this approach performs superior to conventional techniques with no apparent loss of diagnostic information (17). The works of Feng et al (18), Chandarana et al (19), Rosenkrantz et al (20) used a golden-angle radial sampling pattern, compressed sensing, and parallel imaging to achieve a comparable acceleration rate of 19.1 to 28.7. These studies showed improved resolution and reduced motion sensitivity in breast, liver, and prostate DCE-MRI, compared to either parallel imaging alone or coil-by-coil compressed sensing alone. We will refer to these techniques as “indirect” methods, because the anatomical image series are reconstructed first, followed by a separate step for TK parameter fitting on a voxel-by-voxel basis.

In this work, we propose a framework for “direct” estimation of TK parameter maps from fully sampled or undersampled (k,t)-space data. We use a full forward model that converts the TK maps to (k,t)-space, and we pose the estimation of TK maps as an error minimization problem. Our approach is motivated by two factors: 1) Spatial TK parameter maps have much lower dimensionality than those of dynamic image series (two to four parameters, compared to 50–100 time points, per voxel), and 2) TK model-based reconstruction directly exploits what is known about contrast agent kinetics. These allow for robust parameter estimation from an information theoretic perspective and have the potential to provide the most accurate restoration of TK parameter values, and allow for the highest acceleration.
Model-based direct reconstruction has been previously explored in other applications, such as MRI relaxation parameter estimation (21–26), TK parameter estimation in positron emission tomography (PET) (27–29), and TK parameter estimation in MRI (30,31). Notably, for MRI relaxation parameter estimation, Sumpf et al (24) used a model-based nonlinear inverse reconstruction to estimate T2 maps from highly undersampled spin-echo MRI data; Zhao et al (25) estimated T1 parameters directly from undersampled MRI data with a sparsity constraint on the parameter maps. For dynamic PET imaging, Kamasak et al (28) directly estimated TK parameter images from dynamic PET data using a kinetic model-based reconstruction optimization; Lin et al (29) used a sparsity constrained mixture model to estimate TK parameters from dynamic PET data and evaluated with both simulated and experimental PET data.

For DCE MRI parameter mapping, Felsted et al (30) proposed to use a model-based reconstruction algorithm to solve TK parameters directly from undersampled MRI k-space with a modified gradient descent algorithm. An undersampling factor of R = 4 was demonstrated on simulated data; Dikaios et al (31) proposed a Bayesian inference framework to directly estimate TK maps from undersampled MRI data and achieved 8 × acceleration in phantom and in vivo prostate cancer data.

Although the previous studies demonstrate promise, the full potential of TK model-based reconstruction has lacked validation, both at higher undersampling rates or with prospectively undersampled data from patients. In this study, we explore the maximum potential benefit of the direct TK estimation by testing very high undersampling rates. We validate the approach using retrospective undersampling of fully sampled data and using prospectively undersampled DCE data sets from brain tumor patients. Compared to previous work, we are able to demonstrate much higher undersampling rates (up to 100×) in the retrospective study, using a more-efficient, gradient-based algorithm. We use quantitative evaluation (root mean square error [rmSE] in TK parameters) to provide a systematic comparison against a state-of-the-art compressed sensing method that uses spatial and temporal sparsity constraints in 13 brain tumor patients. We also uniquely provide a prospective in vivo study showing that whole-brain coverage with high spatial resolution can be achieved to capture complete pathological information. We demonstrate the potential of direct reconstruction to enable “parameter-free” reconstruction, when no sparsity constraints are added.

**THEORY**

**Direct TK Mapping**

We propose to integrate TK modeling, specifically the Patlak model, into the image reconstruction process. Figure 1 illustrates the forward model that relates TK parameter maps to undersampled (k,t)-space. We use the vector \( \mathbf{r} \in (x,y,z) \) to represent image domain spatial coordinates, \( \mathbf{k} \in (k_x,k_y,k_z) \) to represents k-space coordinates; \( t, c \) are the time and coil dimensions. The variables beneath the arrows of each step are known or predetermined. The steps indicated (above the arrows) in Figure 1 are explained below:

1. **Contrast agent concentration over time** \( C_A(r,t) \) is assumed to follow the Patlak model (Eq. [1]):

   \[
   C_A(r,t) = K^{\text{trans}}(r) \int_0^t C_p(\tau) d\tau + v_p(r)C_p(t)
   \]  

   where \( C_p(t) \) is the arterial input function (AIF). In this study, we used a population-based AIF from Parker et al (32). Notice that the AIF requires specifying a delay time. This is estimated from the k-space origin, which is acquired in every time frame, and has shown to accurately detect the time of contrast bolus arrival (33). We assume that the Patlak is appropriate for all voxels in the imaging volume. We have observed that image regions outside of vessels and tumor typically experience no enhancement during the DCE-MRI acquisition, \( \text{Ktrans} \) which is the contrast agent relaxivity, and \( \text{R1} \) is the relaxation parameter.

2. **Dynamic anatomic images** \( s(r,t) \) are related to \( C_A(r,t) \) by the steady-state spoiled gradient echo (SPGR) signal equation (Eq. [2]):

   \[
   s(r,t) = M_0(r)\sin(1 - \exp(-TRR_1(r,t) + C_A(r,t) t))
   \]

3. **The undersampled raw (k,t)-space data** \( S(k,t,c) \) are related to \( s(r,t) \) by the coil sensitivities \( C(r,c) \) and undersampling Fourier transform \( F_u \) (Eq. [3]):

   \[
   S(k,t,c) = F_u C(r,c) s(r,t)
   \]

In this work, \( C(r,c) \) is estimated from time-averaged data using the standard root sum-of-squares method (14). The image phase information is assumed to be captured by the complex-valued sensitivity maps.

Combining Equations [1] to [3], we reach a general function \( f \) to denote the relationship between TK maps \( K^{\text{trans}}(r) \), \( v_p(r) \) and undersampled (k,t)-space \( S(k,t,c) \) (Eq. [4]):

\[
S(k,t,c) = f(K^{\text{trans}}(r), v_p(r); C_p(t), TR, \alpha, R_1(t,0), M_0(r), r_1, C(r,c))
\]

where \( C_p(t), TR, \alpha, R_1(t,0), M_0(r), r_1, C(r,c) \) are variables that are known or predetermined as mentioned above.
We solve for the unknown $K_{\text{trans}}(r), v_p(r)$ by least-square optimization, formulated as follows (Eq. [5]):

$$(K_{\text{trans}}(r), v_p(r)) = \underset{K_{\text{trans}}(r), v_p(r)}{\text{argmin}} \left| |S(k, t, c) - f(K_{\text{trans}}(r), v_p(r))|^2ight|$$

[5]

This nonlinear optimization problem is solved by a quasi-Newton limited-memory Broyden-Fletcher-Goldfarb-Shannon (L-BFGS) method (36), where $K_{\text{trans}}(r)$ and $v_p(r)$ are solved alternatingly. The details of the optimization algorithm and gradient calculation are provided in Appendix I.

Direct reconstruction by itself is parameter free. This is in contrast to compressed sensing-based algorithms that require tuning of one or more regularization parameters. It is possible to incorporate additional spatial sparsity constraints on the TK maps themselves. In this work, we test the potential value of adding a spatial “db2” wavelet constraint ($\Psi$) to the parameter maps. The optimization problem with sparsity constraint is formulated as follows (Eq. [6]):

$$
(K_{\text{trans}}(r), v_p(r)) = \underset{K_{\text{trans}}(r), v_p(r)}{\text{argmin}} \left| |S(k, t, c) - f(K_{\text{trans}}(r), v_p(r))|^2ight| + \lambda_1\|\Psi K_{\text{trans}}(r)\|_1 + \lambda_2\|\Psi v_p(r)\|_1
$$

[6]

We provide source code, along with sample data sets, and scripts that generate several of the figures from this article (37). Repository: https://github.com/usc-mrel/DCE_direct_recon; Release 1: https://doi.org/10.5281/zenodo.154058.

Indirect TK Mapping

Current state-of-the-art methods for highly accelerated DCE involve reconstructing intermediate images before TK modeling. These indirect methods are the most relevant alternatives to direct TK modeling and serve as a performance benchmark. A basic model for indirect reconstruction solves the minimization problem in Equation [7], where the final image, $s(r, t)$, remains consistent with acquired (k,t)-space data $S(k, t, c)$, yet is sparse in the temporal finite differences ($V$) domain and spatial wavelet domain ($\Psi$).

$$
s(r, t) = \underset{s(r, t)}{\text{argmin}} \left| |S(k, t, c) - F_u C(r, c)s(r, t)|^2\right| + \lambda_1\|Vs(r, t)\|_1 + \lambda_2\|\Psi s(r, t)\|_1
$$

[7]

The image is related to the acquired data using known coil sensitivities $C(r, c)$ and the undersampling Fourier transform, $F_u$. TK modeling (eg, using a Patlak model) is performed in a last step to estimate the spatial TK maps (eg, $K_{\text{trans}}(r)$, $v_p(r)$) from $s(r, t)$. This optimization problem was solved by an efficient augmented Lagrangian method, alternating direction methods of multipliers (38), to get the anatomical images.

METHODS

Digital Phantom

We simulated realistic DCE-MRI data using a digital phantom with known TK parameter maps and using the Patlak TK model. We used a process identical to Zhu et al (39), where the segmentation is extracted from patient data. Realistic sensitivity maps were used, and noise was added to each channel according to noise covariance matrix estimated from the patient data. A pre-contrast white matter signal-to-noise ratio (SNR) level of 20 was chosen to mimic the SNR level in actual DCE data sets.

We retrospectively undersampled (k,t)-space with rates $R$ of $20 \times$ to $100 \times$. Ten noise realizations were generated for each $R$. Undersampling was in the k$_x$-k$_y$ plane,
simulating the $k_x$-$k_z$ plane as in a prospectively undersampled 3D case, using a randomized golden-angle radial sampling pattern (40,41). Detailed description and the videos of the undersampling strategies can be found in the Supporting Information. Direct and indirect methods were used to generate the TK parameters from both fully sampled and undersampled data. TK map rMSEs were computed over a region of interest (ROI) containing the entire tumor boundary.

**In Vivo Retrospective Evaluation**

We reviewed 110 fully sampled DCE-MRI raw data sets from patients with known or suspected brain tumor, receiving a routine brain MRI with contrast on a clinical 3 Tesla (T) scanner (HDxt; GE Healthcare, Waukesha, WI). The data sets were from patients receiving routine brain MRI with contrast (including DCE-MRI) at our institution, and the demographics reflect our local patient population. Our Institution follows standard exclusion criteria for MRI with gadolinium-based contrast (42,43), which includes: a medically unstable, renal impairment, cardiac pacemaker, internal ferromagnetic device that is contraindicated for use in MRI, claustrophobia, and any other condition that would compromise the scan with reasonable safety. The retrospective study protocol was approved by our institutional review board.

The sequence was based on a 3D Cartesian fast SPGR with field of view (FOV): $22 \times 22 \times 4.2$ cm$^3$; spatial resolution: $0.9 \times 1.3 \times 7.0$ mm$^3$; temporal resolution: 5 seconds; 50 time frames; and eight receiver coils. The flip angle is $15^\circ$, echo time is 1.3 ms, and TR is 6 ms. DESPOT1 was performed before the DCE sequence, where three images with flip angles of 2°, 5°, and 10° were acquired to estimate $T_1$ and $M_0$ maps before the contrast arrival. The contrast agent, gadobenate dimeglumine (MultiHance Bracco Diagnostics Inc., Princeton, NJ), which has relaxivity $r_1 = 4.39$ s$^{-1}$ mM$^{-1}$ at 37°C at 3T (44), was administered with a dose of 0.05 mMol/kg, followed by a 20-mL saline flush in the left arm by intravenous injection.

Of the 110 cases, we found 18 that had visible tumor larger than 1 cm by bidirectional assessment. TK parameter maps $K_{trans}$ and $v_p$ were calculated from the fully sampled images, and TK model fitting error was computed by taking the l2 norm between the contrast concentration curves from fully sampled images, $CA(r,t)$, and the fitted concentration curves generated from the TK parameter maps, $GA(r,t)$. We then examined the Patlak modeling error, defined as (Eq. [8]):

$$\frac{\|CA(r,t) - GA(r,t)\|^2}{\|CA(r,t)\|^2} \times 100\% \quad [8]$$

Of the 18 cases with visible tumor larger than 1 cm, 13 cases had Patlak modeling error less than 1%, suggesting that the Patlak model with the population AIF was appropriate. The analysis below was performed on the 13 cases that were fully sampled, had at least one tumor larger than 1 cm, and for whom the Patlak model fitted the fully sampled data with less than 1% error.

For each selected case, three sets of TK maps were generated from: 1) standard Fourier reconstruction of fully sampled data. These served as the gold-standard reference maps. 2) Direct reconstruction method using retrospective undersampling; and 3) Indirect reconstruction using retrospective undersampling. We examined R of $20 \times 100\times$, with increments of $20\times$. For each R, 10 realizations of the sampling pattern were generated using a different initial angle in the randomized golden-angle radial scheme. This effectively creates multiple noise realizations, given that there is almost no overlap in the (k,t) sampling pattern (except for the one sample at the k-space origin, which is included in every sampling scheme at every undersampling factor).

For indirect method (Eq. [7]), the regularization parameters were empirically set as $\lambda_1 = 0.01$ and $\lambda_2 = 0.0001$. These values are motivated by empirical observations made on retrospective undersampling studies on a number of data sets (around 15) based on a criterion of achieving minimal rMSE between the reconstructed dynamic images from subsampled and fully sampled data. Both the regularizations were used in Equation [7], during experimental comparisons against the direct method with spatial sparsity constraint (Eq. [6]). For direct method, regularization parameters were also empirically set as $\lambda_1 = 0.03$ and $\lambda_2 = 0.00001$. For fair comparison, when the direct method had no spatial wavelet constraint (Eq. [5]), $\lambda_2$ in Equation [7] of the indirect method was set to 0.

The quantitative metric, rMSE, was computed on TK parameter maps, within an ROI containing enhancing tumor. Bland-Altman plots were generated using the difference of the reconstructed $K_{trans}$ maps with respect to the fully sampled $K_{trans}$ maps within the ROI to test for any systematic bias.

A two-tailed paired Student t test was performed based on the rMSE of the two methods in the 13 patients. The null hypothesis was equivalence of the two methods, with the null value being zero. The significance criterion was P value less than 0.05. Assumptions of normality were validated using the Shapiro-Wilk test. Bonferroni correction was applied to correct for multiple comparisons, that is, the significance level for each individual test was set to 0.05/13.

For one data set, we applied spatial wavelet sparsity constraints for both methods (Eqs. [6] and [7]) to demonstrate the feasibility and determine any possible improvement.

**In Vivo Prospective Evaluation**

Prospectively undersampled data were acquired in 4 brain tumor patients (65 male [M], 71 M, 46 female [F], and 22 F, all glioblastoma) with Cartesian golden-angle radial k-space sampling (17,41). 3D $T_1$-weighted SPGR data were acquired continuously for 5 minutes. Whole-brain coverage was achieved with an FOV of $22 \times 22 \times 20$ cm$^3$ and spatial resolution of $0.9 \times 0.9 \times 1.9$ mm$^3$. The prospective study protocol was approved by our institutional review board. Written informed consent was provided by all participants.
Five-second temporal resolution was achieved by grouping raw (k,t)-space data acquired within consecutive 5-second intervals. This prospective acquisition undersampled each 5-second temporal frame by 30x2. Note that undersampling is not being used to shorten the scan time, but rather to significantly increase the spatial coverage and spatial resolution. For comparison, the standard clinical protocol at our institution that utilizes Nyquist sampling and 5-second temporal resolution achieves FOV 22x22x4.2 cm3 and spatial resolution 0.9 x 1.3 x 7.0 mm3. For DCE-MRI, the scan time and temporal resolution are kept the same to capture dynamic changes during contrast arrival and washout.

Direct estimation of TK maps was performed using the proposed method. Three-plane and panning volume Ktrans and vp maps for the four data sets are presented for visual assessment. The first frame is necessary in the direct reconstruction process. A detailed description of how to obtain this frame, utilizing the properties of the golden-angle radial sampling pattern, can be found in the Supporting Information. These prospectively undersampled DCE-MRI data were obtained 20 minutes after a standard-of-care conventional DCE-MRI scan; therefore, there was some residual contrast on board.

RESULTS

Figure 2 shows the convergence performance for the direct method at R = 20. Objective function changes are plotted against iteration number, and final results where minimum gradient values reached are shown with different initial conditions. In the experiments, 140 to 180 iterations were needed to reach the stopping criteria. Convergence was achieved regardless of the initial condition. The reconstruction time for indirect and direct method was 265 and 296 seconds, respectively, on a Linux workstation (24 core 2.5 GHz, 192 GB RAM). Ktrans is reported in the units of min−1, and vp is reported as percent fraction. This holds throughout the article.

Figure 3 top shows the phantom results (cropped at the tumor part) of indirect and direct reconstruction of the Ktrans and vp maps at R = 1 (fully sampled data) and R = 60 for both the direct and indirect methods. At R = 1, it is shown that the direct method does not introduce additional error by enforcing the Patlak model into the reconstruction. At R = 60, the direct method performs better than the indirect method for both Ktrans and vp maps, overcoming the large errors and noise introduced by the indirect method. Figure 3 (bottom) shows the rMSE (calculated in tumor boundary ROI) performance across different Rs. Across all Rs tested, the direct method outperformed the indirect method at high undersampling rates.

Figure 4 shows one representative example of the image results of direct and indirect reconstruction. Ktrans and vp maps of a glioblastoma patient at three different undersampling rates obtained from fully sampled data, the proposed direct reconstruction, and by indirect reconstruction are shown. Across all undersampling rates, the direct method qualitatively and quantitatively depicted equal or more-accurate restoration of TK parameter maps compared to the indirect method. At R of 20 x or less, the direct and indirect methods had equivalent performance. At higher undersampling rates, the indirect method failed to capture critical tumor signals and tumor shapes in Ktrans maps and small vessel information in vp maps, whereas the direct method was able to provide accurate restoration. The results with spatial wavelet constraints (only 100x shown here) provide improved noise performance and image quality. It is also worth noting that the indirect method tends to underestimate Ktrans values, whereas the direct method overcomes this underestimation. This is better observed in the Bland-Altman plots in Figure 6.

Figure 5 shows tumor ROI Ktrans rMSE plots for both methods for a range of undersampling rates and all 13 data sets that we used for retrospective evaluation. The error bars show the variance introduced by varying the initial angle of the sampling patterns. The direct method outperformed the indirect method for all the cases at high undersampling factors (>80x). The cases are ordered (left to right) by decreasing performance of direct reconstruction compared to indirect reconstruction. More-detailed analysis can be found in Table 1.
Figure 6 shows the Bland-Altman plots of direct and indirect methods for the $K^{\text{trans}}$ values combining the ROIs of all the 13 cases. The indirect method tended to suppress $K^{\text{trans}}$ values smaller than 0.02 min$^{-1}$ and tended to underestimate values larger than 0.02 min$^{-1}$. This is similar to a soft-thresholding operation and is illustrated by the green line in Figure 6. This may be a side effect of the temporal finite difference constrained reconstruction that suppresses small temporal changes of concentration. Such a trend was not observed in the direct reconstruction, suggesting that tighter integration of the TK model is able to identify and restore low $K^{\text{trans}}$ values.

Table 1 lists the patient demographic information and the performance of direct and indirect methods at $R=60$, evaluated by rMSE for both $K^{\text{trans}}$ and $v_p$ maps. The mean and standard deviation (SD) of rMSE are listed. The order of presentation in this table matches the order of Figure 6. The normality assumptions are met by the Shapiro-Wilk test, and only 4 of 56 cases (13 patients, five undersampling rates) reject the null hypothesis of composite normality assumption at the significance level of 0.05. A two-tailed paired Student t test was performed between the two methods based on rMSE of $K^{\text{trans}}$, and the last two columns show the smallest $R$ where the direct method started to statistically significantly outperform the indirect method ($P < 0.0038$ after Bonferroni correction) and the corresponding $P$ value. The direct method was consistently better than the indirect method across multiple data sets at high undersampling rates, and the difference was statistically significant at $R > 20$ for 6 cases, $R > 40$ for 3 cases, $R > 60$ for 1 case, and $R > 80$ for 3 cases.

Figure 7 illustrates direct reconstruction of $K^{\text{trans}}$ and $v_p$ in two representative prospectively undersampled whole-brain DCE data sets. Panning video TK maps for all 4 cases are provided in the Supporting Information. The whole-brain, high-resolution TK maps enable visualization of the tumor on any arbitrary reformatted plane, providing a complete depiction of the pathological information, and evaluation of narrow enhancing margin and small lesions. The reconstruction time was approximately 10 hours.
pilot study demonstrates the feasibility of applying direct TK parameter reconstruction to whole-brain DCE-MRI.

**DISCUSSION**

We have presented a novel, potentially powerful TK parameter estimation scheme for DCE-MRI, where the TK parameter maps are directly reconstructed from undersampled (k,t)-space. By integrating the full forward model connecting the TK maps to the (k,t)-space data in the reconstruction, this method is able to provide excellent TK map fidelity at undersampling rates up to 100x. Higher rates were not tested. The forward model contained the analytic TK model along with specification of the AIF, coil sensitivity maps, and precontrast $T_1$, as well as $M_0$ maps obtained from prescans. The optimization has the flexibility to incorporate additional spatial sparsity constraints on the TK maps, as demonstrated with a spatial wavelet transform. In the retrospective study, this method outperformed an indirect reconstruction using parallel imaging and compressed sensing. We also uniquely demonstrated the use of this method for prospectively undersampled whole-brain DCE data, where whole-brain TK parameter maps can be produced with excellent image quality.

The proposed method is a parameter-free reconstruction, when no spatial constraints are applied to the TK maps. We demonstrated that by simply enforcing the TK model during reconstruction, performance is improved relative to a state-of-the-art compressed sensing reconstruction, without the need to select a constraint or tune associated regularization parameters. It is straightforward to add sparsity constraints to the optimization problem, as shown in Equation [7]. Such constraints improve the noise performance, but at the expense of tuning regularization parameters. These constraints were found to

![FIG. 4. Retrospective evaluation of direct and indirect reconstruction of $K^{\text{trans}}$ and $v_p$ maps. Both reconstructions are shown without spatial wavelet sparsity constraints in the first three columns and with the sparsity constraint in the last column. By visual inspection, direct reconstruction outperformed indirect results at all undersampling rates. The direct method provided superior delineation of the tumor boundary and other high-resolution features in the $K^{\text{trans}}$ maps than did the indirect method. This was particularly true at the highest undersampling rate (see arrows at 100×). For $v_p$ maps, the direct method better preserved small vessel signals (see arrows at 100×) compared to the indirect method. Spatial wavelet constraints, shown on the rightmost column with 100×, provide additional noise suppression.](image-url)
FIG. 5. Tumor ROI (fully sampled) and rMSE plot for direct and indirect reconstruction results across different R (20× to 100×) and 13 data sets for $K^{\text{trans}}$ values. The error bar indicates the mean and variance of the rMSE for each R where 10 different realizations of sampling patterns were used. The direct method outperformed the indirect method in most cases, especially at $R > 60\times$. The variance for the two methods are comparable.

Table 1
Patient demographic information and the rMSE performance (mean and SDs) of $K^{\text{trans}}$ and $v_p$ for direct and indirect methods at $R = 60\times$

<table>
<thead>
<tr>
<th>No.</th>
<th>Age/Sex</th>
<th>Diagnosis</th>
<th>$K^{\text{trans}}$ rMSE ($\times 10^{-4}$)</th>
<th>$v_p$ rMSE ($\times 10^{-4}$)</th>
<th>$K^{\text{trans}}$ rMSE ($\times 10^{-4}$)</th>
<th>$v_p$ rMSE ($\times 10^{-4}$)</th>
<th>Significantly Different at R &gt;</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>75M</td>
<td>Meningioma</td>
<td>117 ± 2.1</td>
<td>237 ± 4.1</td>
<td>96 ± 2.0</td>
<td>219 ± 6.3</td>
<td>20×</td>
<td>3.72 × 10$^{-7}$</td>
</tr>
<tr>
<td>2</td>
<td>74M</td>
<td>Glioblastoma</td>
<td>107 ± 2.0</td>
<td>250 ± 6.2</td>
<td>76 ± 1.2</td>
<td>188 ± 4.2</td>
<td>20×</td>
<td>6.3 × 10$^{-9}$</td>
</tr>
<tr>
<td>3</td>
<td>73M</td>
<td>Glioma</td>
<td>120 ± 3.8</td>
<td>305 ± 9.5</td>
<td>99 ± 3.2</td>
<td>258 ± 9.4</td>
<td>20×</td>
<td>2.55 × 10$^{-8}$</td>
</tr>
<tr>
<td>4</td>
<td>69F</td>
<td>Meningioma</td>
<td>178 ± 4.5</td>
<td>313 ± 8.3</td>
<td>103 ± 3.4</td>
<td>201 ± 6.2</td>
<td>20×</td>
<td>4.1 × 10$^{-10}$</td>
</tr>
<tr>
<td>5</td>
<td>77M</td>
<td>Glioma</td>
<td>90 ± 4.7</td>
<td>191 ± 8.3</td>
<td>82 ± 5.2</td>
<td>178 ± 7.4</td>
<td>20×</td>
<td>6.86 × 10$^{-6}$</td>
</tr>
<tr>
<td>6</td>
<td>39F</td>
<td>Meningioma</td>
<td>114 ± 5.0</td>
<td>192 ± 10.5</td>
<td>83 ± 4.7</td>
<td>191 ± 8.2</td>
<td>20×</td>
<td>1.75 × 10$^{-6}$</td>
</tr>
<tr>
<td>7</td>
<td>54F</td>
<td>Glioma</td>
<td>167 ± 5.1</td>
<td>358 ± 14.5</td>
<td>156 ± 4.4</td>
<td>376 ± 17.3</td>
<td>40×</td>
<td>2.41 × 10$^{-6}$</td>
</tr>
<tr>
<td>8</td>
<td>44F</td>
<td>Meningioma</td>
<td>155 ± 6.4</td>
<td>327 ± 13.1</td>
<td>123 ± 5.6</td>
<td>281 ± 14.3</td>
<td>40×</td>
<td>6.57 × 10$^{-6}$</td>
</tr>
<tr>
<td>9</td>
<td>60M</td>
<td>Glioblastoma</td>
<td>130 ± 5.0</td>
<td>306 ± 10.2</td>
<td>113 ± 5.5</td>
<td>261 ± 10.1</td>
<td>40×</td>
<td>9.73 × 10$^{-7}$</td>
</tr>
<tr>
<td>10</td>
<td>38F</td>
<td>Glioma</td>
<td>134 ± 5.4</td>
<td>457 ± 6.8</td>
<td>129 ± 5.3</td>
<td>424 ± 6.2</td>
<td>60×</td>
<td>7.42 × 10$^{-5}$</td>
</tr>
<tr>
<td>11</td>
<td>63M</td>
<td>Meningioma</td>
<td>135 ± 5.4</td>
<td>361 ± 10.6</td>
<td>136 ± 6.9</td>
<td>352 ± 22.0</td>
<td>80×</td>
<td>1.25 × 10$^{-6}$</td>
</tr>
<tr>
<td>12</td>
<td>73F</td>
<td>Glioma</td>
<td>171 ± 6.1</td>
<td>436 ± 18.2</td>
<td>171 ± 6.6</td>
<td>433 ± 16.5</td>
<td>80×</td>
<td>3.84 × 10$^{-4}$</td>
</tr>
<tr>
<td>13</td>
<td>79F</td>
<td>Glioma</td>
<td>133 ± 9.5</td>
<td>318 ± 13.0</td>
<td>137 ± 4.7</td>
<td>316 ± 12.2</td>
<td>80×</td>
<td>1.31 × 10$^{-4}$</td>
</tr>
</tbody>
</table>

The patient order is sorted as the direct reconstruction performance degraded (same as Fig. 6). At a significance level of $P < 0.05$ (for individual case, $P < 0.0038$ after Bonferroni correction), the direct method performed better than the indirect method for all the cases, with the cut-off undersampling rates varying between $20\times$ and $80\times$. The P value for the cut-off undersampling rate is shown in the last column.
improve convergence at very high undersampling rates (>50×), where the TK map estimation problem becomes ill-posed.

The prospective study demonstrates that the proposed method can be used to achieve substantially higher spatial resolution and broader spatial coverage DCE-MRI, while maintaining the same temporal resolution and overall scan time. Although not studied in this work, this approach could potentially be used to improve the temporal resolution of DCE-MRI, which is known to provide improvements in patient-specific AIF measurement and TK parameter precision (45).

The proposed method estimates the TK maps directly and rather than reconstructing image time series as an intermediate step. This enables robust parameter estimation and ease of use in clinical application. Clinically, intermediate images (typically 50–100 volumes) are not always viewed. The extracted TK maps are of primary interest given that they succinctly describe the behavior of the intermediate images. It is worth noting that the proposed method can provide intermediate images using the full forward model described in Figure 1. Figure 8 compares synthesized images from reconstructed TK maps against fully sampled anatomical images. The synthesized images show close resemblance to the fully sampled images.

The proposed direct reconstruction scheme requires a priori definition of the AIF. In this work, we used a population-based AIF, and the time of arrival was automatically detected, as described in Lebel et al (33). However, other extensions, which are blind to the choice of AIF, could be explored. For instance, Fluckiger et al (46) proposed a model-based blind estimation of both AIF and TK parameters from DCE images for fully sampled data. This approach may be combined with ours to provide joint reconstruction of AIF and TK parameters directly from undersampled data.

This study has a few important limitations. First, we have thus far only demonstrated effectiveness of this approach using the Patlak model. Patlak was chosen because it is widely used and can be linearized and gradients can be readily computed. We also restricted the retrospective study to data sets that fit the Patlak model. It will be important to develop support for more-sophisticated models and utilize data that do not fit the presumed model to fully characterize failure modes. Use of more-sophisticated models (eg, extended Tofts model or two-compartment exchange model) may fit the data better. Their inclusion will make the reconstruction problem nonlinear and, possibly, nonconvex. Gradient descent algorithms may not be applicable, given that they require analytical solution for the first derivative of CA(r,t) with respect to each TK parameter in the model (step 1 for Patlak model in Fig. 1). Dikaouis et al (47) recently demonstrated the use of a Bayesian formulation of direct TK parameter estimation in DCE. The rationale was to use an optimal model for different tissue types. The additional complexity of more-sophisticated models will necessitate longer reconstruction time, and convergence will require further investigation, and remains as future work.

A second limitation is that the tight integration of TK modeling in our reconstruction could be sensitive to data inconsistencies, such as patient motion. This is equally true for indirect reconstruction. Prospective motion compensation could be added to the proposed model, but the complexities involved and efficacy have not been investigated here.

A third limitation is that the intermediate anatomical images are computed during the reconstruction and thus require a similar amount of memory and computation time as indirect methods. This approach is not currently solving the computational limitation of constrained
reconstructions, but rather provides a framework for improved image quality.

A fourth limitation is that $T_1$ and $M_0$ maps are required in the forward model and were estimated using a separate multiple flip-angle sequence (DESPOT1) performed immediately before the DCE scan. Future work could include joint estimation of the precontrast $T_1$ maps and the TK maps, as suggested by Dickie et al. (48).

We compared the direct method with a state-of-the-art indirect method that utilized a temporal finite difference constraint. Compressed sensing techniques are expected to improve steadily as better constraints are identified (49–51); however, if used for TK parameter estimation, a TK model will be applied to the data, and model inconsistencies introduced by the intermediate sparsity transforms are likely to propagate into the final parameter maps. Although untested, we hypothesize that our direct estimation method is likely to meet or exceed the image quality of any compressed sensing method.

Our proposed direct reconstruction scheme provides a method for highly accelerated DCE. Extremely high acceleration rates have been demonstrated (up to $100\times$), enabling full-brain DCE with high spatial and temporal resolution. Our method is able to provide a parameter-free reconstruction and so avoids the empirical tuning required in other methods. This technique is easily extendable to DCE-MRI in other body parts, such as the breasts, prostate, etc. Future work will include exploration of these additional clinical

FIG. 7. Direct reconstruction of $K^{trans}$ (a,c) and $v_p$ (b,d) maps from two representative prospectively undersampled data. Although lacking the gold standard for the prospective studies, the direct reconstruction provided reasonable $K^{trans}$ values and complete depiction of the entire tumor region. Panning-volume videos for all 4 cases are available in the Supporting Information.
applications, optimization of data-sampling schemes, and integration of more-sophisticated TK models.

CONCLUSION

We have presented a novel and efficient reconstruction scheme to directly estimate TK parameter maps from highly undersampled DCE-MRI data. By comparison with a state-of-the-art indirect compressed sensing method, we demonstrate that the proposed direct approach provides improved TK map fidelity and enables much higher acceleration. With the prospective study, this method is shown to be clinically feasible and provide high-quality, whole-brain TK maps.

ACKNOWLEDGMENTS

We thank Dr Samuel Barnes for useful discussions related to DCE TK modeling. We also thank Dr Meng Law, Dr Mark S. Shiroshi, Mario Franco, and Samuel Valencerina for help recruiting and scanning brain tumor patients. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

APPENDIX I: GRADIENT CALCULATION FOR THE OPTIMIZATION PROBLEM

The optimization problem in Equation [5] is solved alternatively by a quasi-Newton limited-memory Broyden-Fletcher-Goldfarb-Shanno (l-BFGS) method (36). That is, solving one while keeping the other fixed, as the pseudo codes indicated below (Eqs. [A.1] and [A.2]):

Input initial guess $K^{\text{trans}}(r)$ and $v_p(r)$

k=0, while “stopping criteria not met” do {

$K^{\text{trans}}(r)^{(k+1)} = \arg\min_{K^{\text{trans}}(r)} ||S(k, t, c) - y(K^{\text{trans}}(r), v_p(r)^{(k)})||^2_2$ \hfill [A.1]

$v_p(r)^{(k+1)} = \arg\min_{v_p(r)} ||S(k, t, c) - y(K^{\text{trans}}(r)^{(k+1)}, v_p(r))||^2_2$ \hfill [A.2]

k=k+1; }

The gradient of the cost function is evaluated analytically. This can be derived from the model and signal equations. For notational simplicity, the coordinate notations $r$, $k$, and $c$ are neglected ($t$ is preserved to show the difference in dimension between TK parameter maps and dynamic images). For example, $K^{\text{trans}}$ is for $K^{\text{trans}}(r)$ and $S(t)$ is for $S(k, t, c)$.

In Equation [A.1], we denote the cost function as:

$y(K^{\text{trans}}, v_p) = ||S(t) - f(K^{\text{trans}}, v_p)||^2_2$

where for one iteration, $v_p$ and all other known variables are kept constant, and we focus on deriving the gradient of $y$ w.r.t. $K^{\text{trans}}$. We use the derivative chain rule:
\[ \frac{\partial y}{\partial K^{\text{trans}}} = \frac{\partial CA(t)}{\partial K^{\text{trans}}} \frac{\partial s(t)}{\partial t} \frac{\partial y}{\partial t} \]

where,

\[ \frac{\partial y}{\partial s(t)} = C^{H} F_{a}^{H} \left[ k_{a}(t) - f(K^{\text{trans}}, v_{p}) \right] \]

\[ \frac{\partial s(t)}{\partial CA(t)} = r_{1} M_{0} \sin \alpha \frac{TR - e^{-TR R_{1}(t)} (1 - e^{-TR R_{1}(t) \cos \alpha}) - (1 - e^{-TR R_{1}(t)} \cos \alpha) \cdot TR - e^{-TR R_{1}(t) \cos \alpha}}{(1 - e^{-TR R_{1}(t) \cos \alpha})^{2}} \]

\[ \frac{\partial CA(t)}{\partial K^{\text{trans}}} = \sum_{t=0}^{T-1} \left[ C_{p}(\tau) d\tau \right] \]

Sparsity-based constraints can be optionally applied to the TK maps as shown in Equation [6]. In this study, we demonstrate the use of wavelet transform, and we denote the wavelet constrained part as \( y_{1} = \| \psi \|_{1} \). For the evaluation of \( y_{1} \), the \( l_{1} \) norm is relaxed as in Lustig et al [13]:

\[ \frac{\partial y_{1}(K^{\text{trans}})}{\partial K^{\text{trans}}} = \lambda_{1} \psi^{H} W^{-1} \psi^{K^{\text{trans}}} \]

And the \( i^{th} \) diagonal element of \( W \) is calculated as:

\[ W_{i} = \sqrt{\left( \psi K^{\text{trans}} \right)_{i} \cdot \left( \psi K^{\text{trans}} \right)_{i} + \mu} \]

where \( \mu \) is a small relaxation parameter.

The gradient for Equation [A.2] is very similar:

\[ \frac{\partial y}{\partial v_{p}} = \frac{\partial CA(t)}{\partial v_{p}} \frac{\partial s(t)}{\partial t} \frac{\partial y}{\partial t} \]

where all other parts are the same as above except:

\[ \frac{\partial CA(t)}{\partial v_{p}} = C_{q}(t) \]

REFERENCES


