High angular resolution diffusion imaging with stimulated echoes: compensation and correction in experiment design and analysis

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Stimulated echo acquisition mode (STEAM) diffusion MRI can be advantageous over pulsed-gradient spin-echo (PGSE) for diffusion times that are long compared with $T_2$. It therefore has potential for biomedical diffusion imaging applications at 7T and above where $T_2$ is short. However, gradient pulses other than the diffusion gradients in the STEAM sequence contribute much greater diffusion weighting than in PGSE and lead to a disrupted experimental design. Here, we introduce a simple compensation to the STEAM acquisition that avoids the orientational bias and disrupted experiment design that these gradient pulses can otherwise produce. The compensation is simple to implement by adjusting the gradient vectors in the diffusion pulses of the STEAM sequence, so that the net effective gradient vector including contributions from diffusion and other gradient pulses is as the experiment intends. High angular resolution diffusion imaging (HARDI) data were acquired with and without the proposed compensation. The data were processed to derive standard diffusion tensor imaging (DTI) maps, which highlight the need for the compensation. Ignoring the other gradient pulses, a bias in DTI parameters from STEAM acquisition is found, due both to confounds in the analysis and the experiment design. Retrospectively correcting the analysis with a calculation of the full $B$ matrix can partly correct for these confounds, but an acquisition that is compensated as proposed is needed to remove the effect entirely. © 2014 The Authors. NMR in Biomedicine published by John Wiley & Sons, Ltd.

Keywords: diffusion MRI; diffusion tensor imaging; HARDI; STEAM; stimulated echo

INTRODUCTION

Stimulated echo acquisition mode (STEAM) diffusion MRI (1,2) offers advantages over the more common pulsed-gradient spin-echo (PGSE) diffusion MRI when $T_2$ is short compared with the diffusion time and $T_1 > T_2$. The STEAM signal decays with rate $T_1$ during the mixing time $T_m$, which determines the diffusion time, whereas $T_2$ decay occurs throughout the whole PGSE sequence. PGSE is preferred for many standard diffusion-weighted imaging (DWI) applications and avoids the initial 50% signal loss inherent in STEAM, where half of the echo is lost. However, STEAM diffusion MRI is common in tissue with short $T_2$, such as muscle or cartilage (3). STEAM is also used in diffusion-weighted spectroscopy (DWS) of metabolites with short $T_2$ (4). STEAM is also useful for measuring diffusivity in water compartments with significantly lower $T_2$, like myelin water, as previously shown in preclinical settings with PGSE (5).

In general, $T_2$ decreases and $T_1$ increases as field strength increases and early evidence (6) suggests benefits of STEAM for in vivo human-brain diffusion tensor imaging (DTI) at 7T. Ex vivo $q$-space studies of brain tissue (7–9) usually prefer STEAM over PGSE, as fixation and lower temperature reduce diffusivity compared with in vivo studies, which increases the necessary diffusion times (10,11). Similarly, Dyrbys et al. (12) demonstrated the need for long diffusion times to ensure sensitivity to large axon diameter, which is important for microstructure imaging techniques such as ActiveAx (13) and AxCaliber (8).

Although STEAM offers benefits over PGSE in obtaining long diffusion times when $T_2$ relaxation is short, gradients other than the diffusion gradients can significantly degrade its potential by introducing a directional diffusion-weighting bias, if not taken into account. The most significant contributions are usually from the crusher and slice-select gradients, here referred to as butterfly gradients, which add unwanted diffusion-weighting ‘cross terms’ in the $B$ matrix (14). In PGSE, their contribution to diffusion weighting is usually negligible in practice, because the diffusion time for the butterfly gradients is only a few milliseconds (the length of the refocusing pulse). In STEAM, that contribution is typically much more significant, because the butterfly gradients are separated by the diffusion time.

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Abbreviations used: DTI, diffusion tensor imaging; DWI, diffusion-weighted imaging; DWS, diffusion-weighted spectroscopy; FA, fractional anisotropy; HARDI, high angular resolution diffusion imaging; PGSE, pulsed-gradient spin-echo; STEAM, stimulated echo acquisition mode; STEAMCOMP, compensated STEAM acquisition.
approximately $\tau_m$ and that is often much longer than for PGSE. Nevertheless, previous work with STEAM diffusion MRI, such as (8,9), follows standard practice for PGSE and neglects this effect. Mattiello et al. (15) derived the full $B$ matrix accounting for all gradient pulses, which predicts the signal exactly on the assumption of Gaussian dispersion, i.e., the diffusion tensor model. However, additional diffusion weighting from other pulses skews the effective diffusion direction, disrupting the experimental design, i.e., the even distribution and strength of the sampled gradient directions usually acquired in DTI and high angular resolution diffusion imaging (HARDI) techniques. This is a critical issue in brain imaging, where diffusion is often strongly anisotropic and non-uniform gradient orientations can lead to significant orientational bias in the precision of derived diffusion metrics (16).

In this article, we propose a simple compensation for the STEAM sequence, referred to as compensated acquisition, that accounts for the unwanted directional bias caused by the butterfly gradients. The implementation of the compensated acquisitions is simple and only requires a correction of the gradient vectors loaded on the scanner. We show how this compensation effectively cancels the effects of the butterfly gradients, so that the resulting data sets can be treated as if they came from an idealized HARDI protocol, i.e., ignoring the butterfly gradients.

We demonstrate the need and effectiveness of the compensation for STEAM through HARDI-DTI experiments in simulation and on data acquired from a fixed monkey brain. Numerical experiments show that ignoring the butterfly gradients in STEAM leads to severe bias in the fitted diffusion tensor and derived quantities. The full $B$-matrix formulation from (15) reduces the bias, but some still remains because of the disruption to the intended even distribution of gradient directions. However, the compensated acquisition reduces bias even further and simplifies the calculation by allowing us to ignore the cross terms in the $B$ matrix.

**METHODS**

**STEAM pulse sequence**

The compensation in subsequent sections assumes the idealized STEAM pulse sequence in Figure 1. We refer to this figure for nomenclature. The layout is very similar to the conventional PGSE sequence, with diffusion-encoding gradients $G_d$ on each side of the refocusing pulse. All gradients working on the signal pathway in the transversal plane introduce a diffusion weighting. In our case, the major sources, in addition to $G_d$, are a crusher pulse $G_c$ and a slice selection pulse $G_s$. For a conventional sinc RF pulse, the phase modulation corresponds to half the area of the slice selective gradient. In a practical imaging setup, the crusher, or the same effect of the diffusion encoding gradient, is also needed to isolate the original coherence trajectory (17). We will refer to data with $|G_d|=0$ as nominal $b=0$ measurements.

**Diffusion tensor imaging and STEAM**

On the assumption of zero-mean Gaussian particle dispersion, i.e., the diffusion tensor (DT) model, the general formula (18):

$$S = S_0 \exp(-B \cdot D)$$

predicts the signal, where $B = \gamma^2 \int F(t)F^T(t)dt$ is the $B$ matrix (14),

$$F(t) = \int_0^t G(t)dt$$

$G(t)$ is the effective gradient vector at time $t$, $\gamma$ is the gyromagnetic ratio, $G_d$ is a unit vector in the direction of $G_d$ and $S_0$ is the signal with $b=0$. A common approach is to assume that $G_c$ and $G_s$ are negligible, so that we need only consider $G_d$, which reduces Equation [1] to

$$S = S_0 \exp(-b G_d^T DT G_d)$$

where

$$b = (\Delta - \delta_d/3)(\gamma \delta_d |G_d|^2)$$

and

$$\Delta = \tau_m + \delta_d + 2\delta_s + 2\delta_c + \tau_1 + \tau_2$$

is the separation between the diffusion gradients in the STEAM sequence. However, including the contribution from all relevant gradients in Figure 1 gives

$$B = \gamma^2 (\delta_d^2 \tau_{dd} G_d G_d^T + \delta_c^2 \tau_{cc} G_c G_c^T + \delta_s^2 \tau_{ss} G_s G_s^T + \delta_{dc} \tau_{dc} (G_d G_c^T + G_c G_d^T))$$

where

$$\begin{align*}
\tau_{dd} &= \tau_1 + \tau_2 + \tau_m + 2\delta_d + 2\delta_c/3 + 2\delta_s, \\
\tau_{cc} &= \tau_m + 2\delta_c/3 + 2\delta_s, \\
\tau_{ss} &= \tau_m + 2\delta_s/3, \\
\tau_{dc} &= \tau_m + \delta_c + 2\delta_s, \\
\tau_{ds} &= \tau_m + \delta_c, \\
\tau_{cs} &= \tau_m + \delta_s
\end{align*}$$

The expression in Equation [6] is the sum of pairwise interactions between the diffusion, crusher and slice-select pulses, similar to the $B$ matrix for PGSE in (15).
Compensated acquisition correction

The goal of the compensation is to cancel out the effect of the butterfly gradients by modifying the diffusion gradient directions. A simple correction finds the adjustment of $G_d$ that minimizes the diffusion weighting of the nominal primary eigenvector of the separated gradient components with different orientation (15). A1 also uses Equation [3], but with the contributions from $G_i$ and $G_d$ produces the approximated effective gradient vector $G_d'$ close to $G_i$. The simple compensation we propose replaces each $G_d$ in the list of gradient vectors we intend to acquire with

$$G_d' = G_d - \delta_t \tau_d (\delta_d \tau_d)^{-1} G_c - \delta_t \tau_d (\delta_d \tau_d)^{-1} G_c$$  \[8\]

Equation [8] comes from setting $G_d = g G_i + h G_s$ and minimizing the trace of the $B$ matrix in Equation [6] with respect to $g$ and $h$ to obtain $g = -\delta_t \tau_d (\delta_d \tau_d)^{-1}$ and $h = -\delta_t \tau_d (\delta_d \tau_d)^{-1}$. The weightings $g$ and $h$ depend only on the timings of the pulses so are constant within one HARDI shell, but may vary between shells or measurements with different $b$ value or diffusion time.

Another choice of $G_d$ is $(b_1/(\delta_d \tau_d))^{1/2} G_i$, where $G_i$ is the primary eigenvector of the $B$ matrix and $b_1$ is the corresponding eigenvalue. However, the two choices for $G_d$ are very similar in practice. The former is simpler to compute, while the latter applies to more general gradient configurations where the $B$ matrix is more easily assessed by numerical integration.

In practice, implementation of the compensated acquisition simply requires an adjustment to the gradient direction scheme and corresponding $b$ values or gradient strengths uploaded to the scanner. Precise implementations may differ among vendors. Also note that the gradient strength after compensation may exceed the maximum available gradient strength. A solution to this is simply to negate the direction of the intended direction, as the compensation is an additive vector.

Post-processing correction

We consider three approximations to the signal that account for the butterfly gradients in different ways.

- Approximation 1 (A1) is the approach generally used in DTI analysis of PGSE. It ignores butterfly gradients and considers only diffusion gradients.
- Approximation 2 (A2) accounts for the butterfly gradients in a simple but non-trivial way by identifying an effective diffusion gradient $G_d'$ that incorporates the diffusion weighting of the diffusion and butterfly gradients. Our choice comes from inverting Equation [8]:

$$G_d' = G_d - \delta_t \tau_d (\delta_d \tau_d)^{-1} G_c - \delta_t \tau_d (\delta_d \tau_d)^{-1} G_c$$  \[9\]

- Approximation 3 (A3) calculates the full $B$ matrix as in Equation [6] and accounts for cross $b$ does not.

A1 also uses Equation [3] directly to fit the diffusion tensor. A2 also uses Equation [3], but with $G_d'$ from Equation [9] replacing $G_d$. A3 uses the full $B$ matrix, analogous to (15) for PGSE, rather than the single $b$ value and gradient direction in A1 and A2.

By assuming a single $b$ value, A1 and A2 ignore the cross terms in the $B$ matrix, which express the interaction between temporally separated gradient components with different orientation (15). A1 is exact only when $G_c = G_s = 0$. A2 is exact only when $G_d$, $G_c$, and $G_s$ all have the same orientation. A3 accounts for all cross terms, so is always exact for the Gaussian dispersion assumed in DTI. In the absence of butterfly gradients, A3 and A2 are equivalent to A1. An overview of the parameter definitions and signal models is given in Table 1.

EXPERIMENTS

DWI protocols

The experiments compare three kinds of imaging protocol: (i) STEAM protocol, which does not make the adjustment in Equation [8] but uploads the intended list of directions and $b$ values to the scanner; (ii) compensated STEAM acquisition (STEAMCOMP) protocol, which does include the adjustment in Equation [8]; and (iii) PGSE protocol. The PGSE was included to provide principal direction estimates minimally affected by the butterfly gradients for comparison with those from STEAM. All protocols are representative of real-world applications and were in fact optimized for axon diameter mapping of post-mortem brain tissue using ActiveAx; see (19) for details. For the PGSE, two protocols were chosen with high and low gradient strength. Table 2 shows the settings for the preclinical protocols used.

Every image in STEAM and STEAMCOMP has $\delta_1 = 1.5$ms, $G_s = (0, 0, 0.15)\text{mT}\cdot\text{m}^{-1}$, $\delta_2 = 1$ms, $G_c = (0, 0, 0.15)\text{mT}\cdot\text{m}^{-1}$ and $\tau_2 = 0$.

Since $G_s$ and $G_c$ are both along the slice direction (0, 0, 1), the compensation $G_s - G_d$ from Equation [8] is along the negative slice direction; $|G - G_d| = 68.5\text{mT}\cdot\text{m}^{-1}$.

Imaging experiments

We acquire data from a fixed monkey brain, prepared as in (10). The live monkey was handled and cared for on the Island of St Kitts according to a protocol approved by the local ethics committee (The Caribbean Primate Center of St Kitts). Data were acquired on a Varian 4.7T preclinical scanner using a quadrature volume coil. All datasets were acquired at 0.5 mm$^3$ isotropic resolution with a 256 × 128 matrix with 15 contiguous sagittal slices including the mid-sagittal plane.

Simulation experiments

The simulation experiments use two DTs, one with eigenvalues $[0.6, 0.2, 0.2] \times 10^{-9}$ m$^2$ s$^{-1}$, which are typical of coherent white matter in fixed brain tissue at this $b$ value, and the other

| Table 1. Short descriptions of (a) gradient definitions used in Equations [2]–[9] and (b) signal models used in the analysis |
|---|
| Name | Description |
| (a) | $G(t)$ The effective gradient time vector including all gradients. |
| $G_d$ | The applied diffusion encoding gradient vector. |
| $G_c$ | The crusher gradient vector. |
| $G_s$ | The slice gradient vector. |
| $G_l$ | The intended diffusion encoding gradient vector. |
| $G_d'$ | The effective gradient vector that with the contributions from $G_c$ and $G_s$ approaches $G_d$ after compensation. |
| (b) | A1 Assuming diffusion weighting from $G_d$. |
| A2 | Assuming diffusion weighting from $G_d'$. |
| A3 | Full $B$-matrix calculated from $G_d$, $G_s$ and $G_c$. |
CONSIDERATIONS FOR DIFFUSION EXPERIMENTS WITH STEAM MRI

Table 2. (a) PGSE and (b) STEAM protocols. Both come from the experiment design optimisation in (12,25) with $G_{\max} = 300 \text{mTm}^{-1}$. $N$ is the number of diffusion-weighted images in each shell. $K$ is the number of nominal $b = 0$ images associated with each shell. The nominal $b = 0$ images in STEAM have the same $\tau_m$, as the diffusion-weighted images. The compensated STEAM protocol STEAMCOMP follows (b), but replaces each $G_i$ according to Equation [8]. Please refer to Figure 1 for notation.

| $N$ | $K$ | $|G_d|/\text{mTm}^{-1}$ | $\Delta/\text{ms}$ | $\delta_d/\text{ms}$ | $b/\text{smm}^{-2}$ | $\tau_o/\text{ms}$ | $\tau_r/\text{ms}$ |
|-----|-----|----------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| (a) | 103 | 25 | 300.0 | 12.9 | 5.6 | 2243 | 36.8 | 2600 |
| (b) | 106 | 25 | 219.2 | 20.4 | 7.0 | 3084 | 36.8 | 2600 |
| (b) | 108 | 25 | 113.5 | 137.0 | 5.0 | 3425 | 26.0 | 2600 |

The butterfly gradients are weaker than for the ex vivo protocol, because the voxel size is larger (2 mm isotropic). The test DTs have eigenvalues $\{1.7,0.2,0.2\} \times 10^{-9} \text{m}^2 \text{s}^{-1}$ and $\{0.7,0.7,0.7\} \times 10^{-9} \text{m}^2 \text{s}^{-1}$.

Data analysis

We fit the DT to the STEAM, STEAMCOMP and PGSE data using weighted linear least-squares and construct colour-coded FA maps (22). We quantify the orientational similarity between pairs of DT volumes by computing the mean over the brain of the absolute dot product of principal directions weighted by DT linearity (23).

RESULTS

HARDI gradient direction file

Figure 2 shows the distribution of effective gradient directions, i. e. the orientation of $G_d^*$ from Equation [9], for STEAM and STEAMCOMP to illustrate the disruption to the HARDI design. Without compensation, the butterfly gradients skew the effective gradient directions strongly towards the slice direction. The true $b$ values of the uncompensated gradient scheme ranged from 660–8830 s mm$^{-2}$ with a maximum deflection from the intended direction of $36^\circ$ and the mean deflection over all directions was $22^\circ$. With the crusher gradients turned off, the $b$ values still ranged from 2075–5160s mm$^{-2}$ with a maximum deflection of $13^\circ$ and the mean deflection over all directions was $9^\circ$. The compensated protocol has evenly distributed effective gradient directions and a constant $b$ value.

The nominal $b = 0$ were not compensated. As an indication of the impact of butterfly gradients, the $b$ value from them alone, as

Figure 2. Illustration of the target and effective gradient directions in the STEAM protocols using the 108 directions in the STEAM protocol. A black cross marks each direction; shaded crosses are on the far side of the sphere. Panel (a) shows the target set with a cross in both positive and negative gradient directions. Panel (b) shows the set of effective gradient directions, i.e. the direction of $G_d^*$ in Equation [9], without compensation (STEAM); they skew strongly towards the slice direction. Panel (c) shows the effective gradient directions after compensation (STEAMCOMP); these are close to the target set.
in the nominal $b = 0$, is $1316 \text{s mm}^{-2}$ for the preclinical STEAM protocol. The same figure for the PGSE protocol is $17 \text{ s mm}^{-2}$.

Simulation experiments

Tables 3 and 4 list statistics for the fixed-tissue simulations with anisotropic and isotropic DTs, respectively. We show the two extreme cases, when the principal eigenvector of the DT is perpendicular and parallel to the slice direction. Note that perfect compensation makes $A1$ and $A2$ equivalent. Without compensation, $A1$ shows significant bias in FA, $\lambda_1$ and $\mathbf{e}_1$ with both orientations of the anisotropic DT. Bias is most severe for $\mathbf{e}_1$ parallel to the butterfly gradients. In this case, the nominal $b = 0$ acquisition has higher diffusion weighting and larger signal attenuation than the intended diffusion weighting and larger signal attenuation than the intended diffusion-weighted signal. Here, DT estimation with $A1$ completely fails, producing negative eigenvalues and artificially high FA.

Estimates of the isotropic DT show artefactual non-zero FA and significant direction concentration: $\gamma = 1.5$ means 95% of directions are within 6° of the mean. Compensation dramatically improves $A1$. Some downward bias remains in both FA and $\lambda_1$ of the anisotropic DTs, but the bias is similar for both orientations. Compensation largely removes artefactual non-zero FA and orientational bias in the isotropic DT estimates: $\gamma = 0.4$ is typical for a uniformly distributed random sample of 10 000 directions and the 95% angle is over 25°.

Without compensation, $A2$ and $A3$ produce very similar results. Both significantly reduce bias compared with $A1$, although bias remains orientationally dependent and is strongest with parallel $\mathbf{G}_1$ and $\mathbf{G}_2$. Compensation reduces the bias and variance of parameter estimates from $A3$, especially for parallel $\mathbf{e}_1$, and removes orientational dependence. With compensation, $A3$ shows no benefit over $A1$ or $A2$. Tables 5(a) and (b) are equivalent to Tables 3 and 4, but with the crisper gradients turned off. The biases before compensation are smaller but still significant and compensation improves consistency and reduces orientational dependence.

Tables 6 and 7 show the corresponding results from the in vivo human protocol. Without compensation, $A1$ still produces considerable bias, which $A2$ or $A3$ reduces. The compensation provides only minor further improvements with $A3$ compared with $A2$.

Imaging experiments

Figure 3 compares maps from PGSE with STEAM for each approximation qualitatively. The number next to each STEAM map is the orientational similarity with the $b = 3084 \text{s mm}^{-2}$ shell PGSE; higher numbers show greater agreement. The number next to the PGSE map is the orientational similarity of the $b = 2243 \text{s mm}^{-2}$ and $b = 3084 \text{s mm}^{-2}$ shells of PGSE.

For STEAM, $A1$ introduces upward bias in FA in the superior half of the brain, where diffusion should be close to isotropic, such as the area in the cyan box on the PGSE map. The maps also show orientation bias towards the left–right slice direction (the map appears red) due to the directional bias towards the direction of the butterfly gradients shown in Figure 2. The white

| DT orientation | Analysis (FA, $\lambda_1$, $\alpha$, $\gamma$) | A1 | A2 | A3 | A1/A2 | A3 |
|----------------|---------------------------------------------|----|----|----|--------|----|
| FA             | 0.514                                      | 0.884 | 0.573 | 0.495 | 0.573 | 0.495 | 0.575 | 0.574 | 0.575 | 0.574 |
| std            | 0.035                                      | 0.166 | 0.025 | 0.042 | 0.025 | 0.042 | 0.020 | 0.021 | 0.020 | 0.021 |
| $\lambda_1$    | 5.518                                      | 1.522 | 5.278 | 3.377 | 5.278 | 3.377 | 5.566 | 5.544 | 5.566 | 5.544 |
| Std            | 0.262                                      | 0.270 | 0.212 | 0.185 | 0.212 | 0.185 | 0.161 | 0.165 | 0.161 | 0.165 |
| $\alpha$       | 4.670                                      | 63.714 | 2.537 | 5.058 | 2.537 | 5.058 | 1.946 | 1.992 | 1.946 | 1.992 |
| $\gamma$       | 5.309                                      | 1.909 | 6.238 | 5.362 | 6.238 | 5.362 | 6.766 | 6.719 | 6.766 | 6.719 |

Table 3. Statistics from simulations with anisotropic DTs for each approximation using the preclinical STEAM protocol and a SNR of 20. Two simulated datasets were created, one with the DT perpendicular (⊥) and one with the DT parallel (∥) to the butterfly gradient direction. The units of $\lambda_1$ are $10^{-10} \text{m}^2 \text{s}^{-1}$; the units of $\alpha$ are degrees. The true FA is 0.603 and the true $\lambda_1$ is $6 \times 10^{-10} \text{m}^2 \text{s}^{-1}$. Higher $\gamma$ is better in this experiment.

| Analysis (FA, $\lambda_1$, $\alpha$, $\gamma$) | Uncompensated | Compensated |
|---------------------------------------------|---------------|-------------|
| A1/A2 | A3 | A1/A2 | A3 |
| FA    | 0.058 | 0.058 | 0.019 | 0.019 |
| std   | 0.019 | 0.019 | 4.023 | 4.023 |
| $\lambda_1$ | 3.806 | 3.658 | 3.806 | 3.658 |
| std   | 0.305 | 0.171 | 0.171 | 0.058 |
| $\alpha$ | 1.504 | 0.846 | 0.846 | 0.846 |
| $\gamma$ | 1.504 | 0.846 | 0.846 | 0.846 |

Table 4. Statistics, as in Table 3, from simulations with isotropic DTs. The true FA is 0; the true $\lambda_1$ is $4 \times 10^{-10} \text{m}^2 \text{s}^{-1}$. Here $\gamma$ should be zero.

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boxes show bias in anisotropic regions: the left box shows severely biased orientation estimates (some voxels appear green rather than red) in the corpus callosum, where the fibres are parallel to the butterfly gradients; the right box shows less biased orientation estimates in the fornix, which has perpendicular fibres. A2 and A3 are qualitatively indistinguishable from one another and are more consistent with the PGSE map than A1, e.g. in the white boxes. However, they still show upward bias in FA together with consistent artefactual orientation in isotropic regions in the cortex (blue/green colour in cyan box region). Two directions exceeded $G_{\text{max}}$ after compensation and were truncated to $G_{\text{max}}$ by the scanner. This imperfection in compensation leads to the difference between the compensated results in A1 versus A2 and A3.

### Table 5. Statistics from simulations for each approximation using the preclinical STEAM protocol for (a) anisotropic DT and (b) isotropic DT. The experiments are similar to the imaging and simulation experiments presented in Tables 3 and 4, but with the crusher gradients turned off. The biases before compensation are only caused by the slice gradient and thus smaller, but consistency is still improved after compensation.

| Gradients | Analysis | Uncompensated | Compensated |
|-----------|----------|---------------|-------------|
|           | A1       | A2            | A3          | A1/A2     | A3          |
| (a)       |          |               |             |           |             |
| DT orientation |          |               |             |           |             |
| FA        | 0.498    | 0.671         | 0.575       | 0.534     | 0.575       | 0.534       | 0.577       | 0.576       | 0.576       |
| std       | 0.024    | 0.025         | 0.021       | 0.024     | 0.021       | 0.024       | 0.020       | 0.020       | 0.020       |
| $\lambda_1$ | 5.576    | 6.426         | 5.512       | 5.071     | 5.512       | 5.071       | 5.579       | 5.574       | 5.574       |
| std       | 0.181    | 0.333         | 0.173       | 0.175     | 0.173       | 0.175       | 0.160       | 0.160       | 0.160       |
| $\alpha$  | 2.602    | 5.874         | 1.982       | 2.451     | 1.982       | 2.451       | 1.889       | 1.922       | 1.922       |
| $\gamma$  | 6.212    | 6.260         | 6.730       | 6.351     | 6.730       | 6.351       | 6.825       | 6.790       | 6.790       |
| (b)       |          |               |             |           |             |
| FA        | 0.210    | 0.074         | 0.074       |           | 0.057      | 0.057       |
| std       | 0.032    | 0.023         | 0.023       |           | 0.018      | 0.018       |
| $\lambda_1$ | 5.168    | 3.981         | 3.981       |           | 4.025      | 4.025       |
| std       | 0.212    | 0.119         | 0.119       |           | 0.099      | 0.099       |
| $\gamma$  | 3.843    | 0.612         | 0.612       |           | 0.415      | 0.415       |

### Table 6. Simulation statistics for anisotropic diffusion with the human protocol. Two simulated datasets were created, one with the DT perpendicular ($\perp$) and one with the DT parallel ($||$) to the butterfly gradient direction. The true FA is 0.87 and the true $\lambda_1$ is $17 \times 10^{-10}$ m$^2$ s$^{-1}$.

| Gradients | Analysis | Uncompensated | Compensated |
|-----------|----------|---------------|-------------|
|           | A1       | A2            | A3          | A1/A2     | A3          |
| DT orientation |          |               |             |           |             |
| FA        | 0.899    | 0.859         | 0.861       | 0.864     | 0.861       | 0.864       | 0.863       | 0.863       | 0.863       |
| std       | 0.019    | 0.017         | 0.018       | 0.015     | 0.018       | 0.015       | 0.017       | 0.017       | 0.017       |
| $\lambda_1$ | 14.807   | 16.203        | 15.928      | 16.260   | 15.928      | 16.260      | 16.263      | 16.269      | 16.269      |
| std       | 0.581    | 0.533         | 0.610       | 0.548     | 0.610       | 0.548       | 0.532       | 0.521       | 0.521       |
| $\alpha$  | 19.628   | 3.862         | 1.502       | 1.493     | 1.502       | 1.493       | 1.391       | 1.390       | 1.390       |
| $\gamma$  | 7.132    | 7.380         | 7.288       | 7.300     | 7.288       | 7.300       | 7.437       | 7.437       | 7.437       |

### Table 7. Simulation statistics for isotropic diffusion with the human protocol. The true FA is 0 and the true $\lambda_1$ is $7 \times 10^{-10}$ m$^2$ s$^{-1}$.

| Gradients | Analysis | Uncompensated | Compensated |
|-----------|----------|---------------|-------------|
|           | A1       | A2            | A3          | A1/A2     | A3          |
| FA        | 0.356    | 0.098         | 0.098       |           | 0.095      | 0.095       |
| std       | 0.042    | 0.033         | 0.033       |           | 0.031      | 0.031       |
| $\lambda_1$ | 5.433    | 4.313         | 4.313       |           | 4.311      | 4.311       |
| std       | 0.305    | 0.254         | 0.254       |           | 0.238      | 0.238       |
| $\gamma$  | 3.852    | 0.462         | 0.462       |           | 0.414      | 0.414       |
Simulation and fixed-brain data experiments demonstrated that, without the compensation and using the standard DTI model (A1 uncompensated), severe bias can arise in estimated DTI parameters. Accounting for cross terms in the $B$ matrix either exactly (A3 uncompensated) or approximately (A2 uncompensated) reduces that bias considerably; the lack of performance difference between A2 and A3 shows that the minor eigenvalues of the $B$ matrix are negligible. However, including the compensation at acquisition time results in further reductions in bias (e.g., A3 compensated versus A3 uncompensated), highlighting the importance of the compensation. Differences in results from A3 with and without compensation at acquisition show the effect of the experiment design disruption, which is seen in both simulation and real data. The disruption to the experiment design affects parameter estimates most strongly with parallel diffusion and butterfly gradients, because the additional diffusion weighting in the fibre direction pushes parallel signals further into the noise floor. The effect for an arbitrary tensor orientation is in between the two extreme cases explored by the simulation with the tensor aligned parallel or perpendicular to the slice direction. Compensation removes the experiment design disruption and thereby improves the precision in parameter estimates and removes the orientational dependence. Moreover, A1, A2 and A3 after compensation provide similar results, showing that the compensated acquisition largely eliminates the cross terms in the $B$ matrix, thus enabling the use of A1. This is a significant advantage, as A1 is simple to implement and compute and is used in the majority of DTI and HARDI reconstruction software tools. A2 and A3 are useful in their own right for analysis of existing data without the compensation. In addition to HARDI techniques, single-direction model-based STEAM diffusion MRI applications, such as (8,9), are also likely to benefit significantly from the proposed compensation.

In simulations we observe a bias in FA and $\lambda_1$ from A3 with compensation, which is unavoidable, since the model is exact and the experiment design is not disrupted. It comes from Jones’ ‘squashed-peanut’ effect (20): a Rician noise effect as measurements with gradient parallel to $\mathbf{e}_1$ approach the noise floor.

The performance differences are less marked in the human protocol, because the butterfly gradients are smaller due to the lower resolution. However, the values of $\alpha$ between 3.8° and 19.6° that we observe for A1 without compensation are at least as large as the orientational bias incurred by failing to account for small head motions in the $B$ matrix, which (24) finds sufficient to disrupt tractography.

Although differences in DTI maps certainly arise between PGSE and STEAM acquisitions, we expect the similarity in principal directions to be similar enough to highlight gross errors introduced by incorrect modelling or lack of compensation. The butterfly gradients affect the nominal $b=0$ images, as well as the diffusion-weighted images. In the absence of a strong diffusion gradient, the compensation counteracts the effect of crusher gradients, allowing additional echoes to affect the signal and leading to severe image artefacts. Thus the nominal $b=0$ images remain uncompensated with $G_0=0$, but this is solved with a two-point fit to the non-normalized data. The $b$ value in the nominal $b=0$ images defines the minimal diffusion weighting possible with sufficient crusher effect for the given image resolution. For example, in our $b=3425$ s mm$^{-2}$ shell that value is 1316 s mm$^{-2}$ . In the diffusion-weighted images, $G_w$ works in principle as a crusher itself. In the absence of $G_w$, the compensation must only correct for the contribution from $G_0$.
CONCLUSIONS

We demonstrate here that imaging gradients in the STEAM sequence can severely disrupt HARDI experiment design in a way not seen in PGSE and cause bias in parameter estimates. We introduce a compensation in the acquisition by simply subtracting the diffusion weighting of imaging gradients from the diffusion encoding gradients. This is simply implemented by updating the gradient scheme loaded on the scanner. Furthermore, models are presented to handle these problems retrospectively. These methods allow future work on and exploitation of the potential benefits of STEAM, especially for diffusion MRI on high-field scanners, where the lower effective $T_2$ prevents long diffusion times in PGSE.

In particular, they enable us to evaluate STEAM ActiveAx for better sensitivity to large axons, which is the focus of our current work.

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