

Contents lists available at ScienceDirect

Magnetic Resonance Imaging



journal homepage: www.elsevier.com/locate/mri

Chaos and COSMOS—Considerations on QSM methods with multiple and single orientations and effects from local anisotropy



Dimitrios G. Gkotsoulias ^{a,*}, Carsten Jäger ^b, Roland Müller ^a, Tobias Gräßle ^c, Karin M. Olofsson ^d, Torsten Møller ^e, Steve Unwin ^f, Catherine Crockford ^{g,h,i}, Roman M. Wittig ^{g,h,i}, Berkin Bilgic ^{j,k,l}, Harald E. Möller ^a

^a Nuclear Magnetic Resonance Methods & Development Group, Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, Germany

^b Department of Neurophysics, Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, Germany

^c Epidemiology of Highly Pathogenic Microorganisms, Robert Koch-Institute, Berlin, Germany

^d National Veterinary Institute, Uppsala, Sweden

^e Kolmården Wildlife Park, Norrköping, Sweden

^f Wildlife Health Australia, Canberra, Australia

⁸ Department of Human Behavior, Ecology and Culture, Max Planck Institute for Evolutionary Anthropology, Leipzig, Germany

^h The Ape Social Mind Lab, Institut des Sciences Cognitives Marc Jeannerod, Bron, France

ⁱ Taï Chimpanzee Project, Centre Suisse de Recherches Scientifiques, Abidjan, Côte d'Ivoire

^j Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Boston, MA, USA

^k Harvard Medical School, Boston, MA, USA

¹ Harvard/MIT Health Sciences and Technology, Cambridge, MA, United States

ARTICLE INFO

Keywords: Anisotropic magnetic susceptibility COSMOS Gradient-recalled echo Quantitative susceptibility mapping

ABSTRACT

Purpose: Field-to-susceptibility inversion in quantitative susceptibility mapping (QSM) is ill-posed and needs numerical stabilization through either regularization or oversampling by acquiring data at three or more object orientations. Calculation Of Susceptibility through Multiple Orientations Sampling (COSMOS) is an established oversampling approach and regarded as QSM gold standard. It achieves a well-conditioned inverse problem, requiring rotations by 0°, 60° and 120° in the yz-plane. However, this is impractical *in vivo*, where head rotations are typically restricted to a range of $\pm 25^{\circ}$. Non-ideal sampling degrades the conditioning with residual streaking artifacts whose mitigation needs further regularization. Moreover, susceptibility anisotropy in white matter is not considered in the COSMOS model, which may introduce additional bias. The current work presents a thorough investigation of these effects in primate brain.

Methods: Gradient-recalled echo (GRE) data of an entire fixed chimpanzee brain were acquired at 7 T (350 µm resolution, 10 orientations) including ideal COSMOS sampling and realistic rotations *in vivo*. Comparisons of the results included ideal COSMOS, *in-vivo* feasible acquisitions with 3–8 orientations and single-orientation iLSQR QSM.

Results: In-vivo feasible and optimal COSMOS yielded high-quality susceptibility maps with increased SNR resulting from averaging multiple acquisitions. COSMOS reconstructions from non-ideal rotations about a single axis required additional L2-regularization to mitigate residual streaking artifacts.

Conclusion: In view of unconsidered anisotropy effects, added complexity of the reconstruction, and the general challenge of multi-orientation acquisitions, advantages of sub-optimal COSMOS schemes over regularized single-orientation QSM appear limited in *in-vivo* settings.

1. Introduction

Local alterations of iron and myelin in the brain are often linked to

pathologies, making their quantitation an important goal for diagnostic and neuroscientific imaging. Given the distinct magnetic properties of iron-containing compounds (typically paramagnetic) and myelin

* Corresponding author at: Max Planck Institute for Human Cognitive and Brain Sciences, Stephanstraße 1a, 04103 Leipzig, Germany. *E-mail address:* gkotsoulias@cbs.mpg.de (D.G. Gkotsoulias).

https://doi.org/10.1016/j.mri.2024.04.020

Received 22 February 2024; Received in revised form 7 April 2024; Accepted 14 April 2024 Available online 16 April 2024 0730-725X/© 2024 The Author(s). Published by Elsevier Inc. This is an open access article under the

0730-725X/© 2024 The Author(s). Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

(diamagnetic), alterations in their local concentration cause spatial variations of the bulk magnetic susceptibility $\Delta \chi$ [1]. Quantification methods employing magnetic resonance imaging (MRI) to obtain local $\Delta \chi$ estimates have, therefore, been proposed, in particular, quantitative susceptibility mapping (QSM)[2-4]. In QSM, susceptibility is estimated indirectly from frequency shifts measured by the gradient echo (GRE) signal phase φ [5,6]. The processing pipeline includes the estimation of the full phase evolution using phase-unwrapping techniques, the removal of background-field contributions from the local magnetic field variation δB_0 and, finally, a field-to-source inversion method to extract $\Delta \chi$ from δB_0 under the strict assumption that $\Delta \chi$ within a voxel is a scalar, isotropic quantity. Two basic issues arise for QSM: (i) The inverse problem from local tissue phase to susceptibility is intrinsically ill-posed because the dipole response kernel has near-zero values in the vicinity of two conical surfaces in the Fourier domain. Therefore, a direct inversion is ill conditioned at such positions, and hence, additional assumptions are required for avoiding artifacts in the final estimation. (ii) Magnetic susceptibility-especially in white matter (WM)-exhibits anisotropic characteristics as a result of the specific liquid-crystalline arrangement of elongated lipid molecules within the myelin sheaths enveloping axons. This inherent anisotropy is typically ignored in QSM pipelines [6,7].

Most QSM implementations employ post-acquisition regularization solutions to the first problem, for example Morphology Enabled Dipole Inversion (MEDI) [8], FAst Nonlinear Susceptibility Inversion (FANSI) [9] and iLSQR-QSM [10]. iLSQR-QSM is generally considered to be a method that uses minimum a-priory regularizations while still solving residual issues existing in more basic methodologies, such as Thresholded k-space Division (TKD) [11]. Calculation Of Susceptibility through Multiple Orientation Sampling (COSMOS) [2] has been introduced as an acquisition-based solution. It requires sampling at three or more orientations of the object inside the magnetic field. An analytical investigation of the optimal scheme suggests that rotations by $0^\circ,\,60^\circ$ and 120° about the magnet's (physical) x-axis eliminate the chaos $(\chi \dot{\alpha} o \zeta')^1$ of conic surfaces with zeroes and permit the estimation of $\Delta \chi$ without further regularization. With this choice of orientations, COSMOS fully addresses the ill-posed inverse problem yielding susceptibility maps that should be free of streaking artifacts. Considering the second problem, by sampling optimally along a circle, susceptibility anisotropy is partially averaged out (at least along the direction of sample rotation), leading to a more robust $\Delta \gamma$ estimate. Results obtained with COSMOS have, therefore, been used previously as gold standard susceptibility reference in evaluations of QSM processing pipelines [12]. However, the application of COSMOS in human brain imaging in vivo is challenging because achievable head rotation angles are limited to a small range, for example

 $\pm 25^{\circ}$ or less [13–17], which deviates substantially from the ideal rotation scheme. To address the limitation of a restricted range of accessible angles, head rotations about two axes have also been advocated as alternative to the optimal COSMOS scheme [18].

In this work, we used high-resolution 7 T GRE acquisitions in *post-mortem* chimpanzee brain as a versatile and robust setup for a further evaluation of COSMOS [19]. In particular, we compared the optimal scheme to results obtained with a more realistic range of accessible angles *in vivo* in the human brain and with single-orientation iLSQR-QSM. Acquisitions using more than three orientations were also included in these comparisons to evaluate the potential impact from more expanded orientation information (or averaging) in $\Delta \chi$ estimations. Finally, residual effects from susceptibility anisotropy on the obtained maps were evaluated.

2. Methods

2.1. Brain specimen

The specimen used for all acquisitions was obtained post-mortem from a female chimpanzee (Pan troglodytes verus) from Kolmården Wildlife Park, Sweden. The chimpanzee was medically euthanized due to a cervical leiomyoma and secondary hydronephrosis. All procedures followed the ethical guidelines of primatology research at the Max Planck Institute for Evolutionary Anthropology, Leipzig, and were approved by the Ethics Committee of the Max Planck Society. Exhaustive details of the steps involved in the brain extraction, fixation and storage have been published by Gräßle et al. [20] Briefly, a specialized veterinarian performed the brain extraction within 18 h after death. The tissue was preserved in 4% paraformaldehyde (PFA) in phosphatebuffered saline (PBS). For the MRI acquisitions, the PFA was washed out in PBS at pH 7.4. The specimen was then immersed in Fomblin® (Solvay Solexis, Bollate, Italy) and positioned in an individualized, anatomically-shaped, 3D-printed container to avoid gravity-induced non-linear deformations during reorientation. Further details of this setup are given by Gkotsoulias et al., 2023 [21].

2.2. Multi-orientation phase measurements

Complex-valued 3D Fast Low-Angle SHot (FLASH) [22] data were obtained at 7 T on a MAGNETOM Terra (Siemens Healthineers, Erlangen Germany) with a single-channel transmit/32-channel receive head coil (Nova Medical, Wilmington, MA, USA). The acquisition parameters were optimized for high-resolution post-mortem acquisitions (field of view $160 \times 160 \times 145 \text{ mm}^3$, matrix size $458 \times 458 \times 416 \text{voxels}$, 0.35 mm isotropic nominal resolution, transverse orientation) based on prior experience [23,24]. To account for T_1 and T_2^* shortening in fixed tissue, a radiofrequency (RF) pulse flip angle of $\alpha = 28^{\circ}$ was combined with a relatively short repetition time (TR = 26 ms) for efficient sampling (bandwidth 180 Hz/pixel). The echo time of TE = 13 ms was long enough to achieve sufficient phase evolution and, at the same time, a good contrast-to-noise ratio between regions of different microstructural properties. The phase encoding (PE) direction was from right to left (here, along the z-axis of the laboratory frame). GeneRalized Autocalibrating Partially Parallel Acquisitions (GRAPPA) [25] with acceleration factor R = 2 and a partial-Fourier scheme [26] with partial-Fourier factor, $f_p = 7/8$ were employed in PE direction to accelerate the measurements (acquisition time, $TA \approx 34$ min per orientation). A comprehensive list of the scanning parameters is given in Supplementary Table S1. In total, ten measurements were obtained, starting with the natural supine reference position in the scanner (coordinate system with axis-in terms of the subject-from left to right, posterior to anterior, and inferior to superior) and followed by rotating the specimen by angles as given in Table 1 (also shown in Supplementary Fig. S1). After every two scans, the acquisition was paused for approximately 1 h to mitigate effects from potential temperature drifts due to the energy deposited in the specimen by repeated application of RF pulses.

2.3. Image processing

The uncombined image data from the individual receive channels were retrieved from the scanner using an in-house functor providing raw data in nifti format. Multi-channel combination without phase singularities was accomplished using an adaptive-combine algorithm implemented in MATLAB (v. 2022.a; MathWorks, Natick, MA, USA), followed by Laplacian phase unwrapping [27] and V-SHARP background-field removal [28,29] for obtaining the tissue phase volumes of high quality. The phase data for each orientation were registered to the LPI reference employing transformations that were derived by registering the corresponding magnitude volumes using FMRIB's Linear Image

¹ 'Chaos', in Greek mythology the void state preceding the creation of 'cosmos' (the universe).

Table 1

Overview of the sample rotations and the combinations of orientations used for the COSMOS reconstructions. The angles are indicated in degrees, and the x-, y-, and zaxis notation refers to the scanner coordinate system. The reference orientation (#1) was acquired with the specimen in the typical LPI *in-vivo* position. All other reoriented volumes were registered to this orientation.

	1 (LPI)	2	3	4	5	6	7	8	9	10
x-axis rotation	0°	60°	120°	10°	22°	-22°	0°	0°	0°	0 °
y-axis rotation	0°	0 °	0 °	0°	0 °	0 °	22°	-22°	0 °	0 °
z-axis rotation	0 °	0 °	0 °	0°	0 °	0 °	0 °	0 °	30°	-30°
COSMOS-opt	1	1	1							
COSMOS-iv1	1			1	1					
COSMOS-iv2	1				1	1				
COSMOS-iv3	1			1	1	1				
COSMOS-iv4	1			1	1	1	1	1		
COSMOS-iv5	1			1	1	1	1	1	1	1

Registration Tool (FLIRT; FSL 5.0.9) with six-parameter rigid transformations, a normalized Mutual Information (nMI) cost function and spline interpolation [30]. FSL was also used to derive a full-brain mask from the reference volume.

Quantitative susceptibility maps were reconstructed from the tissue phase of the reference orientation using iLSQR [11]. Using MATLAB, the remaining ten registered tissue phase volumes were used in different combinations for a total of six COSMOS reconstructions as indicated in Table 1. These combinations included the optimal COSMOS scheme ("COSMOS-opt" with rotations of 0_r , 60_r and 120_r about the x-axis), two combinations of three rotation angles mimicking the limited range of accessible rotation angles under conditions of in-vivo MRI of the human brain ("COSMOS-iv1" and "COSMOS-iv2" with rotations of 0_r , 10_r , 22_r and of 0_{r}° , 22_{r}° , -22_{r}° about the x-axis, respectively), and three further combinations to explore the potential advantage obtained with more than three orientations about either a single rotation axis ("COSMOSiv3" with rotations of 0_x° , 10_x° , 22_x° and -22_x° about the x-axis) or multiple rotations axes ("COSMOS-iv4" with rotations of 0_{r}° , 10_{r}° , 22_{r}° , -22_{r}° and of 22_{y}° , -22_{y}° about the x- and y-axis, respectively and "COSMOS-iv5" with rotations of 0_x° , 10_x° , 22_x° , -22_x° , of 22_y° , -22_y° and of 45_z° , -45_z° about the x-, y- and z-axis, respectively). The scripts for COSMOS and L2regularized COSMOS were modifications based on openly available MATLAB implementations (https://martinos.org/~berkin/software.ht ml). The statistical analyses included the 3D structural similarity index measure (SSIM) and Pearson's correlation coefficient (R). The measures were computed from a large region of interest (ROI) consisting of gray

matter (GM) and white matter (WM) as previously employed in a QSM reconstruction challenge [12]. Normalized difference maps were obtained by first normalizing the $\Delta \chi$ maps within the range [0,1] and then computing the absolute differences. The Gaussian noise variance was estimated in a 200 × 200 × 200 matrix extending from the center of the specimen [31,32].

3. Results

An example slice (magnitude and phase image) acquired with different specimen orientations and registered to the LPI reference is presented in Fig. 1. All registrations achieved excellent overlap. The transformation matrices indicated rotation angles that were close to the adjusted ones (see Supplementary Fig. S2 and Supplementary Table S2). Phase maps acquired with relatively small rotation angles (e.g., $\pm 10^{\circ}$) are almost indistinguishable from the reference, whereas maps acquired with rotation angles exceeding 20° (e.g., 60° and 120° about the x-axis) show distinct differences. Such differences are particularly prominent in WM regions, where relevant susceptibility anisotropy impacts the local field distribution. This suggests that COSMOS reconstructions from different sets of rotation angles will vary in regions of relevant susceptibility anisotropy. For example, scheme COSMOS-iv1 will be dominated by the measured phase distribution at the reference position (voxel-byvoxel averaging of similar local phase values), whereas the input images to the optimal scheme COSMOS-opt differ considerably in their phase variation (voxel-by-voxel averaging of local phase values distributed over a larger range). This is further evident from a comparison of the

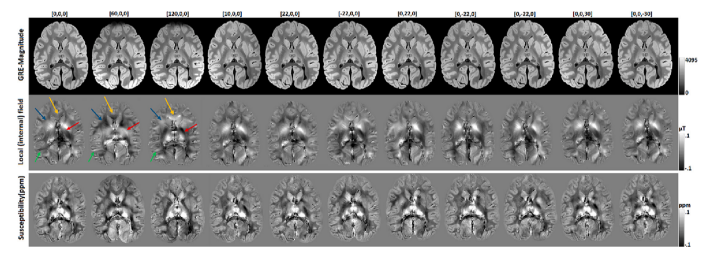


Fig. 1. Arbitrarily selected axial slice of the registered magnitude images (top row) acquired with different specimen orientations. The corresponding phase maps (2nd row) show obvious inconsistencies, presumably resulting from the anisotropic properties of magnetic susceptibility. In the bottom row, the iLSQR QSM calculated for each orientation is presented. Note the significant changes in WM regions—especially in high rotations—presumably resulting from the anisotropic properties of magnetic susceptibility (examples of such regions are indicated by blue, red, green and orange arrows in the internal field images for rotations of 0° , 60° and 120° about the *x*-axis). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

various COSMOS reconstructions with the simple QSM results obtained with iLSQR (Fig. 2). All acquisitions with *in vivo*-feasible rotation angles about a single axis (*e.g.*, *x*-axis) yielded COSMOS reconstructions with residual streaking artifacts. These artifacts were effectively mitigated by L2-regularization in all cases. The optimal COSMOS reconstruction and the *in vivo*-feasible schemes that included rotations about more than one axis appeared artifact-free without regularization.

As expected, the Gaussian noise variance was reduced for COSMOS reconstructions (range from 5×10^{-4} to 8×10^{-4}) compared to iLSQR-QSM (approx. 10^{-3}). Consistent with this observation, the use of only single orientations (*i.e.*, no averaging) appeared to result in less smoothing. We note that the noise variance served as a (relative) metric in the current case as the calculation of the SNR is not straightforward for susceptibility maps.

Fig. 3 presents normalized difference maps of results obtained with COSMOS-opt and COSMOS reconstructions with rotations that are feasible under in-vivo conditions. All difference maps indicated higher values in WM regions, while the COSMOS-inv1,2,3 results indicate also global differences, potentially due to the additional regularization needed to eliminate residual streaking artifacts. Fig. 4 presents normalized difference maps of results obtained with iLSOR-OSM and COSMOS-opt in comparison to corresponding difference maps obtained with iLSQR-QSM and COSMOS-iv2. As expected, the qualitative maps indicate prominent differences between iLSQR-QSM and COSMOS-opt are observed in WM. Differences between iLSQR-QSM and COSMOSiv2 are also evident, however, they appear reduced. The enhanced similarity of the iLSQR-QSM and COSMOS-iv2 results (compared to COSMOS-opt) is particularly evident in WM regions of high anisotropy, such as corpus callosum or internal capsule [7,18], whereas WM regions with more complex fiber dispersion or GM yielded higher agreement of the two COSMOS schemes. Note that the regularization that was included in the COSMOS-iv2 (but not in the COSMOS-opt) reconstruction also adds to these differences in a widespread fashion.

These observations are further corroborated quantitatively by the SSIM and Pearson's correlation coefficient shown in Fig. 5 and Table 2. Using COSMOS-opt as reference, the greatest SSIM and *R* values were obtained with COSMOS-iv2 and schemes with more rotations. This is consistent with the expectations that larger rotation angles are beneficial for COSMOS reconstructions if the angles deviate from the optimal distribution. Acquisitions with further rotations (albeit without increasing the angle span) yielded similar SSIM and *R* values. Compared to the various COSMOS schemes, iLSQR-QSM yielded the lowest SSIM and *R* although its accuracy measures were only moderately lower than those obtained with *in vivo*-feasible COSMOS schemes (Table 2). Of note, quantification of the mean susceptibility in major basal ganglia and

brainstem nuclei (see Supplementary Fig. S3) revealed a trend of slight underestimation of χ with the *in-vivo* feasible schemes when compared to the optimal COSMOS approach (see Supplementary Table S3). iLSQR-QSM results indicated mean values close the *in-vivo* feasible COSMOS data albeit with higher standard deviations, presumably due to omitting averaging.

4. Discussion

This study presents an investigation of effects from susceptibility anisotropy in different implementations of the COSMOS model. It is based *post-mortem* scans that allowed re-orientations replicating the optimal COSMOS reconstruction as well as others that adhere to limitations of physical re-orientations of the head that are feasible *in vivo*. We observed that *in-vivo* feasible and optimal COSMOS yielded highquality susceptibility maps with increased SNR resulting from averaging multiple acquisitions. COSMOS reconstructions from non-ideal rotations about a single axis required additional L2-regularization to mitigate residual streaking artifacts. *In-vivo* feasible COSMOS indicated higher visual resemblance and was statistically more comparable with single-orientation QSM than the optimal COSMOS reconstruction.

Fixed brain specimens can be arbitrarily rotated, overcoming limitations of orientation-dependent in vivo experiments. The chimpanzee brain is smaller in size than a human brain (about 1/3), supporting less time-consuming high-resolution acquisitions while still providing WM architecture morphologically close to the human brain with similar cellular composition, relative volume and complexity [32]. In MRI of cadaver brains, altered diffusion properties, and shorter T_1 and T_2 values are expected, due to autolysis [33,34], which may further depend on the fixative agent [35]. Despite multiple changes after fixation at the molecular level in brain tissue, however, a recent study showed a high correlation between post-mortem and pre-mortem QSM, and susceptibility remained relatively stable throughout the first 6 weeks of fixation in formalin [36]. Furthermore, T_2^* relaxation times decrease with decreasing temperature [37], thus, there may be a slight shift in R_2^* -related metrics, as our scans were performed at room temperature (approx. 18-22 °C).

The accuracy of the rotation-angle adjustment was high (estimated error $\leq 3^{\circ}$ based on the registration results, see Supplementary Table S2), facilitated through a sophisticated setup that was designed to support reorientation experiments [21]. This is different from *in-vivo* scanning of the human brain, where multiple unstable factors impact the robustness of the rotations, such as the subject's head size and shape, fixation within the coil, or head motion. Close-fitting array coils are optimized for high SNR in the LPI position and do not well support

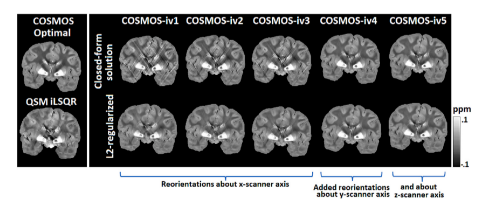


Fig. 2. Susceptibility maps in a central coronal slice obtained with iLSQR-QSM as well as different COSMOS schemes including COSMOS-opt and those with rotations that are feasible under *in-vivo* conditions. The results obtained with COSMOS-iv2 and COSMOS-iv3 appear to have the greatest similarity to those from COSMOS-opt at visual inspection despite remaining differences in WM regions. Note that the results from all *in vivo*-feasible COSMOS acquisitions are also presented after L2-regularization (bottom row) to minimize potential residual streaking artifacts. The reconstructions from COSMOS-opt and from *vivo*-feasible acquisitions that included rotations about more than one axis appeared artifact-free without regularization.

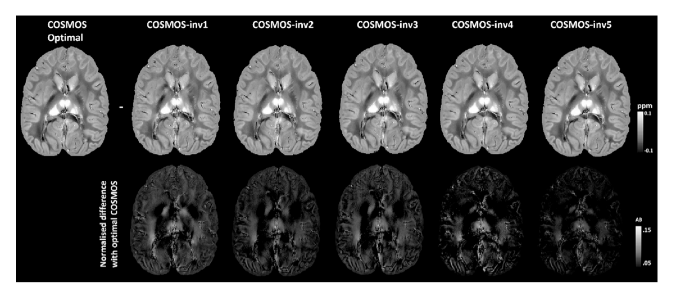


Fig. 3. Top row: Susceptibility maps in a central axial slice obtained with different COSMOS schemes (optimal and with rotations that are feasible under *in-vivo* conditions). Careful observation reveals local differences, which are pronounced in WM. Bottom row: normalized difference maps of results obtained with COSMOS-opt and *in-vivo* feasible COSMOS schemes. Enhanced differences are obtained in WM. The COSMOS-inv1,2,3 schemes also indicate global differences, presumably due to the additional regularization needed to eliminate residual streaking artifacts.

reorientation of the head, which severely limits the range of accessible rotation angles for the majority of adult subjects.

The signal distribution inside the object depends on the specific distance from individual coil elements and might vary upon reorientations. Such inconsistencies lead to SNR fluctuations and inaccuracies in the registration and post-processing pipelines and may, hence, further degrade the quality of COSMOS results. Indications of this bias is evident on the magnitude images after 60° and 120° rotations about the *x*-axis (Fig. 1), where the signal intensity is reduced in the frontal lobe and increased in parieto-occipital brain compared to the LPI reference. This may require further preprocessing before registration, such as N4 bias correction [38], which was not employed in our experiments.

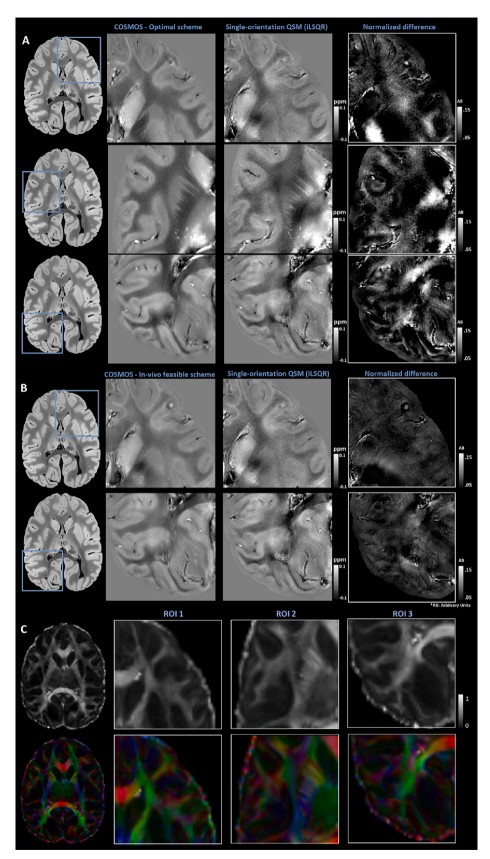
Our results indicate limited performance of the COSMOS model when applied to data with a restricted range of rotation angles as typically encountered *in-vivo*. While the results from suboptimal angle distributions still outperformed single-orientation QSM in comparisons to the optimal COSMOS scheme, the similarity metrics SSIM and *R* for the full-brain data were only <0.9 and < 0.85, respectively. Qualitatively, this is consistent with previous work indicating slightly better performance for multi-angle QSM (with a restricted range of angles) compared to both thresholded or regularized single-angle approaches [13]. We note that our data were recorded under idealized conditions, whereas subtle head motion *in vivo* might degrade multi-orientation acquisitions (requiring longer scan time) more than those with only a single orientation. Extending the rotations beyond one axis relaxed the need for regularization to eliminate streaking artifacts.

An inherent limitation is that COSMOS does not consider susceptibility anisotropy. While this aspect is observed for the early years of QSM development, it has recently received more attention [7,13,39,41,42] whereas previous investigations in COSMOS were typically limited to deep GM structures analysis [13,17]. The anisotropic character of susceptibility in WM leads to variations of the local field distribution upon reorienting the sample in the magnetic field. While structural anisotropy [40] and high-susceptibility source non-local contributions (e.g. deep GM nuclei) might also contribute to susceptibility changes in each measurement, it has been shown that susceptibility anisotropy contributes a fraction to the relative change: in *post mortem* brains leads there are observed contributions of ± 0.01 to 0.03 [42,43]. Along the same lines, in recent post-mortem study, observations in Corpus Callosum and internal capsule (WM ROIs with high degree of anisotropy) showed mean changes of 0.01 to 0.04ppm²¹. Thus, susceptibility anisotropy is a substantial contribution to the estimation, especially when strongly myelinated WM fibers are considered. In the current analysis, the optimal COSMOS scheme (0° , -60° , -120°) appeared to achieve better averaging of anisotropic susceptibility, especially in parts of WM with strong orientational characteristics. In comparison, averaging of anisotropy is limited with the restricted range of angles in-vivo, resulting in enhanced orientation bias. Apart from such performance differences, all COSMOS reconstructions will be impacted by anisotropy-related variability, in particular in WM regions of higher order of the local fiber distributions (Figs. 3-4). Remarkably, the previously suggested χ_{33} component of the susceptibility tensor [12] does not appear to be a better gold standard for scalar QSM estimations, presumably due to unaccounted anisotropic effects contained in offdiagonal susceptibility tensor elements [39]. A better approach to the ground truth may be obtained by projecting the χ_{13} and χ_{23} anisotropic contributions into the apparent scalar susceptibility, which is, however, not trivial.

For COSMOS reconstructions from non-optimal angle distributions limited to a single rotation axis (*e.g.*, the *x*-axis), additional L2regularization was required to mitigate residual streaking artifacts. This is counter-intuitive because COSMOS was introduced as a method for solving the ill-posed inversion problem without regularization—as indeed achieved with COSMOS-opt or upon incorporating rotations about multiple axes. Overall, both *in-vivo* feasible and optimal COSMOS yielded high-quality susceptibility maps with increased SNR resulting from inherent averaging of multiple acquisitions. Nevertheless, in view of unconsidered anisotropy effects, added complexity of the reconstruction, and the general challenge of multi-orientation acquisitions, advantages of sub-optimal COSMOS schemes over regularized singleorientation QSM appear limited.

CRediT authorship contribution statement

Dimitrios G. Gkotsoulias: Writing – review & editing, Writing – original draft, Visualization, Validation, Software, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. Carsten Jäger: Resources, Methodology. Roland Müller: Resources, Methodology. Tobias Gräßle: Resources. Karin M. Olofsson: Resources. Torsten Møller: Resources. Steve Unwin: Resources. Catherine Crockford: Writing – review & editing, Resources. Roman M. Wittig: Writing – review & editing, Resources. Berkin Bilgic: Writing – review



(caption on next page)

Fig. 4. Normalized difference maps of results from (A) iLSQR-QSM and COSMOS-opt and from (B) iLSQR-QSM and COSMOS-iv2 (indicated as "*in-vivo* COSMOS"). Zoomed ROIs are shown for better appreciation of the differences in specific parts of the brain. Pronounced deviations between iLSQR-QSM and COSMOS-opt in WM (*e.g.*, corpus callosum and internal capsule) are less evident between iLSQR-QSM and COSMOS-iv2 in the same regions. (C) Fractional anisotropy (FA) and FA-weighted primary eigenvector (same ROIs as above) from registered diffusion tensor imaging acquired on the same specimen, in a separate scanning session at a different time (Red color denotes Left-Right direction of fibers; Blue color denotes Superior-Inferior direction; Green color denotes Anterior-Posterior direction, always in the object's coordinate system). Increased diffusion anisotropy is evident in regions of enhanced susceptibility differences supporting the assumption that susceptibility anisotropy is related to myelination in these WM regions. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

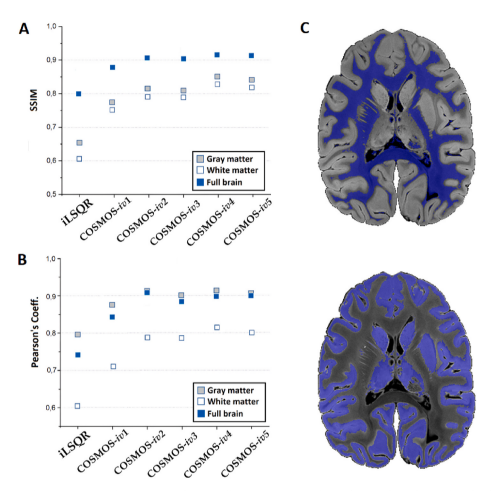


Fig. 5. Structural similarity index measure (A) and Pearson's correlation coefficient (B) as metrics for similarity and correlation of results obtained with iLSQR-QSM and *in vivo*-feasible COSMOS schemes in comparison to COSMOS-opt. Results are shown for masks [shown in blue color in (C)] of total GM (gray squares), total WM (open squares) and the combination of both (blue squares). COSMOS-iv2, yielded the smallest deviations than COSMOS-iv1 (*i.e.*, greater SSIM and *R* values). Adding acquisitions with further rotations did not significantly improve the result. Compared to all COSMOS schemes, iLSQR-QSM yielded the greatest deviation (smallest SSIM and *R* values) from the COSMOS-opt result. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table	2

Values of the	metrics	presented	in	Fig.	5
---------------	---------	-----------	----	------	---

	-	0				
SSIM	iLSQR	inv1	inv2	inv3	inv4	inv5
FULL BRAIN	0.80	0.88	0.91	0.9	0.92	0.92
GM	0.66	0.77	0.82	0.81	0.85	0.84
WM	0.61	0.75	0.77	0.79	0.82	0.81
R	iLSQR	inv1	inv2	inv3	inv4	inv5
FULL BRAIN	0.74	0.84	0.91	0.89	0.90	0.90
GM	0.79	0.87	0.92	0.90	0.92	0.91

& editing, Validation, Software, Resources, Methodology. Harald E. Möller: Writing – review & editing, Writing – original draft, Supervision, Resources, Project administration, Methodology, Funding acquisition, Conceptualization.

Declaration of generative AI and AI-assisted technologies in the writing process

During the preparation of this work the authors did not use any Generative AI and AI-assisting technology tool.

Declaration of competing interest

The authors declare no competing interest.

Data availability

The data that support the findings of this study are available from the corresponding author or the senior author upon reasonable request.

Acknowledgements

This work was funded by a grant to HEM from the European Commission, ITN "INSPIRE-MED" (H2020-MSCA-ITN-2018, #813120). BB acknowledges funding by the National Institutes of Health, grant Nos. R01 EB032378, R01 EB028797, R03 EB031175, U01 EB025162, P41 EB030006, and U01 EB026996, and computing support by the NVIDIA Corporation. We are grateful to Angela D. Friederici and Nikolaus Weiskopf from the Max Planck Institute for Human Cognitive and Brain Sciences and all Principal Investigators of the Evolution of Brain Connectivity (EBC) project for supporting this research through providing access to the brain specimen. We especially thank Ariane Düx and Fabian Leenderz from the Helmholtz Institute for One Health for their involvement in establishing the brain extraction network and the Kolmården Wildlife Park in Kolmården and National Veterinary Institute, Sweden, for executing the brain extraction and providing the tissue.

References

- Möller HE, Bossoni L, Connor JR, Crichton RR, Does MD, Ward RJ, et al. Iron, myelin, and the brain: neuroimaging meets neurobiology. Trends Neurosci 2019; 42:384–401. https://doi.org/10.1016/j.tins.2019.03.009.
- [2] Liu T, Spincemaille P, de Rochefort L, Kressler B, Wang Y. Calculation of susceptibility through multiple orientation sampling (COSMOS): a method for conditioning the inverse problem from measured magnetic field map to susceptibility source image in MRI. Magn Reson Med 2009;61:196–204. https:// doi.org/10.1002/mrm.21828.
- [3] Shmueli K, de Zwart JA, van Gelderen P, Li T-Q, Dodd SJ, Duyn JH. Magnetic susceptibility mapping of brain tissue in vivo using MRI phase data. Magn Reson Med 2009;62:1510–22. https://doi.org/10.1002/mrm.22135.
- [4] Schweser F, Deistung A, Lehr BW, Reichenbach JR. Quantitative imaging of intrinsic magnetic tissue properties using MRI signal phase: an approach to in vivo brain iron metabolism? NeuroImage 2011;54:2789–807. https://doi.org/10.1016/ j.neuroimage.2010.10.070.
- [5] Liu C, Wei H, Gong N-J, Cronin M, Dibb R, Decker K. Quantitative susceptibility mapping: contrast mechanisms and clinical applications. Tomography 2015;1: 3–17. https://doi.org/10.18383/j.tom.2015.00136.
- [6] Deistung A, Schweser F, Reichenbach JR. Overview of quantitative susceptibility mapping. NMR Biomed 2017;30:e3569. https://doi.org/10.1002/nbm.3569.
- [7] Liu C. Susceptibility tensor imaging. Magn Reson Med 2010;63:1471–7. https:// doi.org/10.1002/mrm.22482.
- [8] Liu J, Liu T, de Rochefort L, Ledoux J, Khalidov I, Chen W, et al. Morphology enabled dipole inversion for quantitative susceptibility mapping using structural consistency between the magnitude image and the susceptibility map. Neuroimage 2012;59(3):2560–8. https://doi.org/10.1016/j.neuroimage.2011.08.082.
- [9] Milovic C, Bilgic B, Zhao B, Acosta-Cabronero J, Tejos C. Fast nonlinear susceptibility inversion with variational regularization. Magn Reson Med 2018;80 (2):814–21. https://doi.org/10.1002/mrm.27073.
- [10] Li W, Wang N, Yu F, Han H, Cao W, Romero R, et al. A method for estimating and removing streaking artifacts in quantitative susceptibility mapping. NeuroImage 2015;108:111–22. https://doi.org/10.1016/j.neuroimage.2014.12.043.
- [11] Wharton S, Schäfer A, Bowtell R. Susceptibility mapping in the human brain using threshold-based k-space division. Magn Reson Med 2010;63(5):1292–304. https:// doi.org/10.1002/mrm.22334.
- [12] Langkammer C, Schweser F, Shmueli K, Kames C, Li X, Guo L, et al. Quantitative susceptibility mapping: report from the 2016 reconstruction challenge. Magn Reson Med 2018;79:1661–73. https://doi.org/10.1002/mrm.26830.
- [13] Wharton S, Bowtell R. Whole-brain susceptibility mapping at high field: a comparison of multiple- and single-orientation methods. NeuroImage 2010;53: 515–25. https://doi.org/10.1016/j.neuroimage.2010.06.070.
- [14] Schweser F, Sommer K, Deistung A, Reichenbach JR. Quantitative susceptibility mapping for investigating subtle susceptibility variations in the human brain. NeuroImage 2012;62:2083–100. https://doi.org/10.1016/j. neuroimage.2012.05.067.
- [15] Yoon J, Gong E, Chatnuntawech I, Bilgic B, Lee J, Jung W, et al. Quantitative susceptibility mapping using deep neural network: QSMnet. NeuroImage 2018; 179:199–206. https://doi.org/10.1016/j.neuroimage.2018.06.030.
- [16] Liu T, Liu J, de Rochefort L, Spincemaille P, Khalidov I, Ledoux JR, et al. Morphology enabled dipole inversion (MEDI) from a single-angle acquisition: comparison with COSMOS in human brain imaging. Magn Reson Med 2011;66: 777–83. https://doi.org/10.1002/mrm.22816.
- [17] Bilgic B, Xie L, Dibb R, Langkammer C, Mutluay A, Ye H, et al. Rapid multiorientation quantitative susceptibility mapping. NeuroImage 2016;125:1131–41. https://doi.org/10.1016/j.neuroimage.2015.08.015.

- [18] Wharton S, Schäfer A, Bowtell R. Susceptibility mapping in the human brain using threshold-based k-space division. Magn Reson Med 2010;63:1292–304. https:// doi.org/10.1002/mrm.22334.
- [19] Gkotsoulias DG, Müller R, Schlumm T, Alsleben N, Jäger C, Jaffe J, et al. COSMOSbased susceptibility estimations: Accuracy assessment and comparisons of QSM and multiple-angle acquisitions. In: Proceedings of the 31st Annual Meeting of ISMRM, London, UK; 2022. p. 2370.
- [20] Gräßle T, Crockford C, Eichner C, Girard-Buttoz C, Jäger C, Kirilina E, et al. Sourcing high tissue quality brains from deceased wild primates with known socioecology. Methods Ecol Evol 2023;14:1906–24. https://doi.org/10.1111/2041-210X.14039.
- [21] Gkotsoulias DG, Müller R, Jäger C, Schlumm T, Mildner T, Eichner C, et al. High angular resolution susceptibility imaging and estimation of fiber orientation distribution functions in primate brain. NeuroImage 2023;276:120202. https:// doi.org/10.1016/j.neuroimage.2023.120202.
- [22] Frahm J, Haase A, Matthaei D. Rapid three-dimensional MR imaging using the FLASH technique. J Comput Assist Tomogr 1986;10:363–8. https://doi.org/ 10.1097/00004728-198603000-00046.
- [23] Alkemade A, Pine K, Kirilina E, Keuken MC, Mulder MJ, Balesar R, et al. 7 Tesla MRI followed by histological 3D reconstructions in whole-brain specimens. Front Neuoanat 2020;14:536838. https://doi.org/10.3389/fnana.2020.536838.
- [24] Alkemade A, Bazin P-L, Balesar R, Pine K, Kirilina E, Möller HE, et al. A unified 3D map of microscopic architecture and MRI of the human brain. Sci Adv 2022;8: eabj7892. https://doi.org/10.1126/sciadv.abj7892.
- [25] Griswold MA, Jakob PM, Heidemann RM, Nittka M, Jellus V, Wang J, et al. Generalized autocalibrating partially parallel acquisitions (GRAPPA). Magn Reson Med 2002;47:1202–10. https://doi.org/10.1002/mrm.10171.
- [26] Feinberg DA, Hale JD, Watts JC, Kaufman L, Mark A. Halving MR imaging time by conjugation: demonstration at 3.5 kG. Radiology 1986;161:527–31. https://doi. org/10.1148/radiology.161.2.3763926. Schofield MA, Zhu Y. Fast phase unwrapping algorithm for interferometric applications. Opt. Lett. 2003; 28: 1194–1196. doi:10.1364/OL.28.001194.
- [27] Li W, Wu B, Liu C. Quantitative susceptibility mapping of human brain reflects spatial variation in tissue composition. NeuroImage 2011;55:1645–56. https://doi. org/10.1016/j.neuroimage.2010.11.088.
- [28] Özbay PS, Deistung A, Feng X, Nanz D, Reichenbach JR, Schweser F. A comprehensive numerical analysis of background phase correction with V-SHARP. NMR Biomed 2017;30:e3550. https://doi.org/10.1002/nbm.3550.
- [29] Jenkinson M, Beckmann CF, Behrens TEJ, Woolrich MW, Smith SM. FSL. NeuroImage 2012;62:782–90. https://doi.org/10.1016/j. neuroimage.2011.09.015.
- [30] Immerkær J. Fast noise variance estimation. Comput Vis Image Underst 1996;64: 300–2. https://doi.org/10.1006/cviu.1996.0060.
- [31] Liu W, Liu T, Rong M, Wang R, Zhang H. A fast noise variance estimation algorithm. In: Proceedings of the 2011 Asia Pacific Conference on Postgraduate Research in Microelectronics & Electronics, Macao, China; 2011. p. 61–4. https:// doi.org/10.1109/PrimeAsia.2011.6075071.
- [32] Herculano-Houzel S. The remarkable, yet not extraordinary, human brain as a scaled-up primate brain and its associated cost. Proc Natl Acad Sci U S A 2012;109 (Suppl. 1):10661–8. https://doi.org/10.1073/pnas.1201895109.
- [33] De Barros A, Arribarat G, Combis J, Chaynes P, Péran P. Matching ex vivo MRI with iron histology: pearls and pitfalls. Front Neuroanat 2019;13:68. https://doi.org/ 10.3389/fnana.2019.00068.
- [34] Tatsumi S, Shinohara M, Yamamoto T. Direct comparison of histology of microbleeds with postmortem MR images: a case report. Cerebrovasc Dis 2008;26: 142-6. https://doi.org/10.1159/000139661.
- [35] Birkl C, Soellradl M, Toeglhofer AM, Krassnig S, Leoni M, Pirpamer L, et al. Effects of concentration and vendor specific composition of formalin on postmortem MRI of the human brain. Magn Reson Med 2018;79:1111–5. https://doi.org/10.1002/ mrm.26699.
- [36] Evia AM, Kotrotsou A, Tamhane AA, Dawe RJ, Kapasi A, Leurgans SE, et al. Ex-vivo quantitative susceptibility mapping of human brain hemispheres. PloS One 2017; 12:0188395. https://doi.org/10.1371/journal.pone.0188395.
- [37] Birkl C, Langkammer C, Golob-Schwarzl N, Leoni M, Haybaeck J, Goessler W, et al. Effects of formalin fixation and temperature on MR relaxation times in the human brain. NMR Biomed 2016;29:458–65. https://doi.org/10.1002/nbm.3477.
- [38] Tustison NJ, Avants BB, Cook PA, Zheng Y, Egan A, Yushkevich PA, et al. N4ITK: improved N3 bias correction. IEEE Trans Med Imaging 2010;29:1310–20. https:// doi.org/10.1109/TMI.2010.2046908.
- [39] Milovic C, Özbay PS, Irarrazaval P, Tejos C, Acosta-Cabronero J, Schwesser F, et al. The 2016 QSM challenge: lessons learned and considerations for a future challenge design. Magn Reson Med 2020;84:1624–37. https://doi.org/10.1002/mrm.28185.
- [40] Wharton S, Bowtell R. Effects of white matter microstructure on phase and susceptibility maps. Magn Reson Med 2015;73(3):1258–69. https://doi.org/ 10.1002/mrm.25189.
- [41] Bao L, Xiong C, Wei W, Chen Z, van Zijl PCM, Li X. Diffusion-regularized susceptibility tensor imaging (DRSTI) of tissue microstructures in the human brain. Med Image Anal 2021;67:101827. https://doi.org/10.1016/j.media.2020.101827.
- [42] Li W, Liu C, Duong TQ, van Zijl PC, Li X. Susceptibility tensor imaging (STI) of the brain. NMR Biomed. 2017;30(4). https://doi.org/10.1002/nbm.3540. Lee J, Shmueli K, Fukunaga M, van Gelderen P, Merkle H, Silva AC, Duyn JH. Sensitivity of MRI resonance frequency to the orientation of brain tissue microstructure. Proc Natl Acad Sci U S A 2010 Mar 16;107(11):5130-5. doi:10.1073/pnas.0910222107.
- [43] Liu C, Li W, Johnson GA, Wu B. High-field (9.4 T) MRI of brain demyelination by quantitative mapping of magnetic susceptibility. Neuroimage 2011;56(3):930–8. https://doi.org/10.1016/j.neuroimage.2011.02.024.