# 1 Studying magnetic susceptibility,

## <sup>2</sup> microstructural compartmentalisation and

**3** chemical exchange in a formalin-fixed ex vivo

## 4 human brain specimen

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Running title
Probing white matter phase mechanisms using formalin-fixed tissue
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### 28 Abstract

Purpose: Ex vivo imaging is a preferable method to study the biophysical mechanism of white matter orientation-dependent signal phase evolution. Yet, how formalin fixation, commonly used for tissue preservation, affects the phase measurement is not fully known. We, therefore, study the impacts of formalin fixation on magnetic susceptibility, microstructural compartmentalisation and chemical exchange measurement on human brain tissue.

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Methods: A formalin-fixed, post-mortem human brain specimen was scanned with multiple orientations with respect to the main magnetic field direction for robust bulk magnetic susceptibility measurement with conventional quantitative susceptibility imaging models. Homogeneous white matter tissues were subsequently excised from the whole-brain specimen and scanned in multiple rotations on an MRI scanner to measure the anisotropic magnetic susceptibility and microstructure-related contributions in the signal phase. Electron microscopy was used to validate the MRI findings.

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43 **Results:** The bulk isotropic magnetic susceptibility of ex vivo whole-brain imaging is 44 comparable to in vivo imaging, with noticeable enhanced non-susceptibility contributions. The 45 excised specimen experiment reveals that anisotropic magnetic susceptibility and 46 compartmentalisation phase effect were considerably reduced in formalin-fixed white matter 47 tissue.

48

49 **Conclusions:** Despite formalin-fixed white matter tissue has comparable bulk isotropic 50 magnetic susceptibility to those measured via in vivo imaging, its orientation-dependent 51 components in the signal phase related to the tissue microstructure is substantially weaker, 52 making it less favourable in white matter microstructure studies using phase imaging.

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54 **Keywords:** Quantitative susceptibility imaging, phase imaging, ex vivo imaging, 55 microstructure, white matter

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- 58

### 59 Introduction

Quantitative susceptibility mapping (QSM) is a physics-driven method to the study magnetic properties of biological tissues (1). Some features that differentiated it from conventional MR relaxometry include the field strength independence of the derived maps, relying on spatial deconvolution and its ability to distinguish paramagnetic and diamagnetic substances since they produce opposite contrasts. QSM is commonly performed on gradient echo phase data owing to its direct relationship to magnetic field variations (2,3).

66

One major QSM research challenge is to understand the mechanism of phase evolution in 67 68 white matter (WM) (4,5). In deep grey matter (GM), strong correlations between QSM and iron 69 concentration have been demonstrated (6). Yet, in WM, the abundance of diamagnetic myelin 70 (relative to water) would have suggested a strong QSM contrast relative to cerebrospinal fluid 71 (CSF) (4,5). The lack of this strong contrast has been attributed to various biophysical 72 phenomena (7,8). The lipid-rich myelin bilayer sheath encapsulating the highly-ordered axons 73 in WM results in anisotropic susceptibility (7–12). Additionally, water protons exist in various 74 microstructural environments (13), namely myelin water, intra-axonal water and extra-axonal 75 water, which can have different signal decay rates and frequency shifts depending on the 76 composition of the tissue and the fibre orientation with respect to the main magnetic field  $(B_0)$ 77 (14–17), which also make the signal phase not representing the average magnetic field in a 78 voxel. The chemical exchange of protons between macromolecules and water can also 79 introduce a further frequency shift of the MR signal (18,19). Disentangling the origins of WM 80 phase contrast can improve our understanding of QSM and provide new means to account for 81 their impact in QSM.

82

83 As WM phase contrast is orientation-dependent, studying its properties requires data acquired 84 with different orientations to  $B_0$ . Subject compliance limits the range of angles that can be 85 obtained in vivo (unnatural posture inside the scanner). Experiments with ex vivo samples, on 86 the other hand, do not suffer from this limitation, allowing long scanning sessions without data degradation caused by motion and for histology to be performed as a means of validation of 87 any microstructural findings (20,21). One shortcoming of ex vivo experiments is that MR 88 89 measured parameters in tissues undergone formalin fixation (a common practice to preserve 90 human post-mortem tissue) show substantial differences to those found in vivo. Those 91 differences have been seen both on single- and multi-compartment relaxometry, and diffusion-92 weighted imaging (22,23). Yet, a previous study showed that the bulk magnetic susceptibility 93 of brain tissues measured by QSM did not change significantly between in vivo and ex vivo

94 conditions, and also during a 6-week fixation period (24). This finding is in agreement with the 95 experiment results when studying the microstructural effect in phase imaging (25), where 96 comparable bulk magnetic susceptibilities were observed between fresh and fixed rat optic 97 nerves but the origins of the susceptibility contrast are different.

98

99 In this study, we investigate the magnetic susceptibility, compartmentalisation and chemical 100 shift effects on the MR phase using a formalin-fixed, post-mortem human brain specimen for 101 WM phase-contrast mechanism studies at 3T, providing comprehensive insights for the use 102 of fixed tissue in future QSM methodology studies. We performed multiple orientation 103 experiments in both whole-brain and excised tissue samples (26). This enabled us to obtain 104 both traditional QSM maps and ground truth bulk magnetic susceptibility measurements of the excised samples, as well as a separate estimation of microstructure compartmentalisation 105 106 information. The samples were then studied using electron microscopy (EM) to further 107 evaluate microstructural correlates between MR and histology. 108

### 109 Methods

- 110 The study is divided into 3 parts:
- 1) a post-mortem, formalin-fixed human brain specimen was scanned on an MRI scanner in
- 112 multiple orientations with respect to B<sub>0</sub> for robust magnetic susceptibility measurements;
- 113 2) homogeneous WM specimens were excised from the whole-brain specimen, embedded in
- agar and subsequently scanned again in various orientations in respect of B<sub>0</sub> to measure their
- magnetic susceptibility and microstructure-induced field, similar to the experiment conductedby (26);
- 117 3) the WM specimens used in the second part of the study were imaged by 3D EM, allowing
- 118 MRI data to be compared to histology.
- 119

### 120 **Tissue processing**

A post-mortem human brain specimen from a deceased male (aged 78 years old) with no history of neurological disorder (cause of death: myocardial infarction) was used for this research in accordance with the local ethics committee and the Anatomy Department of Radboud University Medical Center (Radboudumc, Nijmegen, the Netherlands). The brain specimen was immersed in 10% formalin for tissue fixation after being extracted from the skull.

After one month of fixation, the specimen was scanned on an MRI scanner (imaging details in section 2.2) to obtain the brain morphology for creating a tailor-made holder. The holder was made of a stack of 35 4-mm thick, 3D-printed plastic plates having space with the same shape of the specimen in the centre, covering most of the brain (see Figure 1) where the specimen can be fitted tightly inside the holder and a surrounding spherical container. Each plate has a grid layout with 4 mm x 4 mm elements, providing landmarks in MRI images for planning and guidance of tissue excision for the validation experiment in the second MRI session.

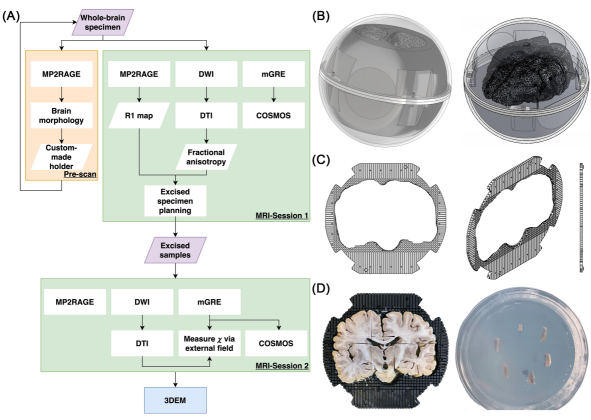
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135 After the first MRI session, 10 homogeneous WM regions of interest (ROIs) were identified (DTI fractional anisotropy  $\geq 0.45$ ) with excisable volume ( $\geq 1$  element of the holder plate grid, 136 137 i.e., 64 mm<sup>3</sup>). These WM tissues and 2 additional deep GM tissues (one from globus pallidus 138 and one from putamen) were then excised and embedded in 1% low-gelling temperature 139 agarose (A9414, Sigma Aldrich, Germany; with demineralised water) to ensure no extra tissue 140 protein denaturation would occur between the two acquisitions. The samples were positioned in a polymethyl methacrylate cylindrical container with the main fibre orientation perpendicular 141 142 to the cylindrical axis. Imaging was performed about 1 day after the excised specimens were 143 acquired and 5 days after the whole brain scan.

#### 144

EM was utilised to provide an additional reference to understand and explain the MRI findings. Two days after the second MRI session, the WM specimens were sectioned to 100  $\mu$ m on a vibratome (VT1000S, Leica Biosystems, Nussloch, Germany) before being immersed in 2.5% glutaraldehyde in 0.1M sodium cacodylate buffer for overnight incubation at 4°C. The specimens were then transferred to 0.25% glutaraldehyde in 0.1M sodium cacodylate buffer for storage at 4°C, and then delivered to the EM facility at the University of Oxford for imaging. The workflow of this study is summarised in Figure 1.





153

154 Figure 1: (A) A summary of this study, consisting of a pre-scan, two MRI sessions and a 3D EM session. 155 (B) Schematic of the experimental setup used in the first MRI session. The setup is made of two parts: 156 an outer transparent sphere allowing free rotation of the specimen and a tailor-made inner holder to 157 ensure the specimen was in a fixed position inside the sphere. (C) Schematic of the plate which forms 158 the inner holder. The centre of the plate is a space with the shape of the specimen, surrounded by a 159 grid structure providing location reference in MRI image (the gaps filled with water while the plate 160 material gave no detectable signal), and guidance in sample excision for the second imaging session. 161 (D) An illustration of how homogenous samples were acquired with the aids of the plate (right) and the 162 excised specimens were embedded in the agar inside the cylindrical container (left).

### 164 **MRI experiments**

#### 165 Data acquisition

The study was approved by the local ethics committee. All MRI data were acquired on a 3T scanner (Prisma, Siemens, Erlangen, Germany) at room temperature (20°C) using a 64channel array head/neck coil (with only 48 head channels were enabled). The experiment consisted of two imaging sessions: the first session was conducted on the whole-brain specimen and the second session was conducted on the excised brain tissues. The following protocol was used for the first session:

- 172 (1) MP2RAGE adopted to sensitise for T<sub>1</sub> values between 250 ms and 1000 ms, 1 mm 173 isotropic resolution, TI1/TI2/TR=311/1600/3000 ms, flip angle ( $\alpha$ ) #1/#2 = 4°/6°, Tacq 174 = 5 min;
- (2) 2D spin-echo EPI DWI, 1.6 mm isotropic, TR/TE=15241/77.6 ms, 2-shell
  (b=0/1250/2500 s/mm<sup>2</sup>, 17/120/120 measurements with 7 b=0 measurements
  collected with reversed phase-encode blips for distortion correction), 20 repetitions,
  Tacq = 5.6 hours;
- 179(3) Monopolar, 3Dmulti-echoGRE,1mmisotropicresolution,180TR/TE1/ $\Delta$ TE/TE6=40/3.45/6.27/34.8 ms,  $\alpha$  =20° (optimised for WM T<sub>1</sub>), 10 orientations181with respect to B<sub>0</sub>, chosen to optimise microstructural information decoding (27); Tacq182= 1.7 hours.
- 183
- 184 In the second scanning session, the excised specimens were scanned with the following185 protocol:
- 186 (1) MP2RAGE sequence with the same parameters as above;
- (2) 2D spin-echo EPI DWI, 1 mm isotropic, TR/TE=15241/77.6 ms, 2-shell
  (b=0/1250/2500 s/mm<sup>2</sup>, 17/120/120 measurements with 7 b=0 measurements), 9
  repetitions, Tacq = 9 hours;
- 190(3) Monopolar, 3Dmulti-echoGRE,0.7mmisotropicresolution,191TR/TE1/dTE/TE6=40/3.45/6.27/34.8ms,FA=15°(optimised for agar T1),10192orientations with respect to B0 (acquisition order was randomised);Tacq = 5.3 hours.
- 193

#### 194 Data processing

195 Each DWI repetition was pre-processed separately with MP-PCA denoising (28),196 susceptibility-induced distortion correction (29,30) and eddy current-induced distortion

197 correction (31). DTI model was then utilised on the DWI data after averaging all repetitions.

198 R<sub>1</sub> maps and DTI results were then linearly registered to the GRE data using ANTs (32).

199

200 GRE data with all orientations were first corrected for gradient nonlinearity induced geometric 201 distortion and then linearly registered to a common space, independent of the experiment 202 orientations.  $R_2^*$  maps were computed using a closed-form solution (33). Field maps were 203 computed using SEGUE (34) spatial phase unwrapping with optimum-weighted echo 204 combination (35) and tissue field maps were computed using LBV (36) in SEPIA (37). For the 205 whole-brain data, bulk isotropic magnetic susceptibility was derived using COSMOS (38). 206 Additionally, the QUASAR algorithm for multi-orientations was also applied to test if the bulk 207 isotropic magnetic susceptibility measurement improved when the field generated by non-208 susceptibility contributions ( $f_{\rho}$ ) were simultaneously estimated (39):

 $f_N = d_N * \chi + f_\rho [Eq. 1]$ 

where  $f_N$  and  $d_N$  are the tissue field and a unit dipole field associated with the acquisition at orientation N and \* is the convolution operator.

212

For the excised specimen data, in addition to the computation of the bulk isotropic magnetic susceptibility using COSMOS, quantification of isotropic and anisotropic magnetic susceptibility ( $\chi_i$  and  $\chi_a$ ) of the sample without confounding with non-susceptibility microstructural contributions was performed by fitting the measured field  $f_N$  on the agar surrounding the specimen in the following manner (26):

218

$$\min_{\boldsymbol{\chi},\boldsymbol{\vartheta},\boldsymbol{\chi}_{-}} \left\| M_{agar}(f_{N} - \chi_{i}\delta f_{i,N} - \chi_{a}\delta f_{a,N} - C_{N}) \right\| [Eq. 2]$$

219 where: M<sub>agar</sub> is the binary mask on agar with inner and outer boundaries 1 and 5 voxels away 220 from the specimen tissue boundary in all directions;  $\delta f_{i,N}$  and  $\delta f_{a,N}$  are the frequency 221 perturbations generated by a homogeneous specimen per unit of isotropic and anisotropic 222 magnetic susceptibility at orientation N, with its orientation of tensor derived from obtained 223 from DTI; and C<sub>N</sub> accounts for any baseline frequency differences in agar due to either 224 chemical exchange in the agar compartment or any residuals remaining after background field 225 removal for a particular orientation. Linear regression analysis was then conducted to compare 226 the magnetic susceptibility measurements between the COSMOS and external field method, 227 and between the two imaging sessions to determine if the excised specimens agree with the 228 ROIs in the whole-brain imaging data.

229

Tissue compartmentalisation contributions to the MR phase can be measured as the residual field ( $f_R$ ) inside the specimen with mask  $M_{specimen}$ :

232 
$$f_{R,N} = \overline{M_{specimen}(f_N - \chi_i \delta f_{i,N} - \chi_a \delta f_{a,N})} [Eq. 3].$$

- 233 which is expected to vary with the angle  $\theta$  between the fibre direction and B<sub>0</sub> as
- 234

 $f_{R,N} = A\sin^2\theta_N + B \left[Eq.4\right]$ 

where A accounts for the microstructure orientation-dependent effect of the specimen and B is orientation-invariant, related to both magnetisation exchange and microstructure (26).

237

### 238 Electron microscopy

Two corpus callosum (CC) specimens from the second MRI session (CC4 and CC5) having the greatest discrepancy of the microstructural compartmentalisation effect underwent 3D EM (40) to provide histology data for the MRI experiment validation. Each of the EM images has a matrix size of 8000×8000 with an in-plane resolution of 13.7×13.7 nm (~0.11x0.11mm FOV) and slice thickness of 100 nm. In total, 651 and 623 slices were acquired for CC4 and CC5 respectively.

245

246 Our EM data showed similar myelin sheath damage (splitting and swelling) as illustrated in 247 (41), resulting in unsatisfactory compartmental classification (intra-axonal, myelin and extra-248 axonal compartments) from standard segmentation tools. To facilitate high-quality 3-249 compartment classification, we first performed a semi-automatic intra-axonal segmentation 250 using ITK-snap (42) on down-sampled 3D EM data (87.7×87.7 nm and 100 consecutive slices). 251 The myelin sheath of each axon was initially defined by expanding the axon assuming a g-252 ratio = 0.5 (for axon diameters <  $1.2\mu$ m) or 0.6 (otherwise), followed by intensity thresholding 253 using the EM images. The resulting myelin mask is clearly influenced by the chosen threshold, 254 therefore, myelin volume fractions (MVF) derived from 5 threshold values range from 125 to 255 145 (step size of 5, most frequent intensities in the myelin mask are 104 (CC4) and 88 (CC5)) 256 were reported. To account for the enlarged myelin volume due to swelling, we also computed 257 the image intensity corrected myelin probability as:

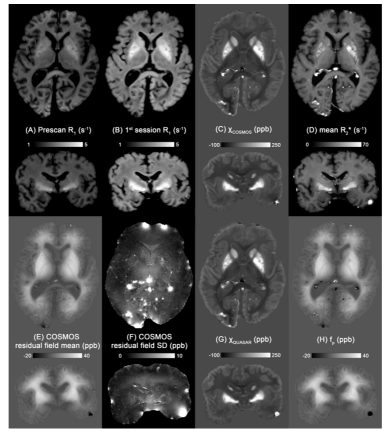
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$$Myelin \ probability = \begin{cases} 1, & I < M\\ \frac{A-I}{A-M}, & M < I \le A \ [Eq. 5]\\ 0, & I > A \end{cases}$$

where A and M are the most frequent values inside the axonal and myelin masks and I is the intensity of a voxel. The axon and myelin masks of each myelinated axon were eventually combined in a single classification map to avoid repeated counts due to overlapping between neighbouring axons.

264 Axonal volume fraction (AVF) and MVF were computed by counting the total number of voxels 265 of each compartment in the classification map, and were used to derive the sample g-ratio (20). Effective axonal diameter was defined as the square root of the product of the 2<sup>nd</sup> and 266 267 3<sup>rd</sup> principal axis lengths of the axons obtained from the regionprop3 function of MATLAB 268 (Mathworks, Natick, US), from which the median and the skewness of the axonal diameter 269 distribution were computed. Fibre dispersion was computed from the axonal volume-weighted 270 average squared dot product between the axon main principal direction and the average 271 orientation of the entire sample (27). To further investigate the effect of realistic myelin sheath 272 geometry on the compartmental frequency shifts, field perturbations induced from myelin  $\chi_i$ 273 was simulated in two scenarios: when the segmented sample axons were parallel or 274 perpendicular to  $B_0$ . The frequency shift distribution in the extracellular space was 275 subsequently analysed.

### 277 **Results**

278 Whole-brain imaging results are shown in Figure 2. The R<sub>1</sub> maps obtained from the first 279 session (5-month fixation) show faster relaxation rates than those from the pre-scan (1-month 280 fixation), with the contrast between WM and cortical GM being clearly reduced, and DGM 281 showing increased R<sub>1</sub> (Figure 2A and 2B). The COSMOS derived magnetic susceptibilities 282 are in reasonable agreement with previously published in vivo data (see supplementary Figure 283 S1), where opposite magnetic susceptibility between WM and GM can be observed (Figure 284 2C). However, the residual field of the COSMOS estimation shows a slowly-varying pattern 285 across the brain that cannot be explained by the isotropic dipole field (Figure 2E) and is 286 relatively stable across orientations (see Figure 2F). This residual map shares similar contrast 287 and values with the QUASAR non-susceptibility contributions map (Figure 2H). Hence, not 288 surprisingly, the bulk magnetic susceptibilities derived from QUASAR (Figure 2G) and 289 COSMOS (Figure 2c) are comparable. Susceptibility tensor imaging was also performed, yet 290 the results beyond the mean susceptibility tensor were found not informative.



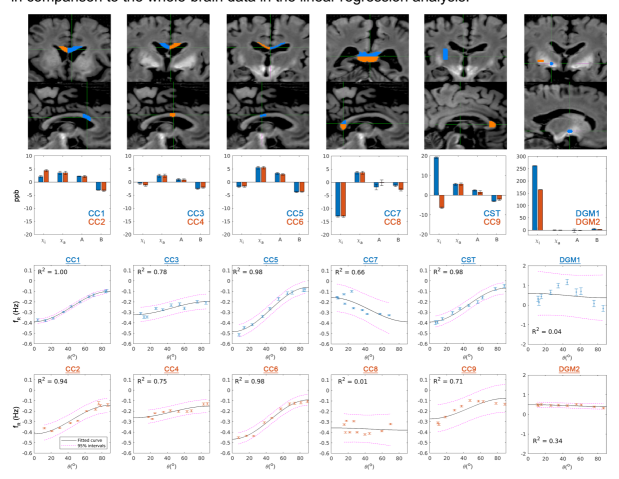
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Figure 2: Quantitative maps of the whole-brain specimens in transverse and coronal directions. (A) R<sub>1</sub>
 map obtained from a pre-scan after 1-month of formalin fixation and (B) R<sub>1</sub> map obtained from the 1<sup>st</sup>
 imaging session while the brain was fixed for 5 months. (C) COSMOS derived bulk magnetic
 susceptibility, (D) mean R<sub>2</sub>\* map across the 10 rotations, (E and F) mean and standard deviation of the

residual fields from COSMOS across orientations, (G and H) QUASAR derived bulk magneticsusceptibility map and non-susceptibility contribution map.

298

299 Figure 3 shows the magnetic susceptibility of the excised specimens measured via the 300 external field on agar with their ROIs illustrated in the whole-brain R<sub>1</sub> map. The mean  $\chi_i$  and 301  $\chi_a$  are -1.17±9.18 ppb and 4.03±1.63 ppb across WM specimens. A relatively strong positive  $\chi_i$  is found in the corticospinal tract specimen (CST; 19.17 ppb), which was found in retrospect 302 303 to be due to some DGM residual in one end of the excised sample. The coefficient A of  $\sin^2\theta$ dependence reflecting the WM microstructure effect has a mean of 1.46±1.55 ppb with a mean 304 intercept B of -2.75±0.79 ppb in WM. However, the R<sup>2</sup> of the specimen residual field fitting 305 306 suggests that not all WM specimens fit the  $\sin^2\theta$  function equally well, particularly for samples 307 obtained from the genu and splenium of the CC (CC7-CC9; R<sup>2</sup> ranging from 0.01 to 0.71). 308 Therefore, we focused on the 6 WM specimens obtained from the body of the CC (CC1-CC6) 309 in comparison to the whole-brain data in the linear regression analysis.



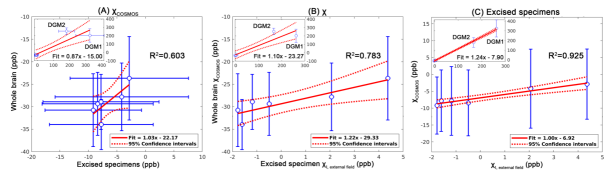
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Figure 3: (Middle row) Barplots of isotropic and anisotropic magnetic susceptibility, and coefficients A and B of the fitting of  $\sin^2\theta$ . (Bottom two rows) Fittings of  $\sin^2\theta$  reflecting microstructure compartmentalisation of the excised specimens. Each column shows the results of two specimens with their ROIs illustrated in the whole-brain R<sub>1</sub> map (top two rows). Error bar indicates the standard error

(except for the coefficients A and B in the barplots; 95% confidence intervals in this case). CC: corpus
 callosum; CST: corticospinal tract; DGM1: globus pallidus; DGM2: putamen.

317

318 Strong linear relations were found in mean susceptibility estimated by COSMOS between the 319 excised specimens and the corresponding ROI in the whole-brain data (cross-session; 320  $R^2$ =0.603, Figure 4A), between the  $\chi_i$  from external field measurement and the mean 321 COSMOS susceptibility in the whole-brain data (cross-session, cross-method; R<sup>2</sup>=0.783, 322 Figure 4B), and between the  $\chi_i$  from external field measurement and the mean COSMOS susceptibility on the excised specimens (cross-method;  $R^2=0.925$ , Figure 4C). All the slopes 323 of the linear regressions are close to 1, whereas the relatively large intercepts in Figure 4A-B 324 reflect the different reference medium in the scans (1<sup>st</sup> session: water; 2<sup>nd</sup> session: 1% agar). 325

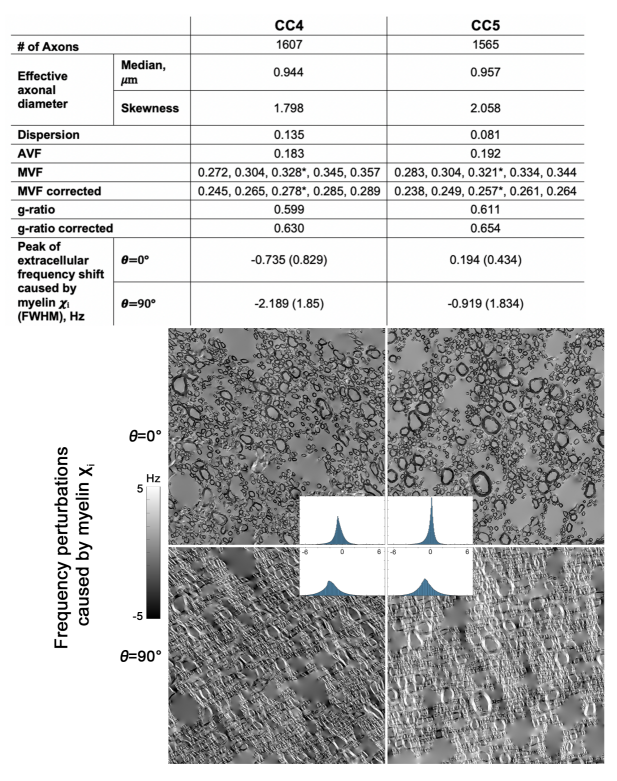


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Figure 4: Linear regression analyses on the 6 WM specimens of (A) the magnetic susceptibility measured between two imaging sessions using COSMOS, (B) between the bulk magnetic susceptibility measured by COSMOS on the first session and the  $\chi_i$  from external field measurement on the second session, and (C) the excised specimens between COSMOS magnetic susceptibility and external field derived  $\chi_i$ . Blue points: measurement data; solid red line: fitted line; dash line: 95% confidence interval; error bar: standard deviation. The subplot of each panel shows the regression result when the DGM specimens were included.

334

The microstructural properties of two WM specimens (CC4: relatively weaker microstructural phase; CC5: strong microstructural phase) derived from 3D EM data are summarised in Figure 5. Both specimens have similar MVF (8% difference), AVF (4% difference) and axonal diameter (1% difference). Noticeable differences are observed in fibre dispersion (50% difference) and in the FWHM of the extra-axonal frequency distributions when the fibre direction is parallel to B<sub>0</sub> (63% difference), the latter being attributed to the differences of both fibre dispersion and the spatial distribution of myelinated axons.



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Figure 5: (Top) Summary of the 3D EM derived CC4 and CC5 microstructural properties. MVF was separately probed using 5 different intensity thresholds (only the middle value indicated by \* was showed in g-ratio). The corrected MVF was derived using Eq. [5]. (Bottom) Frequency induced by the myelin  $\chi_i$  at two orientations to B0 ( $\theta$ =0° & 90°). Sub-figures show the frequency distributions in extracellular space, where the locations and the FWHM of the peaks are shown in the above summary.

### 349 **Discussion**

350 In this study, we examined the magnetic susceptibility and microstructural 351 compartmentalisation effect on MRI phase data on a formalin-fixed, post-mortem human brain 352 specimen. The bulk magnetic susceptibility of the whole-brain specimen shows comparable 353 contrast to those in the previous ex vivo studies (6,24), as well as to in vivo imaging: WM is 354 slightly diamagnetic, whereas cortical and deep GM are paramagnetic. Further investigation 355 reveals the residual fields of COSMOS have a gradient-like appearance varying from the 356 surface toward the centre of the specimen (Figure 2E), which is similar to the expected way 357 of how the solutions (fixative or water) diffused into the specimen. Since these residual fields 358 are relatively constant across different rotations, it is likely to be caused by the exchange effect 359 (17–19). These fields were captured as the non-susceptibility contributions by QUASAR and 360 they did not have a significant impact on the bulk magnetic susceptibility measurement (Figure 361 2C, 2G). This result is distinct from in vivo imaging results (Figure S1), where the susceptibility 362 differences in WM are more noticeable, suggesting that the effect of (sub)cellular structure of 363 WM is considerably reduced and the sphere of Lorentz inclusion utilised in COSMOS is 364 already a good approximation on formalin-fixed tissue.

365

366 While the bulk susceptibility of the WM samples in this study is similar to in vivo imaging, the 367 residual field analysis inside the excised homogenous tissue confirmed that the microstructure 368 compartmental frequency (parameter A in Eq. 4) is notably weaker in our samples than in vivo 369 and reported by others. In a similar experiment (26), the amplitude of the microstructure 370 frequency of a fresh bovine optic nerve at 7T was -18.75 ppb, significantly larger in magnitude 371 and with an opposite sign to what we have obtained in our CC samples, 1.46 ppb. Additional 372 analysis was performed to consolidate this result (see supplementary Figure S2). A reduction 373 of the microstructural compartmentalisation effect had already been reported in the literature 374 when studying fresh vs fixed rat optic nerves (25). One possible explanation is the structural 375 alteration of the myelin sheath in fixed tissues. In our 3D EM images, we observed myelin 376 sheath spitting and swelling in some of the myelinated axons, similar to the observation 377 reported in the previous study (41), and such phenomena appeared more frequently in larger 378 axons than small axons. Based on the general Lorentzian tensor approach (43), the increase 379 of the aqueous space of the myelin sheath can result in the amplitude reduction of the induced 380 frequency shifts inside the myelin sheath and the intra-axonal space. Microstructural 381 differences related to structures (bovine optic nerve vs human CC) and age-associated 382 demyelination (45,46), together with the tissue preparation methods can also contribute to the

differences observed in this study, as all these factors modulate the relative waterconcentration in the three WM compartments.

385

386 All six specimens obtained from the body of the CC have similar magnetic susceptibility 387 anisotropy, suggesting that they have similar MVF based on the HCM approximation (Eq. S25 388 in (17)), and this is supported by the EM analysis (0.278 & 0.257 between the two samples). 389 The amplitude of the residual field inside the specimens is, on the other hand, subject to 390 various properties including MVF, AVF and the aggregate g-ratio of the sample (Eq. A14 in 391 (16)). Interestingly, the realistic geometry of the WM fibre also plays an important role in the 392 compartmental frequency shifts (27,44). This effect is clearly illustrated in the frequency 393 perturbation simulations in Figure 5: not only the centres but also the FWHM of the extra-394 cellular frequency distribution of the two samples are different, despite the two specimens 395 having virtually identical MVF and AVF. The broader frequency spectrum of CC4 induces a 396 faster R<sub>2</sub>\* decay in the extra-axonal space and the specimen also has a more disperse fibre 397 arrangement. These two factors together reduce the amplitude discrepancy between the slow 398  $R_2^*$  (intra- and extra-axonal water) and the fast  $R_2^*$  (myelin water) compartments throughout 399 the echo time as well as the frequency difference between those compartments that could 400 result in a reduced compartmentalization effect.

401

The experimental setup, particularly the 3D-printed holder, is an effective tool also for other ex vivo studies that involve histology. The grid on the 3D-printed plates not only facilitates tissue excision with high precision but also provides landmarks in MRI images for experiment planning and sample matching. The close to unity linear relations of the susceptibility measurements between the two sessions (Figure 4A and 4B) support that the ROIs drawn on the whole-brain images and the excised specimens are corresponding to each other.

408

409 One limitation of this study is the relatively long fixation time of the specimen. The first imaging session happened after 5 months of fixation instead of the scheduled 2 months because of 410 411 the coronavirus measures that took place in the Netherlands. The prolonged fixation time 412 results in a substantial change of specimen  $R_1$  between the pre-scan and the first session 413 (Figure 2A, 2B; average  $R_1$  across the brain increased from 1.95±0.47 s<sup>-1</sup> to 2.67±0.74 s<sup>-1</sup>). 414 The enhanced R<sub>1</sub> in deep GM suggests the contribution of iron to the R<sub>1</sub> was more pronounced. 415 An alternative and more likely explanation would be that the myelin contribution to  $R_1$  could 416 be reduced by the fixation process, which would also explain the diminished cortical GM and 417 WM contrast observed at 5 months. Experiments conducted with a shorter fixation time can 418 potentially reduce the fixation effects on the signal phase. However, progressive changes in

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relaxation parameters begin in the early stage of fixation (23) and it is likely that fixation induced phase differences could also happen simultaneously. Lastly, the EM analysis can be subject to sampling bias due to the limited field-of-view. Preliminary results of a recent study suggest that large axons in the WM tissue can be under-represented in EM compared to light microscopy when larger field of view is available (45). Such under-representation may have an impact on the realistic geometry of the myelinated axons on the phase data.

### 426 Conclusion

427 The contributions of MR phase contrast observed in the formalin-foxed brain specimen are 428 substantially different from fresh tissue, despite the QSM maps derived from in vivo and ex 429 vivo imaging sharing similar contrasts and values. Particularly, the reductions of magnetic 430 susceptibility anisotropy and compartmentalisation are observed in the fixed WM tissue. An increase of non-susceptibility contributions to phase contrast can also be found in fixed tissue, 431 432 which is potentially introduced by formalin fixation. Therefore, WM magnetic susceptibility and 433 microstructural quantification findings in studies using formalin-fixed tissue should be 434 interpreted with care. Our study suggests that the microstructural effects observed in our samples encode information regarding WM arrangements such as dispersion and packing 435 436 while susceptibility anisotropy encodes myelin volume as was predicted from theory.

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