A 48-channel receive array coil for mesoscopic diffusion-weighted MRI of ex vivo human brain on the 3 T connectome scanner

Alina Scholz, Robin Etzel, Markus W. May, Mirsad Mahmutovic, Qiyuan Tian, Gabriel Ramos-Llordén, Chiara Maffei, Berkin Bilgić, Thomas Witzel, Jason P. Stockmann, Choukri Mekkaoui, Lawrence L. Wald, Susie Yi Huang, Anastasia Yendiki, Boris Keil

Institute of Medical Physics and Radiation Protection (IMPS), TH-Mittelhessen University of Applied Sciences (THM), 14 Wiesenstrasse, Giessen 35390, Germany

A.A. Martinos Center for Biomedical Imaging, Department of Radiology, Massachusetts General Hospital, Boston, MA, USA

Harvard Medical School, Boston, MA, USA

Harvard-MIT Division of Health Sciences and Technology, Cambridge, MA, USA

Center for Mind, Brain and Behavior (CMBB), Marburg, Germany

Abstract

In vivo diffusion-weighted magnetic resonance imaging is limited in signal-to-noise-ratio (SNR) and acquisition time, which constrains spatial resolution to the macroscale regime. Ex vivo imaging, which allows for arbitrarily long scan times, is critical for exploring human brain structure in the mesoscale regime without loss of SNR. Standard head array coils designed for patients are sub-optimal for imaging ex vivo whole brain specimens. The goal of this work was to design and construct a 48-channel ex vivo whole brain array coil for high-resolution and high b-value diffusion-weighted imaging on a 3T Connectome scanner. The coil was validated with bench measurements and characterized by imaging metrics on an agar brain phantom and an ex vivo human brain sample. The two-segment coil former was constructed for a close fit to a whole human brain, with small receive elements distributed over the entire brain. Imaging tests including SNR and G-factor maps were compared to a 64-channel head coil designed for in vivo use. There

This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/)

Credit authorship contribution statement

Alina Scholz: Conceptualization, Methodology, Software, Validation, Formal analysis, Investigation, Writing - original draft, Writing - review & editing, Visualization. Robin Etzel: Methodology, Formal analysis. Markus W. May: Methodology, Data curation. Mirsad Mahmutovic: Methodology, Software, Validation. Qiyuan Tian: Data curation, Software, Formal analysis, Visualization. Gabriel Ramos-Llordén: Data curation, Software, Formal analysis, Visualization. Chiara Maffei: Data curation, Software, Formal analysis, Visualization. Berkin Bilgić: Software. Thomas Witzel: Software. Jason P. Stockmann: Methodology. Choukri Mekkaoui: Software, Funding acquisition. Lawrence L. Wald: Resources, Project administration, Funding acquisition. Susie Yi Huang: Investigation, Writing - original draft, Formal analysis, Writing - review & editing, Resources, Project administration, Funding acquisition. Anastasia Yendiki: Conceptualization, Formal analysis, Writing - original draft, Writing - review & editing. Boris Keil: Methodology, Formal analysis, Writing - original draft, Writing - review & editing, Resources, Project administration, Funding acquisition.
was a 2.9-fold increase in SNR in the peripheral cortex and a 1.3-fold gain in the center when compared to the 64-channel head coil. The 48-channel ex vivo whole brain coil also decreases noise amplification in highly parallel imaging, allowing acceleration factors of approximately one unit higher for a given noise amplification level. The acquired diffusion-weighted images in a whole ex vivo brain specimen demonstrate the applicability and advantage of the developed coil for high-resolution and high b-value diffusion-weighted ex vivo brain MRI studies.

**Keywords**
Magnetic resonance imaging; Diffusion-weighted imaging; RF coil; Receive array coil; Brain imaging; Ex vivo brain

1. **Introduction**

Diffusion MRI (dMRI) is a powerful, non-invasive technique for imaging axonal orientations as well as characterizing white and gray matter microstructure (Conturo et al., 1999; Lagana et al., 2010; McNab et al., 2009; Mori and Zhang, 2006; Okano and Mitra, 2015). The basic premise of dMRI in the human brain is that the diffusion of water molecules in white matter is anisotropic, and that its preferential direction is aligned with the orientation of the underlying fibers (Mori and Zhang, 2006). A series of images, each sensitized to diffusion in a different direction, are acquired and used to infer the most likely orientation of water displacement in every voxel (Basser et al., 1994).

There are several requirements that increase the acquisition time needed for whole-brain dMRI. High spatial resolution is desirable for resolving small brain structures. A large number of diffusion directions must be sampled to improve the angular resolution, i.e., the smallest angle between crossing fiber bundles that can be resolved. Finally, advanced dMRI sampling schemes may require images to be acquired with multiple b-values. Satisfying all these requirements would lead to acquisition times that are prohibitive for in vivo imaging in the absence of any image acceleration. As a result, trade-offs must be made that restrict in vivo whole-brain dMRI to the macroscale regime (Okano and Mitra, 2015; Zeng, 2018), with voxel sizes on the order of 1 to 3 mm. Motion artifacts, which are exacerbated by long acquisitions, and distortions near tissue-air interfaces further degrade the effective resolution that is achievable in vivo.

Many of these issues can be circumvented in ex vivo dMRI, which allows for longer acquisition times, absence of motion and significantly reduced susceptibility artifacts with appropriate sample preparation (Roebroeck et al., 2019). Furthermore, ex vivo imaging enables the placement of coil elements closer to the actual brain tissue to maximize sensitivity. Thus, ex vivo imaging can achieve substantially higher spatial and angular resolution, permitting the anatomy and microstructure of complex fiber pathways to be imaged at the mesoscale, sub-millimeter regime, well beyond what is feasible in vivo. The impressive level of anatomical detail that can be resolved by ex vivo dMRI has already been demonstrated on a variety of human tissue samples (Augustinack et al., 2010; Beaujoign et al., 2018; Fritz et al., 2019; Miller et al., 2011; Modo et al., 2016). Ex vivo dMRI, in
combination with optical imaging, is an excellent tool for validating dMRI acquisition and analysis methods in human brain tissue (Jones et al., 2020; Mollink et al., 2017).

However, various challenges arise when acquiring ex vivo dMRI. These primarily include reduced diffusivity and decreased T2, caused mainly by the fixation, tissue dehydration and lower probe temperature (D’Arceuil et al., 2007; Pfefferbaum et al., 2004; Roebroeck et al., 2019). As a result, dMRI data must be acquired with higher b-values ex vivo to achieve similar diffusion contrast as in vivo. Furthermore, when a conventional in vivo head coil is used, it is challenging to center the ex vivo brain in the coil and to ensure that it remains stable throughout the long scan time.

In addition to the above challenges, the higher spatial resolution of post mortem scans comes at the cost of lower signal-to-noise-ratio (SNR). Several strategies for improving SNR in high-resolution ex vivo dMRI have been proposed and tested, mainly focusing on higher magnetic field strengths (Pallebage-Gamarallage et al., 2018; Plantinga et al., 2016), small-bore MRI scanners (Augustinack et al., 2010; Calamante et al., 2012) or high-performance gradient systems (McNab et al., 2013). One of the main innovations introduced by the NIH Blueprint Human Connectome Project was the development of human scanners with ultra-high gradients, which allow high b-values to be achieved without loss of SNR (Setsompop et al., 2013). Initial results have already shown the advantages of a 300 mT/m gradient system for imaging whole post-mortem human brains at 0.6 mm isotropic resolution (McNab et al., 2013), or smaller, non-human primate brain samples at 0.8 mm isotropic resolution (Eichner et al., 2020). Those results were obtained with an in vivo head coil. Dedicated ex vivo brain coils are known to increase signal reception sensitivity, and a few studies have shown the benefits of multi-channel brain array coils for ex vivo tissue imaging applications (Edlow et al., 2019; Roebroeck et al., 2015; Sengupta et al., 2018).

The aim of this study was to push the limits of spatial and angular resolution in ex vivo dMRI by designing, constructing, and validating a 48-channel (48ch) receive array coil for ex vivo whole human brain examinations. The array coil was developed for high spatial resolution and high b-value dMRI acquisitions with long scan times (a few hours to a few days) on the 3 T Connectome scanner (McNab et al., 2013; Setsompop et al., 2013). This work presents high-sensitivity ex vivo diffusion MRI results obtained in a whole human brain specimen at mesoscale resolution (0.73 mm isotropic) using the 48ch receive coil on the 3 T Connectome scanner and expands on preliminary results that were published in conference proceedings (Scholz et al., 2019).

2. Material and methods

2.1. Coil design and construction

To closely cover a whole human brain, we designed an anatomically-shaped ex vivo brain coil former (Fig. 1a and b) based on a nonlinear brain atlas of the International Consortium for Brain Mapping (ICBM). The coil housing was modeled with 3D computer aided design (CAD) software (Rhino3D, Robert McNeel & Associates, Seattle, WA, USA, version 6). It was designed to completely surround the brain with minimal space between the coil elements and imaging volume. The coil former was split into an upper and lower part, such
that a whole brain can be placed inside the coil container. Both coil segments close with an
overlapping rim structure (Fig. 1c). The coil container can accommodate whole brains with
dimensions of 140 mm in the left-to-right direction, an anterior-to-posterior diameter of 182
mm, and a superior-inferior distance of 110 mm. The completed array coil is shown in Fig.
1e–h.

In the bottom coil segment, we incorporated the mechanics for a plugging slide mechanism
(Fig. 1g and h), which directly plugs it into the scanner’s patient bed. The top coil segment
is connected to the scanner using two standard multi-channel coil plugs. The ex vivo coil
container was designed to allow the brain to be placed at the isocenter of the scanner.

The optimum channel count for the constructed ex vivo brain array was determined by (1)
the given area of the coil former’s surface, and (2) a suitable coil element’s Q-ratio for
maintaining sample noise domination. These constraints resulted in a loop count of 48 and a
loop diameter of 54 mm, which comprises an inductance of about 203 nH. The positions of
the 48 coil elements on the outer surface of the coil former were derived from a hexagonal/
pentagonal tiling pattern (Wiggins et al., 2006), with 30 and 18 coil elements located on
the top and bottom segments, respectively (Fig. 1d). The position and outline of all loop
elements, which are decoupled geometrically from neighboring loops by critical overlap
(Roemer et al., 1990), were incorporated in the CAD model. The majority of the loops was
circularly formed, whereas some loops were arbitrarily shaped to fit over the rim structure.
The critical overlap was determined empirically in previously tested bench measurements
and is about 0.27 times the loop diameter. Standoffs for circuit boards and cable routing
were implemented to provide stable mounting positions. The coil former including its cover
were then 3D-printed in polycarbonate (PC) using a 3D printer (Fortus 350, Stratasys, Eden
Prairie, USA).

2.2. Coil circuit

The loop elements were constructed out of 1.3 mm thick tin-coated copper wire. Compared
to flat circuit board copper traces, the wire loops reduce eddy current losses in a high-
density array coil architecture (Kumar et al., 2009). Implemented small bridges in the
conductor enable one loop to cross over another without touching (Keil and Wald, 2013).

Each coil circuit (Fig. 2) consists of a loop with three symmetrically placed
ceramic capacitors (Series 11, Volttronics, Danville, NJ), one variable plastic capacitor
(GFX2700NM; Sprague Goodman, Westbury, New York, USA), a matching network to the
preamplifier (Siemens AG, Healthineers, Erlangen, Germany), and an actively controllable
detuning resonant circuit. A typically redundant passive detuning safety mechanism for in
vivo examinations was omitted for this ex vivo coil.

The variable capacitor $C_T(3–33 \text{pF})$ was used to fine-tune the loop resonance to the Larmor
frequency at 3T (123.25 MHz). $C_2$ and $C_3$ create a capacitive voltage divider. The variable
capacitor $C_M(3–33 \text{pF}, \text{GFX2700NM}; \text{Sprague Goodman, Westbury, New York, USA})$
provides impedance matching of the loop output to a 50 $\Omega$ noise matched condition needed
by the preamplifier to operate at the lowest noise figure at 123.25 MHz (Reykowski et
al., 1995). To ensure accurate detuning of the loop elements, an active detuning circuit
was implemented. It consists of one of the voltage dividing capacitors $C_3$ and a variable inductor $L$ (Coilcraft Inc., 25–32 nH, 165-02A06L, Cary, IL, USA) in series to a PIN diode $D$ (MA4P4002B-402; Macom, Lowell, MA, USA) (Edelstein et al., 1986). During transmit, a DC current is applied to forward bias the PIN diode. This in turn activates the detuning resonant circuit at the Larmor frequency and generates a high impedance in the loop to suppress current flow. The RF-choke $L_{RFC}$ (Coilcraft Inc, 2.7 μH, 1812CS-333XJLC Cary, IL, USA) and $C_4$ block the RF signal to prevent passing into bias source.

While nearest neighbors use geometrical decoupling, next-nearest neighbors and further coil elements are decoupled by the impedance transformation of the input of the preamplifiers (Roemer et al., 1990). The capacitors $C_2$ and $C_M$ and the preamplifier’s input impedance form a resonant circuit, which enables a voltage-source measurement setup, where RF current flow is minimized. As a consequence, inductive coupling across elements is highly reduced and all coil elements receive independently, while maintaining a 50 Ω output impedance.

Both the matching and detuning network of the coil element are placed on the preamplifier’s daughter board, rather than soldering these components directly to the coil former. Therefore, the daughter board is a part of the coil element. The printed circuit board (PCB) daughter board is connected to the loop with an intermittent pin connector. This setup allows a fast construction process of dense array coils.

According to the RF scanner architecture, pre-amplified signals from two loops elements are multiplexed onto one output coaxial cable. The bundled output cables are passed through cable traps to prevent RF common mode currents on the shield of the coaxial cable (Peterson et al., 2003). The cable traps comprise a wounded coaxial cable bundle, which form an inductance ($\approx$109 nH), and a parallel ceramic high power capacitor (15.2 pF, Series 25, Voltronics, Danville, NJ), which resonates at Larmor frequency. Two traps are incorporated into the cables of the upper array coil segment and one cable trap is located directly in the bottom coil housing part.

### 2.3. Coil bench measurements

For bench measurements during the construction process, a custom-made coil plug simulator was used. It provides voltage for the preamplifiers (3 V) and the opportunity to apply a DC current (100 mA) to bias manually each PIN diode forward, which allows for active detuning of single coil elements. To gather information about bench level metrics, e.g. transmission and reflection measurements, a vector network analyzer (VNA) (ENA series, Agilent Technologies, Santa Clara, CA) and custom-built RF tools such as single / double probes and sniffer probes were used. These measurements included tuning to Larmor frequency, active detuning, preamplifier decoupling and geometrical nearest neighbor decoupling of each coil element.

The loops were tuned under a $S_{21}$ control with a 50 Ω dummy load plugged into the preamplifier socket, while all other elements of the array were detuned. Active detuning was performed by using $S_{21}$ measurement with the double-probe for each loop, while all other coil elements were detuned and the relevant loop under test was switched between the
tuned and detuned state. The difference of both states at the Larmor frequency indicates the magnitude of active detuning. A similar $S_{21}$ double-probe measurement was carried out to determine the effectiveness of the implemented preamplifier decoupling, first by plugging the preamplifier into the socket on the PCB and second by terminating the socket with a load impedance of $50 \, \Omega$. Again, all but the loop element to be tested were detuned.

Coupling of nearest neighbor elements was measured with direct $S_{21}$ VNA measurement by using coaxial cables, which were directly plugged into the preamplifier sockets. During this measurement, all other coil elements were detuned. This measurement configuration was also used to verify $50 \, \Omega$ coil impedance matching using $S_{11}$ and $S_{22}$ measurements (Keil and Wald, 2013; Reykowski et al., 1995).

Furthermore, unloaded-to-loaded coil quality factor ratio ($Q_U/Q_L$) of one representative coil element was measured within the populated but detuned array assembly, using the $S_{21}$ double-probe method (Hoult, 1978). As a load, a fixed tissue brain sample in periodate-lysine-paraformaldehyde (PLP) solution was used.

### 2.4. MRI data acquisition and analysis

Imaging metrics were acquired on a clinical 3T MRI scanner (MAG-NETOM, Skyra, Tim 4G, Dual Density Signal Transfer, Siemens AG, Healthineers, Erlangen, Germany), equipped with a customized gradient coil (AS302 CONNECTOM 1.0 gradient) with a maximum gradient strength of 300 mT/m and a maximum slew rate of 200 T/m/s.

For evaluating the developed ex vivo whole brain array coil, we constructed a human-brain-shaped phantom using a 3D printer (Objet30 Pro, Stratasys, Eden Prairie, USA). The phantom was filled with agarose and dielectrically tuned to match the RF coil’s loading condition with the PLP-packed ex vivo brain. The corresponding quantities were: 830 ml distilled H$_2$O, 29 g NaCl, 12.5 g of agar powder (Sigma-Aldrich Corp., St. Louis, MO) and 936 g sugar. The matched loading condition was validated via an $S_{11}$ measurement on the VNA’s smith chart at Larmor frequency. The dielectric values of the phantom were measured to be $\sigma = 0.49 \, \text{S/m}$ and $\varepsilon_r = 66.3$ with a VNA equipped with a dielectric probe kit (85070E kit, Agilent Technologies, Santa Clara, CA).

#### 2.4.1. Array coil characterization

For determining SNR and G-factor, the phantom was scanned with a proton density (PD)-weighted FLASH sequence (repetition time (TR) = 200 ms, echo time (TE) = 4.8 ms, flip angle (F) = 15°, matrix (M): 192×192 (SNR) and 64×64 (G-factor and SNR in parallel imaging), field of view (FOV): 256×256 mm$^2$, slice thickness: 8 mm, bandwidth (BW): 200 Hz/pixel). Information about noise correlation was obtained with the same sequence but without RF excitation. The coil sensitivities for the G-factor calculations were derived from a pre-scan before the actual MRI data acquisition. This scan provided a low-resolution full FOV image of the phantom for each coil element, which was used to estimate the sensitivity profiles of the individual receiver coil.

---

1 under development and not commercially available in the U.S. and its future availability cannot be assured.
Pixel-wise SNR maps were calculated using the noise-covariance-weighted, root sum-of-squares image reconstruction method from Kellman and McVeigh (2005). To evaluate the array coil’s encoding capability for parallel imaging, SENSE G-factor maps were computed using the acquired noise correlation matrix and complex sensitivities of the coil elements (Pruessmann et al., 1999). The FOV of the G-maps was tightly enclosed to the phantom, in order to enhance the aliasing pattern inside the imaging object.

A valuable metric is the remaining image SNR after the parallel imaging acceleration has been performed. We calculated the remaining SNR by dividing the SNR globally by the square root of the reduction factor $R$ and further locally with the noise amplification given by the G-factor.

For further characterization of the coil performance, we examined the encoding power for simultaneous multislice (SMS) acquisitions with blipped-controlled aliasing in parallel imaging (Feinberg et al., 2010; Larkman et al., 2001; Setsompop et al., 2012). To assess the encoding capability of combined SMS and in-plane acceleration, a reduction factor of $R = 2$ and a slice acceleration factor from $MB = 4$ up to $MB = 8$ with a 1/3 FOV shift were evaluated. Noise correlation and SNR and G-factor maps of the 48ch ex vivo brain coil were compared to a customized 64-channel (64ch) whole head receive array coil (Keil et al., 2013) with identical acquisition parameters.

In addition, time course stability of each coil element was measured with a single-shot, gradient-echo, echo planar imaging (EPI) sequence (time points: 500, $TR = 1000 \text{ ms}$, $TE = 30 \text{ ms}$, $F = 90^\circ$, $M: 64 \times 64$, FOV: $200 \times 200 \text{ mm}^2$, slices: 16 slices of 15 mm, BW: 2298 Hz/pixel) with the brain phantom. This scan was repeated 16 times without pause, resulting in a 2 h stability scan protocol. The average intensity of a 15-pixel square region of interest (ROI) in the phantom center was detrended with linear and quadratic temporal trends and plotted. The stability was calculated as the variation of signal intensity from peak-to-peak as a percentage from the average signal intensity (Weisskoff, 1996).

### 2.4.2. Ex vivo brain dMRI

High-resolution (0.73 mm isotropic) diffusion imaging was performed on a whole ex vivo human brain packed with paraformaldehyde-lysine-periodate (PLP) in a tight-fitting, sealed plastic bag. The brain had been excised from a male who had died of non-neurological causes, and had been placed in fixative (10% formaldehyde) for 90 days before being transferred to PLP solution for long-term storage. Diffusion-weighted images were acquired using the same imaging protocol on the 48ch whole brain ex vivo coil and 64ch in vivo head coil to enable comparisons between the two coils. We used a 3D diffusion-weighted spin-echo segmented EPI sequence (imaging parameters: $TR = 500 \text{ ms}$, $TE = 65 \text{ ms}$, echo spacing: $1.22 \text{ ms}$, $M: 160 \times 268 \times 208$, FOV: $118 \times 196 \times 152 \text{ mm}^3$, BW: 1244 Hz/pixel, 16 shots, EPI factor = 10, no partial Fourier). A multi-shell sampling scheme was used that included 18 non-collinear diffusion encoding directions with $b = 4000 \text{ s/mm}^2$ (gradient strength of 91 mT/m, $\delta = 16.1 \text{ ms}$, $\Delta = 27.6 \text{ ms}$), and 36 non-collinear diffusion encoding directions with $b = 10000 \text{ s/mm}^2$ (gradient strength of 133 mT/m, $\delta = 16.1 \text{ ms}$, $\Delta = 27.6 \text{ ms}$). A total of 9 $b = 0$ volumes were acquired, interleaved every 6 diffusion-weighted volumes. The total acquisition time for the $b = 4000 \text{ s/mm}^2$ scan was 10.3 h. The total acquisition time for the $b = 10000 \text{ s/mm}^2$ scan was 20.6 h. The
phase-encoding direction was anterior-posterior when considering the conventional sagittal plane. Since the brain in the constructed coil was rotated compared to the usual orientation of a patient, the anatomical axis of the phase-encoding direction was inferior-superior.

The image acquisition parameters were chosen so as to maximize SNR while avoiding prohibitively long scan times. The use of a 3D read-out with excitation of the entire imaging volume per TR enabled the use of a relatively short TR of 500 ms while preserving SNR efficiency (Miller et al., 0000). Due to the reduction of $T_1$ and $T_2$ values in fixed human brain tissue (340 ms and 45 ms, respectively (McNab et al., 2009)) compared to in vivo values, a relatively short TE of 65 ms was chosen to preserve SNR. To achieve the TE and $b$-values used here, the maximum gradient strength was calculated to be on the order of 90–130 mT/m based on the sequence design, which seeks to maximize gradient strength while keeping the diffusion time $\Delta$ as short as possible.

Diffusion-weighted volumes were corrected for eddy current distortions with the *eddy* tool from FSL (Andersson and Sotiropoulos, 2016). The multi-shot acquisition mitigated EPI distortions and thus no further distortion correction was performed prior to image analysis. We performed a diffusion tensor imaging (DTI) analysis (Basser, 1995) on the images from the lower shell, and a diffusion kurtosis imaging (DKI) analysis (Jensen et al., 2005) on the images from both shells. Finally, the full data set was used to fit a fiber orientation distribution (FOD) at each voxel with multi-shell, multi-tissue, constrained, spherical deconvolution (MSMT-CSD) (Dhollander et al., Montreal, 2019). Probabilistic tractography was then performed on these FODs (Tournier et al., Stockholm, 2010), with a minimum FOD amplitude of 0.2, a maximum bending angle of 45 degrees, and a minimum streamline length of 20. We seeded at every voxel in a white-matter mask, which we extracted by thresholding the average of the diffusion-weighted volumes to remove the PLP background.

### 3. Results

#### 3.1. Coil bench measurements

The $Q_{U}/Q_L$-ratio of a 54 mm loop element was measured to be 233/46=5.1 with six surrounding but non-resonant neighboring loops. Thus, the array’s loop elements operate in the sample noise dominated regime. The geometrical decoupling of nearest neighbors was $S_{21}$ measured with an average value of −16 dB and ranged from −14 dB to −18 dB. Non-adjacent and thus non-overlapping coil elements, which are primarily decoupled via preamplifier decoupling, obtained an average decoupling value of −18 dB with a range from −17 dB to −19 dB. The isolation between tuned and detuned states caused by the active detuning circuit reached an average value of 42 dB.

#### 3.2. Image performance

Time course stability tests show a peak-to-peak variation of 0.36% over 8,000 time-points EPI sequence measured in a ROI comprising 15×15 pixels.

**3.2.1. Signal-to-noise-ratio in unaccelerated images**—Fig. 3 shows the noise correlation matrix of the 48ch ex vivo coil and that of the 64ch in vivo coil. The ex vivo
array has a range of noise correlations from 0.02% to 35.8% with an average value of 7.5%, while the in vivo array has noise correlations from 0.12% to 53.8% with an average value of 7.1% for the off-diagonal elements.

Fig. 4 compares the SNR maps from the newly developed 48ch ex vivo brain coil to that of the existing, custom 64ch whole head coil, in different planes of the agar phantom. For both coils, the measured SNR is highest in the outer periphery and decreases towards the center. The SNR gain of the newly constructed 48ch array coil reaches over the whole brain volume. It outperforms the larger 64ch in vivo head coil by a factor of 2.5, when the average SNR over the whole brain phantom is considered. The highest gain is found in the periphery of the phantom, especially in the regions where the 48ch brain coil has a substantially closer proximity to the sample. In the periphery and in the center of the phantom, a 2.9-fold and 1.3-fold SNR gain was measured, respectively.

Examples of SNR profiles can be found in Fig. 5. In all three slices, the SNR of the 48ch ex vivo brain coil exceeds that of the 64ch in vivo head coil over the entire profile.

3.2.2. Signal-to-noise-ratio and G-factor in parallel imaging—Fig. 6 shows the SENSE inverse G-factor maps in a representative coronal plane of the brain phantom for both one-dimensional and two-dimensional acceleration obtained from the 48ch ex vivo brain coil and the 64ch in vivo head coil. The newly constructed 48ch coil provides significant improvement compared to the 64ch head coil for both in-plane acceleration types. Both coils show minimal noise amplifications for acceleration factors of $R = 2$, $R = 3$ and $R = 2 \times 2$. However, for higher accelerations ($R > 3$) the 48ch ex vivo coil provides favorable encoding capabilities when compared to the 64ch in vivo head coil. At $R = 4$, the 48ch coil shows on average a 16% lower G-factor than the 64ch head coil. When comparing the peak G-factors between both, the 48ch coil shows a 21% improvement. The enhanced encoding power of the 48ch coil becomes even more apparent when very high acceleration factors are compared. The improved average and peak G-factor for $R = 7$ was measured to be 35% and 41% lower. At $R = 5 \times 5$ the noise amplifications could be reduced on average by 43%, while the peak G-factor decreased by 53%.

A more meaningful figure of merit is the SNR obtained from the accelerated image, where both the under-sampled k-space trajectory and the local noise amplification were taken into account. Fig. 7 illustrates the accelerated SNR for both coils using box plots. Since the constructed 48ch coil provides both a higher baseline SNR and lower G-factors, it highly outperforms the 64ch head coil across all acceleration scenarios. The average SNR from the 64ch coil only reaches the lower 25th percentile of the 48ch ex vivo coil. Further, it should be noted that the relative gain in average SNR increases with higher acceleration factors (e.g., factor 2.4 for $R = 2$ and 3.9 for $R = 7$) for both one-dimensional and two-dimensional acceleration. For a direct comparison between 1D and 2D accelerations, when the reduction factor $R$ was matched ($R_{1D} = 4$ and $R_{2D} = 2 \times 2 = 4$), the average SNR of the 48ch ex-vivo coil was measured to be 600 and 657, respectively. For the 64ch head coil, these numbers were 210 and 268, respectively. Thus, both coils showed overall less noise amplification in the 2D-acceleration scheme.
3.2.3. Signal-to-noise-ratio and G-factor in simultaneous multislice imaging imaging—Fig. 8 compares the inverse G-factor maps for the SMS image reconstruction technique from a coronal slice of the brain phantom. Compared to the 64ch head coil, the constructed 48ch coil indicates overall substantially lower noise amplification for the SMS examination, as well as for combined SMS and in-plane acceleration. At a multi-band factor of $MB = 4$, the 48ch coil generates negligible noise amplifications ($g_{\text{mean}} = 1.0002$ and $g_{\text{max}} = 1.0569$), while the 64ch head coil shows substantial noise gains $g_{\text{mean}} = 1.1218$ and $g_{\text{max}} = 1.6345$. Furthermore, the dedicated 48ch ex vivo brain coil achieves similar to slightly better encoding capabilities at $MB = 8$ as the 64ch head coil at $MB = 4$ ($g_{\text{mean48ch}} = 1.0047$ vs. $g_{\text{mean64ch}} = 1.1317$ and $g_{\text{max48ch}} = 1.2913$ vs. $g_{\text{max64ch}} = 1.2913$). Therefore, the 48ch coil allows the application of a slice acceleration factor of $MB = 8$ with negligible noise gain.

To assess the accelerated SNR during SMS acquisitions, Fourier averaging needs to be taken into account: In the case of the $MB = 8$ acceleration, eight times more $^1$H spins are simultaneously excited compared with a single-slice acquisition. Thus, for a multiband factor $MB$, the SNR efficiency can be improved up to a factor of $\sqrt{MB}$, if the imaging parameters between the non-accelerated and the SMS-accelerated case remain identical. Under these constant circumstances, the Fourier averaging translates to an SNR increase by a factor of up to $\frac{\sqrt{8}}{g_{\text{max}} - 48} = 2.2$, when compared to a commonly used consecutive single-slice acquisition schemes. The $MB = 8$ achievable SNR obtained from the 64ch is only increased by a factor of up to $\frac{\sqrt{8}}{g_{\text{max}} - 64} = 2.2$. In direct comparison, when the baseline SNR, Fourier averaging, and G-factors are taken into account, the 48ch coil achieves up to a 4.5-fold SNR improvement at $MB = 8$ compared to the 64ch head coil.

3.2.4. Diffusion imaging in ex vivo brain—Fig. 9 shows various maps obtained from the multi-shell dMRI scan of a post mortem human brain, acquired at 0.73 mm isotropic resolution with the 48ch ex vivo brain coil. The six columns show: (i) a $b = 0$ image, (ii) a diffusion-weighted image from the lower shell ($b = 4000$ s/mm$^2$), (iii) the fractional anisotropy (FA) map, (iv) the FA map color encoded by the principal eigenvector of the diffusion tensor, (v) a diffusion-weighted image from the higher shell ($b = 10000$ s/mm$^2$, and (vi) the mean kurtosis map.

For each map in the figure, an axial view is shown in row (a) and a coronal or sagittal view is shown in row (c). Row (b) shows magnified regions of interest that highlight fine anatomical detail in the striatum (red box, columns i-ii and iv-v) or in the basal ganglia and thalamus (green box, columns iii and vi). Note that the exquisite mean kurtosis contrast allows a clear delineation of internal structures such as the putamen, caudate nucleus, internal and external global pallidus, and thalamus. Row (d) shows a magnified coronal view of the primary motor cortex (blue box, columns i-ii and iv-v), highlighting the presence of radial fibers, and a magnified sagittal view of the hippocampus (yellow box, columns iii and vi). These maps illustrate that, by combining the high gradient strengths available on the 3 T Connectome scanner with the high sensitivity of our ex vivo coil, we can collect dMRI data with high spatial resolution and high SNR, and map detailed gray and white matter anatomy, both in deep brain and near the cortical surface.
Fig. 10 compares the FODs and probabilistic tractography obtained from the multi-shell dMRI data collected with the 48ch ex vivo brain array and the 64ch in vivo head array. The higher SNR achieved by the newly developed ex vivo coil, in both superficial and deep brain areas, yielded less noisy FODs that better captured the course of the underlying fiber bundles. This resulted in higher-quality tractography. For example, the tractogram from the 48ch brain array data included fuller bundles of trans-callosal streamlines, as well as subcortical U-shaped streamlines. The lower SNR of the 64ch head coil led to more spurious peaks in the FODs, which had a negative impact on the ability of tractography to reconstruct these bundles.

4. Discussion

We designed, constructed, and evaluated a 48ch ex vivo brain array receive coil for high-resolution and high b-value dMRI of a whole ex vivo human brain on the 3 T Connectome scanner (McNab et al., 2013; Setsompop et al., 2013). The coil was characterized by both bench tests and image metrics. Bench tests included element measurements of the coil quality factor $Q$, active detuning, geometrical decoupling, and preamplifier decoupling. MRI evaluations included measurements of the noise correlation, pixel-wise SNR, and G-factor, as well as time course stability using a brain shaped agar phantom. We demonstrated the coil’s performance in achieving high SNR with the acquisition of multi-shell 0.73 mm isotropic resolution diffusion-weighted MR images of a whole ex vivo brain.

4.1. Array coil characterization

In many applications, large channel count arrays with relatively small loop sizes such as the 54 mm loops used here are necessary to increase both reception sensitivity and encoding power. However, very small loop elements quickly lose their sample noise dominance. Under these circumstances, small elements do not translate to higher SNR acquisitions anymore. For in vivo imaging at 3T, this critical size is reached at about 60 mm diameter (Keil et al., 2011). In ex vivo brain imaging, however, loop sizes can be made substantially smaller than for in vivo imaging. This is attributed to the brain fixation medium, which has a higher conductivity compared to in vivo tissue and thus provides a higher fraction of sample noise. While the noise increases in the ex vivo sample, the electronic noise can be decreased by omitting in vivo human safety features in the coil element circuitry, such as passive detuning and RF-fuses. This condition results in an enhanced $Q_L / Q_U$-ratio when using small receiver elements. Therefore, the implemented loop size of $\approx 54$ mm provides a relatively high $Q_L / Q_U$-ratio of 5.1, outperforming most coils optimized for in vivo applications with loop diameters ranging from 50 mm to 65 mm from our previous studies (Janssens et al., 2012; Keil et al., 2011; 2013). As a consequence, the minimum loop diameter at which sample noise dominance is maintained decreases for imaging fixed tissue brain samples in PLP solution, allowing us to contemplate very high-density arrays for ex vivo sample examinations.

Despite RF electrical optimizations, the mechanical coil former is an important and critical design aspect for ex vivo imaging. To improve SNR, the loops were populated very close to the sample, maximizing signal reception. Thus, the completely brain-enclosing coil former
with uniformly distributed loop elements guarantees nearly omni-directional signal reception from the sample (decreased sensitivity was observed with some loops aligned to be almost parallel to the magnetic field \( B_0 \)). However, an entirely surrounding coil array requires a split housing mechanism, which disturbs the loop layout and makes it difficult to maintain geometric decoupling at the split housing edge. Therefore, an overlapping edge structure was implemented, enabling adjacent loop elements to be geometrically decoupled across the two housing segments, while the overall array coil structure remains self-contained.

In array coil design, the central ultimate SNR is already approached with only 12 surrounding coil elements at 3 T (Wiesinger et al., 2005). Implementing higher loop element counts only yields SNR improvements at the periphery for a given geometry. Nevertheless, relative central SNR gains are achievable with tightly fitting array coils. Due to the lack of dedicated \textit{ex vivo} receiver arrays, \textit{in vivo} head coils are commonly used in many \textit{ex vivo} brain studies (Iglesias et al., 2018; McNab et al., 2009; Shatil et al., 2016; 2018). However, these coils are not well suited in terms of sample fitting and SNR performance. Optimizing both the mechanical features for close fitting of samples and the RF circuitry can thus result in significant SNR gains in the brain. This implementation provides a 30% SNR increase of the 48ch coil at the phantom center when compared to the larger 64ch head coil. In addition, in the peripheral regions of the brain phantom, the tight-fitting form factor also provides favorable SNR gains, as evidenced by an almost 3-fold SNR improvement over the 64ch coil. The high SNR can be exploited to reduce the voxel size, enabling high spatial resolution MR imaging of a whole \textit{ex vivo} brain.

The average noise correlation of 9% indicates a well decoupled array and highly independent operating receiver loops. Adjacent loops show much higher coupling values up to 36%, which can be attributed to insufficient overlap, resulting in a remaining mutual inductance and shared resistance especially in the sample voxels beneath the overlapping loop regions.

The constructed 48ch \textit{ex vivo} brain coil shows remarkably better encoding performance when compared to the 64ch head coil. The encoding power of the 48ch coil enables approximately one additional acceleration unit, for both one-dimensional and two-dimensional accelerations, with the same noise amplification as the 64ch head coil. Improvements in G-factors are usually achieved by implementing higher channel counts on a given geometry. However, when comparing array coil formers of different sizes, similar improvements in G-factors can be achieved by (1) reducing the diameter of the coil elements at constant or even lower channel counts, and (2) positioning the coil elements in close proximity to the sample. The tight-fitting, smaller loop elements of the constructed 48ch coil provide an overall stronger spatial modulation in the signal sensitivity’s magnitude and phase. Consequently, this coil arrangement allows favorable encoding capabilities for unaliasing folded images (SENSE method) or synthesizing spatial harmonics (GRAPPA or SMASH methods). Additionally, the entirely enclosed \textit{ex vivo} coil former of the 48ch coil leads to better spatial coverage for the aliased pixels when compared to a head array coil, which obviously has limited coverage along the inferior aspect and in the area covering the face.
When the total acceleration factor $R$ is the product of the acceleration factors from two individual orthogonal phase-encoding directions, the images show less amplified noise in comparison to $R$-fold 1D accelerations. This can be attributed to the embedded sensitivity variations of the array coil being efficiently exploited in two spatial dimensions, allowing overall more favorable encoding capabilities in the image reconstruction.

Reducing scan time using parallel imaging techniques is not strictly essential when constraints on acquisition time are lifted for ex vivo examinations. On the other hand, 2D acquisitions are still often used despite their SNR inefficiency per unit time (Eichner et al., 2020; Miller et al., 2011). For example, mapping tissue microstructural features throughout the whole human brain involves measurements at multiple $b$-values (Huang et al., 2020), and protocol optimization may be facilitated by 2D scans acquired at resolutions on the order of 0.8 to 1 mm isotropic. For such 2D acquisitions, slice acceleration enables the excitation and measurement of multiple slices (Feinberg et al., 2010; Setsompop et al., 2018; 2012). Unlike conventional parallel imaging, which requires under-sampled data acquisition, these techniques provide acceleration by exciting the spins in multiple slices at the same time using multi-band radiofrequency pulses. These newer multi-band MR acquisitions have the SNR advantages of 3D sampling based on Fourier averaging (Larkman et al., 2001; Setsompop et al., 2012). Therefore, SNR efficiency can be improved by up to a factor of $\sqrt{MB}$, if the imaging parameters remain the same. In practice, however, the SNR gain is reduced locally by the SMS G-factor of the coil and globally due to changes in the sequence parameters. At higher SMS acceleration factors, shorter repetition times are employed, which decreases the level of steady-state longitudinal magnetization. Therefore, the optimal SMS acceleration factor for an MRI study is a trade-off between the benefits of the simultaneously acquired volume and the negative impact from local noise amplifications and the chosen repetition times.

The SNR recovery achieved by the SMS method is highly advantageous for dMRI, which normally suffers from low signal strength. Therefore, it is advantageous for ex vivo array coils to provide a high encoding capability for SMS in order to accommodate modern acquisition techniques. Commonly used in vivo head coils do not optimally fulfill this requirement for SMS ex vivo scans, as they lack enough elements in the $z$-direction. The radially surrounding, $z$-directional, stacked elements of the constructed coil provide favorable spatial coverage for SMS image encoding, allowing the separation of multiple collapsed slices. In the case of an $MB = 8$ acceleration scheme, the combination of the enhanced SMS encoding power and the increased baseline SNR of the 48ch coil, would provide an up to 4.5-fold SNR improvement when compared to the 64ch head coil.

4.2. **Diffusion imaging in ex vivo brain**

Previous work comparing ex vivo dMRI to optical imaging suggests that high spatial resolution (1 mm or higher) improves the accuracy of dMRI-derived axonal orientation estimates, and may have a greater impact than high angular resolution or ultra-high $b$-values (Jones et al., 2020). While dMRI acquisitions with sub-mm resolution have been feasible in pre-clinical scanners, for smaller samples, they have remained a challenge for whole human brains. The 48ch ex vivo brain coil that we have developed here has enabled us to collect
high-resolution (0.73 mm isotropic) dMRI data from a whole ex vivo human brain. We have shown that this coil achieves higher SNR than a 64ch in vivo head coil throughout the brain, leading to improved delineation of brain circuitry with dMRI tractography. The 48ch brain coil exhibits the highest SNR gains in the peripheral area of the imaged sample, which will be of particular utility for imaging fiber architecture in the cortex with high precision. Examples of this in our results are the radial fibers in the primary motor cortex in Fig. 9 (column iv, row d), as well as the subcortical U-shaped tracts in Fig. 10. Future work in laminar microstructure in the cortex will greatly benefit from this coil. The high sensitivity of the coil enables high spatial resolution in combination with very high b-values, as we have shown with our preliminary results at $b = 10000 \text{s/mm}^2$. Indeed, mean kurtosis maps obtained from the acquired data reveal exquisite contrast that captures the internal structure of the basal ganglia and hippocampus.

The coil presented here paves the way for sub-mm resolution ex vivo dMRI on whole human brains at the high $b$-values accessible on the 3 T Connectome scanner. This capability will allow us to map the connectational anatomy and microstructure of the human brain at unprecedented resolutions, as well as provide reference data for evaluating in vivo dMRI scans to gain deeper insight into human brain structure at multiple scales. Currently, 3D EPI suffers from ghosting artifacts when both spatial resolution and gradient amplitude are increased up to the limit of the Connectome scanner’s capacity. In future work, we will address this issue by combining appropriate $k$-space reconstruction techniques (Ramos-Llorden et al., 2021) and the use of an additional field monitoring camera system (Mahmutovic et al., 2021). We expect this novel coil design, in combination with the current 3 T Connectome scanner equipped with 300 mT/m gradient strengths and next-generation gradient system planned for the Connectome 2.0 project (Yendiki et al., 2020), to advance our understanding of human brain circuitry in health and disease.

### 5. Conclusion

A 48ch close-fitting receive array coil for dMRI of whole ex vivo human brains at 3 T was designed, constructed, and tested with a brain-shaped phantom and an ex vivo brain. We characterized the coil with unloaded-to-loaded $Q$-ratio, noise correlation, SNR, G-factor, SMS G-factor and stability measurements in comparison to a 64ch whole-head in vivo coil. Compared to in vivo array coils, smaller loop sizes can be used for ex vivo brain samples due to increased loading characteristics of the fixed brain tissue. This allows the design of high-channel count arrays, improving both peripheral SNR and encoding performance for accelerated imaging. Due to the high SNR and parallelism, the designed coil is well-suited for high-resolution, high $b$-value ex vivo dMRI acquisitions and will enable to map the connectomics and microstructure of the human brain at multiple scales.

### Acknowledgments

Funding: This work was supported by the National Institutes of Health [R01EB021265, 1U01EB026996, R01HL131635, 1P41EB030006]; and the Federal Ministry of Education and Research Germany (BMBF) [IN2016-2-226].
**Abbreviations:**

- 48ch: 48-channel
- 64ch: 64-channel
- ABS: acrylonitrile butadiene styrene
- BW: bandwidth
- CAD: computer aided design
- CSD: constrained spherical deconvolution
- dMRI: Diffusion MRI
- DKI: diffusion kurtosis imaging
- DTI: diffusion tensor imaging
- DWI: Diffusion-Weighted Imaging
- DWMRI: Diffusion-weighted Magnetic Resonance Imaging
- EPI: echo planar imaging
- EPROM: Erasable Programmable Read-Only Memory
- F: flip angle
- FA: fractional anisotropy
- FOD: fiber orientation distribution
- FOV: field of view
- M: matrix
- MRI: Magnetic Resonance Imaging
- MSMT-CSD: multi-shell, multi-tissue, constrained, spherical deconvolution
- PC: polycarbonate
- PCB: printed circuit board
- PD: proton density
- PLP: periodate-lysine-paraformaldehyde
- RF: radio frequency
- ROI: region of interest
- SMS: simultaneous multislice
- SNR: signal-to-noise-ratio

*Neuroimage. Author manuscript; available in PMC 2021 September 14.*
**References**

Andersson JL, Sotiropoulos SN, 2016. An integrated approach to correction for off-resonance effects and subject movement in diffusion MR imaging. Neuroimage125, 1063–1078. doi:10.1016/j.neuroimage.2015.10.019. [PubMed: 26481672]

Augustinack JC, Helmer K, Huber KE, Kakunoori S, Zöllei L, Fischl B, 2010a. Direct visualization of the perforant pathway in the human brain with ex vivo diffusion tensor imaging. Front. Hum. Neurosci4, 42. doi:10.3389/fnhum.2010.00042. [PubMed: 20577631]

Basser PJ, 1995. Inferring microstructural features and the physiological state of tissues from diffusion-weighted images. NMR Biomed8 (7), 333–344. doi:10.1002/nbm.1940080707. [PubMed: 8739270]

Basser PJ, Mattiello J, LeBihan D, 1994. MR Diffusion tensor spectroscopy and imaging. Biophys. J66 (1), 259–267. doi:10.1016/S0006-3495(94)80775-1. [PubMed: 8130344]

Beaujoin J, Palomero-Gallagher N, Boumezbeur F, Axer M, Bernard J, Poupon F, Schmitz D, Mangin J-F, Poupon C, 2018. Post-mortem inference of the human hippocampal connectivity and microstructure using ultra-high field diffusion MRI at 11.7 T. Brain Struct. Funct223 (5), 2157–2179. doi:10.1007/s00429-018-1617-1. [PubMed: 29387938]

Calamante F, Tournier J-D, Kurniawan ND, Yang Z, Gyengesi E, Galloway GJ, Reutens DC, Connelly A, 2012. Super-resolution track-density imaging studies of mouse brain: comparison to histology. Neuroimage59 (1), 286–296. doi:10.1016/j.neuroimage.2011.07.014. [PubMed: 21777683]

Conturo TE, Lori NF, Cull TS, Akbadak E, Snyder AZ, Shimony JS, McKinstry RC, Burton H, Raichle ME, 1999. Tracking neuronal fiber pathways in the living human brain. Proc. Natl. Acad. Sci96 (18), 10422–10427. doi:10.1073/pnas.96.18.10422. [PubMed: 10468624]

Dhollander T, Mito R, Raffelt D, Connelly A, Montreal, 2019. Improved white matter response function estimation for 3-tissue constrained spherical deconvolution. In: Proceedings of the 27th Annual Meeting of ISMRM, p. 555.

D’Arceuil HE, Westmoreland S, de Crespigny AJ, 2007. An approach to high resolution diffusion tensor imaging in fixed primate brain. Neuroimage35 (2), 553–565. doi:10.1016/j.neuroimage.2006.12.028. [PubMed: 17292630]

Edelstein W, Hardy C, Mueller O, 1986. Electronic decoupling of surface-coil receivers for NMR imaging and spectroscopy. J. Magn. Reson67 (1), 156–161. doi:10.1016/0022-2364(86)90421-X.

Edlow BL, Mareyam A, Horn A, Polimeni JR, Witzel T, Tisdall MD, Augustinack JC, Stockmann JP, Diamond BR, Stevens A, Tirell LS, Folkert RD, Wald LL, Fischl B, Van der Kouwe A, 2019. 7 Tesla MRI of the ex vivo human brain at 100 micron resolution. Sci. Data6 (1), 1–10. doi:10.1038/s41597-019-0254-8. [PubMed: 30647409]

Eichner C, Paquette M, Mildner T, Schlumm T, Pléh K, Samuni L, Crockford C, Wittig RM, Jäger C, Möller HE, Friederici AD, Anwander A, 2020. Increased sensitivity and signal-to-noise ratio in diffusion-weighted MRI using multi-echo acquisitions. Neuroimage221, 117172. doi:10.1016/j.neuroimage.2020.117172. [PubMed: 32682095]

Feinberg DA, Moeller S, Smith SM, Auerbach E, Ramanna S, Glasser MF, Miller KL, Ugurbil K, Yacoub E, 2010. Multiplexed echo planar imaging for sub-second whole brain FMRI and fast diffusion imaging. PLoS ONE5 (12). doi:10.1371/journal.pone.0015710.

Fritz F, Sengupta S, Harms R, Tse D, Poser B, Roebroeck A, 2019. Ultra-high resolution and multi-shell diffusion MRI of intact ex vivo human brains using kT-dsteam at 9.4 t. Neuroimage202, 116087. doi:10.1016/j.neuroimage.2019.116087. [PubMed: 31408716]

Hoult D, 1978. The NMR receiver: a description and analysis of design. Prog. Nucl. Magn. Reson. Spectrosc12 (1), 41–77. doi:10.1016/0039-6565(78)80002-8.
Huang SY, Tian Q, Fan Q, Witzel T, Wichtmann B, McNab JA, Bireley JD, Machado N, Klawiter EC, Mekkaoui C, Wald LL, Nummenmaa A, 2020. High-gradient diffusion MRI reveals distinct estimates of axon diameter index within different white matter tracts in the in vivo human brain. Brain Struct. Funct. 225, 1277–1291. doi:10.1007/s00429-019-01961-2. [PubMed: 31563995]

Iglesias JE, Insausti R, Lerma-Usabiaga G, Boccetta M, Van Leemput K, Greve DN, Van der Kouwe A, Fischl B, Caballero-Gades C, Paz-Alonso PM, 2018. A probabilistic atlas of the human thalamic nuclei combining ex vivo MRI and histology. Neuroimage 183, 314–326. doi:10.1016/j.neuroimage.2018.08.012. [PubMed: 30121337]

Janssens T, Keil B, Farivar R, McNab JA, Polimeni JR, Gerits A, Arsenault J, Wald LL, Vanduffel W, 2012. An implanted 8-channel array coil for high-resolution macaque MRI at 3 T. Neuroimage 62 (3), 1529–1536. doi:10.1016/j.neuroimage.2012.05.028. [PubMed: 22609793]

Jensen JH, Helperna J, Ramani A, Lu H, Kaczynski K, 2005. Diffusional kurtosis imaging: the quantification of non-gaussian water diffusion by means of magnetic resonance imaging. Magn. Reson. Med 53 (6), 1432–1440. doi:10.1002/mrm.20508. [PubMed: 15906300]

Jenssens T, Keil B, Farivar R, McNab JA, Polimeni JR, Gerits A, Arsenault J, Wald LL, Vanduffel W, 2012. An implanted 8-channel array coil for high-resolution macaque MRI at 3 T. Neuroimage 62 (3), 1529–1536. doi:10.1016/j.neuroimage.2012.05.028. [PubMed: 22609793]

Jones R, Grisot G, Augustinack J, Magnain C, Boas DA, Fischl B, Wang H, Yendiki A, 2020. Insight into the fundamental trade-offs of diffusion MRI from polarization-sensitive optical coherence tomography in ex vivo human brain. Neuroimage 214, 116704. doi:10.1016/j.neuroimage.2020.116704. [PubMed: 32151760]

Keil B, Alagappan V, Mareyam A, McNab JA, Fujimoto K, Tountcheva V, Triantafyllou C, Dilks DD, Kanwisher N, Lin W, Grant PE, Wald LL, 2011. Size-optimized 32-channel brain arrays for 3 T pediatric imaging. Magn. Reson. Med 66 (6), 1777–1787. doi:10.1002/mrm.22961. [PubMed: 21656458]

Keil B, Blau JN, Biber S, Hoecht P, Tountcheva V, Setsompop K, Triantafyllou C, Wald LL, 2013. A 64-channel 3T array coil for accelerated brain MRI. Magn. Reson. Med 70 (1), 248–258. doi:10.1002/mrm.24427. [PubMed: 22851312]

Keil B, Wald LL, 2013. Massively parallel MRI detector arrays. J. Magn. Reson. 229, 75–89. doi:10.1016/j.jmr.2013.02.001. [PubMed: 23453758]

Kellman P, McVeigh ER, 2005. Image reconstruction in SNR units: a general method for SNR measurement. Magn. Reson. Med 54 (6), 1439–1447. doi:10.1002/mrm.20713. [PubMed: 16261576]

Kumar A, Edelstein WA, Bottomley PA, 2009. Noise figure limits for circular loop mr coils. Larkman DJ, Hajnal JV, Herlihy AH, Coutts GA, Young IR, Ehrholm G, 2001. Use of multicoil arrays for separation of signal from multiple slices simultaneously excited. J. Magn. Reson. Imaging 13 (2), 313–317. doi:10.1002/1522-2586(200102)13:2<313::AID-JMRI1045>3.0.CO;2-W. [PubMed: 11169840]

Mahmutovic M, Scholz A, Kutscha N, May M, Schlumm T, Möller R, Pine K, Edwards L, Weiskopf N, Brunner D, Möller H, Keil B, 2021. A 64-channel brain array coil with an integrated 16-channel field monitoring system for 3T MRI. In: Proceedings of the 29th Annual Meeting of ISMRM, p. 0623.

McNab JA, Edlow BL, Witzel T, Huang SY, Bhat H, Heberlein K, Feiweier T, Liu K, Keil B, Cohen-Adad J, Tsidall MD, Folkert RD, Kinney HC, Wald LL, 2013. The human connectome project and beyond: initial applications of 300 mT/m gradients. Neuroimage 80, 234–245. doi:10.1016/j.neuroimage.2013.05.074. [PubMed: 23711537]

McNab JA, Jbabdi S, Deoni SC, Douaud G, Behrens TE, Miller KL, 2009. High resolution diffusion-weighted imaging in fixed human brain using diffusion-weighted steady state free precession. Neuroimage 46 (3), 775–785. doi:10.1016/j.neuroimage.2009.01.008. [PubMed: 19344686]

Miller KL, Stagg CJ, Douaud G, Jbabdi S, Smith SM, Behrens TE, Jenkinson M, Chance SA, Esiri MM, Voets NL, Jenkinson N, Aziz TZ, Turner MR, Johansen-Berg H, McNab JA,. Diffusion imaging of whole, post-mortem human brains on a clinical MRI scanner. NeuroImage.

Miller KL, Stagg CJ, Douaud G, Jbabdi S, Smith SM, Behrens TE, Jenkinson M, Chance SA, Esiri MM, Voets NL, Jenkinson N, Aziz TZ, Turner MR, Johansen-Berg H, McNab JA, 2011. Diffusion

Neuroimage. Author manuscript; available in PMC 2021 September 14.
imaging of whole, post-mortem human brains on a clinical MRI scanner. Neuroimage57 (1), 167–181. doi:10.1016/j.neuroimage.2011.03.070. [PubMed: 21473920]

Modo M, Hitchens TK, Liu JR, Richardson RM, 2016. Detection of aberrant hippocampal mossy fiber connections: ex vivo mesoscale diffusion MRI and microtractography with histological validation in a patient with uncontrolled temporal lobe epilepsy. Hum. Brain Mapp37 (2), 780–795. doi:10.1002/hbm.23066. [PubMed: 26611565]

Mollink J, Kleinijenhuis M, van Cappellen van Walsum A-M, Sotiropoulos SN, Cottaar M, Mirfin C, Heinrich MP, Jenkinson M, Pallebage-Gamarallage M, Ansorge O, Jbabdi S, Miller KL, 2017. Evaluating fibre orientation dispersion in white matter: comparison of diffusion MRI, histology and polarized light imaging. Neuroimage157, 561–574. doi:10.1016/j.neuroimage.2017.06.001. [PubMed: 28602815]

Mori S, Zhang J, 2006. Principles of diffusion tensor imaging and its applications to basic neuroscience research. Neuron51 (5), 527–539. doi:10.1016/j.neuron.2006.08.012. [PubMed: 16950152]

Okano H, Mitra P, 2015. Brain-mapping projects using the common marmoset. Neurosci. Res93, 3–7. doi:10.1016/j.neures.2014.08.014. [PubMed: 25264372]

Pallebage-Gamarallage M, Foxley S, Menke RA, Huszar IN, Jenkinson M, Tendler BC, Wang C, Jbabdi S, Turner MR, Miller KL, Ansorge O, 2018. Dissecting the pathobiology of altered MRI signal in amyotrophic lateral sclerosis: a post mortem whole brain sampling strategy for the integration of ultra-high-field MRI and quantitative neuropathology. BMC Neurosci19 (1), 11. doi:10.1186/s12868-018-0416-1. [PubMed: 29529995]

Peterson DM, Beck BL, Duensing GR, Fitzsimmons JR, 2003. Common mode signal rejection methods for MRI: reduction of cable shield currents for high static magnetic field systems. Concepts Magn. Reson. Part B Magn. Reson. Eng19 (1), 1–8. doi:10.1002/cmr.b.10090.

Pfefferbaum A, Sullivan EV, Adalsteinsson E, Garrick T, Harper C, 2004. Post-mortem MR imaging of formalin-fixed human brain. Neuroimage21 (4), 1585–1595. doi:10.1016/j.neuroimage.2003.11.024. [PubMed: 15050582]

Plantinga BR, Roebroeck A, Kemper VG, Uludağ K, Melse M, Mai J, Kuijf ML, Herrler A, Jahanshahi A, ter Haar Romeny BM, Temel Y, 2016. Ultra-high field MRI post mortem structural connectivity of the human subthalamic nucleus, substantia nigra, and globus pallidus. Front. Neuroanat10, 66. doi:10.3389/fnana.2016.00066. [PubMed: 27378864]

Preussmann KP, Weiger M, Scheidegger MB, Boesiger P, 1999. SENSE: Sensitivity encoding for fast MRI. Magn. Reson. Med42 (5), 952–962. doi:10.1002/(SICI)1522-2594(199911)42:5<952::AID-MRM16>3.0.CO;2-S. [PubMed: 10542355]

Ramos-Llorden G, Lobos RA, Kim TH, Tian Q, Touretki S, Witzel T, Keil B, Yendiki A, Bilgic B, Haldar JP, Huang S, 2021. Improved multi-shot EPI ghost correction for high gradient strength diffusion MRI using structured low-rank modeling k-space reconstruction. In: Proceedings of the 29th Annual Meeting of ISMRM, p. 1346.

Reykowski A, Wright SM, Porter JR, 1995. Design of matching networks for low noise preamplifiers. Magn. Reson. Med33 (6), 848–852. doi:10.1002/mrm.1910330617. [PubMed: 7651124]

Roebroeck A, Miller KL, Aggarwal M, 2019. Ex vivo diffusion MRI of the human brain: technical challenges and recent advances. NMR Biomed32, e3941. doi:10.1002/nbm.3941. [PubMed: 29863793]

Roebroeck A, Sengupta S, Bastiani M, Schillak S, Tramm B, Waks M, Lataster A, Herrler A, Tse D, Poser B, 2015. High resolution MRI neuroanatomy of the whole human brain post mortem with a specialized 9.4 T RF-coil. In: Proceedings of the Organization for Human Brain Mapping, p. 1856. doi:10.13140/RG.2.2.21380.07041.

Roemer PB, Edelstein WA, Hayes CE, Souza SP, Mueller OM, 1990. The NMR phased array. Magn. Reson. Med16 (2), 192–225. doi:10.1002/mrm.1910160203. [PubMed: 2266841]

Scholz A, May M, Etzel R, Mahmutovic M, Kutscha N, Wald LL, Yendiki A, Keil B, 2019. A 48-Channel ex vivo Brain Array Coil for Diffusion-Weighted MRI at 3T. In: Proceedings of the 27th Annual Meeting of ISMRM, p. 1494.
Sengupta S, Fritz F, Harms R, Hildebrand S, Tse D, Poser BA, Goebel R, Roebroeck A, 2018. High resolution anatomical and quantitative MRI of the entire human occipital lobe ex vivo at 9.4 t. Neuroimage168, 162–171. doi:10.1016/j.neuroimage.2017.03.039. [PubMed: 28336427]

Setsompop K, Fan Q, Stockmann J, Bilgic B, Huang S, Cauley S, Nummenmaa A, Wang F, Rathi Y, Witzel T, LL W, 2018. High-resolution in vivo diffusion imaging of the human brain with generalized slice dithered enhanced resolution: simultaneous multislice (gslider-SMS). Magn. Reson. Med141–151. doi:10.1002/mrm.26653. [PubMed: 28261904]

Setsompop K, Gagoski BA, Polimeni JR, Witzel T, Wedeen VJ, Wald LL, 2012. Blipped-controlled aliasing in parallel imaging for simultaneous multislice echo planar imaging with reduced g-factor penalty. Magn. Reson. Med67 (5), 1210–1224. doi:10.1002/mrm.23097. [PubMed: 21858868]

Setsompop K, Kimmlingen R, Eberlein E, Witzel T, Cohen-Adad J, McNab JA, Keil B, Tisdall MD, Hoecht P, Dietz P, Cauley S, Tountcheva V, Matschl V, Lenz V, Heberlein K, Pothen A, Thein H, Van Horn J, Toga A, Schmitt F, Lehne D, Rosen B, Wedeen V, Lawrence L, 2013. Pushing the limits of in vivo diffusion MRI for the human connectome project. Neuroimage80, 220–233. doi:10.1016/j.neuroimage.2013.05.078. [PubMed: 23707579]

Shatil AS, Matsuda KM, Figley CR, 2016. A method for whole brain ex vivo magnetic resonance imaging with minimal susceptibility artifacts. Front. Neuro7, 208. doi:10.3389/fneur.2016.00208. [PubMed: 27965620]

Shatil AS, Uddin MN, Matsuda KM, Figley CR, 2018. Quantitative ex vivo MRI changes due to progressive formalin fixation in whole human brain specimens: longitudinal characterization of diffusion, relaxometry, and myelin water fraction measurements at 3T. Front. Med5, 31. doi:10.3389/fmed.2018.00031.

Tournier J-D, Calamante F, Connelly A, Stockholm, 2010. Improved probabilistic streamlines tractography by 2nd order integration over fibre orientation distributions. In: Proceedings of the 18th Annual Meeting of ISMRM, p. 1670.

Weisskoff RM, 1996. Simple measurement of scanner stability for functional NMR imaging of activation in the brain. Magn. Reson. Med36 (4), 643–645. doi:10.1002/mrm.1910360422. [PubMed: 8892220]

Wiesinger F, De Zanche N, Pruessmann K, 2005. Approaching ultimate SNR with finite coil arrays. In: Proceedings of the 13th Annual Meeting of ISMRM, p. 672.

Wiggins GC, Triantafyllou C, Potthast A, Rekowskia A, Nittka M, Wald L, 2006. 32-Channel 3 tesla receive-only phased-array head coil with soccer-ball element geometry. Magn. Reson. Med56 (1), 216–223. doi:10.1002/mrm.20925. [PubMed: 16767762]

Yendiki A, Witzel T, Huang SY, 2020. Connectome 2.0: cutting-edge hardware ushers in new opportunities for computational diffusion MRI. In: Bonet-Carne E, Hutter J, Palombo M, Pizzolato M, Sepehrband F, Zhang F (Eds.), Computational Diffusion MRI. Springer, Switzerland, pp. 3–12. doi:10.1007/978-3-030-52893-5_1.

Zeng H, 2018. Mesoscale connectomics. Curr. Opin. Neurobiol50, 154–162. doi:10.1016/j.conb.2018.03.003. [PubMed: 29579713]
Fig. 1.
48ch ex vivo brain coil (a-c) Computer aided design model of the coilformer with graved loops and standoffs for preamplifier boards. (a) Top coilformer part. (b) Bottom coilformer part. (c) Inner side of both coilformer parts with overlapping frames to allow geometrical decoupling of the loops from top and bottom part. (d) Placement of the 30 top loops (blue) and the 18 bottom loops (green) around the brain (gray). (e-h) Completely constructed coil consisting of the top part (e and f) and bottom part (g).
Fig. 2.
Circuit schematic for one coil element. Each loop consists of three fixed capacitors ($C_1$–$C_3$) and one variable capacitor ($C_T$). $C_T$ fine-tunes the resonant frequency of the coil to Larmor frequency corresponding at 3T. $C_2$ and $C_3$ create a capacitive voltage divider. $C_3$ is part of the active detuning circuit (blue) together with the variable inductor $L$ and the PIN diode $D$. $C_2$ and $C_M$ (green) provide both impedance matching of the loop and impedance transformation to establish preamplifier decoupling. Typical values for the components are: $C_1 = 33 \text{ pF}$, $C_2 = 56 \text{ pF}$, $C_3 = 56 \text{ pF}$, $C_4 = 2.2 \text{ nF}$, $C_T \approx 18 \text{ pF}$, $C_M \approx 18 \text{ pF}$, $L \approx 24.5 \text{ nH}$, $L_{RFC} = 2.7 \text{ \mu H}$. 

Neuroimage. Author manuscript; available in PMC 2021 September 14.
Fig. 3.
Noise correlation matrix of the 48ch ex vivo brain coil and the 64ch in vivo head coil with the scale normalized to 1.
Fig. 4.
Comparison of the SNR, normalized to 100, of a transverse (left), coronal (middle) and sagittal (right) slice of the brain phantom with the 48ch ex vivo brain coil (top row), the 64ch head coil (middle row) and ratio maps between the two coils (bottom row). The 48ch ex vivo brain coil shows a 1.3-fold SNR gain in the center and a 2.9-fold SNR improvement in the peripheral regions when compared to the 64ch head coil.
Fig. 5.
SNR profiles of the 48ch ex vivo brain coil (blue) and the 64ch in vivo head coil (green) through the center of a transverse (top), a coronal (middle), and a sagittal (bottom) slice. The dedicated 48ch ex vivo coil shows substantial SNR gains at the periphery in the brain phantom. Due to the close-fitting coil array with omnidirectional signal reception, even SNR improvements at the center of the brain phantom of \( \approx 30\% \) are feasible.
Fig. 6.
Comparison of inverse phantom G-factor maps between the 48ch ex vivo brain coil (top row) and the 64ch head coil (middle row) for different acceleration factors ($R$) obtained from a representative coronal slice. The bottom row shows the ratio maps of the inverse G-factors obtained from both coils. The G-Factors from the 48ch ex vivo brain coil show overall lower noise amplification, when compared to the 64ch head coil.
Fig. 7.
Parallel imaging accelerated SNR as a function of acceleration factor ($R$) from the 48ch brain coil and the 64ch head coil for one-dimensional (top) and two-dimensional (bottom) accelerations. The box plots represent median (horizontal line), average (cross mark) lower/upper quartiles and minimum-maximum range (whiskers) without outliers. The constructed 48ch coil shows higher accelerated SNR in the entire range of acceleration factors.
Fig. 8. Comparison of inverse G-factor maps of the brain phantom for accelerated imaging with SMS technique. The 48ch brain coil shows overall considerable lower noise amplification in comparison to the 64ch head coil.
Fig. 9.
High-resolution DTI and DKI results at 0.73 mm isotropic resolution with \( b = 0 \) images (column i), diffusion-weighted images (DWI) acquired at \( b = 4000 \text{ s/mm}^2 \) along left-right diffusion-encoding direction (column ii), diffusion-weighted images (DWI) acquired at \( b = 10000 \text{ s/mm}^2 \) along left-right diffusion-encoding direction (column iii), fractional anisotropy (FA) maps color encoded by the primary eigenvectors (V1) from DTI (column iv), FA maps (column v) and mean kurtosis maps (column vi). Row (a) shows axial views of each map. Row (b) shows enlarged regions of interest in the internal capsule (red box) and basal ganglia (green box). Row (c) shows coronal or sagittal views for each map. Row (d) shows enlarged regions of interest in subcortical white matter (blue box) and hippocampus (yellow box). The images highlight the fine-scale gray and white matter anatomy captured by the data, both in deep and superficial brain areas.
Fig. 10.
Comparison of FODs and probabilistic tractography in the same ex vivo human brain, between data acquired with the 48ch ex vivo brain coil and the 64ch head coil. A whole-brain, coronal color map of the principal diffusion directions as obtained by CSD illustrates the location of the selected regions of interest. For the three smaller boxes in the corpus callosum (1), superior frontal gyrus (2), and middle frontal gyrus (3), we compare FODs obtained with the 48ch ex vivo brain coil (left) and the 64ch head coil (right). For the larger box centered over the corpus callosum (outlined in blue dashed lines), we show tractography results obtained with the two coils (see blue dashed insets at the bottom of the figure). We show tractography streamlines from the corpus callosum (in red) and adjacent U-fibers (blue, light blue, green). The higher SNR achieved by the 48ch ex vivo brain coil resulted in less noisy FODs, which in turn improved the quality of tractography in these bundles.