

# 7 Tesla MRI of the *ex vivo* human brain at 100 micron resolution

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## 21 Abstract

22 We present an ultra-high resolution MRI dataset of an *ex vivo* human brain  
 23 specimen. The brain specimen was donated by a 58-year-old woman who  
 24 had no history of neurological disease and died of non-neurological causes.  
 25 After fixation in 10% formalin, the specimen was imaged on a 7 Tesla MRI  
 26 scanner at 100  $\mu\text{m}$  isotropic resolution using a custom-built 31-channel  
 27 receive array coil. Single-echo multi-flip Fast Low-Angle SHot (FLASH) data  
 28 were acquired over 100 hours of scan time (25 hours per flip angle), allowing  
 29 derivation of a T1 parameter map and synthesized FLASH volumes. This  
 30 dataset provides an unprecedented view of the three-dimensional  
 31 neuroanatomy of the human brain. To optimize the utility of this resource, we  
 32 warped the dataset into standard stereotactic space. We now distribute the  
 33 dataset in both native space and stereotactic space to the academic  
 34 community via multiple platforms. We envision that this dataset will have a  
 35 broad range of investigational, educational, and clinical applications that will  
 36 advance understanding of human brain anatomy in health and disease.

37

<b>Design Type(s)</b>	Single measure design
<b>Measurement Type(s)</b>	Nuclear magnetic resonance assay
<b>Technology Type(s)</b>	7 Tesla MRI scanner
<b>Factor Type(s)</b>	
<b>Sample Characteristic(s)</b>	Homo sapiens • brain

38

## Background & Summary

Postmortem *ex vivo* MRI provides significant advantages over *in vivo* MRI for visualizing the microstructural neuroanatomy of the human brain. Whereas *in vivo* MRI acquisitions are constrained by time (i.e. ~hours) and affected by motion, *ex vivo* MRI can be performed without time constraints (i.e. ~days) and without cardiorespiratory or head motion. The resultant advantages for characterizing neuroanatomy at microscale are particularly important for identifying cortical layers and subcortical nuclei<sup>1-5</sup>, which are difficult to visualize even in the highest-resolution *in vivo* MRI datasets<sup>6,7</sup>. *Ex vivo* MRI also provides advantages over histological methods that are associated with distortion and tearing of human brain tissue during fixation, embedding, and slide-mounting.

As the field of *ex vivo* MRI has developed over the past two decades, several laboratories have focused on imaging blocks of tissue from human brain specimens using small-bore scanners<sup>2,8</sup> and specialized receive coils<sup>9-11</sup>. This approach allows for spatial resolutions of up to 35–75 microns for analyses of specific neuroanatomic regions<sup>9,11-13</sup>. However, ultra-high resolution imaging of whole human brain specimens at high magnetic field strengths has been far more challenging, due to the need for multi-channel receive coils and large-bore clinical scanners that can accommodate a whole-brain specimen. Whole-brain imaging is required to observe neuroanatomic relationships across distant brain regions, as well as to provide a complete view of human neuroanatomy in standard stereotactic space.

Here, we report the results of a multidisciplinary effort to image a whole human brain specimen *ex vivo* at an unprecedented spatial resolution of 100

64  $\mu\text{m}$  isotropic. Central to this effort was the construction of an integrated  
 65 system consisting of a custom-built 31-channel receive array coil and volume  
 66 transmit coil, which was designed to accommodate and tightly enclose an ex  
 67 vivo human brain<sup>14</sup>. The scans were performed on a 7 Tesla whole-body  
 68 human MRI scanner using four single-echo spoiled gradient-recalled echo  
 69 (SPGR/GRE) or Fast Low-Angle SHot (FLASH) sequences. We used varying  
 70 flip-angles (FA15°, FA20°, FA25°, FA30°) to generate multiple synthesized  
 71 volumes, each of which provides a different tissue contrast. The scans,  
 72 performed over ~100 hours (~25 hours per FA), generated an ~8 TB dataset  
 73 (~2 TB per flip angle) that required custom-built computational tools for offline  
 74 MRI reconstruction and creation of the synthesized volumes. Offline MRI  
 75 reconstruction considerably reduces the data amount. We release the  
 76 resulting FA25° acquisition, as well as the synthesized FLASH25 volume here,  
 77 both in native space and coregistered to standard stereotactic space, for use  
 78 by the academic community. We envision a broad range of investigational,  
 79 educational, and clinical applications for this dataset that have the potential to  
 80 advance understanding of human brain anatomy in health and disease.

81

## 82 **Methods**

### 83 **Specimen acquisition and processing**

84 A 58-year-old woman with a history of lymphoma and stem cell  
 85 transplantation, but no history of neurological or psychiatric disease, died in a  
 86 medical intensive care unit. She was initially admitted to the hospital for  
 87 fevers, chills, and fatigue, and then was transferred to the intensive care unit  
 88 for hypoxic respiratory failure requiring mechanical ventilation. Her hospital  
 89 course was also notable for a deep venous thromboses and a pulmonary

90 embolism. The cause of her death on hospital day 15 was determined to be  
91 hypoxic respiratory failure due to viral pneumonia. At the time of her death,  
92 her surrogate decision-maker provided written informed consent for a clinical  
93 autopsy and for donation of her brain for research, as part of a protocol  
94 approved by our Institutional Review Board.

95 At autopsy, her fresh brain weighed 1,210 grams (normal range = 1,200  
96 to 1,500 grams). The brain was fixed in 10% formalin 14 hours after death.  
97 Gross examination revealed a normal brain (Fig. 1), without evidence of mass  
98 lesions or cerebrovascular disease. To ensure adequate fixation and prevent  
99 specimen flattening (which can prevent specimens from fitting into custom ex  
100 vivo MRI coils), we followed a series of standard specimen processing  
101 procedures, as previously described<sup>15</sup>.

102

### 103 **Specimen preparation for scanning**

104 After remaining in fixative for 35 months, the brain specimen was transferred  
105 to Fomblin Y LVAC 06/6 (perfluoropolyether, Solvay Specialty Polymers USA,  
106 LLC, West Deptford, NJ), which is invisible to MR and reduces magnetic  
107 susceptibility artifacts. The specimen, immersed in Fomblin, was then  
108 secured inside a custom-built, air-tight brain holder made of rugged  
109 urethane<sup>16</sup>. The brain holder contains degassing ports for removal of air  
110 bubbles, which further reduces magnetic susceptibility artifacts.

111

### 112 **Construction of a receive array coil and transmit volume coil for *ex vivo*** 113 **imaging of the whole human brain**

114 We built a receive coil apparatus consisting of a 31-channel surface coil loop  
115 array with two halves. The apparatus was fabricated using a 3D printer of

116 slightly larger dimensions than the brain holder, which slides inside the single-  
117 channel birdcage volume transmit coil (Fig. 2). The brain holder is an oblate  
118 spheroid (16 × 19 cm) that conforms to the shape of a whole brain (cerebral  
119 hemispheres + cerebellum + brainstem)<sup>16</sup> (Fig. 2d). It is made of two separate  
120 halves that can be sealed together with a silicone gasket after packing the  
121 brain inside. This holder must also withstand the degassing process when  
122 under vacuum pressure. Degassing is performed in three steps: 1) introducing  
123 vacuum suction into the container with the brain inside, which allows the  
124 bubbles to expand under decreased pressure and exit tissue cavities; 2)  
125 opening the valve to fill the holder with fomblin and then sealing off the fill  
126 valve; and 3) continuation of vacuum suction with low-amplitude vibration of  
127 the holder for 2-6 hours. The vibration facilitates the removal of bubbles from  
128 tissue cavities. All three steps are performed inside a fume hood.

129 The coil former (Fig. 2c) consists of two halves and encloses the brain  
130 holder. The receive array coil consists of 31 detectors (Fig. 2a), with 15  
131 elements on the top half (diameter = 5.5 cm) and 16 on the bottom half  
132 (diameter = 8.5 cm). Coil elements were constructed using 16 AWG wire  
133 loops<sup>17</sup>, each with four or five evenly spaced capacitors (Supplementary Fig.  
134 1). All elements were tuned to 297.2 MHz and matched to a loaded  
135 impedance of 75 Ω to minimize preamplifier noise. Preamplifier decoupling  
136 was achieved with a cable length of 6 cm. Preamplifiers were placed directly  
137 on the coil elements, yielding a substantial reduction in cable losses compared  
138 to a previous 30-channel *ex vivo* brain array<sup>18</sup>. The active detuning circuit was  
139 formed across the match capacitor using an inductor and PIN diode.

140 Tuning, matching, and decoupling of neighboring elements was  
141 optimized on the bench with a brain sample immersed in periodate-lysine-

142 paraformaldehyde (PLP) solution. Because coil loading varies with the fixative  
143 used, the coil must be tuned and matched on the bench using a brain sample  
144 with the correct fixative. (For example, testing can be performed with a brain  
145 sample immersed in PLP or formalin, but not the regular loading phantom  
146 comprised of water and salt). Loops tuned/matched on PLP showed  
147 unloaded-to-loaded quality factor ratio (Q-ratio) of  $Q_{UL}/Q_L = 210/20 = 10.5$ ,  
148 corresponding to an equivalent noise resistance of 11 ohms for the loaded coil  
149 ( $Q = \omega L/R$ ). By contrast, formalin is a less lossy fixative, giving a coil Q-ratio  
150 of  $Q_{UL}/Q_L = 210/60 = 3.5$ , corresponding to an equivalent noise resistance of 4  
151 ohms.

152 A shielded detunable volume coil (Fig. 2) was built for excitation, with  
153 the following parameters and features: band-pass birdcage, diameter 26.7 cm,  
154 and an extended length of 32 cm to accommodate brain samples of larger  
155 dimensions. For the detuning circuit we used diodes in every leg of the  
156 birdcage. These diodes are powered with the high-power chokes, which can  
157 withstand high voltage and short duration inversion pulses.

158 In summary, this coil system incorporates an improved mechanical  
159 design, preamps mounted at the coil detectors, and an extended transmit coil  
160 design capable of producing high-power pulses.

161

## 162 **7 Tesla MRI data acquisition**

163 The brain specimen was scanned on a whole-body human 7 Tesla (7T)  
164 Siemens Magnetom MRI scanner (Siemens Healthineers, Erlangen,  
165 Germany) with the custom-built coil described above. We utilized a GRE  
166 sequence<sup>19</sup> at 100  $\mu\text{m}$  isotropic spatial resolution with the following acquisition  
167 parameters: TR = 40 msec, TE = 14.2 msec, bandwidth = 90 Hz/px, FA = 15°,

168 20°, 25°, 30°. Total scan time for each FA was 25:01:52 [hh:mm:ss], and each  
 169 FA acquisition generated 1.98 TB of raw k-space data. To improve the signal-  
 170 to-noise ratio (SNR) and optimise T<sub>1</sub> modelling, we collected FLASH scans at  
 171 four FAs: 15°, 20°, 25°, 30° (Fig. 3). Accounting for localizers, quality  
 172 assurance (QA) scans, and adjustment scans, the total scan time was 100  
 173 hours and 8 minutes, and we collected nearly 7.92 TB of raw k-space data.

174

# **175 MRI data reconstruction**

176 The size of the k-space data exceeded the storage capacity of the RAID  
 177 provided by the scanner image reconstruction computer. The image  
 178 reconstruction also required more RAM than what was available. We  
 179 therefore implemented software on the scanner to stream the data directly  
 180 via TCP/IP to a server on an external computer added to the scanner network,  
 181 which saved the data as they were received. Because of additional limitations  
 182 related to the total size of the raw data for any single scan, as dictated by the  
 183 imager RAID size, we also divided each acquisition into segments. The  
 184 server on the external computer stored the data as they were acquired,  
 185 creating date stamps for every k-space segment.

186 After the scan was completed, the streamed k-space data were  
 187 transferred to a computational server where we ran custom software to stitch  
 188 together the segments, reconstruct the images for each channel (via a 3D FFT  
 189 on each volume per channel<sup>20</sup>), and combine the images derived from the 31  
 190 channels via the root-sum-of-squares of the signal magnitudes at each voxel.  
 191 These signal magnitudes were channel-wise decorrelated using a covariance  
 192 matrix of the channels' thermal noise. The output from coil combination was  
 193 the final acquired image (Data Citation 1; Videos 1, 2 and 3).



194

## 195 **MRI data processing**

196 The acquired data underwent a series of processing steps, culminating in the  
197 creation of a  $T_1$  parameter map and synthesized FLASH volumes (Fig. 3 and  
198 Fig. 4; Videos 4, 5, and 6; Data Citations 1 and 2). The volumes were  
199 estimated directly from the four FLASH acquisitions using the DESPOT1  
200 algorithm<sup>19,21</sup> with the program 'mri\_ms\_fitparms' distributed in FreeSurfer  
201 (<http://surfer.nmr.mgh.harvard.edu>) to quantify tissue properties independent  
202 of scanner and sequence types. This algorithm fits the tissue parameters (i.e.  
203  $T_1$ ) of the signal equation for the FLASH scan at each voxel using multiple  
204 input volumes. The volumes at the originally acquired TRs and flip angles  
205 were then regenerated from the parameter maps by evaluating the FLASH  
206 signal equation. In principle, a volume with any TR and flip angle combination  
207 could be synthesized. These synthesized volumes are created from all the  
208 acquired data, and therefore have better SNR than the individually acquired  
209 input volumes. We choose to release the 25 degree synthetic volume as it  
210 has maximal SNR and the best apparent contrast for cortical and subcortical  
211 structures<sup>9</sup>.

212 Of note, *ex vivo* MRI of the fixed human brain yields a different contrast  
213 than *in vivo* MRI, mainly from a shortened  $T_1$ , but also from a decrease in  $T_2^*$ ,  
214 both of which are related to formalin fixation<sup>22</sup>. The predominant source of  
215 signal contrast in *ex vivo* MRI is likely myelin<sup>23</sup> and/or iron<sup>24</sup>. Specifically,  
216 myelin appears to be a source of  $T_1$  contrast, while cortical iron appears to be  
217 a source of  $T_2^*$  contrast<sup>25</sup>.

218

## 219 **Coregistration of the dataset to standard stereotactic space**

220 The dataset was spatially normalized into the MNI ICBM 2009b NLIN ASYM  
 221 template<sup>26</sup> (Supplementary Fig. 2a). This template constitutes the newest  
 222 version of the “MNI space” and is considered a high-resolution version of MNI  
 223 space because it is available at 0.5 mm isotropic resolution. To combine  
 224 structural information present on T<sub>1</sub> and T<sub>2</sub> versions of the template, we  
 225 created a joint template using PCA, as previously described<sup>27</sup>. The four  
 226 synthesized FLASH volumes (FA15, FA20, FA25, and FA30) were  
 227 downsampled to isotropic voxel-sizes of 0.5 mm for spatial normalization and  
 228 initially registered into template space in a multispectral approach using  
 229 Advanced Normalization Tools (ANTs; <http://stnava.github.io/ANTs/>; <sup>28</sup>). This  
 230 multispectral approach simultaneously accounts for intensity data in all four  
 231 volumes. The initial normalization was performed in four stages (rigid body,  
 232 affine, whole brain SyN and subcortically focused SyN) as defined in the  
 233 “effective: low variance + subcortical refine” preset implemented in Lead-DBS  
 234 software ([www.lead-dbs.org](http://www.lead-dbs.org/); <sup>29</sup>).

235 To refine the warp, we introduced fiducial regions of interest (ROI)  
 236 iteratively using a tool developed for this task (available within Lead-DBS).  
 237 Specifically, we manually drew line and point fiducial markers in both native  
 238 and template spaces (Supplementary Fig. 2b). In addition, we manually  
 239 segmented four structures in native space (subthalamic nucleus, internal and  
 240 external pallidum and red nucleus). The three types of fiducials (line ROI,  
 241 spherical ROI and manual segmentations of key structures) were then added  
 242 as “spectra” in subsequent registration refinements (Supplementary Fig. 2c).  
 243 Thus, the final registration consisted of a large number of pairings between  
 244 native and template space (the first four being the actual anatomical volumes,  
 245 the subsequent ones being manual segmentations and paired helper

fiducials). To achieve maximal registration precision, the warp was refined in over 30 iterations with extensive manual expert interaction, each refinement continuing directly from the last saved state. We used linear interpolation to create the normalized files in the data release (Data Citations 1 and 3).

250

## 251 **Code availability**

Neuroimaging data were processed using standard processing pipelines (<http://surfer.nmr.mgh.harvard.edu/>, <https://github.com/freesurfer/freesurfer>). All code used for registration of volumes into standard stereotactic space are available within the open-source Lead-DBS software (<https://github.com/leaddbs/leaddbs>). Because registration involved multiple manual user interface steps, no ready-made code is provided, but the process can be readily reproduced with the provided data and software.

259

## 260 **Data Records**

The native space FA25° acquisition and synthesized FLASH25 volume are available for download at <https://datadryad.org> (Data Citation 1). Additional synthesized volumes are available upon request to the corresponding author. Axial, coronal, and sagittal videos of the native space FA25° acquisition (Videos 1, 2, and 3) and synthesized FLASH25 volume (Videos 4, 5, and 6) are also available at the Dryad data repository (Data Citation 1). The synthesized FLASH25 volume is available for interactive, online viewing at <https://histopath.nmr.mgh.harvard.edu> (Data Citation 2). The normalized FLASH25 volume in standard stereotactic space is available at the Dryad data repository (Data Citation 1) and is hosted on [www.lead-dbs.org](http://www.lead-dbs.org) (preinstalled as part of the LEAD-DBS software package; Data Citation 3).

272

## 273 **Technical Validation**

### 274 **Coil signal-to-noise ratio (SNR) measurements**

275 The receive coil has a  $Q_{UL}/Q_L$  ratio that ranged from 6 in the top half elements  
276 to 8 in the bottom half elements due to larger coil diameter. The  $S_{12}$  coupling  
277 between neighbouring elements, measured with all other coils active detuned,  
278 ranged from  $-10.9$  to  $-24$  dB. All individual elements had  $S_{11} < -20$  dB and  
279 active detuning of  $> 30$  dB. We evaluated the performance of the transmit coil  
280 by examining the  $B_1^+$  profile<sup>14</sup>, which shows the efficiency throughout the  
281 entire spatial distribution of the brain specimen. The efficiency was greatest in  
282 the center of the specimen and fell off gradually towards the edges, as  
283 expected for a whole brain specimen at 7T.

284 We compared the SNR of the 31-channel *ex vivo* array to that of a  
285 standard 31-channel 7T head coil and a 64-channel 3T head coil. SNR maps  
286 were computed following the method of Kellman & McVeigh<sup>30</sup>. We calibrated  
287 the voltage required for  $180^\circ$  pulse using a  $B_1^+$  map (estimated with the AFI  
288 method)<sup>31</sup> with an ROI of 3-cm diameter at the center of the brain. We  
289 estimated array noise covariance from thermal noise data acquired without RF  
290 excitation. The SNR gain with the 31-channel *ex vivo* array was 1.6-fold  
291 versus the 31-channel 7T standard coil and 3.3-fold versus the 64-channel 3T  
292 head array (Fig. 5). The noise coupling between channels was 11% for the  
293 31-channel *ex vivo* array, a 2-fold improvement relative to our previous  
294 array<sup>18</sup>.

295

### 296 **Coregistration accuracy**

297 We assessed the neuroanatomic accuracy of the final registration results (i.e.  
 298 the fit between structures on the normalized FLASH volumes versus the high-  
 299 resolution MNI template) by visual inspection using a tool specifically designed  
 300 for this task (implemented in Lead-DBS). An example of this visual inspection  
 301 assessment for the subthalamic nucleus and globus pallidus interna is  
 302 provided in Supplementary Fig. 3. The final maps are stored in NIfTI and mgz  
 303 files in isotropic 150  $\mu\text{m}$  resolution (Data Citation 1). The normalized  
 304 FLASH25 volume is additionally distributed pre-installed within Lead-DBS  
 305 software and can be selected for visualization in the 3D viewer (Data Citation  
 306 3). Fig. 6 shows an example in synopsis with deep brain stimulation electrode  
 307 reconstructions in a hypothetical patient being treated for Parkinson's  
 308 Disease.

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336

## 337 **Author contributions**

338 B.L.E. designed the study, analyzed the data, and prepared the manuscript.

339 A.M. built the coil, acquired and analyzed the data, and contributed to the

340 manuscript.

341 A.H. created the warp from native space to standard stereotactic space,

342 performed the coregistration for Lead-DBS implementation, and contributed to

343 the manuscript.

344 J.R.P. designed the study, acquired and analyzed the data, and contributed to

345 the manuscript.

346 T.W. designed the study, acquired the data, and contributed to the

347 manuscript.

348 M.D.T. acquired and analyzed the data, and contributed to the manuscript.

349 J.A. designed the study, acquired and analyzed the data, and contributed to

350 the manuscript.

351 J.P.S. advised on the building and testing of the coil, and contributed to the

352 manuscript.

353 B.R.D. analyzed the data and contributed to the manuscript.

354 A.S. acquired and analyzed the data, and contributed to the manuscript.

355 L.S.T. processed and analyzed the data, and contributed to the manuscript.

356 R.D.F. performed the pathological assessment and contributed to the

357 manuscript.

358 L.L.W. supervised the building of the coil and contributed to the manuscript.

359 B.F. supervised and designed the study, analyzed the data, and contributed to

360 the manuscript.



361 A.v.d.K. supervised and designed the study, acquired and analyzed the data,  
362 and contributed to the manuscript.

363 **Additional Information**

364 **Competing interests**

365 None of the authors has a conflicting financial interest. Dr. Fischl and Mr.  
366 Tirrell have financial interest in CorticoMetrics, a company whose medical  
367 pursuits focus on brain imaging and measurement technologies. Their  
368 interests were reviewed and are managed by Massachusetts General Hospital  
369 and Partners HealthCare in accordance with their conflict of interest policies.

## 370    **Figures**

371

372    **Figure 1. Human brain specimen.** The human brain specimen that  
373 underwent *ex vivo* MRI is shown from inferior (**a**), superior (**b**), right lateral (**c**)  
374 and left lateral (**d**) perspectives. Gross pathological examination of the brain  
375 was normal.

376

377    **Figure 2. Receive array coil and transmit volume coil for *ex vivo* imaging**  
378 **of the whole human brain.** (**a**) The 31-channel receive array has 15  
379 elements on the top half (with a diameter of 5.5 cm) and 16 on the bottom half  
380 (with a diameter of 8.5 cm), each made of 16 AWG wire loops with four or five  
381 evenly spaced capacitors. All elements are tuned to 297.2 MHz. (**c**) The coil  
382 former has slightly larger dimensions than the brain holder, which slides inside  
383 a volume coil (**b**). (**d**) A custom air-tight brain holder was designed to conform  
384 to the shape of a whole human brain. The brain holder is an oblate spheroid  
385 container (16 x 19 cm) with degassing ports that are used to apply a vacuum  
386 suction, thereby reducing air bubbles in the specimen and surrounding fomblin  
387 solution.

388

389    **Figure 3. Comparison of FA25° acquisition and synthesized FLASH25**  
390 **volume.** Representative images from the FA25° acquisition (left column) and  
391 the synthesized FLASH25 volume (right column) are displayed in the sagittal  
392 (top row), coronal (middle row) and axial (bottom row) planes. These images  
393 provide a qualitative comparison of the respective signal-to-noise properties of

the FA25° acquisition (~25 hours) and the synthesized FLASH25 volume (~100 hours). All images are shown in radiologic convention.

**Figure 4. Delineation of subcortical neuroanatomy.** Representative axial sections from the synthesized FLASH25 volume are shown at the level of the rostral pons and caudal midbrain (**a-c**, see inset in panel **c**). Zoomed views of the brainstem, medial temporal lobe, and anterior cerebellum (within the white rectangles in **a-c**) are shown in the bottom row (**d-f**). The anatomic detail that can be visualized in this *ex vivo* 100  $\mu$ m resolution MRI dataset is beyond that which can be seen in typical *in vivo* MRI datasets. All images are shown in radiologic convention. Neuroanatomic abbreviations: Amg = amygdala; Cb = cerebellum; CP = cerebral peduncle; MB = mammillary body; P = pons; SCP = superior cerebellar peduncle; VTA = ventral tegmental area; xSCP = decussation of the superior cerebellar peduncle; Th = thalamus.

**Figure 5. Signal-to-noise ratio (SNR) analysis of coil performance.**

Representative SNR maps are shown in the sagittal (top row), coronal (middle row) and axial (bottom row) planes for a test brain sample immersed in periodate-lysine-paraformaldehyde. The maps demonstrate an SNR gain of 1.6-fold for the 31-channel 7 Tesla (7T) *ex vivo* coil (left column) compared to the 31-channel 7T standard coil (middle row), and a gain of 3.3-fold compared to the 64-channel 3T head coil (right column). The noise coupling between channels was 11% for the 31-channel *ex vivo* coil array, a 2-fold improvement relative to our previous array<sup>18</sup>.

419 **Figure 6. Normalization of the *ex vivo* MRI dataset into standard**  
 420 **stereotactic space and integration into the Lead-DBS software platform.**  
 421 **(a)** Exemplary use-case of the normalized FLASH25 volume in deep brain  
 422 stimulation (DBS). DBS electrodes are visualized for a hypothetical patient  
 423 using Lead-DBS software (<https://www.lead-dbs.org>)<sup>29</sup>. An axial image from  
 424 the normalized scan, at the level of the rostral midbrain, is shown as a  
 425 backdrop, with 3D-structures defined by the DISTAL atlas<sup>32</sup> (right subthalamic  
 426 and left red nucleus hidden for optimal visualization of the underlying  
 427 anatomy). Panels **(b)** and **(c)** show zoomed views of key DBS target regions:  
 428 the left globus pallidus interna (GPi in **b**) and the subthalamic nucleus (STN in  
 429 **c**). The images in **(b)** and **(c)** are shown in radiologic convention.

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## 552 **Videos**

553

554 **Video 1. Axial images from the FA25° acquisition.** These images were  
555 acquired in ~25 hours of scan time. The images are shown in radiologic  
556 convention.

557

558 **Video 2. Coronal images from the FA25° acquisition.** These images were  
559 acquired in ~25 hours of scan time. The images are shown in radiologic  
560 convention.

561

562 **Video 3. Sagittal images from the FA25° acquisition.** These images were  
563 acquired in ~25 hours of scan time.

564

565 **Video 4. Axial images from the synthesized FLASH25 volume.** These  
566 images were acquired in ~100 hours of scan time. The images are shown in  
567 radiologic convention.

568

569 **Video 5. Coronal images from the synthesized FLASH25 volume.** These  
570 images were acquired in ~100 hours of scan time. The images are shown in  
571 radiologic convention.

572

573 **Video 6. Sagittal images from the synthesized FLASH25 volume.** These  
574 images were acquired in ~100 hours of scan time.



a



b



c

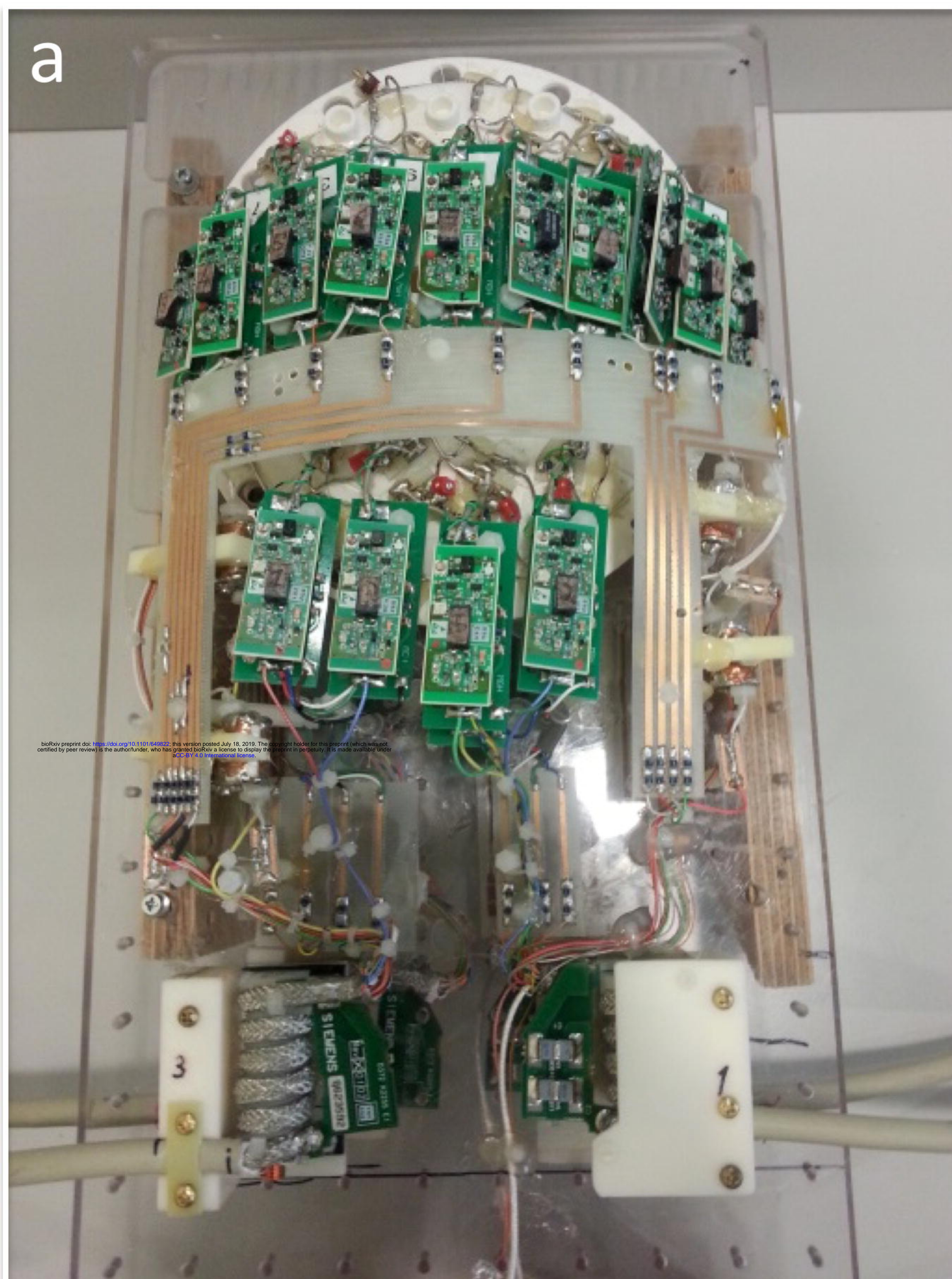


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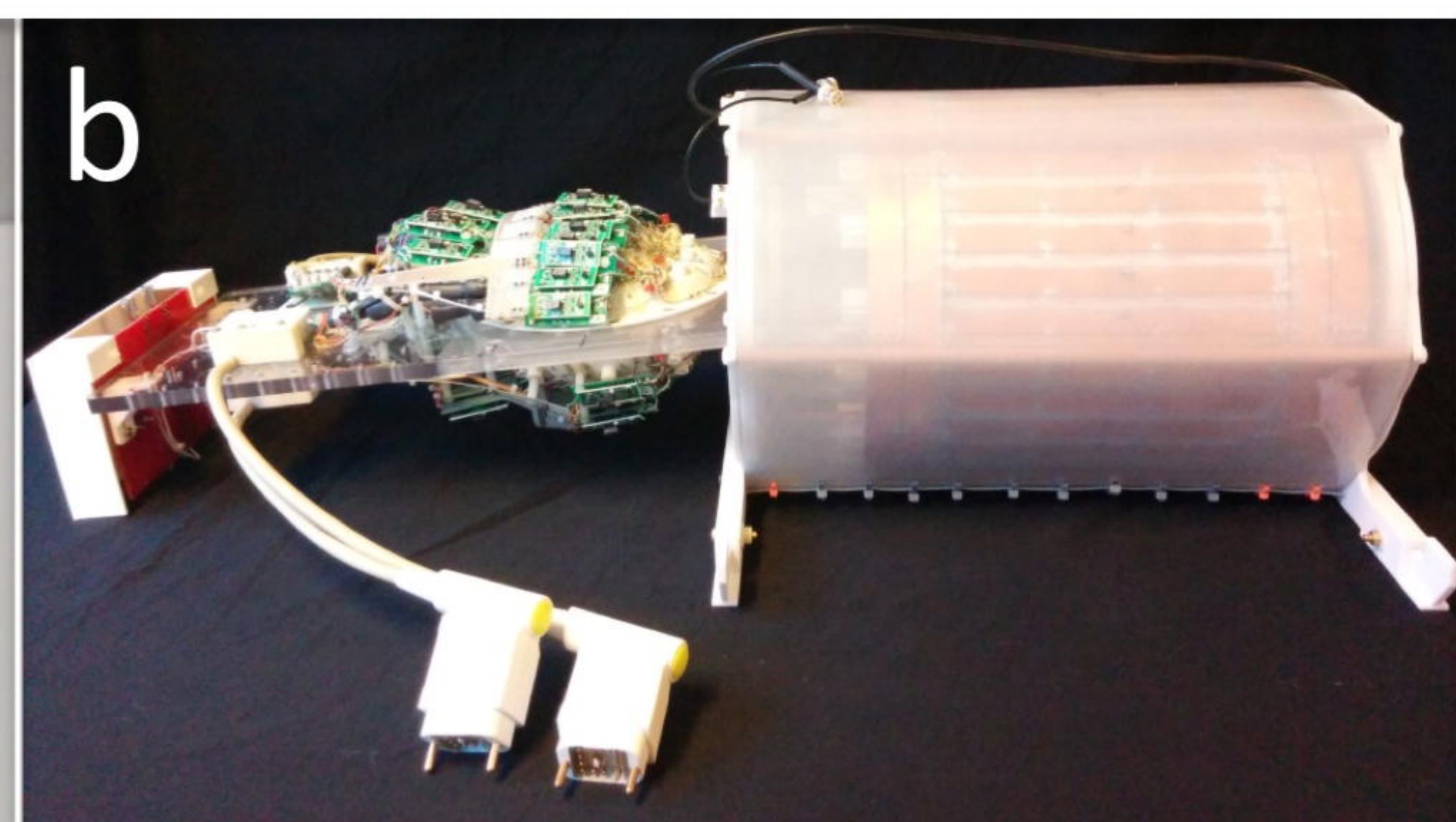




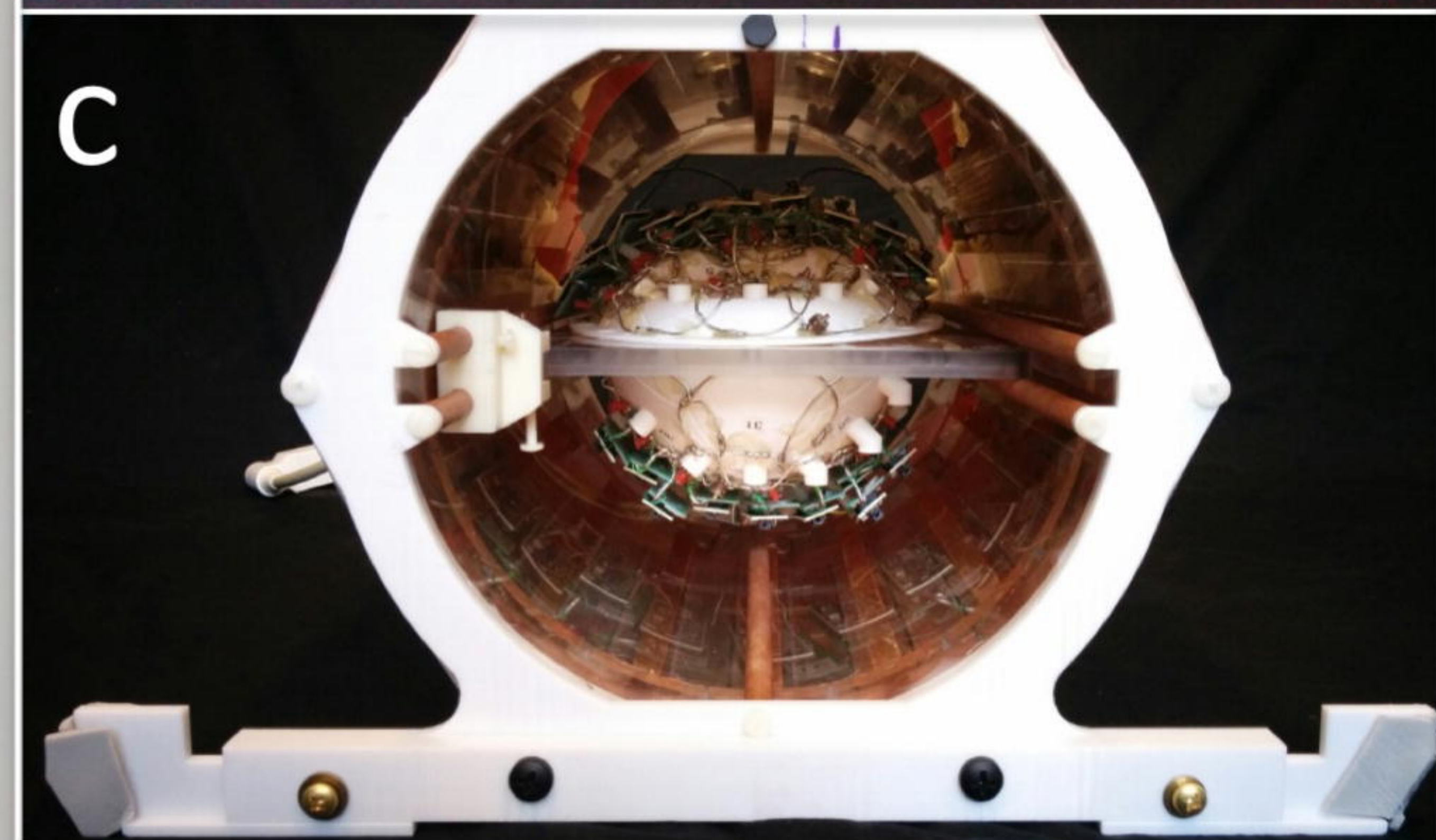
a



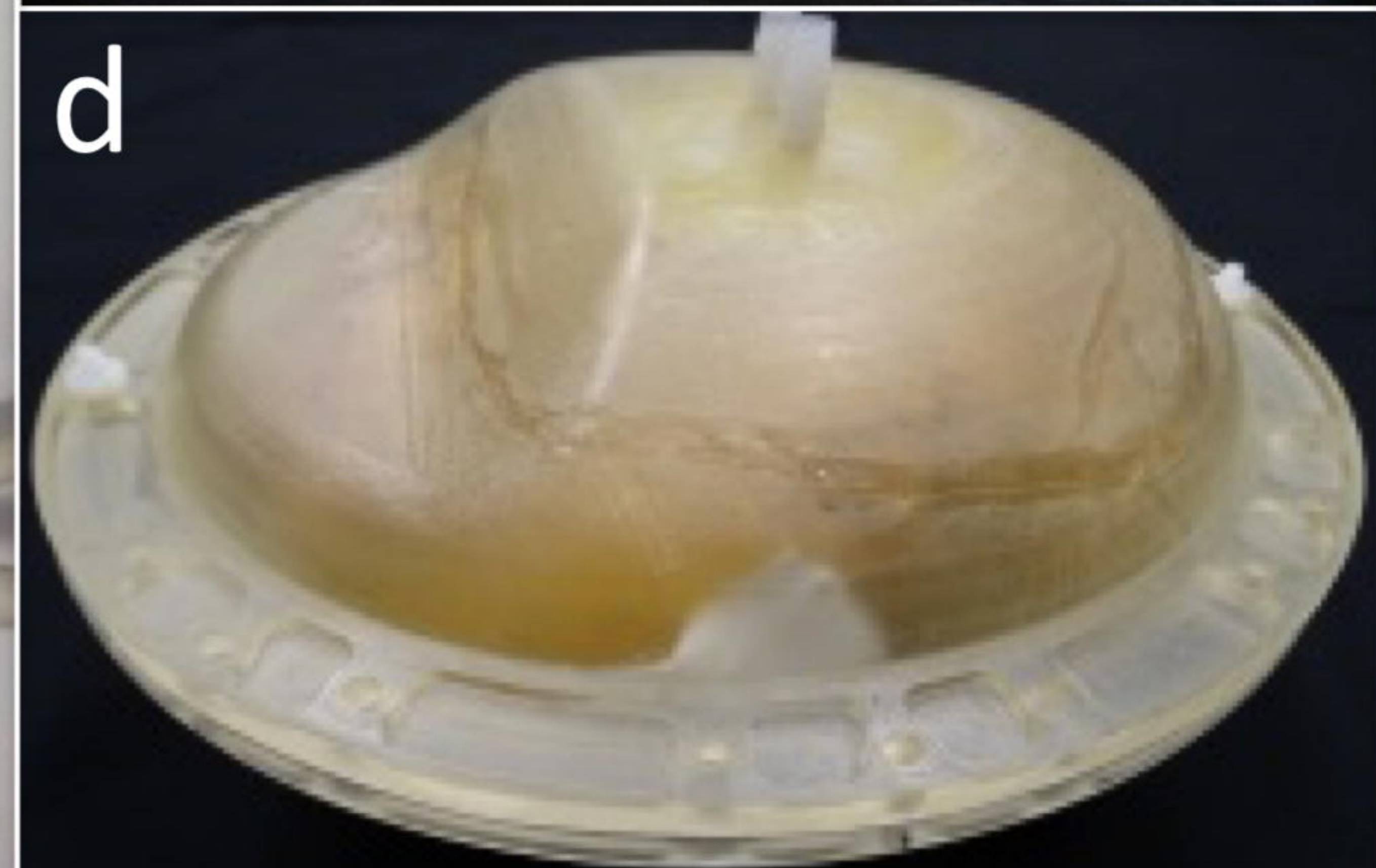
b



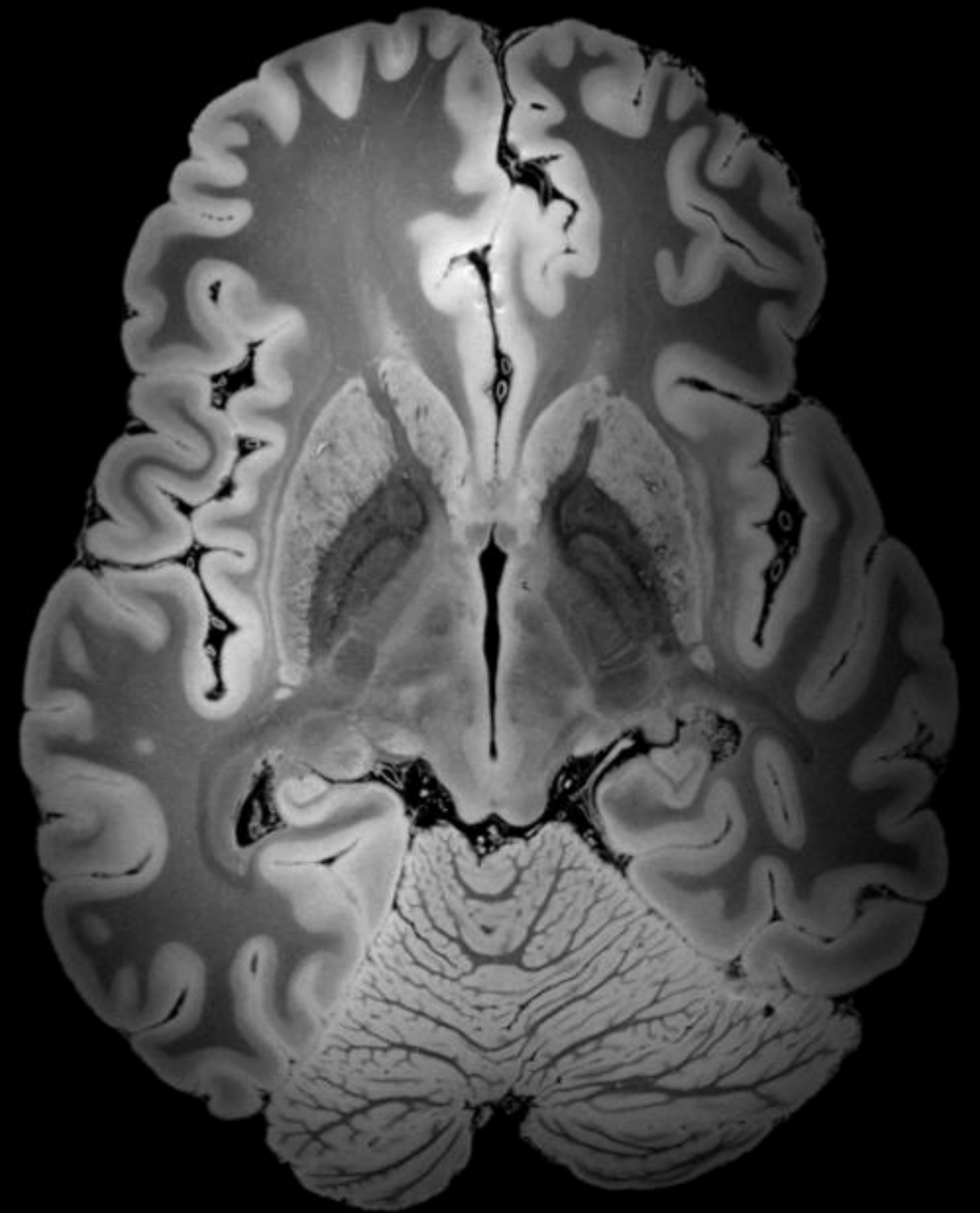
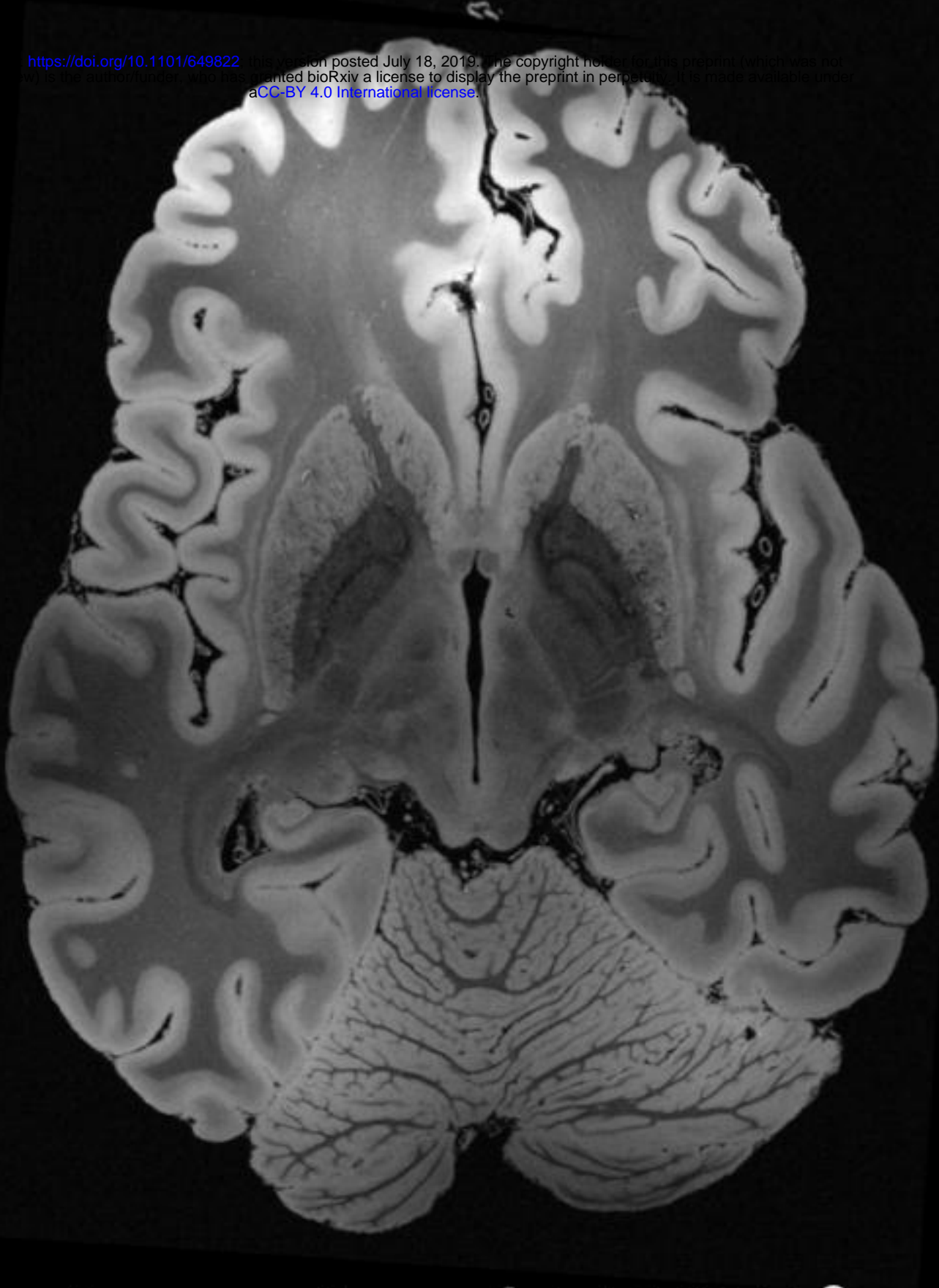
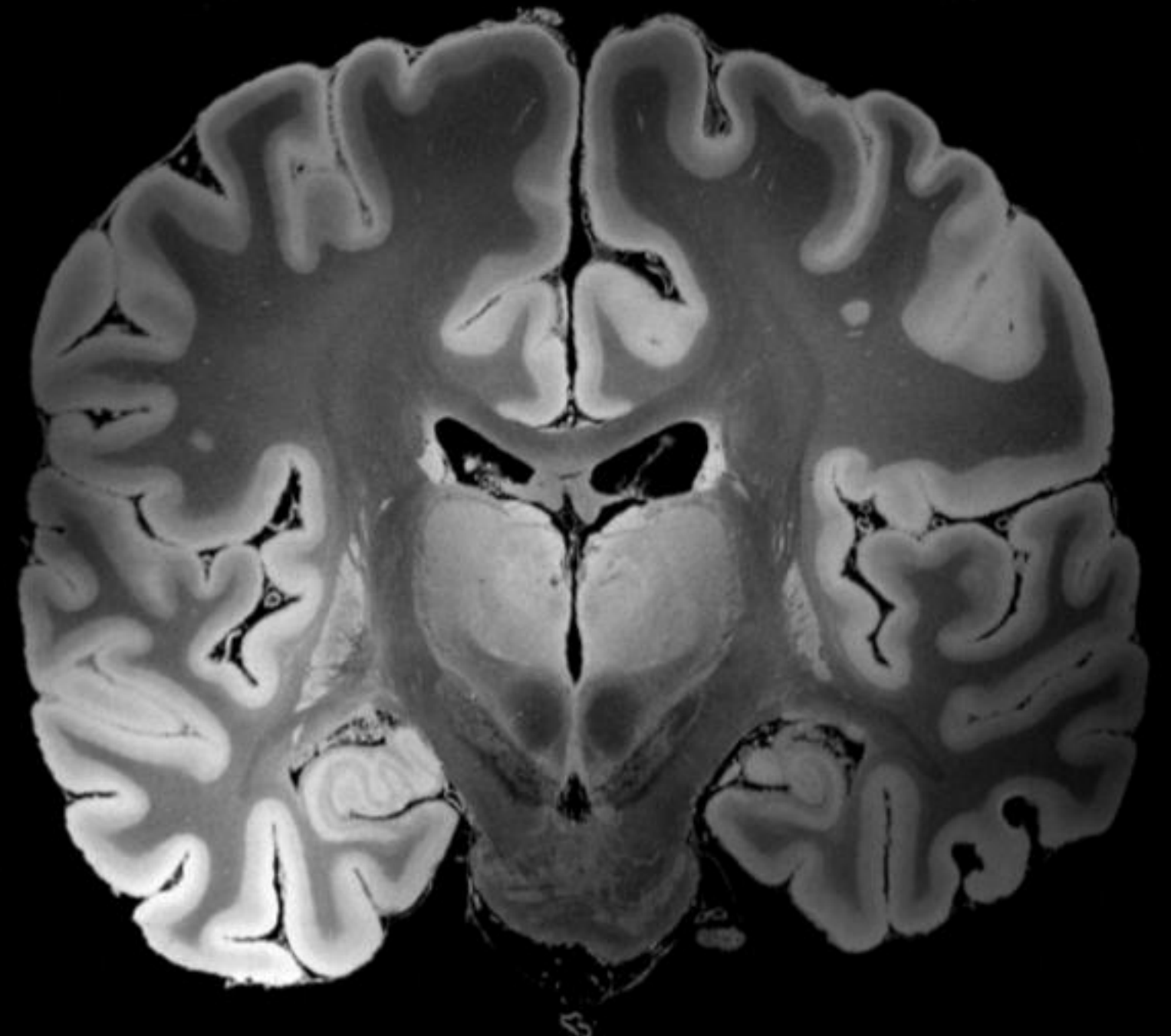
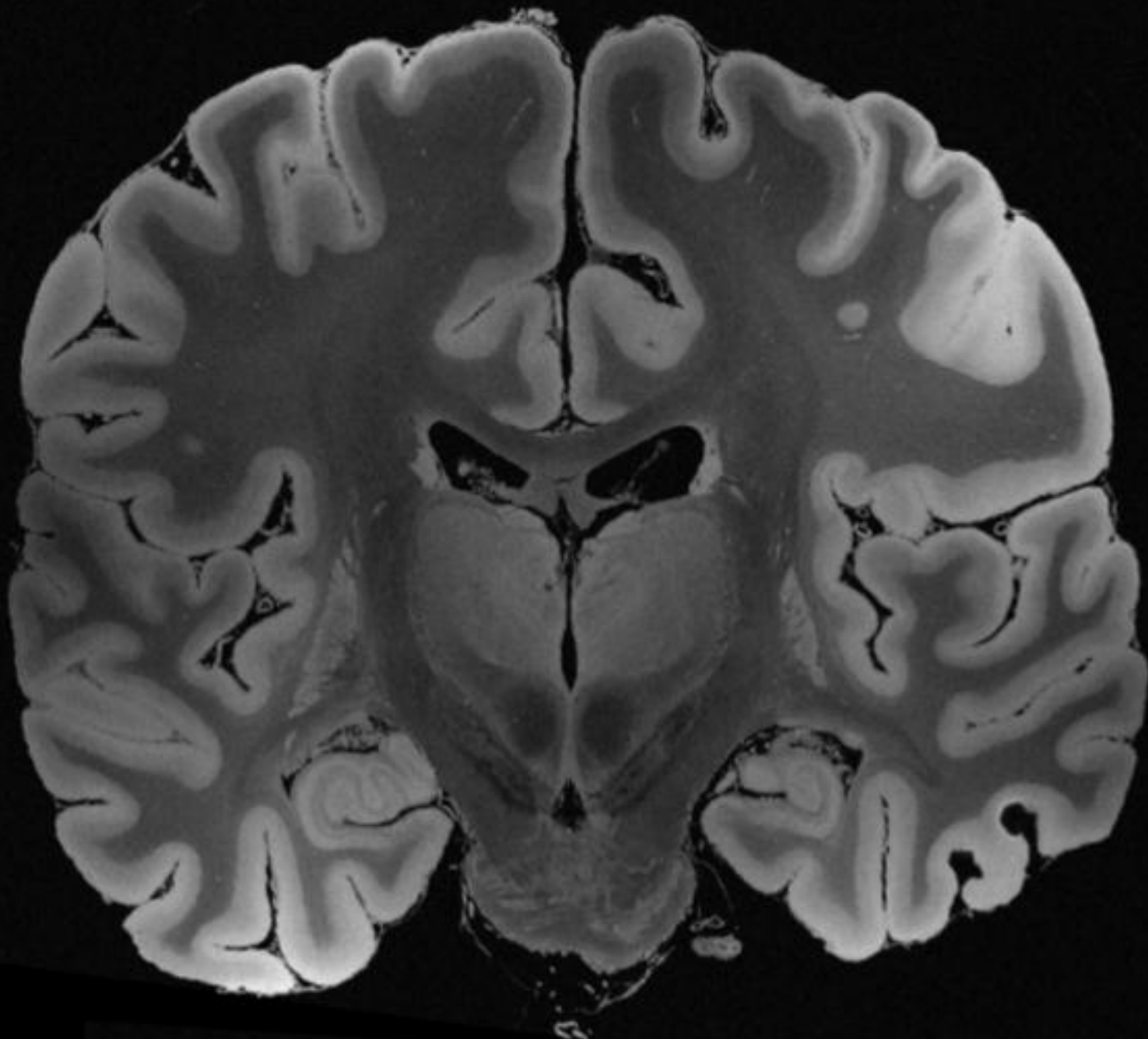
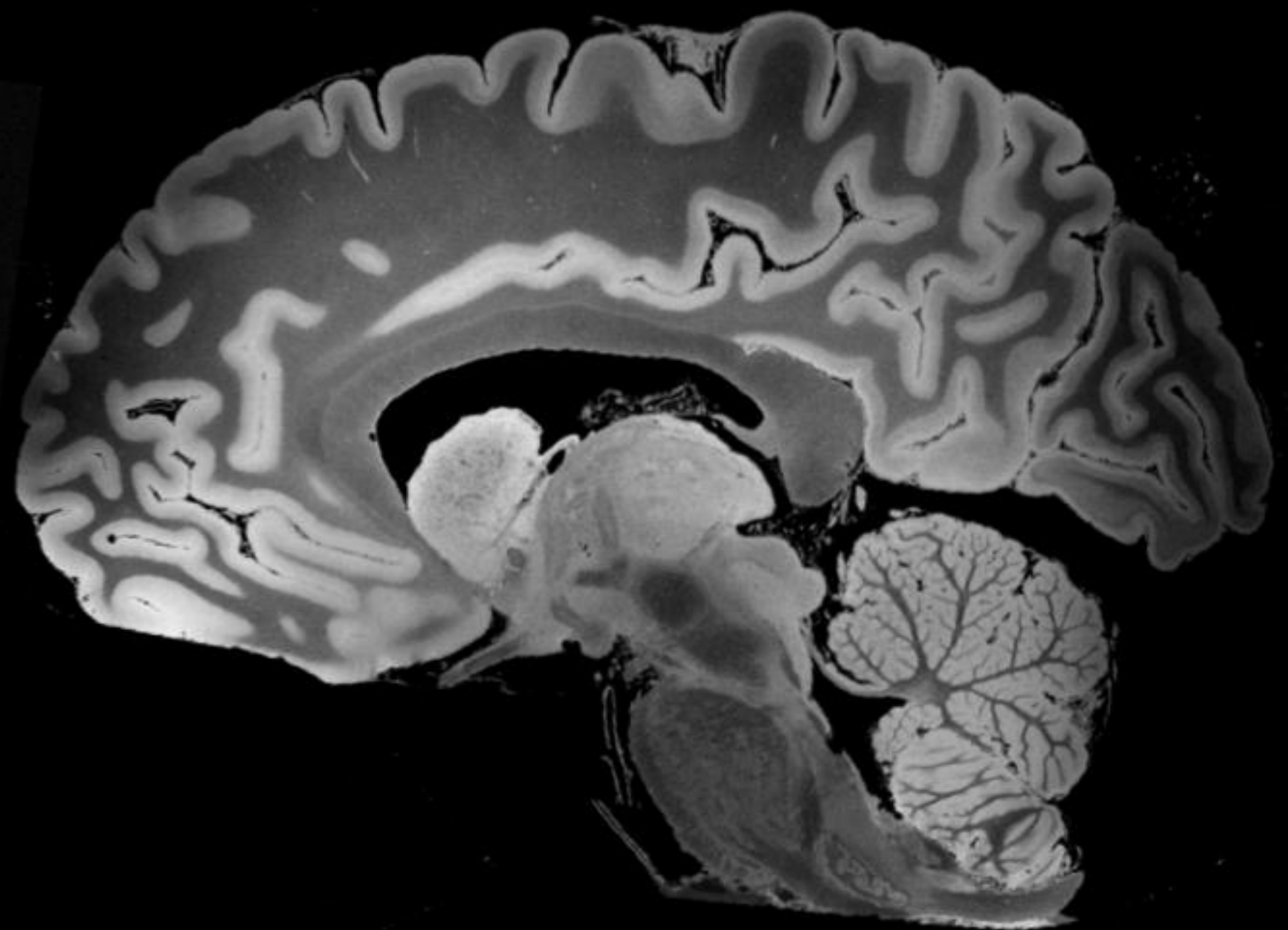
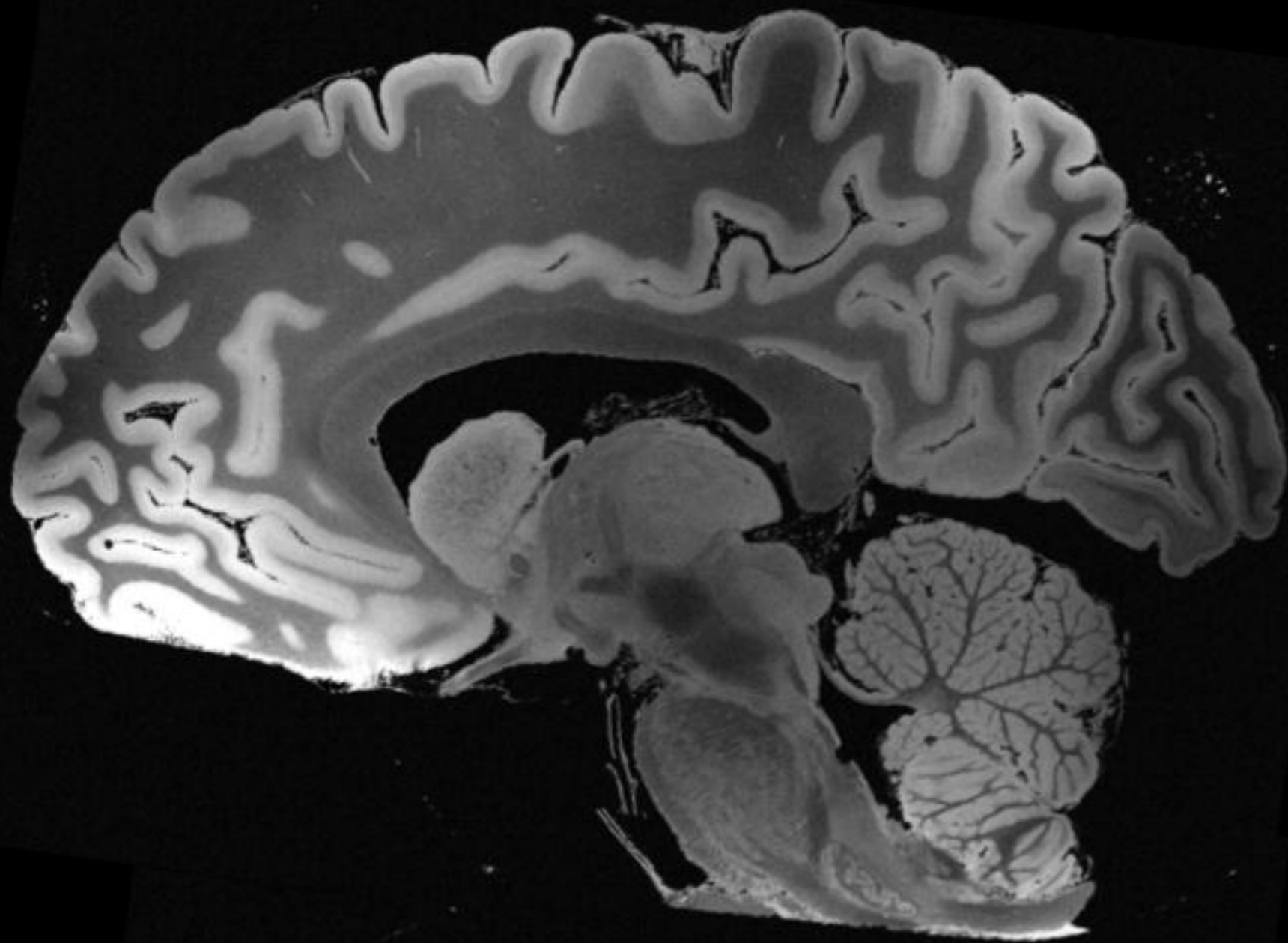
c



d



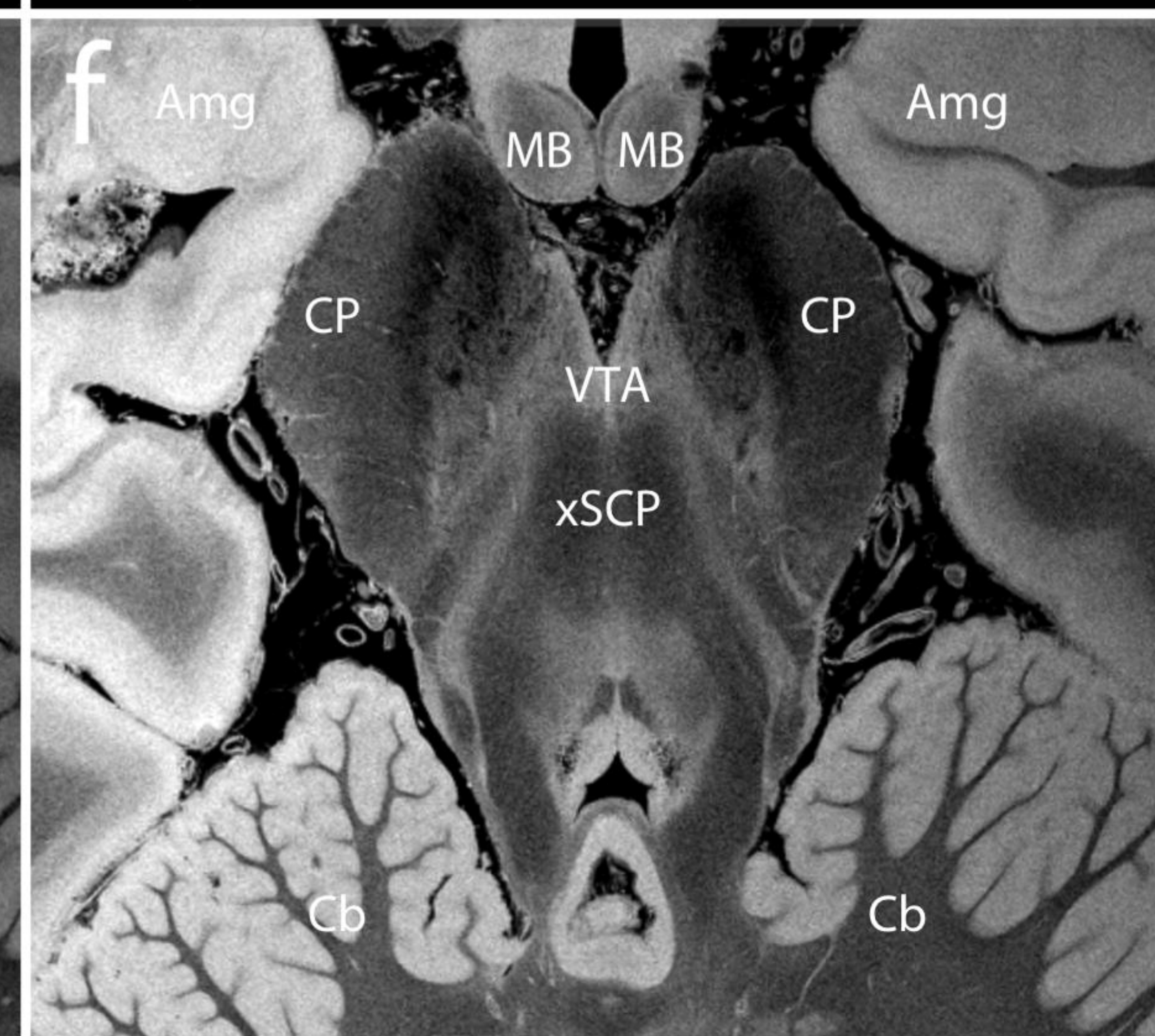
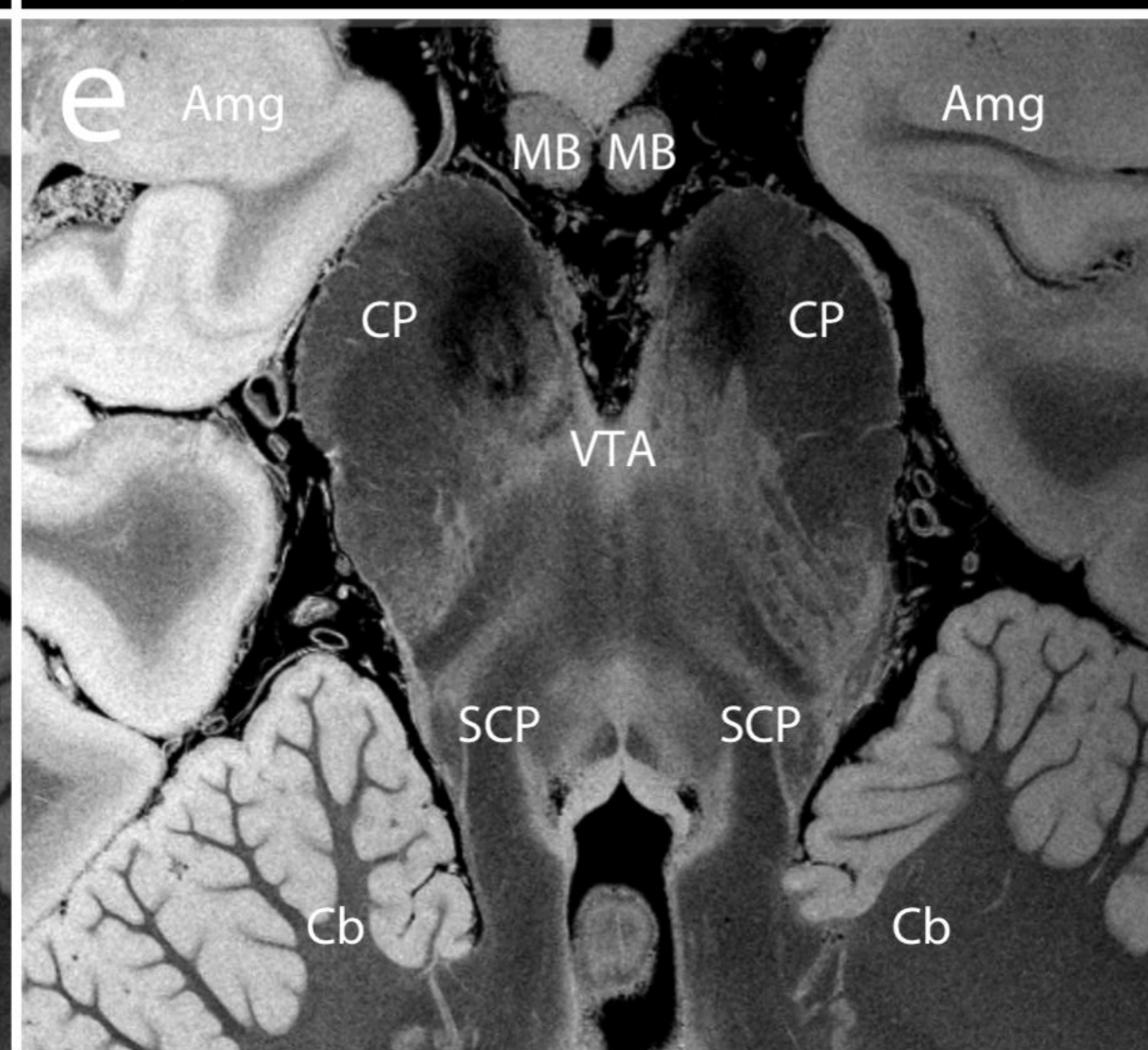
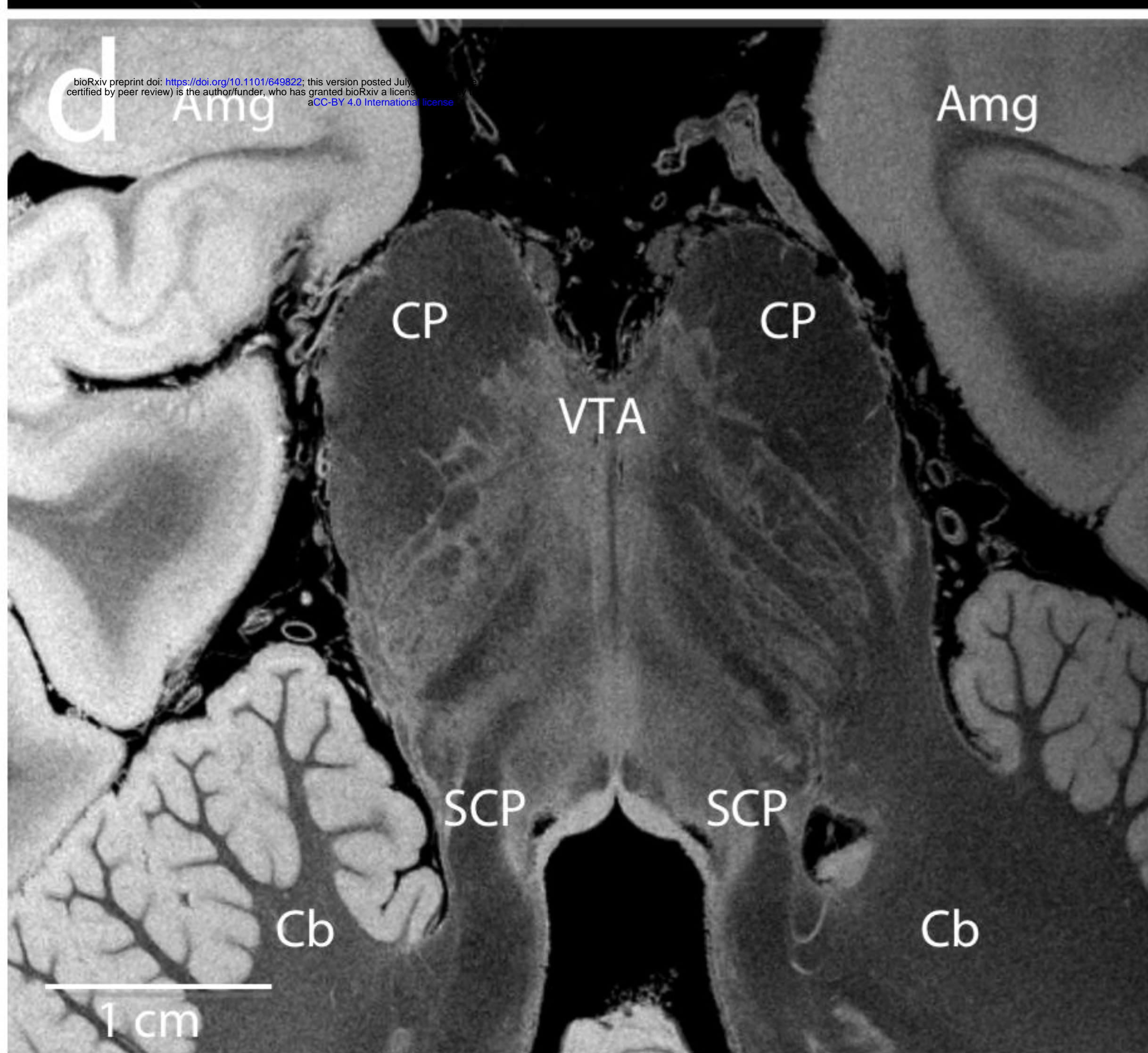
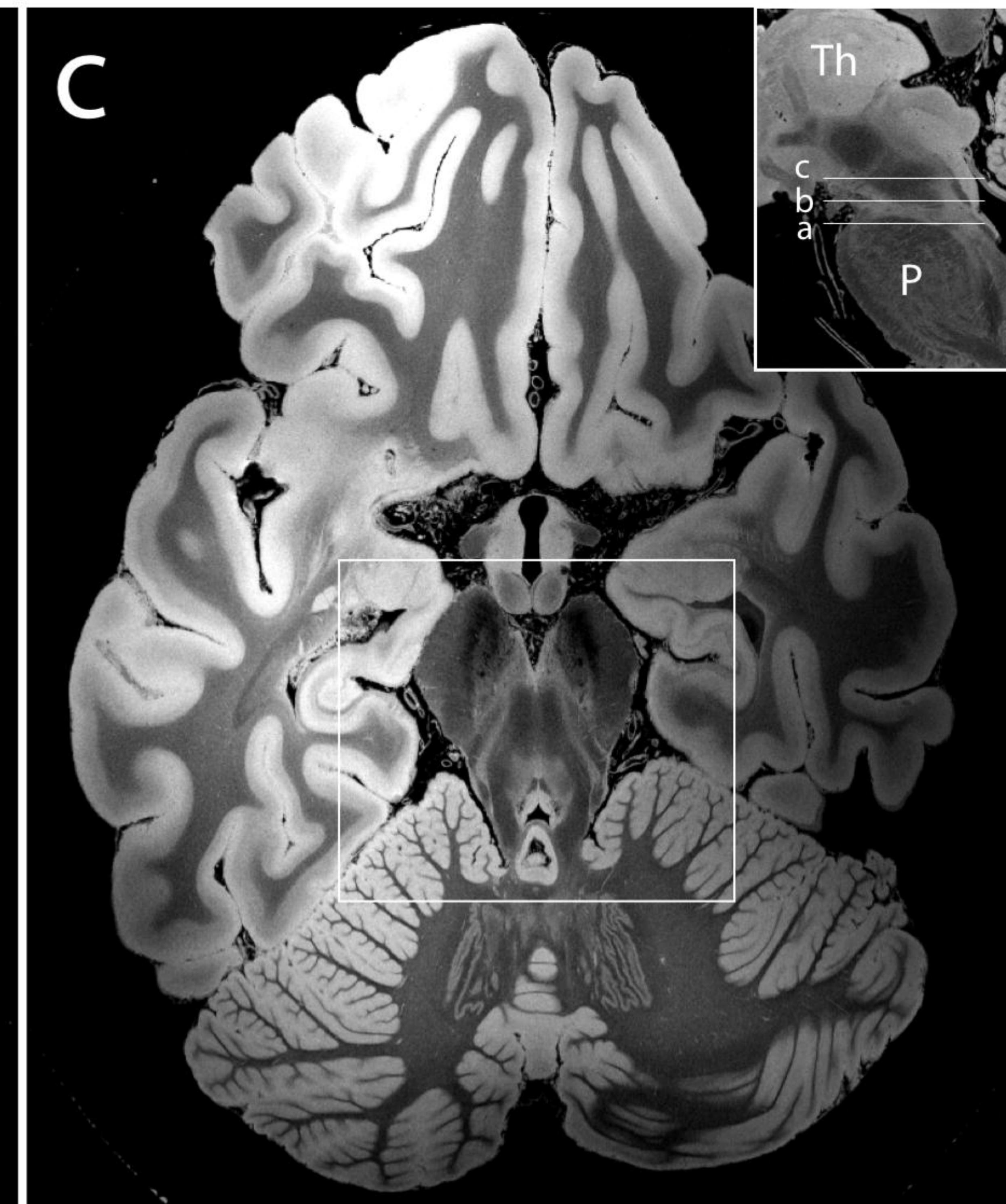




Acquired FA25°  
(~25 hours)

Synthesized FLASH25  
(~100 hours)







7T 31Ch  
*ex vivo* coil

7T 31Ch  
standard coil

3T 64Ch  
head coil

Sagittal

Coronal

Axial

