The combination of optical tissue transparency with immunofluorescence allows the molecular characterization of biological tissues in 3D. However, adult human organs are particularly challenging to become transparent because of the autofluorescence contributions of aged tissues. To meet this challenge, we optimized SHORT (SWITCH—H₂O₂—antigen Retrieval—TDE), a procedure based on standard histological treatments in combination with a refined clearing procedure to clear and label portions of the human brain. 3D histological characterization with multiple molecules is performed on cleared samples with a combination of multi-colors and multi-rounds labeling. By performing fast 3D imaging of the samples with a custom-made inverted light-sheet fluorescence microscope (LSFM), we reveal fine details of intact human brain slabs at subcellular resolution. Overall, we proposed a scalable and versatile technology that in combination with LSFM allows mapping the cellular and molecular architecture of the human brain, paving the way to reconstruct the entire organ.
olumetric reconstruction with a cellular resolution of samples, such as the human brain, is difficult to obtain due to the geometrical, structural, and biological composition of the organ itself. To overcome some of these issues, sections with a thickness less than 100 μm are used to investigate the human cortical cytoarchitecture. However, this strategy lacks three-dimensionality and can decontextualize the section from the surrounding environment, introducing artifacts. Several volumetric imaging technologies like confocal, multiphoton, and light sheet-fluorescence microscopy (LSFM) have been employed for visualizing deeper inside the tissue. To perform 3D reconstructions, these optical techniques require an optimized clearing protocol, which preserves the proteins of interest and removes the components that affect the imaging (generally lipids and chromophores) by producing a refractive index mismatching that is correlated with an increase in light scattering and absorption by chromophores and pigments. By removing such undesirable biological components, two main important features can be obtained: first, the refractive index can be easily matched, allowing efficient light penetration and imaging at depth; second, the absence of biological barriers like lipid bilayers allows performing deep exogenous labeling of the tissue. Although the advances in tissue clearing and labeling techniques coupled with fast optical techniques have enabled access to the sensory–motor cortex, and superior frontal cortex areas (i.e., hippocampus, Broca’s area, motor cortex, precentral cortex, and superior frontal cortex) at subcellular resolution and the neuronal and myelin densities, are factors that can compromise the staining and the clearing efficiency in the human and non-human primate cerebral cortex. Recent applications of optical clearing methods to the human brain provide new details for structure-function relationships in healthy and pathological conditions. However, such methodologies have some drawbacks and limitations. Most of them are compatible with specific samples such as fresh-frozen samples, pediatric tissue, fetal brains, and specimens with controlled post-mortem conditions. Also, the immunostaining is usually performed with small dyes few antibodies and thinner sections. The advent of the tissue-hydrogel engineering methodologies has extended the utility and labeling efficiency in human samples. In particular ELAST converts the tissue into an elastic hydrogel allowing homogenous staining of a 1 cm-thick section with several antibodies. However, the sample preparation requires long processing times and advanced custom-made equipment. Additionally, the recent advent of expansion microscopy and its variants grants super-resolution imaging by expanding the hybrid sample-hydrogel, increasing the sample mesh size and improving the accessibility and density of labeling for proteins even within a dense specimen region. However, these methodologies require strong denaturation processes, which can induce epoxide damaging and alteration in the tissue architecture, making challenging the detection of multiple markers. Small dye based on oligonucleotides (aptamers) or nanobodies can be the best choice for achieving a 3D homogenous staining. However, the production of these probes for different epitopes could be demanding and time-consuming, and the biochemical complexity of such samples may interfere with the staining process, generating a superficial probe accumulation. Moreover, the inter- and intramolecular binding effects generated by formaldehyde and amino groups of proteins during a prolonged paraffin fixation could induce deep epitope changes with a masking effect. Also, the time extension for achieving sufficient transparency can affect the staining efficiency, and several antibodies can fail to work following a tissue transformation protocol in the grey as well as white matter. For instance, CLARITY procedures strongly reduce the staining efficiency for specific epitopes in human formalin-fixed specimens. Due to these problems, large-scale reconstruction in the white and grey matter of long-storage formalin-fixed samples, with different neuronal markers and cell types with molecular details remains an unmet goal in the human brain. Based on the physicochemical properties of human brain samples, we developed a protocol compatible with specimens that show different tissue characteristics and postmortem treatments. Such optimization consists of a simple sequence of treatments that enable to clear and decolorize the sample, as well as to increase the sensitivity of reactions between antibodies and antigens in a large portion of the human brain cortex to allow high-resolution volumetric acquisition with LSFM. In particular, optimal transparency and access of external probes is enhanced by the clearing process, the autofluorescence of the fixed tissue is bleached, and the unmasked effect of antigen retrieval is performed. This method, termed SHORT (SWITCH—H$_2$O$_2$—antigen Retrieval—TDE) employs the tissue ultrastructure preservation of the SWITCH protocol with a series of conventional buffers used in the histological treatments for performing cm-size mesoscopic analysis of 500 μm-thick slices with light-sheet fluorescence microscopy (LSFM). Using our custom-made LSFM, we evaluated the staining efficiency in the white and grey matter, and we assessed the 500-μm thickness as a limit for obtaining a homogenous labeling for different antibody staining (both neuronal, glial, and vascular). This result increases by 5 times the slice thickness treated with the classic SWITCH technique allowing to reduce the cutting artifact and the acquisition times. Additionally, we optimized multiple staining and multiple round procedures to characterize with several markers the same tissue, obtaining comprehensive molecular phenotyping of the sample. Finally, we demonstrated the widespread applicability of the method by coupling SHORT with LSFM to investigate the 3D cytoarchitecture of different human brain areas (i.e., hippocampus, Broca’s area, motor cortex, precentral cortex, and superior frontal cortex) at subcellular resolution and with a fast acquisition rate.

**Results**

**Autofluorescence reduction characterization of cleared tissues.** To evaluate the effect of tissue clearing on human brain tissue we examined the autofluorescence (AF) signal of transformed tissues. To this aim, we selected two different tissue transformation clearing approaches: SWITCH and SHIELD to process human 500-μm thick slices. The family of these transformation tissue methods consists in the control of the chemical reaction via specific buffers, termed “system-wide control of interaction time and kinetics of chemicals (SWITCH)-Off buffer”. In such a solution, the intramolecular and intermolecular cross-link reactions are completely suppressed, to inhibit the fixative molecules depletion (glutaraldehyde for SWITCH or polyglycerol-3-polyglycylcyl for SHIELD) on the surface and diffuse them throughout the tissue. By moving the specimens to a SWITCH-On buffer, the fixative initializes a uniform and global

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fixation/gelation. This process transforms the tissue into a heat- and chemical-resistant hybrid, which allows rapidly clearing at high temperatures and staining with several probes, including antibodies and fluorescent dyes. The advantage of such procedures is their compatibility with post-mortem fixation of the specimens without the need for a perfusion step. The delipidated specimens were then equilibrated with TDE to reach the final transparency, as previously used, and examined by LSFM using four excitation wavelengths at 405, 488, 561, and 638 nm.

The natural AF at 405 and 488 nm in both protocols is likely attributable to the normal presence of the reduced form of nicotinamide adenine dinucleotide (NAD(P)H), the oxidized form of flavins, and vessels in the human cortex. However, as shown in Fig. 1a, the crosslinking agent glutaraldehyde in SWITCH induces a higher AF signal compared to SHIELD at 488 nm. Moreover, in the red-shifted spectrum, lipofuscin pigments show highly intensity signals, approximately at 561 and 638 nm, localized primarily to the neuronal soma. In this spectral range, further AF signal is especially marked in the SWITCH processed tissue, and it can be imputable to the crosslink reaction between lysine residues and glutaraldehyde, which produces a strong yellow and red emission.
To decrease the AF introduced by the tissue transformation treatments, seven approaches, previously reported to reduce the AF of several tissue types and cells were tested: hydrogen peroxide (H$_2$O$_2$), Quadrol, ascorbic acid, hydroquinone, sodium borohydride (NaBH$_4$), copper sulfate (CuSO$_4$), and Sudan Black (SD). These treatments were tested for their easy preparation combined with rapid decolorization procedures. As SHIELD showed low background autofluorescence across the visible spectral range, we used this clearing method as the reference standard for probing the AF decrease in SWITCH processed slices (Fig. 1a, b).

In addition to classic AF-quenching methods, we also investigated the role of antigen retrieval (AR) treatments on transformed tissues. Indeed, although AR is not implicated in the quenching process, we wanted to integrate it in the clearing procedure for its epitope-unmasking effects to increase immunofluorescence compatibility. In agreement with previous works, the alkaline AR solutions (including Tris-EDTA buffer) showed the highest retrieved antigen immunoreactivity against most of the antibodies tested relatively to the AR solutions at pH 6.0 (i.e., sodium citrate buffer).

We found that high-temperature AR for a short time using tris-EDTA buffer (see recipe in Methods), the reducing agent’s ascorbic acid and hydroquinone showed no significant changes in the overall AF spectrum in SWICH processed human cortex slices (Fig. 1a, b). Also, Quadrol showed a red-shifted quenching effect, while AR presented a reduction of the intensity at 638 nm, likely attributable to the reduction of lipopigments in the human cortex. Although low concentration for a short time of oxygen peroxide did not significantly modify the AF signal, 30% H$_2$O$_2$ significantly diminished the tissue AF at 405, 488, 561, and 638 nm excitation light (Fig. 1a). Also, NaBH$_4$, CuSO$_4$, and SB (see Autofluorescence Elimination Reagent in Methods) were the most effective autofluorescence quenchers, reducing the AF signal about twofold compared to SWITCH and SHIELD processed slices (Fig. 1b and Fig. S1). For these reasons, 30% H$_2$O$_2$, NaBH$_4$, and CuSO$_4$ were selected for their capability for diminishing the AF on human cortex tissue and used for the analysis of immunostaining penetration depth discussed below.

**Optimization of SHORT immunostaining.** As a prerequisite step to performing the molecular phenotyping of cleared human brain specimens, we analyzed the diffusion capability of different antibodies to select the most effective ones. After the staining, the penetration of the antibodies deep inside the sample was evaluated by imaging it with LSFM throughout the 500-μm thickness of the slice. We found that the first 200 μm shows a high fluorescence signal compared to the middle and the opposite side. Indeed, several physical processes, such as light scattering and absorption, optical aberration and the tissue properties (like low transparency) can induce signal decay and contrast loss.

To address this issue, we tested immunofluorescence staining on 500-μm thick SWITCH- and SHIELD-processed tissue using different AF elimination treatments, AR, antibodies incubation temperature, and buffers. In particular, we tested several combinations of different AF elimination reagents, unmasking effects of AR (see above), temperature (37 °C and 4 °C) (Fig. 1c, d), buffers (PBST, PBST + BSA, HEPES buffer supplemented with Quadrol and urea; Fig. S2d) and antibody concentrations, to determine the conditions that permitted the best antibody penetration efficiency (Fig. S3a–f). For the analysis we used an anti-NeuN antibody (a neuron-specific nuclear protein) to validate the protocol with a high-density epitopes staining. Results showed that the fluorescence intensity displays a rimmed pattern, marked at 4 °C, with an intense peak at 50 μm depth in SWITCH and SHIELD (Fig. 1c, d). Instead, high-temperature incubation (37 °C) generates an almost homogenous probe distribution in human SWITCH-processed slices treated with H$_2$O$_2$ and AR. These variations upon the original SWITCH protocol clearly demonstrate their effectiveness and motivate the naming of the improved protocol as SHORT. Additionally, SHORT induces a more intense signal and better S/N ratio than the SWITCH classic version (Fig. 1c). As shown in figure S3c, the combined effect of H$_2$O$_2$ and AR enhances the labeling process with respect to the single treatments (either H$_2$O$_2$ or AR). At this point, we concluded that 37 °C was the best temperature condition for probing the staining efficiency. Treatments like NaBH$_4$, CuSO$_4$, which strongly reduced autofluorescence in SWITCH-processed slices (Fig. 1a, b and Fig. S1), induced a low-intensity peak in the first 50-100 μm and an emission signal comparable to the natural autofluorescence along with the whole tissue thickness at 638 nm (Fig. 1e, f). We also evaluated the CuSO$_4$ treatment before and after the immunostaining process, finding no improvement of the signal (Fig. S2e). This is attributable to the deposition of Cu$^{2+}$ ions which increases the scattering process and/or the epitope damaging of these treatments. While optimizing the primary and secondary concentrations of primary anti-NeuN and Alexa Fluor 647 antibodies, we demonstrated that the best dilutions are 1:50 and 1:500, respectively (Fig. S3b). On the other hand, SHIELD-processed slices showed an evident probe accumulation in the first 100 μm at different temperatures (4 °C and 37 °C) (Fig. 1d). Such results were also demonstrated by combining SHIELD with other autofluorescence elimination reagents and unmasking techniques (Fig. S2a, b, d). Finally, both the SHIELD and SWITCH protocol did not show efficient labeling in HEPES buffer, HEPES buffer supplemented with Quadrol and Urea, and PBST supplemented with BSA (Fig. S2b). d). The overall assessment of the tissue-processing procedures resulted in the optimization of SHORT (autofluorescence characteristics in Fig. S4): a combination of the SWITCH technique with H$_2$O$_2$/AR treatments, high-temperature incubation of antibodies, and TDE refractive index matching, to enable homogeneous labelling of human brain samples.

Figure 2a, b show the complete pipeline for cutting, processing, and acquiring 500-μm thick human brain slices. Briefly, SHORT is based on the SWITCH chemical principles to generate meshgel polymers from inside the tissue, to provide structural and biomolecular preservation (Fig. 2). After the glutaraldehyde fixation, lipids (responsible for preventing the staining to deep structures and inducing the scattering process during imaging) can then be removed without losing native tissue components using a clearing solution (200 mM SDS, 10 mM lithium hydroxide, 40 mM boric acid) at 55 °C (Fig. 2a). Next, SHORT requires the incubation in 30% vol/vol in H$_2$O$_2$ for 1 h at RT, and Tris-EDTA for 10 min at 95 °C (Fig. 2b). The resulting treatments show an efficient staining for different biological features by incubating the antibodies at 37 °C. After staining, the tissue-gel hybrid is immersed in a refractive index matching (TDE/PBS) and enclosed in a sandwich holder to stabilize and preserve the specimens (Fig. 2a). Also, a multi-round of staining, stripping, and acquisition on SHORT-process specimens can be performed for proteome investigation (Fig. 2a, b) as presented in the following paragraph.
like parvalbumin (PV) and calretinin (CR), indeed we observed that the S/B (signal/background) measured overall the 500 μm depth of the sample, is sufficient to detect the labelled neurons (Fig. 3a). We verified the compatibility of SHORT with different nuclear stains, including Sytox Green, DAPI, and Propidium Iodide (Fig. S6; Fig. 4k). Also, to verify the compatibility of SHORT with lipophilic dyes, we tested the DiD staining and we visualized the myelinated fiber pathways in a 100-μm human superior frontal cortex slice (Fig. S7). Then, we optimized different strategies to co-stain the tissue up to three different markers and perform multiple rounds of labelling. To avoid cross-recognition between the antibodies we selected different species both for the primary and the secondary antibodies. In particular, we developed costaining protocols for: NeuN and glutamic acid decarboxylase (GAD67) (Fig. 4f and Fig. S8); NeuN and CR; NeuN and somatostatin (SST) (Fig. 3b); vasoactive intestinal peptide (VIP) with SST and PV; NeuN with CR and SST; glial fibrillary acidic protein (GFAP) with SST and vimentin (VIM) (Fig. 3c). All the antibodies and dyes used in the study are summarized in supplementary Table S1.

To access the phenotyping analysis potential of SHORT, we generated multi-round labelling of the tissue by removing the antibodies and restained the sample with other markers. To remove the staining completely, the antibodies were stripped off from the tissue by soaking the stained sample in the clearing buffer at 80 °C for 4 h. We observed that the stripping process reduces the autofluorescence signal of lipopigments (i.e., lipofuscin) by about 2 times (Fig. S9). We were able to perform three consecutive rounds of staining in 500 μm-thick slices of the human brain cortex, confirming effective antibody removal, preservation of neuronal morphology as well as re-staining capability. In particular, we first costained the tissue with anti-VIP, anti-SST, and anti-PV antibodies. We then stripped out the antibodies and we restained the tissue and repeated the imaging process twice, labelling for the second and third round, respectively: SST, CR, NeuN and SST, GFAP, VIM, using SST as a fiducial marker (Fig. 3c). We achieved multiplexed profiling of 7 different targets in the same piece of human superior frontal cortex by multispectral imaging of 3 antigens in each round. To evaluate if the stripping process affects the staining of the tissue, we quantified the signal-to-noise ratio (SNR) of SST immunostaining at rounds 1, 2 and 3, (N = 30 SST-immunoreactive neurons for each round), finding a loss of 30% protein signal during the first round (Fig. 3d). Our data shows that the SNR of SST immunostaining in rounds 1, 2, and 3 decreases from round 1 to round 2. This suggests that the amount of protein lost in the stripping process decreases after every round, but antigens are still preserved and they can be detected using SHORT. To verify the absence of distortion associated with the high-temperature incubation during the stripping process at cellular level, we quantified the structural similarity index (SSIM index54) of 3 different SST-immunoreactive neurons and their dendritic branches tortuosity (n = 6) before and after stripping. High SSIM index value (0.9 ± 0.06) (Fig. S10a) was found and no significant structural change in the branching patterns (Fig. S10b) was observed.

### Volumetric 3D reconstruction of SHORT cleared samples with light-sheet fluorescence microscopy

The investigation of the 3D molecular tissue phenotyping of large human brain samples was performed by coupling SHORT with a custom-made LSFM that enables imaging of large samples up to 30 × 30 cm² thanks to a large sample accommodation chamber (see Methods for further details). Tissue blocks of the hippocampus, precentral gyrus, prefrontal cortex, motor cortex, and Broca’s area (Table S2) of approximately 3 cm³ in size were cut with a vibratome (Comprestome VF-900-0Z) into slices up to 500 μm of thickness and treated with SHORT for tissue clearing and labelling. Samples were mounted in a custom-made sealed sample holder (termed “sandwich”, Fig. S11) suitable for both imaging and long-term storage. Indeed, we observed a preservation of fluorescence signals of SHORT-processed slices stored in the sandwich for 1 year. We compared the signal obtained from a first acquisition of a
Round 2

Round 1

Round 3

employing an isotropic resolution of 3.3 × 3.3 × 3.3 µm³, enables using the high-throughput capability of our LSFM which, reconstruction of human cortical cytoarchitecture was performed routine acquisition and long storage of labeled specimens. The 3D

Fig. 3 Multiplexed staining and antigen preservation of SHORT-processed tissue. a Labelling efficiency of two neuron markers, PV and CR, acquired by LSFM. Mean and SD were calculated by taking the signal of 8 different neurons and background in three different regions: 0–240 µm corresponding to the maximal light exposure (0 µm), 240–470 µm corresponding to the middle of the slices (350 µm), and 490–700 µm which was opposite to the excitation side (700 µm). Abbreviations: S, signal; B, background. High-resolution images, scale bar – 100 µm. b Different examples of optimized co-staining in SHORT processed slices. Maximum intensity projection (MIP) of 20 slices. Downscaled images, scale bar – 100 µm. c Three distinct rounds of immunostaining for probing neurons (NeuN), inhibitory markers (Vip, SST, PV, CR) vasculature (VIM) and glial cells (GFAP). MIP of 70 slices. Downscaled images neuron, scale bar – 100 µm. d SNR was quantified for SST immunolabeling in round 1, round 2, and round 3 (n = 30 neurons). MIP of 70 slices. Downscaled images, scale bar – 100 µm.

Broca Area labeled with CR, SST, and NeuN, with the one acquired after one year, detecting a reduction of 51% for Alexa Fluor 488; 24% for Alexa Fluor 568; and 14% for Alexa Fluor 647 (Fig. S12) (no photobleaching process due to re-acquisition was detected). In addition to RI matching, TDE also leads to tissue dehydration, which reduces slightly the specimen volume (Fig. S13) and preserves the tissue, a highly beneficial effect for routine acquisition and long storage of labeled specimens. The 3D reconstruction of human cortical cytoarchitecture was performed using the high-throughput capability of our LSFM which, employing an isotropic resolution of 3.3 × 3.3 × 3.3 µm³, enables to resolve single neuronal subtypes and soma morphology features of their individual subpopulations (Fig. 4b–f). The clearing process was highly effective in the gray and white matter (Fig. 4a) and the antibody labeling was homogenous along with the entire depth of the samples (Fig. 4f). Although we obtained a good probe penetration depth at different excitation wavelengths (488, 561, and 638 nm) (Fig. S12 and S14; Supplementary Movie 4), we encountered difficulties in achieving a similar degree of staining homogeneity and light penetration depth at 405 nm with the nuclear staining DAPI. This effect is known and termed the inner filtering effect in the literature and will require further work to be overcome.

We first used NeuN combined with GAD67 to perform multiple labelling throughout the 1.5 × 1.5 × 0.5 cm³ hippocampus human brain slices (Fig. 4b–f). Figure 4d, e show the feasibility of such methodology by imaging hippocampal neurons of the CA1 as well as the human dentate gyrus SHORT-processed slices. Furthermore, the multiple staining approach was also performed on human brain slices from Broca’s area (Fig. 4g–j), and prefrontal cortex (Fig. 4k–n), using three different antibodies against different neuron types, including SST, CR, as well as NeuN. It is known that CR- and SST-immunoreactive inter-neurons were distributed in all cortical layers but, there is little information in the literature on the distribution of SST-expressing neurons in white matter. Our measurements show that white matter regions are rich in SST-expressing neurons (Fig. 4o) in the prefrontal cortex.

In addition, the multi-round characterization enabled by SHORT allowed us to obtain homogeneous labeling of seven different markers, including all neurons (NeuN), inhibitory neuron markers (SST, CR, PV, VIP), glial cells (GFAP), and vasculature (vimentin, VIM), as demonstrated by the successful imaging of a slide from the inferior frontal pole (Fig. 5a, b). Thanks to the specific mounting in the sandwich holder (Fig. S11), the tissue orientation could be maintained through the acquisitions, allowing a good coregistration of the different acquisitions and facilitating future quantitative interrogation of the data.

Discussion

Although several brain tissue-clearing and labeling methods are available, few of them were originally optimized on the human cortex. Indeed, prolonged formalin fixation and variable post-mortem conditions—such as pH variation and tissue oxygenation—could affect the quality of immunostaining due to the epitope masking effect and antigen damage. Using as a starting point the SWITCH/TDE method and coupling it with conventional buffers
used in classical immunocytochemistry, we developed and optimized SHORT, an efficient clearing and labelling method for 3D human cortex molecular probing with LSFM. In general, SWITCH allows preserving the endogenous biomolecules, while TDE performs sample clearing. However, we encountered several hurdles for performing volumetric reconstruction of the human cortex. The first challenge is the long fixation time, which could cause a variety of changes in the 3D structure of proteins, decreasing the possibility to detect specific epitopes for phenotyping and connectomes analyses.\textsuperscript{37,38} Also, the presence of lipopigments—such as lipofuscin and neuromelanin—\textsuperscript{59–61} and blood, have remained an insuperable hurdle during the acquisition process, causing spurious signals that make cell identification and counting more complex. In addition, aldehyde fixation using formaldehyde and glutaraldehyde leads to a wide emission spectrum resulting in high background fluorescence that limits the achievable SNR. Indeed, the generation of the double bonds between the crosslinker and proteins increases the autofluorescence, making the sample unsuitable for immunofluorescence assay. To overcome this limitation, SHORT was engineered to be capable of decolorizing such autofluorescence, blood-infused tissue and lipopigments, in combination with epitope restoration. This strategy allowed to improve the image quality, the antibody’s penetration, and the SNR of reconstructed large portions of human brain slices. Also, we investigated several autofluorescence elimination reagents commonly used in conventional immunocytochemistry. Some of them, such as SB\textsuperscript{45,46}, NaBH\textsubscript{4}\textsuperscript{3,43,44}, and CuSO\textsubscript{4}\textsuperscript{43,45,46}, significantly reduced the autofluorescence signal, but they were not compatible with our method, because they increase the scattering process during the acquisition by forming stable complexes (such as Cu\textsuperscript{2+} and lipofuscin), or causing epitope damage. For these reasons, the best option for tissue structural preservation and autofluorescence quenching capability was provided by peroxyde hydrogen,
coupled with the unmasking effect of antigen retrieval, a combination that suggested naming our method SHORT (SWITCH—H₂O₂—antigen Retrieval—TDE). We established that our approach allows staining 500-µm-thick human slices with different ages and fixation times and performing volumetric acquisition using LSFM. Although LSFM lacks axial resolution power, it is the best optical imaging technique for probing multiple markers in large specimens. Indeed, LSFM offers fast acquisition time and low photobleaching with respect to other conventional optical techniques, such as confocal and two-photon fluorescence microscopy. The probe penetration efficiency with antibodies against a variety of antigens, including NeuN, SST, and CR, which could be masked following SDS treatment, was investigated by LSFM. Although SHIELD was fully applied to clear formalin-fixed 2-mm-thick coronal blocks of the human brain (9 × 5.5 × 0.2 cm³) and label with different markers, our replication of this approach showed a non-homogeneous labelling for high density-epitopes. This is likely attributable to the dense epoxy meshgel and antibody choice, which can induce a probe accumulation in the first 100 µm of the slice. With SHORT, volumetric blocks of the human brain can be reconstructed by cutting the area of interest in slices of 500 µm of thickness. In this paper, we showed the reconstruction of 500 µm-thick slices of various areas of the brain demonstrating the versatility of the method. In addition, SHORT was applied in Costantini et al. demonstrating how this approach can be used to reconstruct a portion of Broca’s area of 1.5 × 1.3 × 0.8 cm³ cutting it in only 16 slices and allowing the cell census of this volumetric portion of the cortex. The entire dataset is available on the DANDI archive (https://gui.dandiarchive.org/#/dandiset/000026).

Exploiting the structural and epitope preservation of such methods, we then attempted and successfully performed multi-rounds of immunostaining and stripping in the SHORT-processed human cortex. Indeed, optical microscopy is limited by the visible light bandwidth and the host of animal species in the classic immunocytochemistry, which makes the detection of more than four different markers challenging without resorting to complex multiplexing techniques that require computational spectral disentangling. The recent advantages of the multiplexed tissue strategies, including MAP, SWITCH, SHIELD, and miriEx have allowed bypassing this limit. These works demonstrated that specific antigens can be repeatedly labeled without loss of antigenicity in 100-µm thick human and mouse brain samples. In agreement with these results, we optimized the possibility to perform multi-rounds in SHORT-processed 500-µm-thick human brain slice, by labeling neuronal (VIP, SST, NeuN, PV), glial (GFAP), and vasculature markers (VIM) in three distinct rounds of immunostaining and stripping. The simultaneous acquisition of three different markers for each round makes easier the co-registration of different channels in a large portion of the human cortex. We showed robust preservation of epitopes following clearing, antigen retrieval, and stripping cycles in a processed large human portion.

We also performed staining using DiD on 100 µm-thick samples of superior frontal cortex, demonstrating the compatibility of SHORT with lipophilic tracing for the rapid visualization of human cytoarchitecture using other methods than immunohistochemistry. In future, the advantages of SHORT may allow combining molecular details obtained with antibody staining and cellular connectivity tranced with lipophilic dyes from diverse cell types within the same tissue.

Although SHORT allows simultaneous detection of multiple markers and multi-round cycles without compromising the tissue integrity, such protocol still presents potential limitations. In particular, the decreased penetration efficiency at 405 nm for nuclear stains like DAPI and the substantial attenuation of the excitation beam in such samples, termed the inner filter effect, should also be improved in the future. To facilitate sample mounting and obtaining human slices with a flat surface we developed a sample “sandwich” holder that also permits long-term storage of the specimens. Although our pipeline allows modulating the XY dimension of the samples using appropriate sandwiches that are size-adjustable, we are still developing specific software for aligning and stitching non-rigidly subsequent the human cortex slices imaged with the LSFM to obtain consolidated 3D maps and reconstructions of the sample’s volume.

In conclusion, we presented a new method, named SHORT, able to preserve the tissue integrity with epitope restoring in centimeter-sized human brain specimens for molecular and cellular phenotyping. Such methodology allowed homogenous tissue

Fig. 5 SHORT allows multiple rounds of staining, imaging, and stripping of large portions of human brain slices. a Seven different antibodies were used across 3 sequential rounds of immunostaining, with SST employed as fiducial markers in each round. Round 1: PV—SST—VIP; round 2: CR—SST—NeuN; round 3: VIM—SST—GFAP. Scale bar = 1 mm. b The merged image shows a particular of the white and grey matters labeled with VIP, SST, PV, CR, NeuN, GFP, VIM. MIP of 70 slices. Scale bar = 100 µm.
labeled of different markers and clearing of human samples. Also, we implemented a new pipeline compatible for long-term tissue storage of such precious samples. Thus, in combination with fast light-sheet microscopy systems, SHORT could be a key technology to map large human brain regions, up to the whole organ, accelerating the understanding of its structural and cellular characteristics, as well as of physiological and pathological conditions underlying such a complex system.

Methods

Samples. Brain tissue specimens were collected from 3 different human donors with no known neuropsychiatric disorders that were obtained from the body donation program of the Université de Tours and from the Massachusetts General Hospital (MGH). The tissue donors provided their informed and written consent to the donation of their body for research purposes. We used 5 different cortical regions from these donors: the precentral cortex from donor 1 was used for autofluorescence analysis, the NeuN depth analysis, and stripping efficiency validation; hippocampus from donor 1, prefrontal cortex from donor 2, motor cortex, and Broca’s area from donor 3 were used for antibody validation, stripping process, and volumetric acquisitions (Table S2). All cases were neurotypical and had no neuropathological diagnosis of neurodegenerative disorder. Tissue sections of 450 ± 50 μm thickness were obtained using a custom-made vibratome (Compresstome VF-900-0Z). The slices were stored at 4 °C in PBS 0.02% NaN3.

SHORT protocol for human brain slices. An adapted version of the SWITCH/TDE protocol from Costantini et al. was used to clear large portions of the human brain samples. The specimens were incubated in a Switch-Off solution, consisting of 50% PBS titrated to pH 3 using HCl, 25% 0.1 M HCl, 25% 0.1 M potassium hydrogen phthalate (KHP) and 4% glutaraldehyde. After 24 h, the solution was replaced with the SWITCH-On solution, containing PBS pH 7.4 with 1% glutaraldehyde. After 3 washes for 2 h each in PBS at RT, the specimens were inactivated by overnight incubation in a solution consisting of 4% glycine and 4% acetic acid at 37 °C. Following inactivation, the samples were extensively washed in PBS and then incubated in the clearing solution containing 200 mM SDS, 10 mM lithium hydroxide, 40 mM boric acid for 2-4 d at 55 °C depending on the sample size. After the clearing process, the samples were washed 3 times in PBS + 0.1% Triton X-100 (PBST) at 37 °C for 24 h. Next, the slices were treated with 30% H2O2 for 1 h and, after 3 washes for 10 min each, antigens were retrieved with the preheated tris-EDTA buffer (10 mM Tris base (v/v), 1 mM EDTA solution (v/v), 0.05% Tween 20 (v/v), pH 9) for 10 min at 95 °C in a volume of 20 ml. This step allows increasing the sensitivity of reaction of antibodies directed to specific targets. For reaching a good sensitivity, the cleared samples must remain in this solution for about 5-24 h. After cooling down to RT, the specimens were washed 3 times in DI water for 5 min each and then equilibrated with PBS for 1 h.

SHIELD/TDE protocol for human brain slices. The reagents for SHIELD were provided by LifeCanv. Briefly, the tissue slices were incubated for 24 h at 4 °C with shaking for 1 day in the SHIELD-Off solution (10 ml SHIELD-Epoxy Solution, 5 ml SHIELD-Buffer Solution, 5 ml DI water). Next, the samples were incubated at RT with shaking for 1 day using SHIELD-On Buffer and SHIELD-Epoxy Solution mixed in a ratio of 1:1 (final volume of 40 ml). Both SHIELD-Off and SHIELD-On were freshly prepared and not used after two months from the opening. Finally, the samples were cleared for 3 days at 55 °C using 200 mM NaCl, 10 mM lithium hydroxide, 40 mM boric acid and washed 3 times in PBST at 37 °C for 24 h.

Alternative decolorization treatments (autofluorescence analysis). After clearing, small portions of the prefrontal cortex (99-year-old female, 6 months in formalin, donor 1, Table S2) were used for investigating the optimal autofluorescence masking treatments. Such treatments were probed on SWITCH/SHIELD treated tissue slices (N = 4 independent experiments for each autofluorescence masking treatment were performed in the prefrontal cortex, donor 1). To promote and enhance their effects, some of these reagents were combined together.

- Oxygen peroxide: the tissue slices were soaked in 0.6%, 15%, and 30% (v/v) H2O2 diluted in DI water for 1 h at RT. Sections were then washed with PBS 3 times, for 10 min each.
- Sodium borohydride: the tissue slices were immersed in 0.1%, 1% and 2% (w/v) NaBH4 in DI water for 1 h at RT. Next, the samples were washed with PBS 3 times, for 10 min each.
- Copper (II) Sulfate (CuSO4): this reagent was applied before and after the immunostaining protocol. Sections were treated with 1 mM and 10 mM CuSO4 in 50 mM ammonium acetate for 1 h, then washed with PBS for 3 times.

For all the treatments, the sample tests were then equilibrated with 30% TDE/PBS (v/v) for 1 h and 68% TDE/PBS (v/v) for, at least, 1 h, before the placement in the sandwich and acquired using our LSFM.

Immunolabeling protocol. After clearing, small portions of the prefrontal cortex (99-year-old female, 6 months in formalin, Table S2) were used for investigating the optimal probe penetration using different autofluorescence masking treatments, temperatures, and dilution buffers (N = 3 independent experiments were performed for each treatment, temperatures, and dilution buffer in the prefrontal cortex, donor 1). For the optimization of the immunolabeling protocols, the antibiotic incubations were performed at 37 °C and 4 °C, using the following buffers: HEPES buffer 1X (HEPES pH 7.5, 10 mM v/v, Triton X 100, 200 mM NaCl, stock solution 2X), HEPES buffer supplemented with 2.5% Quadrol and 0.5 M urea. For the NeuN depth test, the primary antibody was incubated for 3 d at the dilution 1:50 and 1:100. Afterwards, the anti-NeuN stained samples were incubated for 1 day with the secondary antibody (dilution 1:200, 1:500, 1:1000). These protocols were performed for all the autofluorescence masking treatments described previously (n = 3) using the prefrontal cortex (99-year-old female, 6 months in formalin).

For the double and triple staining, the primary antibodies (anti-NeuN, anti-SST, anti-CR diluted at 1:50, 1:200 and 1:200, respectively) were incubated for 7 days at 37 °C. After 3 washes for 30 min each at 37 °C with PBST, the stained sample was incubated with the secondary antibodies alpaca-antirabbit IgG AF 488, donkey anti-rat IgG AF 568, goat antichicken IgY AF 647 (see Table S1), for 5 days at 37 °C and, then, washed 3 times for 1 h each at 37 °C. Such optimized staining protocol was performed on the superior frontal cortex (Fig. 3 and 4k-q; donor 2), Broca’s area (Fig. 4g-i; donor 3). For the hippocampus slice (Fig. 4b-d; donor 1), the primary antibodies (anti-NeuN, anti-GAD67 diluted at 1:50, 1:200, respectively) were incubated for 7 days at 37 °C. After 3 washes for 30 min each at 37 °C with PBST, the stained sample was incubated with the secondary antibodies goat-anti chicken IgG AF 488 and donkey anti-mouse IgG AF 647 (see Table S1), for 5 days at 37 °C and, then, washed 3 times for 1 h each at 37 °C.

For the multi-round experiments, the tissue was processed and stained as described previously. After acquisition, the samples were removed from the sandwich holder, equilibrated in 30% TDE/PBS (v/v) for 1 h and washed 3 times in PBS for 1 h each. Then, the specimens were delabeled with the clearing solution (200 mM SDS, 10 mM lithium hydroxide, 40 mM boric acid) at 4 °C for 1 h. Subsequently, the samples were extensively washed with PBST for 24 h. The stained proteins were: PV, SST, and Vip, labeled with bovine anti-goat IgG AF 488, donkey anti-rat IgG AF 568, goat antichicken IgY AF 647 (see Table S1), for 5 days at 37 °C and, then, washed 3 times for 1 h each at 37 °C. Such optimized staining protocol was performed on the superior frontal cortex (Fig. 3 and 4k-q; donor 2), Broca’s area (Fig. 4g-i; donor 3). For the hippocampus slice (Fig. 4b-d; donor 1), the primary antibodies (anti-NeuN, anti-GAD67 diluted at 1:50, 1:200, respectively) were incubated for 7 days at 37 °C. After 3 washes for 30 min each at 37 °C with PBST, the stained sample was incubated with the secondary antibodies goat-anti chicken IgG AF 488 and donkey anti-mouse IgG AF 647 (see Table S1), for 5 days at 37 °C and, then, washed 3 times for 1 h each at 37 °C.

Lipophilic dye staining with DiD. Slices (100 μm-thick) of superior frontal cortex from donor 2 were processed with SHORT and then incubated in 0.5% DiD (1,1'-dioctadecyl-3,3,3',3'-tetramethylindodicarbocyanine, 4-chlorobenzenesulfonyl salt) (stock solution 1 mg/ml, dilution 1:2) dissolved in 10% SDS (stock solution 20%, dilution 1:2) at 37 °C for 2-3 d. Samples were then soaked in a solution of 0.5% DiD in PBS and incubated in PBS for 1 h at RT. After 3 washes for 30 min each with PBS (v/v) supplemented with DAPI diluted 1:1000 for 1 day. Afterwards, the samples were placed in the sandwich with 68% TDE supplemented with DAPI at the dilution of 1:2000.
The tissue sections were imaged with our custom-made inverted light-sheet fluorescence microscope (LSFM). The illumination and detection objectives are a pair of 10× objectives (Nisle LMD 405, MLD 633) selected to maximize the magnification, NA 0.53, WD 10 mm, with a correction collar for refractive index matching with the immersion solution. The objectives are orthogonal and tilted at 45° in respect to the sample holder plane to avoid placing mechanical constraints on the lateral extension of the samples. Additionally, the set-up includes 4 over-lapped fluorescent filters (Colonial Scientific) with excitation at 488 nm and emission at 561 nm. Laser power 2 mW.

Imaging

Statistics and Reproducibility

For tissue autofluorescence characterization, cleared samples were imaged at 405, 488, 561, and 638 nm using our custom-made LSFM. To quantify the fluorescence intensities in each channel, 20 regions of interest (ROI) of 20 × 500 μm² were selected randomly in the grey matter of volumetric reconstruction of −3 × 3 × 0.5 mm³, using Fiji®. The 20 ROIs were analyzed for each treatment and tissue transformation protocol on consecutive small pieces of cortex (precentral cortex) from patient donor “1” (Table S2). We decided to characterize the effect of the various reagents on the brain of a very old patient (99 years old) as it presented the more severe presence of lipofuscin, whose accumulation increases during aging leading to an enhancement of the autofluorescence signals. The laser power at 405, 488, 561, and 638 nm was set at 5 mW. Means and standard deviations were plotted using Origin (OriginLab Corporation, Origin 2019b). The significance of the difference in mean values was determined using the Mann-Whitney test (Origin 2019b). The statistical significance was set as an α level = 0.05 (*P < 0.05; **P < 0.01; ***P < 0.001). To guarantee the reproducibility of the findings, 4 independent experiments were performed on different slices.

For the NeuN depth analysis, treated SWITCH/SHIELD processed slices were acquired using our LSFM at 638 nm and n = 3 of 200 × 700 μm² ROIs were selected for each treatment by Fiji, and mean and standard deviation were plotted using Origin. For quantifying the antibodies removal, ROIs of 6 × 6 pixels of images acquired at 561 nm using confocal microscopy were selected by Fiji, their mean and SD were plotted using Origin. For the shrinking quantification, the sample area before and after the clearing process was calculated by Fiji (N = 3).

For signal retention, 10 neurons CR-, SST-, and NeuN-immunoreactive in a region of 500 × 500 μm² were selected in the following downsampled reconstructions: after the immunostaining process (column 1; mean and standard deviation (SD)), after 1 year in the sandwich holder (column 2; mean and SD), and then reacquired after another week for the quantification of the bleaching process (3rd column; mean and SD; see Fig. S12). The signal retention was calculated as signal on the background taking 10 ROIs corresponding to the same area (10 × 10 μm²) and 10 ROIs of the background (10 × 10 μm²) using Fiji, and plotted with Origin.

The degree of the deformation of the tissue caused by SHORT during the stripping process was performed on the high-resolution LSFM images by using the structural similarity index (SSIM index) and the morphological deformation of the branching patterns at single-cell level in SST-immunoreactive neurons. For the SSIM index, 3 different pairs of neurons were cropped and scaled up by a factor of 4 to obtain a minimum number of 256 × 256 pixels. Then, the images were processed using the plugin SSIM index, to quantify the structural similarity. For the branching pattern tortuosity, the minimum distance and the real branch length of 6 different dendrites (see Fig. S10) was calculated using Fiji and plotted with Origin.

For visualization of the acquired data, the tiled raw images of extended samples acquired in a snake-like pattern by our LSFM are stitched into a 3D dataset. First, an affine transformation is applied to each image stack to resample the images and convert them from the objective reference frame to the sample reference frame. This is necessary to compensate for the fact that the objectives are tilted at 45° while the sample moves horizontally, i.e. parallel to the ground, during the acquisition. This transformation effectively “re-slices” the image stacks, optionally combining a downsampling factor in a single transformation matrix. Afterimage reslicing, the resolution of the resulting images is isotropic at 3.3 μm. Assuming Z is the thickness axis, X is the direction along which the sample moves and Y is the transversal axis, the resliced stacks are then stitched together along the Y axis using ZetaStitcher [https://github.com/lens-biophotonics/ZetaStitcher], a Python package designed to stitch large volumetric images such as those produced by LSFM. For visualization only purposes, a custom illumination intensity homogenization algorithm is applied to the stitched dataset or to its maximum intensity projection in order to equalize variations in the laser beam power that may occur within the objective field of view and/or between different image stacks. For each image, independent of the wavelength independently, the algorithm simply averages the observed intensity along the X axis to create a smoothed intensity profile along Y that is used as a flat field reference to even out illumination intensity artifacts occurring across the transversal sample extension. After compression of the tiled raw images to JPEG2000 format and stitching the downscaled rows, the 3D downscaled rendering of these planes was performed by Fiji. Maximum Intensity Projection of 20-150 slices were used in Figs. 3, 4, 5, S6, S7, S12, and S14.

As example: the acquisition of a 500-μm-thick slice of a Broca's area of 2 × 2 cm² stained with 2 different dyes requires 30 min and generates 0.4 terabyte. To efficiently store the 3D subvolumes (termed tiles) acquired by LSFM, the tiles were compressed to JPEG2000 format, which allows a reduction of the large-scale data by 60 times. To process and extract biological meaning from the acquired image data, the tiled raw images were resliced and downscaled with an isotropic resolution of 3.3 × 3.3 × 3.3 μm³ (starting resolution: 0.5 × 0.5 × 3.3 μm³). Next, the multi-tiles downsampled images were fused to perform a downscaled reconstruction of the whole slice. Such image processing allows converting the stained Broca’s area of 2 × 2 cm² in a Z-stack of 6 gigabyte (3 for each color) which can be visualized and analyzed by Fiji.

Data availability

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Raw data other than the representative images are available from the corresponding author upon reason the request.

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Author contributions
L.P., A.L. optimized the clearing and decolorization protocols; L.P., I.C., A.L. tested compatibility with immunolabeling. L.P., M.S. tested the different temperature for the antibody penetration and performed multi-staining and multi-rounds; M.S., L.P., and G.S. optimized the sample holder sandwich. L.P., V.G., and N.B. performed light-sheet imaging and confocal imaging. L.P., G.M., and V.G. contributed to the analysis and image processing. C.D. prepared the human brain slices. V.G., M.S., P.R.H, L.S. wrote parts of the manuscript. I.C., L.P., F.S.P conceived the study, supervised the study, analyzed results, and wrote the manuscript. All authors read and approved the manuscript.

Competing interests
The authors declare no competing interests.

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