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Comparative analysis reveals Ce3D as optimal clearing method for in toto imaging of the mouse intestine.

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ABBREVIATIONS

2D: two-dimensional
3D: three-dimensional
BABB: benzyl alcohol/benzyl benzoate
BABB-D15: BABB/diphenyl ether (1:15)
BSA: bovine serum albumin
DBE: dibenzyl ether
DCM: dichloromethane
DMSO: dimethyl sulfoxide
DPE: diphenyl ether
ENS: enteric nervous system
GFP: green fluorescent protein
GI: gastrointestinal
HE: hematoxylin-eosin
NHS: normal horse serum
PBS: phosphate-buffered saline
PBST: PBS containing 0.2% Triton X-100 and 20% DMSO
PFA: paraformaldehyde
PTwH: PBS/0.2% Tween-20 with 10 mg/ml heparin
RI: refractive index
RIMS: refractive index matching solution
RT: room temperature
SEM: scanning electron microscope
THF: tetrahydrofuran
V: volume
Key Points

1. Visikol®, ScaleS- and CUBIC-cleared mouse intestinal samples yield low to modest optical penetration depth, not allowing whole-thickness imaging of the intestinal wall and resulting in substantial structural deformation of the mucosal (sub)epithelial layer.

2. The 3DISCO, iDISCO+ and uDISCO clearing protocols render 1.5-mm-thick samples of the intact mouse intestinal wall fully transparent but equally suffer from destructive tissue effects (e.g. shrinkage and damage to tissue morphology).

3. The hydrophyllic clearing protocols Ce3D, SeeDB2 and UbasM result in excellent GFP retention and preservation of morphology as well as antigenicity and fluorescent staining. Only Ce3D renders full-thickness tissue homogeneously transparent.

4. Ce3D emerged as the most efficient, low-toxic and least time-consuming, cheap protocol for full-thickness 3D mapping of the gastrointestinal wall.
ABSTRACT

**Background:** The intestinal wall has a complex topographical architecture. The multi-layered network of the enteric nervous system and its intercellular interactions are difficult to map using traditional section-based or whole-mount histology. With the advent of optical clearing techniques, it has become feasible to visualize intact tissue and organs in 3D. However, as yet, a gap still needs to be filled in that no in-depth analysis has been performed yet on the potential of different clearing techniques for the small intestine.

**Aim:** The goal of this study was to identify an optimal clearing protocol for *in toto* imaging of mouse intestinal tissue.

**Methods:** Five aqueous-based clearing protocols (SeeDB2, CUBIC, ScaleS, Ce3D and UbasM) and four organic reagent-based clearing protocols (3DISCO, iDISCO+, uDISCO and Visikol®) were assessed in segments of small intestine from CX3CR1GFP/GFP and wild-type mice. Following clearing, optical transparency, tissue morphology, green fluorescent protein (GFP) fluorescence retention and compatibility with (immuno-)labeling were analysed.

**Key results:** All organic reagent-based clearing protocols – except for Visikol – rendered tissue highly transparent but led to substantial tissue shrinkage and deformation. Of the aqueous-based protocols, only Ce3D yielded full-thickness tissue transparency. In addition, Ce3D displayed excellent GFP retention and preservation of tissue morphology.

**Conclusions:** Ce3D emerged as a most efficient protocol for enabling rapid full-thickness 3D mapping of the mouse intestinal wall.

**Key words:** optical clearing, tissue transparency clearing capability, tissue morphology, GFP preservation, Ce3D, in toto imaging.
INTRODUCTION

Until recently, the topographical architecture of the enteric nervous system (ENS) and its interactions with other intramural components such as the mucosal barrier, the muscle and immune cells could only be visualized with 2D histological techniques, based on thin cross-sections or whole-mount preparations. However, 2D techniques are invasive, artefact-prone, and complicate the interpretation of complex 3D architectures (1-7). In recent years, tissue clearing has re-emerged as a powerful tool for bypassing tissue sectioning. This has led to the development of a plethora of tissue clearing protocols that reduce optical scattering and photon absorption by extracting the lipid fraction and/or by refractive index matching thereby yielding full-thickness tissue transparency (4, 7-14). These protocols allow detailed 3D visualization of intact organs by confocal, light-sheet or two-photon microscopy, but the appropriateness of each method is organ-specific and depends on the aspired goal in terms of clearing capability, preservation of fluorescent reporter protein signal, compatibility with immunolabeling and nuclear staining, morphological tissue integrity, complexity and cost (6, 14).

Despite the rapid and efficient clearing of large samples, the use of organic solvent-based clearing methods such as BABB (15), 3DISCO (16, 17), Visikol (18) and iDISCO(+) (19, 20) is typically associated with tissue shrinkage and quenching of fluorescent proteins (4). Moreover, the toxic nature of these organic solvents requires careful sample handling and limits the compatibility with immersion objectives. Although advances were made to overcome these disadvantages (20, 21), dehydration methods still suffer from sample shrinkage and ultrastructural damage remains unaddressed. To overcome these limitations, a novel category of clearing approaches based on aqueous solutions has been developed: SeeDB/SeeDB2 (22, 23), ClearT/ClearT2 (24), CUBIC and derived protocols (25, 26) including UbasM (11), Scale/ScaleS (27, 28) and Ce3D
Whereas the earlier protocols (SeeDB, ClearT2 and Scale) were still hampered by modest clearing capability and GFP preservation (7, 29), CUBIC and its modifications yielded better clearance (26). The more recently developed protocols (SeeDB2, ScaleS, Ce3D and UbasM) have been reported to show superior outcomes in terms of clearing capability, GFP retention and morphological preservation (11, 14, 23, 28). CLARITY and derived protocols compose a third category of clearing approaches, which involves the embedding of intact tissue into a hydrogel (30-33). Although these protocols aim to further improve tissue transparency with minimal protein loss, they are more time-consuming or require additional infrastructure (7, 29).

As far as the GI tract is concerned, a systematic benchmark of different clearing methods is lacking. Clearing of gut tissue has been performed with variable results and for specific aims using different approaches or modifications of existing protocols including FocusClear (1-3, 34-38), BABB-based methods (39), urea-based ScaleA2 and SeeDB (38), the PACT variant of CLARITY (32) and RIMS (40). Therefore, the aim of this study was to evaluate and directly compare the performance of the most effective, state-of-the-art clearing protocols on intestinal tissue. The outcome of this study provides a methodological guideline for researchers interested in exploring and visualizing connections and interactions in the intestinal wall through 3D image reconstruction and analysis.
MATERIALS AND METHODS

Tissue collection

All animal handling and housing procedures were conducted in accordance with European directive 86/609/EEC. The small intestine was obtained from adult CX3CR1<sup>GFP/GFP</sup> (41) and wild-type C57BL/6 mice. Animals were housed in controlled environmental conditions of temperature (22 °C) and illumination (12/12h light-dark cycle). All mice received food and water ad libitum. Animals were sacrificed by cervical dislocation and the ileum was collected and washed in ice-cold Krebs solution (117 mM NaCl, 5 mM KCl, 2.5 mM CaCl<sub>2</sub>·2H<sub>2</sub>O, 1.2 mM MgSO<sub>4</sub>·7H<sub>2</sub>O, 25 mM NaHCO<sub>3</sub>, 1.2 mM NaH<sub>2</sub>PO<sub>4</sub>·2H<sub>2</sub>O, and 10 mM glucose; pH 7.4). Afterwards, intestinal samples were opened along the mesenteric border and fixed with pins in a Sylgard-coated Petri dish. Subsequently, tissues were fixed in 4% paraformaldehyde (PFA) for 2h at 4 °C, washed with 0.01M phosphate-buffered saline (PBS; pH 7.4), and cut into segments of 1 cm length for tissue clearing.

Immunostaining and nuclear staining of intestinal whole-mounts

In general, the immunolabelling procedure preceded the clearing protocols, except for CUBIC, UbasM and Ce3D (see below). Nuclear counterstaining with TO-PRO-3 or DAPI was used for assessment of optical penetration depth across the intestinal layers. For immunofluorescent staining, a single neuronal immunostaining (βIII-tubulin-Cy3-immunostaining) or double staining with either an endothelial marker (CD31) or a fibroblast-like cell marker (PDGFR-α) was performed for all clearing protocols to test its compatibility with immunostaining. Additionally, the compatibility with the different clearing protocols of frequently used neurochemical markers used for subtyping cells in the intestinal wall were tested as well. In general, after an overnight blocking step,
intestinal samples were incubated in primary antibody dilutions for 2 days, rinsed in between and then followed by an overnight incubation in the secondary antibody solutions. The exact protocols of immunostaining procedures were slightly adapted to the clearing protocol used and are described in more detail in the respective paragraphs. The primary and secondary antibodies used in this study are listed in Table 1.

**Tissue permeabilization**

Tissue permeabilization for 3DISCO, iDISCO+, uDISCO and Visikol protocols was performed including either sample pretreatment with methanol or an alternative pretreatment protocol without methanol according to Renier and colleagues (19). In the first method, intestinal samples were rinsed 3x30 min in 0.01M PBS (pH 7.4), followed by permeabilization in a graded series of 20%, 40%, 60%, 80% and 2x 100% methanol on a shaker at RT (1h for each step). The samples were then rehydrated in a reversed methanol series (80%, 60%, 40%, 20%) and further permeabilized in PBS containing 0.2% Triton X-100 and 20% DMSO (PBST) for 2x1 h. In the methanol-free method, intestinal tissues were rinsed 3x30 min in PBS and then 2x1 h in PBST, followed by overnight incubation in PBST at 37 °C. Subsequently, the tissues were incubated in PBS containing 0.1% Tween-20, 0.1% Triton X-100, 0.1% deoxycholate, 0.1% NP40 and 20% DMSO, at 37 °C overnight, and then rinsed 2x1 h in PBS, containing 0.2% Triton X-100 for 1 h before the onset of further immunostaining procedures.

**Immunostaining and nuclear staining**

The immunostaining protocol for 3DISCO, iDISCO+, uDISCO was performed according to Renier and colleagues (19). To block any residual active aldehyde groups, intestinal samples were incubated overnight in PBS/0.2% Triton X-100/20% DMSO/0.3
M glycine at 37 °C and then blocked for 1 day in PBS/0.2% Triton X-100/10% DMSO/6% Normal Horse Serum (NHS) at 37 °C. After blocking, intestinal samples were rinsed twice in PBS/0.2% Tween-20 with 10 mg/ml heparin (PTwH) for 1 h, followed by incubation for 2 days in primary antibody solution (PTwH, containing 5% DMSO and 3% NHS), at 37 °C. Ileal segments were then rinsed 5x30 min in PTwH and incubated overnight in secondary antibody dilutions in PTwH containing 3% NHS at 37 °C. Samples were finally washed 5x30 min in PTwH prior to counterstaining with TO-PRO-3 (1:500 in PTwH) for at least 5 h. Finally, tissues were washed 5x30 min in PTwH. After the staining procedure, clearing was performed as detailed below.

**Tissue clearing**

The 3DISCO clearing protocol was carried out at RT under continuous gentle shaking (16). For iDISCO+ and uDISCO clearing we used the protocols described by Renier and colleagues (19) and Pan and colleagues (21), respectively. The Visikol clearing procedure was performed according to manufacturer’s instructions (Visikol, Whitehouse Station, NJ, USA).

For ScaleS and SeeDB2 clearing the protocols of Hama and colleagues (28) and Ke and colleagues (22) were applied, respectively. The CUBIC and UbasM clearing procedures were carried out according to the protocols described by Susaki and colleagues (26) and Chen and colleagues (11), respectively.

The Ce3D procedure involved incubation of intestinal samples for 8 h in a blocking solution containing 0.01M PBS/1% NHS/1% BSA/0.3% Triton X-100. For immunostaining, tissues were incubated in the primary antibody solution (0.01M PBS/1% NHS/ 1% BSA/0.3% Triton X-100/primary antibody) and then washed in 0.01M PBS/0.2% Triton-X100/0.5% 1-thioglycerol (Ce3D washing solution) for 3x10 min.
Subsequently, the intestinal tissues were incubated in the secondary antibody solution, which contained 0.01M PBS/0.3% Triton X-100/secondary antibody. The samples were then again rinsed in Ce3D washing solution for 3x10 min and then counterstained with either DAPI or TO-PRO-3 (1:500) for at least 1 h. Subsequently, ileal segments were again washed in Ce3D washing solution for 5x30 min. Stained tissues were immersed in the Ce3D medium for 12 h at RT and imaged in Ce3D medium (14). Note that the N-methylacetamide that is used in the Ce3D solution has high reproductive toxicity and that the N-methylacetamide produces toxic nitrogen oxide fumes when heated.

Macroscopic evaluation of transparency and tissue shrinkage

For macroscopic estimation of the degree of tissue transparency/opaqueness and tissue shrinkage/swelling 4% PFA-fixed ileal tissues, approximately 1 cm in length, were photographed in a transparent Petri dish above a standard grid allowing comparison of the transparency and tissue morphology achieved before and after the clearing procedures. Pictures were analyzed using Image-Pro Plus version 4.5.0.29 (Media Cybernetics, Silver Spring, MD, USA). Data were expressed as mean ± standard deviation (SD).

Confocal microscopy and data acquisition

Fixed intestinal samples, opened along the mesenteric attachment were visualized with a Leica SP8/LMS confocal microscope (Leica Microsystems CMS GmbH, Germany). Samples were positioned in a glass bottom Petri dish, submerged in their respective RI matching solution and covered by a golden ring with nylon mesh to prevent the tissue from floating. To evaluate the optical penetration depth, 300 image stacks (z
step size ~ 1.5 µm) of TO-PRO-3 counterstained samples with a 512x512 pixel resolution
(X and Y pixel size is 1.12 µm) at an excitation wavelength of 640 nm were recorded
using a 20x/0.75 HC PL Apo objective lens with a working distance of 600 µm. Immersion oil and glycerol solution (80%) were used for the four organic reagent-based
clearing protocols and the five aqueous-based clearing protocols, respectively. The
correction collar of this multi-immersion objective was adjusted accordingly to match the
refractive index of oil (RI~1.52) or 80% glycerol (RI~1.46). Thirty high-resolution image
stacks (z step size ~ 1 µm) with 1024x1024 pixel resolution were collected for imaging
of the specific layers of the intestinal whole-mounts.

**GFP fluorescence quantification**

After immunostaining and clearing, the retention of GFP fluorescence intensity in
CX3CR1-positive mononuclear phagocytes of intestinal whole-mounts of CX3CR1-GFP
transgenic mice was measured using ImageJ image processing freeware (National
Institutes of Health, Bethesda, MD, USA). In brief, Z-stacks of microvilli were projected
along the axial axis according to the maximum pixel intensity. Subsequently, a fixed
intensity threshold was set, just above background fluorescence, and the average intensity
of the objects detected above this threshold was measured. The average object intensity
was expressed relative to the background threshold value (i.e. GFP intensity ratio). Non-
cleared samples were used as controls to evaluate the GFP preservation of each clearing
protocol. Statistical analyses were performed using GraphPad Prism 6 software. All data
were expressed as mean ± standard error (SEM). A non-parametric t-test was used and
values with p < 0.05 were considered statistically significant.
Morphological damage assessment by hematoxylin-eosin (HE) staining and scanning electron microscopy (SEM)

To assess morphological damage at the light microscopical level, organic solvent-based cleared tissues were washed in 100% ethanol for 3x10 min and were then further processed for paraffin embedding, sectioned (5 µm) and stained for HE. Hydrophilic reagent-based cleared tissues had to be dehydrated first in a graded alcohol series before paraffin-embedding, subsequent sectioning and HE staining. For SEM, cleared samples, previously fixed in 4% PFA, were washed 3x30 min in 0.01M PBS and additionally fixed for 72 h at 4°C in 2.5% glutaraldehyde in 0.1 M Na-cacodylate buffer, dehydrated, critical point dried, gold-sputtered and viewed in a Quanta FEG250 SEM system (ThermoFisher Scientific).
RESULTS

Morphological preservation

Macroscopic images from intestinal samples before and after clearing taken on a mm-grid show that the mm-grid is less visible in the images of Visikol and ScaleS. All other protocols resulted in full transparency of the tissue. The highest shrinkage rates were observed for the organic solvent-based DISCO clearing procedures (3DISCO 57.16 ± 9.8%; iDISCO+ 32.25 ± 20.5%; µDISCO 43.60 ± 10.5% and Visikol 18.18 ± 9.8%), whereas much less shrinkage was observed after ScaleS and Ce3D clearing (6.65 ± 7.3, 5.40 ± 6.0% respectively) (Fig. 1). UbasM-, SeeDB2- and CUBIC-based clearing procedures resulted in a substantial (UbasM) or slight (SeeDB2 and Cubic) increase of tissue area (41.09 ± 7.1%, 4.32 ± 3.8% and 4.35 ± 2.6% respectively) (Fig. 1). In addition, in contrast to the 3DISCO and Ce3D procedures, CUBIC, iDISCO+ (not shown) and uDISCO produced significant destructive effects on the mucosal villi as observed on the SEM and HE images (Fig. 3). Tissue structural deformations induced by dehydration and delipidation procedures were also seen upon macroscopic examination, mainly for the 3DISCO- and uDISCO-cleared samples (Fig. 1).

Clearing capacity and optical penetration depth

Optical penetration depth analysis showed that only 3DISCO, iDISCO, uDISCO and Ce3D achieved full-thickness optical transparency of the intestinal whole-mounts (Fig. 2). The clearing capacity of the UbasM procedure yielded comparable results to the procedures mentioned above in specific areas but appeared not homogeneous for all different layers in the intestinal whole-mounts resulting in inhomogeneous fluorescent staining (Fig. 2 and Suppl Fig.).

GFP fluorescence preservation/quantification
For organic solvent-based clearing protocols, two distinct pretreatments of intestinal tissues have been reported for tissue permeabilization, based on either a dehydration/rehydration procedure with methanol or alternative pretreatment using an aqueous solution that contained PBS, detergent series and DMSO (19). The earlier reported significant GFP quenching effect of methanol (4, 42, 43) was also confirmed in our study. Consequently, we selected the alternative pretreatment without methanol for comparative GFP quantification. Estimates of GFP fluorescence, based on comparison of maximum intensity projection images of the different procedures, showed that Ce3D and SeeDB2 led to significantly lower GFP quenching compared to 3DISCO, uDISCO, iDISCO+, Visikol, CUBIC and ScaleS (Fig. 4).

**Compatibility of the clearing protocols with immunostaining**

Double immunostainings with the neuronal marker βIII-tubulin and either the endothelial marker CD31 or the fibroblast-like cell marker PDGFR-α were performed in combination with the nuclear stain DAPI on intestinal whole-mounts of CX3CR1-GFP transgenic mice (Fig. 5). Ce3D clearing allowed the simultaneous visualization of both the submucosal and myenteric enteric nerve network at the distinct topographical levels and its close interactions and associations with epithelial, immune and vascular components (Fig 5; Suppl video). In addition to the above-mentioned markers, other frequently used neurochemical markers (S100, GFAP, nNOS, NF200, Vimentin, c-kit, calretinin and calbindin) were tested as well (see Table 1; data not shown) and all these antibodies showed compatibility with the Ce3D clearing protocol.

**Comparison of the clearing protocols in terms of duration, handling and safety**

Additional comparison of the remaining protocols in terms of clearing time, technical handling and safety requirements revealed that SeeDB2 and Ce3D were the preferable methods (see Table 2, Suppl Fig.). In contrast to organic solvent-based clearing
protocols, these water-based clearing procedures involve the use of reagents that are non-irritant and less non-toxic and require less immersion steps, although it should be noted that one component in the Ce3D solution, i.e. N-methylacetamide, does have toxic features and might produce toxic nitrogen oxide fumes when heated (which is required in the protocol). Compared to the other protocols requiring a minimum incubation time ranging from a few days to 1 week, Ce3D required the shortest clearing time (12h) with good morphological preservation and compatibility with GFP and different antibodies tested.
DISCUSSION

The applicability of 9 clearing techniques (4 organic solvent-based and 5 aqueous-based protocols) was evaluated for 3D visualization of the entire thickness of the intestinal wall in terms of the following parameters: tissue clearing efficiency, preservation of tissue morphology and GFP fluorescence, immuno- and nuclear staining compatibility, clearing protocol time and safety of the chemical reagents used. CLARITY-based protocols have not been performed in this study, since these are protocols that are more time-consuming or require additional dedicated (commercially available) equipment and have been reported to lead to GFP quenching, ultrastructural changes and compromised structural integrity due to aggressive lipid removal from the biological samples (7, 23, 29).

In line with data reported in other organs (6, 14, 16, 17, 19, 21, 29, 44), of all 9 optical clearing techniques compared in this study, 3DISCO, iDISCO+, uDISCO and Ce3D showed the highest and most homogeneous clearing capacity. In terms of visualization of full-thickness intestinal samples, the Ce3D protocol equally yielded better clearing results compared to SeeDB2 (23). Compared to the original CUBIC protocol, its improved version UbasM did result in higher transparency of the intestinal tissue and less morphological damage (similar to the Ce3D protocol), but suffered from inhomogeneous clearing throughout the intestinal wall. The inferior results in clearing capacity for ScaleS and in particular for Visikol, observed in our experiments on mouse intestine, indicate that its clearing performance is tissue-specific, since better results with these protocols were obtained in brain tissue (28, 45). This tissue dependence might be explained by differences in lipid content, pigment, or the degree of inhomogeneity in refractive index (4).
Despite having excellent clearing capacity and providing good results in the brain, the distinct DISCO protocols are less suitable to obtain optimal images of the full-thickness intestinal wall due to the destructive effects they have on intestinal samples (e.g. damage in mucosal epithelium and/or membrane integrity, and substantial tissue shrinkage). It has been reported that the potent clearing properties of 3DISCO, iDISCO+ and uDISCO, which might be an advantage for brain tissue, may lead to destructive effects on biological (sub)cellular structures, due to the aggressive properties of a.o. methanol, THF and DCM (10, 11, 45). Similarly, in line with earlier reports on other tissues (8, 11, 28, 35, 46) the CUBIC protocol affected the morphology of the intestinal samples, limiting compatibility of these protocols for post-hoc correlative ultrastructural studies.

Substantial or even complete loss of GFP fluorescence occurred in intestinal samples of CX3CR1 transgenic mice after 3DISCO, iDISCO+, uDISCO and Visikol clearing, which is in line with previous findings in other tissues (11, 13, 14, 28, 29, 46, 47). Therefore, taken together with their destructive effects on morphology, these protocols were not further considered for testing their compatibility with immunostaining with intestinal neurochemical markers. Of the 5 aqueous-based protocols, urea-based methods such as ScaleS and CUBIC equally induced substantial GFP quenching, probably due to the presence of the high detergent concentration and the denaturating properties of urea (7, 29, 46, 48). In this study, only UbasM (using much lower detergent concentrations compared to CUBIC), SeeDB2 and Ce3D (both using iohexol solutions) met the requirements in terms of GFP signal retention, as was also found for other fluorescent reporters (23, 49). The latter three protocols were also compatible with immunostaining procedures enabling the visualization of the intrinsic enteric network and its interaction with blood vessels, immune cells and the distinct components of the
mucosal barrier. The aqueous-based clearing procedures were also superior to the non-aqueous organic solvent-based protocols in terms of long-term storage due to the more aggressive solvents which have a strong negative impact on antigenicity (29).

In conclusion, Ce3D emerged as the preferred clearing protocol for 3D imaging of full-thickness intestinal segments, since it is a non-destructive, cost-effective and fast clearing technique with excellent GFP signal preservation and immuno- and nuclear counterstaining compatibility. Thus, it provides a promising tool for studying the distinct intramural components in the GI tract in healthy and diseased conditions. This 3D imaging approach of full-thickness samples opens avenues for detailed mapping and distributional analysis of distinct subsets of cells of the neuro-immune-vascular interactive network within the entire GI wall, thereby leading to a better understanding of the pathophysiological changes occurring in specific GI disorders.
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COMPETING INTERESTS

The authors declare no competing interests.

AUTHOR CONTRIBUTION

GB, IP and ST performed experiments and participated in data analysis. RB and JPT designed the study. GB, RB and JPT wrote the largest part of the manuscript. JD, IP, JZ, RB, WDV and JPT critically revised the manuscript. RB, WDV and JPT provided essential research tools, research infrastructure and funding for this study.
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**LEGENDS**

**Figure 1.** Macroscopic images from intestinal samples before and after clearing, taken on a mm-grid and showing the degree of opaqueness and shrinkage/swelling. The mm-grid is less visible in the images of Visikol and ScaleS compared to the other protocols which all led to full transparent tissue. The three DISCO protocols and Visikol showed large shrinkage of the tissue while UbasM resulted in a substantial swelling. The graph represents the average projected area change (%) for all studied protocols. Mean data ± SD are plotted. iDISCO+, uDISCO, ScaleS (n = 5); 3DISCO, Visikol, SeeDB, CUBIC and UbasM (n = 4) and Ce3D (n = 3).

**Figure 2.** TOPRO3 staining of full-thickness intestinal samples represented as yz images illustrating the level of transparency. Images were taken with an inverted microscope and the serosa was positioned towards the bottom of the petri-dish. The white dots indicate the borders formed by the serosal layer (z = 0) and the top of the villi (z = 1). Full-depth imaging was achieved with the DISCO protocols and Ce3D. This is also evident from the corresponding graphs that present the changes in intensity of the fluorescence (normalized) over the length of the imaged tissue (z normalized for all graphs). The z
value that corresponded with the maximal intensity of fluorescence was set as zero; only z values above this point are shown in the graph. The curve of ScaleS is used as a reference (red line). Scale bar = 50 μm

Figure 3. Representative images of HE-stained sections and SEM images of non-cleared control and cleared ileal segments. (A), (C), (E), (G), and (I) display HE-stained sections from non-cleared control, 3DISCO, uDISCO, CUBIC and Ce3D, respectively. (B), (D), (F), (H) and (J) show SEM images corresponding to the HE sections. The images of HE-stained sections and SEM unambiguously illustrate the damage to the villus membrane integrity or epithelial mucosa of the villi after clearing with uDISCO and CUBIC (E, F, G, H). On the other hand, the tissue is less affected by 3DISCO and Ce3D procedures as seen in (C), (D), (I) and (J). Inserts depict a more detailed image of the area delineated with the dashed line, corresponding with the top of the villi. Scale bars of the overview images = 100 μm; scale bars of the inserts = 50 μm

Figure 4. Graph representing the degree of GFP preservation of each clearing protocol. To calculate the GFP intensity ratio the average object intensity was expressed relative to the background threshold value. Non-cleared samples were used as controls. This estimate of GFP fluorescence confirmed that the GFP signal was less affected by Ce3D and SeeDB2 protocols. Imaging was performed immediately after finishing the clearing procedure. Mean data ± SEM are plotted. Control (n = 9); iDISCO+, uDISCO, Visikol, SeeDB2, CUBIC, ScaleS and Ce3D (n = 10); 3DISCO, UBASM (n = 5).
Figure 5. Full-depth 3D observation showing the epithelium, the enteric nerve network (βIII-tubulin) and its close association with intestinal mononuclear phagocytes (CX3CR1), fibroblast-like cells (PDGFRα) and the vascular network (CD31) in Ce3D-cleared intestinal segments. (A) Full-depth 3D distribution of the intestinal villi and epithelium visualized by DAPI nuclear staining (at the left side of the dotted line). At the right side of the dotted line, the DAPI channel is removed allowing the visualization of the underlying structures, i.e. the underlying neuronal and vascular network as well as CX3CR1+ cells across the distinct intestinal layers. A more detailed view of these underlying structures is shown in (B). (C) Bottom view of the myenteric plexus with the deeper located CX3CR1-GFP cells and blood vessels. (D) Similar image as (B) but now showing the fibroblast-like cell network instead of blood vessels. (E) Optical cross-section of an intestinal villus illustrating the intimate network of enteric nerves, blood vessels, immune cells and fibroblast-like cells. Scale bars (A – D) = 100 μm; Scale bar (E) = 50 μm

TABLES

Table 1. Antibodies used for immunostaining.

<table>
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<tr>
<th>Epitope</th>
<th>Host</th>
<th>Dilution</th>
<th>Supplier</th>
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<tbody>
<tr>
<td>βIII tubulin</td>
<td>Rabbit</td>
<td>1:500</td>
<td>Biolegend; Poly18020</td>
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<tr>
<td>S100</td>
<td>Rabbit</td>
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<td>Dako; Z0311</td>
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<td>NF200</td>
<td>Rabbit</td>
<td>1:500</td>
<td>Sigma-Aldrich; N4142</td>
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<tr>
<td>Vimentin</td>
<td>Goat</td>
<td>1:50</td>
<td>Millipore; AB1620</td>
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<tr>
<td>Clearing protocol</td>
<td>Preservation efficiency</td>
<td>IHCa</td>
<td>Handling</td>
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<tr>
<td><strong>Aqueous solution-based</strong></td>
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<tr>
<td>CUBIC</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>SeeDB2</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>ScaleS</td>
<td>+</td>
<td>nd</td>
<td>+</td>
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<tr>
<td>Ce3D</td>
<td>+++</td>
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<tr>
<td>UbasM</td>
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<tr>
<td><strong>Hydrophobic solution-based</strong></td>
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<tr>
<td>3DISCO</td>
<td>+</td>
<td>+</td>
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<tr>
<td>iDISCO+</td>
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<tr>
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</tr>
<tr>
<td>Visicol</td>
<td>-</td>
<td>nd</td>
<td>+</td>
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</table>

+++ = good; ++ = medium; + = weak; -- = not present; *= immunostaining; nd = not determined given the low clearing capacity; *= only incubation time of tissue permeabilization and/or tissue pretreatment and clearing procedures was considered without immunolabeling.

**Supplementary material**

**Supplementary figure.** Similar images as shown in Figure 5, displaying the enteric nerve network (βIII-tubulin), the mononuclear phagocytes (CX3CR1), and the vascular
network (CD31) in UbasM-cleared (A, B and E) and SeeDB2-cleared intestinal segments (D, C). Note the inhomogeneous clearing capacity (asterisks) and the slightly lower immunostaining quality for the UbasM protocol (A,B). (C-E) Cross-sections of intestinal villi of CX3CR1 transgenic mice after SeeDB2 and UbasM clearing and immunostaining, respectively. Scale bar (B) = 100 μm, Scale bar (C, D, E)= 50 μm

Supplementary Video. 3D observation of the arrangement of the neurovascular network and the CX3CR1+ mononuclear phagocytes throughout the full thickness of the intestinal wall. Red: blood vessels stained by the endothelial marker CD31; green: GFP-labeled CX3CR1-positive mononuclear phagocytes; blue: the enteric nerve network visualized by means of βIII-tubulin immunostaining; purple: nuclear (DAPI) staining.