Panoptic imaging of transparent mice reveals whole-body neuronal projections and skull–meninges connections

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Analysis of entire transparent rodent bodies after clearing could provide holistic biological information in health and disease, but reliable imaging and quantification of fluorescent protein signals deep inside the tissues has remained a challenge. Here, we developed vDISCO, a pressure-driven, nanobody-based whole-body immunolabeling technology to enhance the signal of fluorescent proteins by up to two orders of magnitude. This allowed us to image and quantify subcellular details through bones, skin and highly autofluorescent tissues of intact transparent mice. For the first time, we visualized whole-body neuronal projections in adult mice. We assessed CNS trauma effects in the whole body and found degeneration of peripheral nerve terminals in the torso. Furthermore, vDISCO revealed short vascular connections between skull marrow and brain meninges, which were filled with immune cells upon stroke. Thus, our new approach enables unbiased comprehensive studies of the interactions between the nervous system and the rest of the body.

Most diseases, even when arising in a specific site, eventually affect the entire organism. Histological techniques developed in the last century have been the standard procedure for charting pathology, yet a more complete understanding of biological mechanisms requires an unbiased exploration of the whole organism, not just pre-defined tissues. However, mammalian tissues are naturally opaque, hindering high-resolution imaging in any tissue deeper than a few hundred micrometers, a major reason why sectioning is needed for histological examination of target organs. Recent innovations in tissue clearing technology now allow three-dimensional (3D) histological examination of intact organs. Tissue clearing is a chemical process aiming to match refractive indices throughout intact tissues, yet a more complete understanding of bio-process aiming to match refractive indices throughout intact tissues now allow three-dimensional (3D) histological examination of intact organs. For the first time, we visualized whole-body neuronal projections in adult mice. We assessed CNS trauma effects in the whole body and found degeneration of peripheral nerve terminals in the torso. Furthermore, vDISCO revealed short vascular connections between skull marrow and brain meninges, which were filled with immune cells upon stroke. Thus, our new approach enables unbiased comprehensive studies of the interactions between the nervous system and the rest of the body.

vDISCO, a pressure-driven, nanobody-based whole-body immunolabeling technology to enhance the signal of fluorescent proteins using nanobodies, which consist of the variable domain of heavy chain antibodies (V₃₅H). This technology can enhance fluorescent signals more than 100 times and thereby allows head-to-toe light sheet microscopy scanning of transparent mice (panoptic imaging) and quantification of subcellular details. We used this technology to construct the first neuronal projection map of an adult mouse and to reveal far-reaching changes in neuronal projections and inflammatory processes following acute CNS injuries. Panoptic imaging also revealed short vascular connections between the skull marrow and the rest of the body.

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and meninges (both at the brain surface and sagittal sinus), which may serve as immune gateways following stroke.

Results

vDISCO principles and signal enhancement. Quantitative assessments of intact transparent mouse bodies remained a major challenge impeding the study of neurological diseases at the whole-body scale. We reasoned that fluorescent proteins such as EGFP and red fluorescent protein (RFP) are relatively dim compared with bright fluorochromes (for example, Alexa and Atto dyes) after tissue clearing, making them difficult to detect in centimeters-thick mouse bodies, especially through intact thick bones (for example, skull and vertebrae) and skin. Therefore, we set out to enhance the signal of fluorescent proteins via whole-body immunolabeling with brighter and more stable fluorescent dyes to provide a much higher contrast (signal-to-background ratio) in cleared mice. Furthermore, using fluorescent dyes in the far-red spectrum could help to overcome tissue autofluorescence, thus allowing reliable detection of subcellular details in all tissues (Supplementary Fig. 1). We reasoned that nanobodies are particularly suited to achieve thorough immunolabeling throughout entire adult mouse bodies because of their small molecular weight (12–15 kDa) compared with that of conventional antibodies (~150 kDa). Indeed, nanobodies were more efficient for labeling large tissues compared with conventional antibodies (Supplementary Figs. 2 and 3).

First, we tested the signal quality of nanobody labeling in dissected mouse brains from Thy1-GFP-M mice, in which a subset of neurons express EGFP. Following the nanoboosting, the tissues were cleared using organic solvents. We found that nanoboosting enhanced the signal intensity by one to two orders of magnitude compared with direct imaging of fluorescent proteins (Fig. 1a–i). In the nanoboosted samples, fine details of neurons were evident even in the deep brain regions of Thy1-GFP-M mice (compare Fig. 1b,c with Fig. 1e,f). We also obtained a similar signal intensity increase in the cerebellum of Thy1-GFP-M brains (compare Fig. 1g with Fig. 1h), which is notoriously difficult to clear due to the high lipid content.

A major aim of tissue clearing approaches is to perform automated quantifications in large imaging datasets in an unbiased and timely way. Towards this goal, we used the NeuroGPS-Tree algorithm, a robust, automated neuron-tracing tool that was recently developed for tracing neurons in scans of cortical regions obtained by high-resolution confocal microscopy. We found that virtually all of the neuronal cell bodies and neurites were detected and linked to each other as complete neurons upon vDISCO nanoboosting in light-sheet microscopy scans (Supplementary Figs. 4 and 5). In contrast, in unboosted samples or in standard immunoglobulin G antibody-boasted samples, many fine extensions of neurons were not identified or not connected to somas and neuronal trees (Supplementary Fig. 4). Nanoboosting allowed imaging not only of neuronal details but also of smaller individual cells such as microglia cells and immune cells (Fig. 1j–m and Supplementary Fig. 6). Compared with unboosted samples, we could resolve fine details of microglia cells in intact transparent brains of CX3CR1^GFP^+ mice using light-sheet microscopy (Fig. 1j–m and Supplementary Video 1). Nanoboosting also enabled automated quantification of CX3CR1 GFP^+ cells using the ClearMap algorithms: we found approximately 2.3 million microglia cells in the adult mouse brain (Supplementary Fig. 7a). We also automatically quantified microglia in all brain regions annotated by the Allen Brain Atlas. Thereby we found, for example, ~150,000 microglia in the hippocampus and ~50,000 in the thalamus of adult mice (Supplementary Fig. 7b–k). Furthermore, we found no significant decrease in signal intensity over time of the same samples, suggesting that vDISCO nanoboosting stabilizes the fluorescent signal (Fig. 1n,o), allowing long-term preservation and re-imaging of the samples. Thus, owing to the enormous enhancement and stabilization of fluorescent signals using nanobodies, we could use available computational tools to automatically trace neurons and count cell numbers in images acquired by light-sheet microscopy.

vDISCO allows panoptic imaging of intact adult mouse bodies. To apply whole-body clearing technology at the systems level, it is critical to obtain information not only from dissected organs but also from intact transparent bodies. Towards this goal, we aimed at establishing an approach to achieve nanoboosting in the entire mouse body. We reasoned that vDISCO amplification of the fluorescent signals would allow reliable quantification of cells through intact thick bones and highly autofluorescent muscles. To ensure homogeneous delivery of nanobodies into all body regions, we used high-pressure cardiac perfusion with two- to three-fold increased pressure compared with standard perfusion protocols (160–230 mmHg compared with 70–110 mmHg (ref. 25)). We also illustrated the effect of high-pressure delivery compared with standard pressure using methylene blue. This novel high-pressure delivery approach facilitated a consistent distribution of the dye into deep tissues (Supplementary Fig. 8). Assessing the tissue integrity upon high-pressure delivery, we did not observe any structural changes (Supplementary Fig. 9). Furthermore, we used a permeabilization solution containing Triton X-100, methyl-β-cyclodextrin (to extract the cholesterol from membranes) and trans-1-acetyl-4-hydroxy-l-proline (to loosen the collagen network)16. To further reduce the background signal caused by the residual blood and to decalcify the bones, we treated whole mouse bodies with aminoalcohols14 and EDTA15 before the whole-body immunolabeling step. Combination of these diverse improvements allowed us to achieve whole-body immunolabeling in adult mice with a high degree of transparency (Fig. 2a and Supplementary Fig. 10) and to obtain quantitative information on single cells in deep brain regions through the intact skull16,17 (Supplementary Figs. 11 and 12). Here, we used anti-XFP nanobodies conjugated with bright Atto dyes (called nanoboosters) to boost the fluorescent protein signal in transparent mice. Moreover, we used anti-vimentin nanobodies to label activated glia cells in the brain upon trauma, demonstrating that our approach also works with nanobodies targeting endogenously expressed proteins (Supplementary Fig. 13).

In addition to the specific boosted signal, we also visualized and segmented other major tissues in the transparent body: muscles by their autofluorescence within the blue-green spectrum, and bones and internal organs by propidium iodide (PI) labeling (Fig. 2b,c and Supplementary Fig. 14a–d). Usage of organic solvents, which induce isotropic tissue shrinkage14, allowed us to perform head-to-toe panoptic imaging of entire mice by light-sheet microscopy.

Being able to image subcellular details of neurons through intact bones and highly autofluorescent muscles in the whole mouse body using vDISCO panoptic imaging, we constructed the first whole-body neuronal projection map of a Thy1-GFP-M transgenic mouse (Fig. 2d,e, Supplementary Fig. 14 and Supplementary Video 2). We noticed that in the peripheral nervous system, mainly axons innervating muscles were labeled in this mouse line, which enabled us to investigate the neuromuscular junctions in greater detail. We also observed fluorescent labeling of the internal organs, such as kidneys of Thy1-YFP-H mice18 (Supplementary Fig. 15). Owing to the great increase in signal-to-background ratio by vDISCO, we could readily visualize axonal projections from the spinal cord through intact vertebrae to their terminals into the muscles, even in very remote locations such as toes (Fig. 2f–i and Supplementary Videos 3 and 4). Panoptic imaging of individual neuronal projections through intact bones can now provide more details on neuroanatomy. For example, how axons coming from consecutive roots enter the spinal cord—that is, in an overlapping or non-overlapping manner—has been unclear26. Using our technology, we observed that in mice, the
Fig. 1 | Enhancement and permanent preservation of the fluorescence signal with vDISCO. a–f, Signal quality comparison between vDISCO-boosted (a) and unboosted (d) half-brain samples coming from the same 11-month-old Thy1-GFP-M mouse. To achieve the best comparison between the two procedures, we divided the mouse brains in two halves for light-sheet microscopy imaging (one hemisphere boosted and imaged in the far-red channel, the other hemisphere unboosted and imaged in the green channel for endogenous EGFP). The boosted hemisphere showed highly distinguishable cellular details (b, c) such as axonal projections not visible in the unboosted hemisphere (e, f), especially in the regions with dim GFP labeling in Thy1-GFP-M mice, such as mid-brain (b, e) and cerebellum (c, f) (representative images, repeated at least on 3 different samples). g, h, Comparison of signal quality in cerebellum from the boosted (g) and unboosted (h) samples (repeated with at least 3 mice per group). i, Plots of signal intensity profiles from boosted samples (left) versus unboosted samples (right) along the blue and orange lines in g and h, respectively; n = 3 brains for each group (2- to 6-month-old Thy1-GFP-M mice). j–m, Light-sheet microscopy images (×4 and ×25 magnification) of the microglia from CX3CR1<sup>gfp</sup> mice in boosted (j, k) versus unboosted samples (l, m) showing the fine details of microglia ramifications obtained with vDISCO boosting (similar results repeated with at least 3 mice per group). n, Representative light-sheet microscopy images of 1 of 3 CX3CR1<sup>gfp</sup> mouse brains at 0, 2, 12 and 18 months after boosting, showing the preservation of the fluorescence signal over 18 months. o, Fluorescence level quantifications, in arbitrary units (a.u.), in CX3CR1<sup>gfp</sup> brains after boosting at different time points postclearing (n = 3 mice, 2–6 months old).
axonal bundles coming from different roots enter the spinal cord at non-overlapping territories (Supplementary Fig. 16). Thus, the vDISCO approach provides a holistic view of the intact mouse, which should lead to novel discoveries on how interconnected organ systems function in health and what happens during their perturbation in disease.

Clearing of the whole body without removal of the skin. The skin is the largest organ, forming ~15–20% of the body. Among many other components, it consists of nerve endings, glands, immune cells, blood and lymphatic vessels, signifying its diverse biological roles in addition to being a physical barrier against the external environment\(^2\). However, clearing and imaging whole mice with skin has not been reported. Towards this goal, we adopted the vDISCO protocol to clear and image mice with skin. We observed that vDISCO could efficiently clear whole nude mice without removal of the skin (Fig. 3a,b). In addition, we cleared C57BL/6j (BL6) background mice after shaving the fur. In CX3CR1\(^{GFP^+}\) mice, we could readily image
Fig. 3 | vDISCO panoptic imaging on mice with the intact skin. a, b, Adult NMRI nu/nu nude mouse, 3–4 months old (a), cleared with the vDISCO method (b): the insets show close-up views of the cleared lungs (magenta) and skin (cyan) in b (similar results were observed from 3 independent NMRI nu/nu mice). c–h, vDISCO on 6-week-old CX3CR1GFP/+ mouse with intact skin. c, Cleared body, with the inset (cyan) showing visible inguinal lymph nodes (black arrowhead) through the transparent skin. d, Light-sheet images of CX3CR1GFP/+ mouse boosted with anti-GFP 647-nanobooster (cyan) and labeled with PI (magenta), showing CX3CR1 GFP+ cells (cyan) densely located in the lymph node and sparsely located in and under the skin, such as around hair follicles (magenta). e–h, Confocal 1-mm z-scan of the skin and lymph node in d with visible CX3CR1 GFP+ cells (cyan) and nuclei (magenta). e, Three-dimensional rendering of the stack showing the skin and the lymph node beneath. f, Two-dimensional image at the level of the skin with individual CX3CR1 GFP+ cells in the tissue and in the hair follicles (yellow arrowheads). g, The fine details of the ramifications of the immune cells at the surface of the skin are visible (green arrowheads). h, Two-dimensional image of the lymph node with visible immune cells (yellow arrowheads) in the organ. See also Supplementary Video S. i–k, vDISCO on 2-month-old Thy1-GFP-M mouse with intact skin. i, Cleared body, with the inset (magenta) showing a visible inguinal lymph node through the skin. j, Light-sheet images of Thy1-GFP-M mouse boosted with anti-GFP 647-nanobooster (green) and labeled with PI (magenta), showing Thy1-GFP-expressing immune cells (white arrowheads) and neuronal projections coming from the spinal cord and projecting to the rest of the body, including the skin (green); the insets show close-up details of a neuronal projection (green fibers and yellow arrowheads) projecting into the skin. k, High-magnification confocal image of the region in j (yellow dashed rectangle) showing details of the immune cells (white arrowheads) and neuronal projections (green and yellow arrowheads) into the skin of the animal. All the results from c to k were similarly observed in 2 independent mice for each mouse line.
immune cells through the intact skin owing to the increase of fluorescent signal by vDISCO (Fig. 3c–h and Supplementary Video 5). In Thy1-GFP-M mice, we observed the axonal innervations in the skin in addition to Thy1 GFP+ immune cells located under the skin (Fig. 3i–k). These results demonstrate that vDISCO allows detailed imaging of intact mice with skin, which may be of great value for studying skin-related pathologies.

Degeneration of peripheral nerve terminals after brain injury. Traumatic brain injury (TBI) is a major cause of death and disability, and currently there is no disease-modifying treatment available to combat especially chronic sequelae. In addition to its acute consequences, TBI often leads to chronic focal and global neurological impairments, such as dementia, epilepsy, progressive motor decline, peripheral neuropathy and dystonia36,37. These impairments could be due to degeneration of the corresponding brain regions and/or the dysfunction of nerves outside of the brain. However, so far, the impact of TBI on long-range neuronal projections outside of the brain has not been assessed. For example, while TBI-induced sensory-motor dysfunction has been observed in both animal models and human pathologies31–34, the underlying mechanisms remain unknown, hampering the development of effective therapeutic approaches. Here, imaging the whole CNS, we found extensive neurodegeneration of the descending motor axons in the brainstem and spinal cord upon TBI over the right somatosensory and motor cortices (Supplementary Fig. 17). However, how TBI alters neuronal projections in the peripheral nervous system remains elusive. We used panoptic imaging to examine neuronal changes throughout the body more than a month after the brain injury (chronic stage) in comparison with control animals. As mainly axons innervating muscles are labelled in Thy1-GFP-M mice, we primarily focused on these nerve terminals. Our quantifications conducted in intact cleared mouse bodies demonstrated that the complexity of nerve terminals at the neuromuscular junctions was reduced after TBI, especially in the upper torso, compared with unlesioned control mice (Fig. 4a–d). We observed that nerve endings were reduced in complexity at the contralateral body regions, with fewer axonal ramifications left, implying partial degeneration of these axon terminals (Fig. 4e,f). Interestingly, we also observed changes at the ipsilateral axons, albeit to a smaller degree with respect to the contralateral body region (Fig. 4e). Overall, these data demonstrate that light-sheet microscopy imaging of entire transparent mouse bodies by vDISCO holds great promise for discovering hitherto unknown biological and pathophysiological processes.

Visualizing new routes for immune cell trafficking in the brain and spinal cord. The lymphatic system, which connects various lymphatic organs in the body, is crucial for immune responses. Until recently, the brain was considered to be devoid of any lymphatic vessels. In recent years, brain lymphatic vessels were re-discovered and are considered to play critical roles in brain pathologies35–38. As these lymphatic vessels are located between the brain and the skull39, their connections are largely destroyed when the brain is harvested for standard histology.

We utilized panoptic imaging to overcome this hurdle and imaged details of meningeal vessels in intact transparent mice. We used Prox1-EGFP reporter mice, in which lymphoid endothelial cells express EGFP40. We readily observed previously described brain lymphatic structures, particularly along the sagittal sinus, pterygopalatine artery and transverse sinus (Fig. 5a–e and Supplementary Fig. 18) in addition to many other body regions (Supplementary Fig. 19 and Supplementary Video 6). Next, we imaged multiple subtypes of immune cells within and outside the meningeal vessels using the CX3CR1-YFP+/×CCR2-RFP+/mice by multiplexing with two different nanoboosters (anti-GFP conjugated to Atto-647N and anti-RFP conjugated to Atto-594N). The CX3CR1 GFP+ microglia cells in the brain parenchyma (Fig. 5fg) and CCR2 RFP+ peripheral immune cells in the meningeal vessels were clearly identifiable (Fig. 5gh and Supplementary Video 7). As expected, we also observed CX3CR1 GFP+ immune cells in meningeal vessels, which likely represent meningeal monocytes and macrophages (Fig. 5h). It remains unclear how the immune cells within meningeal vessels contribute to the pathology of acute brain injuries, as they may be in an advantageous position to invade the brain. To start addressing this query, we used the middle cerebral artery occlusion (MCAO) model of stroke, which was performed in LysM-EGFP mice, a transgenic mouse line expressing EGFP in myeloid cells (mainly in neutrophils and monocytes) but not in microglial cells41. The mice with MCAO showed an invasion of LysM GFP+ cells into the brain parenchyma, especially in the peri-infarct region, compared with sham-operated controls (Fig. 5j). Numbers of LysM GFP+ cells also increased in the meningeal structures after MCAO (Fig. 5i), suggesting that the meninges might play a role as entry and/or exit routes of the brain in addition to the disrupted blood–brain barrier and the choroid plexus42. Thus, vDISCO panoptic imaging of transparent mice is a powerful tool to study the anatomy and cellular repertoire of meningeal vessels in health and disease.

Next, we aimed at investigating vascular connections between the skull and the meninges in greater detail. Towards this goal, we injected a cerebrospinal fluid (CSF) tracer (fluorophore-conjugated ovalbumin) into the cisterna magna in VEGFR3-YFP mice, another commonly used reporter line for meningeal vessels. We observed that the CSF tracer was quickly transported into the VEGFR3+ meningeal vessels (Fig. 6a–d). Interestingly, we also observed that the tracer-labeled vessels extended into the skull (Fig. 6b,c). We further investigated the details of these new connections using lectin, another vessel tracer binding to endothelial cells. We found that these vascular connections are mostly located between the skull marrow and the outer surface of the meninges (both at the brain and sagittal sinus interfaces; Fig. 6c,f, Supplementary Fig. 20 and Supplementary Videos 8 and 9). We measured that they have an average size of 46.9 ± 5.8 µm in length and 19.2 ± 2.5 µm in width at the brain interface and of 83.7 ± 0.3 µm in length and 28.5 ± 1.8 µm in width at the sagittal sinus interface (mean ± s.e.m., n=3 mice; Supplementary Fig. 20); therefore, we named them as short skull–meninges connections (SMCs). They also occasionally contain cells (Fig. 6f,e and Supplementary Fig. 20). To address whether the SMCs might play a role in neuropathological conditions, we studied the distribution of LysM GFP+ immune cells in these connections after MCAO. The mice with MCAO showed significantly more LysM GFP+ cells in the SMCs compared with sham-operated controls (Fig. 6g–j and Supplementary Video 10), suggesting that they might represent an entry route for myeloid cells from the skull marrow to the meningeal compartments after brain injury. Thus, vDISCO imaging revealed connections between skull and meninges, which might be important for immune cell trafficking, especially during brain pathologies.

Finally, we explored the immune cell trafficking after spinal cord injury using vDISCO panoptic imaging. To this end, we induced a spinal cord injury in CD68-EGFP transgenic mice expressing EGFP in monocytes and macrophages43. Upon spinal cord injury, we observed an influx of CD68 GFP+ cells at the lesion site as well as a rather widespread increase in CD68 GFP+ cells around the spinal cord (Supplementary Fig. 21 and Supplementary Video 11). Indeed, only 3.5% of these phagocytes were seen in the injured spinal cord, while ~96.5% of the CD68 GFP+ cells were located in the surrounding tissues of the spinal cord including adjacent muscles, spinal cord roots and meningeal vessels, similar to sham mice (Supplementary Fig. 22). Thus, vDISCO is a powerful tool to image and quantify the trafficking of immune cells to and within the injured CNS.
Panoptic imaging of transparent adult mouse bodies from head to toe holds the promise of providing an unbiased and highly resolved view of entire organ systems in health and disease. Here, we developed a new whole-body nanobody labeling method in conjunction with whole-body tissue clearing, enabling a reliable visualization and quantification of subcellular details throughout centimeters-thick tissues of intact mouse bodies. This new panoptic imaging technology is straightforward in application and is suitable for systemic analysis of a vast range of biomedical inquiries, as demonstrated here in visualization of the neuronal projections in whole mouse bodies, the description of vascular pathways between the skull and meninges, and the visualization of remote neuronal and immune consequences of acute CNS injury models. While we developed vDISCO for panoptic imaging of the whole mouse body, it is readily applicable for individual organs, using a simplified protocol to drastically increase and stabilize the signal contrast, as we show for mouse brains imaged more than a year after boosting and clearing (Supplementary Video 12 and Supplementary Table 1).

The panoptic imaging of nervous and immune systems in intact mouse bodies was achieved owing to the massive enhancement of the signal intensity by one to two orders of magnitude via whole-body immunolabeling employing nanobodies conjugated to bright dyes in the far-red spectrum. Therefore, the strong autofluorescence of tissues in shorter-wavelength spectra, in which traditional fluorescent proteins such as GFP are excited and imaged, is avoided. Here, we primarily used Atto dyes at 647 nm, and additional Atto 594 nm for multiplexing two different cell types. In the future, usage of nanobodies conjugated to dyes that emit in the near-infrared range can further facilitate multiplexed detection of more than two targets. While the number of commercially available nanobodies is still limited, the nanoboosters used in this study can stain a broad selection of 21 different fluorescent proteins including EGFP, YFP, Venus, mCherry, and mRFP. Our method can also be used with nanobodies against endogenous proteins, as we demonstrated using anti-vimentin nanobodies labeling activated glia cells in the brain upon trauma. Currently, nanobodies are developed mainly for in vivo applications due to their small size (for example, for crossing the blood–brain barrier); therefore, they are mostly selected for...
in vivo epitopes but not for fixed tissues. We believe that our study will encourage the production of diverse nanobodies for deep-tissue immunolabeling.

Here, we used transgenic mice endogenously expressing fluorescent proteins; however, the labeling can also be achieved by rabies virus retrograde tracing, systemic adeno-associated virus injections or transplantation of genetically engineered cells (such as stem cells or adoptive transfer of immune cells). Another key advantage of vDISCO panoptic imaging is that upon staining with Atto dyes through nanoboosting, the signal becomes permanent, permitting long-term imaging sessions and future re-analysis of the same sample if needed. This also enables high-resolution imaging.

**Fig. 5 | Visualizing meningeal vessels through intact skull by vDISCO panoptic imaging.** a, A representative 4-week-old transparent Prox1-EGFP mouse head from 5 independent animals, showing the labeled vessels underneath the skull. b–e, The Prox1-EGFP mouse head, showing the brain lymphatic vessels along the sagittal sinus, pterygopalatine artery and transverse sinus (yellow, white and green arrowheads in c, d, and e, respectively). Bone structures become prominent with PI labeling (red). Single experiment. f, Three-dimensional visualization of prefrontal cortex (PFC) and olfactory bulb (OB) in a 2-month-old CX3CR1<sup>GFP</sup>/+ (cyan) and CCR2<sup>RFP</sup>/+ (red) double-transgenic mouse (similar results were observed from 3 independent double-transgenic mice). g, h, High-magnification image of marked region in f, showing CCR2<sup>RFP</sup> cells (green arrowheads) and CX3CR1<sup>GFP</sup> cells (yellow arrowheads) in meningeal vessels (MV). See also Supplementary Video 7. i, j, LysM-EGFP transgenic mice (6 months old) with MCAO versus unlesioned control, showing the infiltration of immune cells in MCAO (similar results were observed from 4 independent mice per group). LysM<sup>GFP</sup> cells are shown in red and nucleus labeling by PI in cyan. Immune cells in the meningeal vessels of the injured mice were observed (yellow arrowheads). All images were obtained by light-sheet microscopy.
Fig. 6 | Uncovering SMCs through intact skull by vDISCO panoptic imaging. a–d. A 6-month-old VEGFR3-YFP mouse head in sagittal (a,b) and axial (c,d) views, showing that injected CSF tracer (ovalbumin–Alexa647 (OV-AF647); red) fills not only meningeal vessels but also some short SMCs between skull marrow and brain surface (b,c; yellow arrowheads). Representative images, single experiment. e,f. Details of SMCs after whole mouse body PI and lectin labeling of a 3-week-old BL6 mouse processed by vDISCO: the SMCs from skull marrow to brain surface with a funnel-shaped opening are marked with dashed lines in f. Cells in the channel are indicated by white arrowheads. Similar results were observed from 5 independent mice. g,h. LysM-EGFP transgenic mice (6 months old) after MCAO (g) versus sham control (h), showing increased numbers of LysM GFP+ cells (yellow arrowheads) in the SMCs (marked with dashed lines) in MCAO. Similar results were observed from 3 independent mice per group. LysM GFP+ cells are shown in red and nucleus labeling by PI in cyan. See also Supplementary Video 10. i. Quantification of LysM GFP+ cells in the SMCs in MCAO versus sham mice (mean ± s.e.m., n = 3 animals per group; statistical significance (**P = 0.003) was assessed by two-tailed t test).
of desired body parts by confocal or two-photon microscopy. Indeed, the samples from this study are available to the scientific community upon request for further studies. We believe making vDISCO-treated samples accessible to other labs for further studies can save time and resources, reduce the number of mice used in experimental research and create more powerful datasets by allowing direct correlation of various inquiries by multiple studies from the same animals.

A current limitation is the absence of light-sheet microscope systems that could image the entire body of a mouse in one session without any displacement and turning of the sample. Construction of such new light-sheet microscopes should further decrease the time needed for data acquisition and simplify data processing for 3D reconstruction. In fact, as shown here, optics and light sources of current commercial light-sheet microscopes are sufficient to image intact mouse bodies at subcellular resolution, although the whole body has to be manually moved in the imaging chamber to achieve the scanning of different body regions. In addition, development of machine learning-based (artificial intelligence) algorithms to automatically identify and quantify cellular changes on the whole-organism scale will be important to further scale up this technology in the future.

Using panoptic imaging, we constructed a neuronal projection map for the Thy1-GFP-M transgenic line, showing subcellular details of long-range neuronal connections from the CNS to the distal extremities. Here, we visualized neuronal details in Thy1-GFP-M mice, in which only a subset of CNS neurons and peripheral nervous system neurons innervating skeletal muscles are labeled. In the future, panoptic imaging of other transgenic lines labeling a greater and more diverse subset of neurons will allow the exploration of other biological systems such as the autonomic innervation of internal organs.

In recent years, researchers have been providing more evidence on post-TBI changes that seem to form the basis of chronic complications, such as epilepsy, neuropsychiatric disorders, dementia and progressive motor decline. However, the effects of a localized brain lesion on the rest of the body have been poorly understood, mainly due to the technical challenges in studying long-range neuronal projections. Because vDISCO fluorescent signal boosting allows both imaging and quantifications of light-sheet microscopy images of transparent mouse bodies, we identified TBI-induced changes at the peripheral axonal projections innervating the skeletal musculature of the torso. These data are in line with recent work by others and us demonstrating the functional deficits in fine motor movements in mice and the presence of sensory-motor dysfunctions in TBI patients. Our data provide a possible cellular mechanism for these functional impairments, suggesting that they could arise due to degeneration of long-range axons throughout the spinal cord and peripheral nervous system. This discovery opens alternate avenues for the identification of therapeutic targets. For instance, recent literature showed that microtubule stabilization is a powerful way to block degeneration of axons in spinal cord injury. In this regard, it would be important to assess whether microtubule stabilization would also counteract axon degeneration outside of the brain and circumvent the development of chronic sensory-motor dysfunctions. Here, we also demonstrated that meningeal vessels extending between the cranium and CNS tissue and their cellular content can readily be imaged by vDISCO panoptic imaging in naïve animals and acute CNS injury models. Importantly, we observed short vascular connections between the skull and meninges both at the brain interface (recently seen also by others) and at the sagittal sinus, which were filled with immune cells following ischemic stroke injury. Most of the SMCs seem to be open both at the skull marrow and meningeal ends without being connected to the vascular system of the brain/head (Supplementary Video 9).

In the future, it will be important to identify to which meningeal compartment or compartments (subdural, subarachnoid or subpial) the SMCs are connected. In addition, due to the present lack of appropriate reporter mice, it was not possible to distinguish the type of SMCs (blood or lymphatic vessels) with increased LysM+ cells after stroke. In the future, vDISCO imaging of immune cells in a lymphatic vessel reporter mouse line with different fluorescent proteins (for example, LysM-EGFP + Prox1-RFP double reporter) could reveal more details of this new route for immune cell infiltration. As neuroinflammation is a major determinant of neuronal function and survival following CNS injuries of various etiologies, our technology should accelerate investigation of these novel immune cell infiltration routes in various CNS pathologies.

In conclusion, vDISCO enhancement and stabilization of fluorescent proteins combined with signal acquisition in far-red spectra should facilitate discovery of previously inaccessible biological information. For example, imaging and quantification of neuronal projections, vascular organizations and immune cell populations in the entire mouse body should contribute to a more comprehensive understanding of the initiation, progression and extent of neurological conditions.

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Any methods, additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at https://doi.org/10.1038/s41593-018-0301-3.

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Methods

Animals. We used the following mixed-gender animals in the study: CX3CR1-YFP+, CD68-EGFP, and Thy1-YFP-H−. Prox1-EGFP (Tg(Prox1-EGFP);K221Gsat/Mmucd; Mutant Mouse Resource and Research Centers strain code 031006-UCD), VEGFR3-YFP, CXCR4-EGFP, xCR2-YFP−. (B6.129-C–Ctbr2m1.1Ilc/J; Jackson Laboratory strain code 017586), LysM-EGFP (Ly2tm1.1Iox; Mouse Genome Informatics strain code 2654931), CD68-EGFP (C57BL/6-tdG-ln(CD68-EGFP)1Drg/j; Jackson Laboratory strain code 026827), C57BL/6J, and NMRI nu/nu mice. The animals were housed under a 12:12 h light/dark cycle. The animal experiments were conducted according to institutional guidelines (Klinikum der Universität München/Ludwig Maximilian University of Munich), after approval of the ethical review board of the government of Upper Bavaria (Regierung von Oberbayern, Munich, Germany) and the Animal Experiments Council under the Danish Ministry of Environment and Food (2015-15-2021-00553), and in accordance with the European directive 2010/63/EU for animal research. Ages of the animals are indicated in the figure legends. All data are reported according to the Animal Research: Reporting of In Vivo Experiments (ARRIVE) criteria. Sample sizes were chosen on the basis of prior experience with similar models and are specified in figure legends. Within each strain, animals were randomly selected. Animals that resulted negative for the expression of fluorescent proteins by genotyping were excluded from the study. No statistical methods were used to determine sample sizes, but our sample sizes are similar to those reported in previous publications.

TBI. TBI was performed using a controlled cortical impact device (Leica Benchmark Stereotaxic Impactor, 39463923). At 30 min before surgery, we administered carprofen (4 mg per kg body mass) and butorphanol (0.05 mg per kg) to the animals via subcutaneous injection. Then, anesthesia was induced in animals with 4% isoflurane in a N2O/O2 (70%/30%) mixture and afterwards maintained with 1.5% isoflurane in the same mixture for the whole surgery. As soon as the animals did not show any pedal reflex, they were placed in the stereotaxic frame by the zygomatic arch, with the head slightly tilted to form an angle of 20° in relation to the body. The head and neck regions were shaved to expose the neck muscles, which were bluntly dissected to expose the cisterna magna. Cannulas composed of a dental needle (SOPIRA Carpule 30 G 0.3×10 mm) were inserted into the cisterna magna with the aid of an injection pump (LEGAZIO 130 Syringe pump; KD Scientific, 788130), 10 μl of CSF tracer was injected into the cisterna magna at a rate of 1 μl min−1. At the end of the injection, CSF tracer was allowed to circulate in the subarachnoid and paravascular spaces for 1 h. Mice were then transcardially perfused according to the ‘Perfusion and tissue preparation’ section below.

Perfusion and tissue preparation. Mice were deeply anesthetized using a combination of midazolam, medetomidine and fentanyl (MMF; 1 ml per 100 g body mass for mice; intraperitoneal). As soon as the animals did not show any pedal reflex, they were intracardially perfused with heparinized 0.1 M PBS (pH 7.4; 0.05% sodium azide). Mice were deeply anesthetized by intraperitoneal injection of a combination of midazolam (5 mg per kg), medetomidine (0.5 mg per kg) and fentanyl (0.05 mg per kg). The mid-thoracic spinal cord of anesthetized CD68-EGFP mice was surgically exposed by a dorsal laminectomy as previously described. A hemisection of the spinal cord was performed using fine-tip surgical scissors (Fine Science Tools 15000-08 spring scissors, 2.5 mm cutting edge). For sham surgery, the dorsal laminae of mid-thoracic vertebrae were surgically exposed in anesthetized CD68-EGFP mice. After spinal cord injury or sham surgery, muscle tissue and skin were sutured with a surgical thread (Ethilon suture 6-0, 667 H) and animals were allowed to recover on a heating pad. After 48 h, mice were transcardially perfused according to the ‘Perfusion and tissue preparation’ section below.

Spinal cord injury model. Mice were deeply anesthetized by intraperitoneal injection of a combination of midazolam (5 mg per kg), medetomidine (0.5 mg per kg) and fentanyl (0.05 mg per kg). The mid-thoracic spinal cord of anesthetized CD68-EGFP mice was surgically exposed by a dorsal laminectomy as previously described. A hemisection of the spinal cord was performed using fine-tip surgical scissors (Fine Science Tools 15000-08 spring scissors, 2.5 mm cutting edge). For sham surgery, the dorsal laminae of mid-thoracic vertebrae were surgically exposed in anesthetized CD68-EGFP mice. After spinal cord injury or sham surgery, muscle tissue and skin were sutured with a surgical thread (Ethilon suture 6-0, 667 H) and animals were allowed to recover on a heating pad. After 48 h, mice were transcardially perfused according to the ‘Perfusion and tissue preparation’ section below.

Clearing of unboosted samples. For the quantification of fluorescence signal after clearing, without boosting, we followed the uDISCO passive clearing protocol as described by Pan et al.1. In brief, dissected brains were placed in 5 ml tubes (Eppendorf, 0030 119.401) and covered with 4.5 ml of clearing solution. All incubation steps were performed in a fume hood with gentle shaking or rotation, with the samples covered with aluminum foil to keep them in darkness. To clear the samples, we first pre-warmed them in a gradient of tert-butanol (Sigma-Aldrich, 360538): 30 vol%, 50 vol%, 70 vol%, 80 vol%, 90 vol%, 96 vol% (in distilled water), and 100 vol% at 35°C for 12 h each step followed by immersion in dichloromethane (Sigma-Aldrich, 270997) for 45–60 min at room temperature and finally incubation with the refractive index matching solution BABB-D15, containing 15 parts BABB (benzyl alcohol + benzyl benzoate 1:2; Sigma-Aldrich, 21122 and W213802), 1 part diphenyl ether (Alfa Aesar, A15791) and 0.4 vol% vitamin E (2,6,α-tocopherol; Alfa Aesar, A17039), for at least 6 h at room temperature until transparency was achieved.

Clearing of boosted samples. For the quantification of fluorescence signal after clearing, with boosting, we followed the dDISCO active clearing protocol as described by Leica Benchmark Stereotaxic Impactor, 39463923). At 30 min before surgery, we administered carprofen (4 mg per kg) once per day for 4 d and sacrificed at >1 month post-injury by transcardial perfusion according to the ‘Perfusion and tissue preparation’ section below.

Cisterna magna injection for meningeval vessel labeling. For the cisterna magna injections, mice (VEGFR3-YPF, 6 months old) were anesthetized with a mixture of 60% isoflurane and 40% O2 (0.05 mg per kg) to the animals via subcutaneous injection. After toe pinch reflexes ceased, mice were fixed in a stereotaxic frame by the zygomatic arch, with the head slightly tilted to form an angle of 20° in relation to the body. The head and neck regions were shaved to expose the neck muscles, which were bluntly dissected to expose the cisterna magna. Cannulas composed of a dental needle (SOPIRA Carpule 30 G 0.3×12 mm; Kelzer, and stainless steel tubing (outside diameter 0.024 in ID (interior diameter); Scandicraft, PE10-CL-500) were used to perform the cisterna magna injections. A cannula filled with CSF tracer (2% ovalubinin 45 kDa conjugated to Alexa Fluor 647 (Thermo Fisher Scientific, O37487), diluted in artificial CSF (126 mM NaCl, 2.5 mM KCl, 1.25 mM NaH2PO4, 2 mM MgSO4, 2 mM CaCl2, 10 mM glucose, 26 mM NaHCO3; pH 7.4 when gassed with 95% O2 and 5% CO2) was inserted into the cisterna magna. With the aid of an injection pump (LEGAZIO 130 Syringe pump; KD Scientific, 788130), 10 μl of CSF tracer was injected into the cisterna magna at a rate of 1 μl min−1. At the end of the injection, CSF tracer was allowed to circulate in the subarachnoid and paravascular spaces for 1 h. Mice were then transcardially perfused according to the ‘Perfusion and tissue preparation’ section below.

Validation of nanobooters for dDISCO. Post-fixed brains from fluorescent protein-expressing mice (for example, Thy1-GFP-M) were cut into 400 μm slices using a vibratome (Virto +.1200; sliding knife, 750 μm). The slices were imaged before and after a Z-axis Axiostar EM535/Sycop fluorescence stereomicroscope to make sure that the samples have fluorescent protein expression. All the following steps were done with gentle shaking. The slices were first incubated for 3 h in a 24 well Multicell (Falcon,
tube (Falcon, 352070) filled with the same permeabilization solution containing β-cyclodextrin (Sigma-Aldrich, 332615), 0.2% l-proline (Sigma-Aldrich, 441562) and 0.05% sodium azide (Sigma-Aldrich, 71290) in 0.1 M PBS. Subsequently, they were incubated overnight at 37°C with the same permeabilization solution, and the nanobooster of interest was added with dilution 1:500 considering final volume = 500 μL. The multwell was covered with aluminum foil to keep the slices in dark and was sealed well with parafilm. To prevent the solution from evaporation, we kept as negative control where there was no adding of nanoboster. After the incubation, slices were washed with the washing solution (1.5% goat serum, 0.5% Triton X-100, 0.05% sodium azide in 0.1 M PBS) at room temperature for 15 min 4 times and then washed with 0.1 M PBS at room temperature for 10 min 3 times. Then the slices were imaged again after staining with the AxioZoom microscope to check if the staining worked. In the same way, the slices were moved to 3 DISCO protocol: first they were incubated at room temperature in the following gradient of tetrahydrofuran (THF; Sigma-Aldrich, 186562) in distilled water (45 min for each step): 50 vol% THF, 70 vol% THF, 80 vol% THF, 100 vol% THF and 100 vol% THF, after dehydration samples were incubated for 15 min in dichloromethane, and finally in BABB until transparency. Finally, the slices were imaged again with the AxioZoom microscope to make sure that the staining worked even after clearing. For more details of the AxioZoom imaging setup, see the ‘Fluorescence stereomicroscopy imaging’ section below.

vDISCO whole-body immunostaining, PI labeling and clearing. In order to remove remaining blood and heme after PFA perfusion, and to decalify the bones, the animals were subjected to perfusion with decalcification solution and decalcification solution before immunostaining. The decalcification solution was made with 25–30 vol% dilution of CUBIC reagent 1 in 0.1 M PBS. CUBIC reagent 1 was prepared with 25 wt% urea (Carl Roth, 3941.3), 25 wt% N,N,N’,N’-tetrakis (2-hydroxypropyl)ethylenediamine (Sigma-Aldrich, 122262) and 15 wt% Triton X-100 in 0.1 M PBS. The decalcification solution consisted of 10 wt/vol% EDTA (Carl Roth, 170229685) in 0.1 M PBS, adjusting the pH to 8–9 with sodium hydroxide (Sigma-Aldrich, 71687).

The solutions for the immunolabeling pipeline were pumped inside the body of the animal by transcardial-circulatory perfusion, exploiting the same entry point hole into the heart created during the PBS and PFA perfusion step (see above) and following the procedure described by Pan et al. In brief, the mouse body was placed in a 300 ml glass chamber (Omnilab, 5163279) filled with 250–300 ml of appropriate solution, which covered the body completely. Next, the transcardial-circulatory system was established involving a peristaltic pump (ISMATEC, REGLO Digital MS-4/8 ISM 834; reference tubing, SC0266), keeping the pressure up to 26–28 °C. With this setting, the animals were perfused for 6 d at 160–230 mmHg (45–60 r.p.m.). One channel from the pump, made by a single reference tube, was set for circulation of the solution through the heart into the vasculature: one ending of the tube was connected to the tip of a syringe (cut with extended PVC tubing. The pumping channel was set in the standard way as described above in this section for both nanobody and antibody experimental groups. The only exception was that for the antibody group, in the immunostaining step, we replaced the nanobooster with Alexa647-conjugated anti-GFP antibody (Invitrogen, A31852) with the same concentration (see above).

Pumping pressure measurement. A Kkmoon Digital Manometer Pressure Gauge Manometer (HT-1891) was used to measure the pressure. To achieve one-to-one signal connection with the pump, a cable was connected to the two-channel connector (B.Braun Disconet C Dreieinwahlen, 169494) was inserted into the pumping channel, and the second channel was further connected to the manometer with extended PVC tubing. The pumping channel was set in the standard way as described including a transcardiac perfusion needle. First, the switch on the two-head connector was set without connecting the measuring tubing, and pumping was performed until all the air was released and the pressure was measured. Subsequently, the switch was set to connect the measuring tubing while pumping was continued, and the pressure was measured when the manometer readouts stabilized. Three measurements were made at different time points including morning, afternoon and evening to exclude the variation of room temperature, and the data were quantified by averaging.

Comparison of high-pressure versus low-pressure perfusion with methylene blue. The diffusion of methylene blue (Sigma-Aldrich, M9140), perfused into the body of the animals, was used to compare the efficiency of high-pressure versus low-pressure perfusion (Supplementary Fig. 8). For this purpose, we used 3 Bl6 animals (4 months old) per experimental group: post-fixed whole bodies were perfused for 15 min with 0.05% methylene blue in 0.1 M PBS, at either 50 r.p.m. (~180 mmHg) for the high-pressure group or 25 r.p.m. (~70 mmHg) for the low-pressure group. The same pump setting explained in the section ‘vDISCO whole-body immunostaining, PI labeling and clearing’ was used, including the usage of superfusate to fix the needle in place inside the hole in the heart for 20 min until the superfusate was ~20–35 μg in 250 ml (0.08–0.14 mg·mL⁻¹), 1,700 in dilution (stock concentration 0.5–1 mg·mL⁻¹; the amount of nanoboster was adjusted depending on the expected presence of fluorescent protein in the mouse body) and 290 μL of PI (stock = 10 mg·mL⁻¹) was added with working the iDISCO protocol (3 DISCO protocol (3 DISCO protocol). Next, we removed protocol (Supplementary Fig. 3a–d). This was done performing the antibody validation step, which tested the compatibility of the antibody with the methanol pretreatment required by iDISCO+. To do so, we followed the original iDISCO+ publication 1 and the latest protocol updates from https://idisco.info. Briefly, post-fixed brains from Thy1-GFP-M and CXCR1(+/-)
mice (2–3 months old) were cut into 600 μm slices with a vibratome. First, the slices were incubated overnight in 100% methanol (Roth, 4627.6) at room temperature; then, after rehydrating and washing with 0.1 M PBS, the iDISCO+ immunostaining protocol was done as visually using 30% as permeabilization time, 6 h as blocking time and overnight as immunolabeling time, adding 1:80 of Alexa467-conjugated anti-GFP antibody (already used in the previous section) in 500 μl of illumination solution. Non-methylene-treated slices were used as positive controls. In the end, the slices were imaged using the Zeiss AxioZoom stereo microscope (see ‘Fluorescence stereomicroscopy imaging’ section below).

iDISCO+ on dissected brains. After having verified that the previously mentioned conventional anti-GFP antibody is compatible with iDISCO+, we performed comparison of staining efficiency between iDISCO with nanobody and iDISCO+ with conventional antibody (Supplementary Fig. 5e–i). For this purpose, 3 CX3CR1<sup>+</sup> post-fixed brains (6–7 months old) were used per experimental group. The staining procedure was divided into the iDISCO + protocol was done as normally using 3 h as permeabilization time, 6 h as blocking time, then, after rehydrating and washing with 0.1 M PBS, the iDISCO + protocol was adjusted considering the size of a whole body compared with a dissected brain (for example, during methanol pretreatment on the animals followed by sequential active perfusion iDISCO with conventional antibody, (4) high-pressure active perfusion vDISCO with conventional antibody, and (5) high-pressure active perfusion iDISCO+ with nanobody. Group 1 was processed using the above-described vDISCO whole-body immunostaining and clearing method; the group 2 was processed following all the steps of the vDISCO whole-body immunostaining and clearing protocol described above, but replacing all the original steps of immunostaining by the same vDISCO whole-body immunostaining and clearing protocol, but replacing the nanobuster with Alexa467-conjugated anti-GFP antibody at the same concentration as the nanobuster during the staining step; groups 4 and 5 were processed performing the iDISCO+ methanol pretreatment on the brains followed by sequential perfusion with all the necessary ingredients (except the clearing with Alexa467) to iDISCO+; the size of a whole body compared with a dissected brain (for example, during the methanol dehydration, each step was increased to 5 h versus 1 h for dissected brains). The antibody used for group 4 is the Alexa647-conjugated anti-GFP antibody mentioned above.

Previously published whole-body clearing methods for comparison. For PARS<sub>+</sub> antibody mentioned above.

Different whole-body labeling protocols comparison. To show that vDISCO with nanobody is the only method that can efficiently boost the GFP signal in mouse whole body samples, we performed the following comparison using 3 GFP-M animals (2–3 months old) per experimental group: (1) high-pressure active perfusion vDISCO with nanobody, (2) passive incubation vDISCO with nanobody, (3) high-pressure active perfusion vDISCO+ with conventional antibody, (4) high-pressure active perfusion iDISCO+ with conventional antibody, and (5) high-pressure active perfusion iDISCO+ with nanobody. Group 1 was processed using the above-described vDISCO whole-body immunostaining and clearing method; the group 2 was processed following all the steps of the vDISCO whole-body immunostaining and clearing protocol described above, but replacing all the original steps of perfusion by passive incubation; group 3 was processed using the same vDISCO whole-body immunostaining and clearing protocol, but replacing the nanoboster with Alexa467-conjugated anti-GFP antibody at the same concentration as the nanoboster during the staining step; groups 4 and 5 were processed performing the iDISCO+ methanol pretreatment on the brains followed by sequential perfusion with all the necessary ingredients (except the clearing with Alexa467) to iDISCO+; the size of a whole body compared with a dissected brain (for example, during the methanol dehydration, each step was increased to 5 h versus 1 h for dissected brains). The antibody used for group 4 is the Alexa647-conjugated anti-GFP antibody mentioned above.

Light-sheet microscopy imaging. Single-plane illuminated (light-sheet) image stacks were acquired using an Ultramicroscope II (LaVision BioTec), featuring an axial resolution of 4 μm with the following filter sets: ex 470/40 nm, em 535/50 nm; ex 545/25 nm, em 605/70 nm; ex 560/30 nm, em 609/54 nm; ex 580/25 nm, em 625/30 nm with the following filter sets: ex 470/40 nm, em 535/50 nm; ex 545/25 nm, em 605/70 nm; ex 560/30 nm, em 609/54 nm; ex 580/25 nm, em 625/30 nm; ex 640/40 nm, em 690/50 nm. For low-magnification whole-body imaging of the Thy1<sup>+</sup> mice on a 5.6 mm diameter yield at xenon light source, we used a x2 objective with a zoom range from 0.63 up to 2.5 cm, covering the entire width of the mouse body. Tile scans with 60% overlap along the longitudinal y axis of the mouse body were obtained from ventral and dorsal surfaces up to 13 mm in depth, covering the entire volume of the body used a z-step of 8 μm. Exposure time was 120 ms, laser power was adjusted depending on the intensity of the fluorescent signal (in order to never reach the saturation) and the light-sheet width was kept at maximum. After tile imaging of the sample within the entire field of view, already scanned regions were cut using a thin motorized diamond blade (0.2 mm; Dremel 8200) for further imaging. After low-magnification imaging, the whole body, or a forelimb of the Thy1<sup>+</sup> mouse was imaged with a x2 objective (Olympus MVPLAPO2XC/0.5 NA (WD = 6 mm)) coupled with the same Olympus MVX10 zoom body at zoom magnification x1.6. Moreover, the same x2 objective with different zooms was used to perform high-magnification imaging of specific body regions (for example, the back of the animal at the level of lumbar vertebra, the inguinal lymph node area or the head). Individual organs (e.g. brain, lungs, intestine and thyroid) were imaged using a high-magnification objectives: x2 objective (Olympus MVPLAPO2XC/0.5 NA (WD = 6 mm)) coupled with the same Olympus MVX10 zoom body, x4 objective (Olympus XFLUOR X4 corrected/0.28 NA (WD = 10 mm)), x25 objective (Olympus XLPLN X25/0.95 NA (WD = 4 mm)) and x20 objective (Zeiss x20 Crl Plan-Neofluar/0.1 NA (WD 5.6 mm) coupled to an Olympus revolving zoom body (TU-VCAT) kept at x1. High-magnification tile scans were acquired using 8–30% overlap, and the light-sheet width was reduced to obtain maximum illumination in the field. For the specific imaging settings used in each figure panel, see Supplementary Table 2.

Fluorescence stereomicroscopy imaging. For images in Supplementary Fig. 3, brain slices were kept immersed in PBS. For nanobody validation, cleared slices were put on a glass lid and kept immersed in BABB. The imaging was performed with Zeiss AxiosZoom EMSy/ScyCoP3 fluorescence stereomicroscopy using Zen 2 software (v.2.0.0.0.0; Carl Zeiss AG) using a x1 long-working-distance air objective lens (Plan Z x1, 0.25 NA, WD = 56 mm; Supplementary Table 2). For the comparison of YFP-H mouse tissues under high-pressure pumping versus low-pressure pumping in iDISCO+ (Supplementary Fig. 9), the mouse brains were cut into 1 mm slices using a vibratome and imaged with magnification ×7 for exposure to BABB. To make sure that the imaging region was immersed in BABB. Closing or sealing of the Petri dish was not necessary because BABB has a high evaporation point and doesn't dry out for several days. The only precaution taken was to check the resistance of the dish against BABB, by testing that the glue sealing the glass to the dish and the plastic parts were not dissolved after several hours of exposure to BABB.

The imaging was done using a x40 oil-immersion objective lens (Zeiss, ECP-Plan-Neofluar X40/1.30 oil DIC M27, 1.3 NA, WD = 0.21 mm) and a x25 water-immersion long-working-distance objective lens (Leica, 0.95 NA, WD = 2.5 mm) mounted on a custom mounting thread. See Supplementary Table 2.

Reconstructions of whole-mouse body scans. We acquired light-sheet microscope stacks using ImSpector (v.5.295, LaVision BioTec) as 16-bit grayscale TIFF images for each channel separately. In each pair of neighboring stacks, alignment was done by manually selecting 3 to 4 anatomic landmarks from the overlapping regions, and then the stitching was done sequentially with the Scope Fusion module of the Vision4D (v.2.12.6 x 64, Arisvox) software. Landmarks were mainly chosen from the skull bones or occasionally from the neuronal structures on the basis of visual inspection of the anatomical features. After completing the 3D reconstructions, the data visualization was done with Amira (v.6.3.0), FEl Visualization Sciences Group, Imaris (v.9.1, Bitplane) and Vision4D in both volumetric and maximum-intensity projection color mapping. Depth coding was done using the Temporal-Color Code plugin in Fiji (ImageJ2, v.1.51, https://fiji.sc/).

Image processing. Processing, data analysis, 3D rendering and video generation for the rest of the data were done on an HP workstation Z840, with 8 core Xeon processor, 196 GB RAM, and Nvidia Quadro K5000 graphics card and HP workstation Z840 dual Xenon 256 GB DDR4 RAM, nVidia Quadro M5000 8GB graphic card. We used Imaris, Amira and Fiji (ImageJ2) for 3D and 2D image visualization. Tile scans were stitched by Fiji's stitching plugin49. Stitched images were saved in TIFF format and optionally compressed in LZW format to enable fast processing. We removed tiles with acquisition errors using Fiji’s TrakEM2 plugin and ImLigh2Library. In case of tiling errors in the z dimension, we used TeraStitcher (v.1.10; https://abria.github.io/TeraStitcher/) with its default settings to globally optimize the tiled volumes and reconstruct the entire dataset. To increase the quality of the images, we used the following functions in Fiji: to enhance the contrast of microglia cells (Fig. 1n and Supplementary Fig. 2a–f), we used ‘Enhance Local Contrast (CLAHE)’; to equalize the images (Fig. 4ab and Supplementary Fig. 3g–i), we used ‘Pseudo Flat Field Correction’, to enhance the contrast over the background of the axonal terminals in Fig. 6cd, we used the custom-made macro for Fiji that we utilized to generate the pre-processed data for NeuroGPS-Tree (see the section ‘Neuron tracing below’).

Neuron tracing. For automated neuron tracing in our light-sheet datasets obtained with Zeiss x20 Crl Plan-Neofluar/0.1 NA (WD = 5.6 mm) objective, we used the NeuroGPS-Tree algorithm2. NeuroGPS-Tree was developed for tracing relatively small volumes of confocal microscopy data; therefore, we initially reduced the file size to under 1 GB (approximate maximum data size for NeuroGPS-Tree computation) by using the Fiji scale function. Due to high signal-to-noise ratio discrepancies across different animals and conditions, we then pre-processed the data with a custom-made macro in Fiji (available upon request), which consisted of background removal, pseudo-background correction, noise filtering...
and sharpening. Next, the pre-processed data were loaded and analyzed first in NeuroGPS for soma detection and later in NeuroGPS-Tree for neurite detection (both steps are part of the same algorithm package). The best parameters for soma and neurite detection were chosen following the original publication. To quantify the features (such as the number of neurites per cell and the number of somas with at least one neurite) of these detected neuronal cells, we used Amira software: we chose the 10 neurons with the biggest file size per group and analyzed them using the Spatial Graph Statistics function of the software.

Quantifications. All the details of imaging settings used to acquire the data for the quantifications described in this section are available in Supplementary Table 2.

Analysis of fluorescence signal profiles from light-sheet images. The fluorescence signal profiles from each channel (excitation 470 nm, 560 nm and 647 nm) were plotted in the same z-stack and normalized as percentage over the maximum peak using Fiji (Supplementary Fig. 1). To create the VDSCO+ and unboosted protocols and consequently the reduction of the background and the improvement of the signal over background ratio in the far-red channel, we analyzed neurons and axonal bundles expressing GFP imaged with excitation at 470 nm, and neurons and axonal bundles labeled with anti-GFP nanobody conjugated with Atto647N imaged with excitation at 640 nm at the same anatomic region. First, the scans were taken with the light-sheet microscope using the x4 objective (Olympus XFLUOR x4 corrected/0.28 NA (WD = 10 mm)). Then, the signal profiles were analyzed in Fiji and measured from a defined straight line covering the neuronal structure and surrounding tissue background, and the normalized plots of the signal profile (Fig. 1i) were calculated by normalizing the plots of neuronal structures obtained as described above of all the fluorescence levels of the respective surrounding background. Each experimental group consisted of 3 animals, and for each animal at the same anatomic region we plotted 3 profiles.

The same analysis of signal profile was used to compare the labeling efficiency of VDSCO with nanobody and VDSCO+ with antibody (Supplementary Fig. 3g–i): in this case, the signal of the confocal microscopy. Each experimental group consisted of 3 animals, and for each animal at the same anatomic region we plotted 3 profiles from 3 different microglia. The same analysis of signal profile was used to compare different clearing methods in Supplementary Fig. 12.

Fluorescence level. Fluorescence level quantification was expressed as a signal-to-background ratio and was calculated using Fiji as a ratio of signal over background at the following time points: 0, 2, 3, 4.5, 5.5, 12 and 18 months after nanoboosting (Fig. 1n,o). Each x4 light-sheet microscopy brain scan was taken with the same imaging parameters, and an image in TIF format of the same anatomic region for all the samples was quantified. The mean value of the background for each image was obtained by averaging the background values of 12–40 regions from equally sized areas of the image in regions of the sample without signal. To calculate the mean value of the signal for each image, we used the threshold function of the software: the threshold was adjusted to consider the fluorescence signal visible in the image. After adjusting the threshold, only the sharp signal from specific cellular structures was analyzed per image. To this end, we used Fiji’s ‘analyze particles’ function to measure the signal intensity only of particles sized between 5–10 and 100–150 pixels (visible fluorescent cells) and calculated the average value from all the particles. Next, this value was divided by the mean value of the background of the respective image, obtaining the fluorescence level over the background. The corresponding images, visually showing the preservation of signal over time in relation to the respective fluorescence levels, were processed using the Enhance Local Contrast (CLAHE) function in Fiji to increase the contrast of fluorescent cells in the tissue.

Quantification of deep-tissue staining efficiency of nanobody versus antibody. To quantify the depth of staining and therefore to compare whole organ staining efficiency of nanobody versus antibody labeling (Supplementary Fig. 2), we used CX3CR1GFP+ animals (3–7 months old), 3 whole brains per experimental group. The tile scans of the brains were taken with light-sheet microscopy in axial view using the x4 objective (Olympus XFLUOR x4 corrected/0.28 NA (WD = 10 mm)). After stitching, we chose for the z-plane the same anatomic structure for each animal along the white matter. To compare the original publication to our double imaging of whole brains were analyzed in Fiji; the distance of staining depth was quantified using the ‘line’ function in the z-plane at the greatest width of the brain, measuring from the surface until the last visible boosted microglia. Then, for each image, this staining depth value was normalized over the total thickness of the brain at the considered z-level, measuring a line drawn from the same point of the surface until the axial middle line of the brains.

Quantification of whole-body labeling efficiency of different whole-body labeling protocols. To quantify the rate of staining of the different whole-body labeling protocols in Supplementary Fig. 11, the tile scans of 3 whole heads per method were taken with light-sheet microscopy in the x2 objective (Olympus XFLUOR x2 corrected/0.28 NA (WD = 10 mm)) and using the ‘freehand selections’ function in Fiji, following the brain outline. Statistical comparison was performed considering each set of corresponding regions from the different samples separately.

ClearMap. To quantify microglia distribution, we used ClearMap. Since the script was originally developed for quantification of the cFos+ cells, to comply with the offered method, we did the following pre-processing steps on our microglia data used before ClearMap:

- Background equalization to homogenize intensity distribution and appearance of the microglia cells over different regions of the brain, using pseudo-flat-field correction function from Bio-Voxeel toolbox.
- Convolved background removal, to remove all particles bigger than relevant cells. This was done with the median option in the Bio-Voxeel toolbox.
- Two-dimensional median filter to remove remaining noise after background removal. The filter radius was chosen to ensure the removal of all particles smaller than microglia cells.
- Unshapen mask to amplify the high-frequency components of a signal and increase overall accuracy of the cell detection algorithm of ClearMap.

After pre-processing, ClearMap was applied by following the original publication and considering the threshold levels that we obtained from the pre-processing steps. As soon as the quantification was completed, the data was exported as an Excel file for further analysis. For example, the cellular density per brain region was obtained considering the absolute number of cells detected by ClearMap and the volume of that specific brain region, which was calculated using a custom script (available upon request) based on ClearMap (Elastix registration).

Quantification of peripheral neuronal degeneration in acute brain injury. Peripheral neuronal degeneration in TBI animals versus unlesioned control animals was done in the Thy1-GFP-M line and was assessed considering the complexity of axonal ramifications that projected from the left (left = contralateral side) cervical and thoracic vertebra to the left muscles of the back at the level of the torso of the animal. To quantify density, the following procedure was applied. The thickness of axonal terminations, we used the ‘point tool’ function and the same ‘ROI manager tool’ to detect the cell numbers separately, the border of the spinal cord was marked by using the ‘free hand line’ function in Fiji. Then, the analysis was done over the maximum intensity projections of the tile scans with Fiji software. To measure the length of the ramifications, we used the ‘free hand line’ function and the ‘ROI manager tool’ of Fiji to record all the traced axonal ramifications of interest, which were calculated from a manually segmented axonal density. For each spinal cord, the total length of all of the recorded ramifications with the ‘measure’ function of Fiji. To count the nerve terminals, we used the ‘point tool’ function and the same ‘ROI manager tool’ to record all the visible axonal endpoints protruding from the traced ramifications. The analysis was performed over 2–4 branches from the same anatomic region for each animal, in 6 animals (3–6 months old) per experimental group. The same quantification procedure was followed to analyze the complexity at the equivalent ipsilateral (right side from dorsal view) side of the animals, in 6 and 5 animals for the TBI and unlesioned groups, respectively.

Quantification of CGG8 GEP+ cells with spinal cord injury. The spine areas of interest from CD68+GEF pig line mice with spinal cord injury or sham surgery and from naïve animals were imaged with light-sheet microscopy using the x2 objective described in the ‘Light-sheet microscopy imaging’ section with a total magnification of x3.2 and with 2 x 3 tile z-stacks. After stitching, the entire z-stacks of scans were loaded in Fiji software with the ‘use virtual stack’ mode. To segment the spinal cord tissue from surrounding tissue, including bone and muscles, and count the cell numbers separately, the border of the spinal cord was marked by using the ‘free hand line’ function in Fiji as a manual selection. These manual selections were done in every 6–9 image slices and were recorded by the ‘ROI manager tool’ function. The complete selections for the entire z-stacks were then achieved by using the ‘interpolate ROIs’ function and were saved as zip files. Then, those z-stacks of scans were loaded in Fiji software again, without selecting the ‘use virtual stack’ mode. By loading the complete selections generated from former steps respectively, each spinal cord was segmented by using a custom-made macro with the ‘clear outside function’ and saved as a new ‘image sequence’. To maintain the integrity of the raw data, one should not save the changes when closing the workspace. After running the clearMap (clearMap) and the volume of that specific brain region, which was calculated using a custom script (available upon request) based on ClearMap (Elastix registration).
the ‘clear’ function. All the custom-made macros are available upon request. Next, the segmented image stacks of spinal cord or surrounding tissue were loaded with Imaris respectively, and the ‘surface’ function was used to segment the CD68 GFP+ cells. By generating the ‘surface’ selection, the ‘statistic’ function was available to count the number of cells in the region of interest.

Quantification of short SMCs with MCAO. The transparent skulls from LysM-EGFP line mice with MCAO or sham were imaged with intact heads from the sagittal view with light-sheet microscopy using the ×2 objective described in the ‘Light-sheet microscopy imaging’ section with a total magnification of ×32 and with 2 × 4 tile z-stacks. After stitching, the entire z-stacks of scans were loaded in Imaris, and the length, width and cell numbers in the SMCs were manually calculated by using the ‘measure-line’ function. Then, the length and width of the channels were corrected by a shrinkage rate of DISCO clearing.

Statistical analysis. Data collection and analysis were not performed blind to the conditions of the experiments. Data distribution was assumed to be normal, but this was not formally tested. Data are presented as mean ± s.d. except for in Fig.6 and Supplementary Fig. 22, where they are presented as mean ± s.e.m. Sample sizes are indicated in the figure legends. Statistical analysis was performed using Prism GraphPad software v.6 with 95% confidence interval. P values were calculated using two-tailed unpaired t test to compare data between two groups. P values were calculated using one-way ANOVA followed by Dunnett’s post hoc test to compare data in Supplementary Figs. 4 and 11. For CD68-EGFP quantification in Supplementary Fig. 22, P values were calculated using one-way ANOVA followed by Tukey multiple comparison test to compare data between SCI (spinal cord injury), sham and naïve groups. A P value of <0.05 was considered statistically significant. In ANOVA analysis the reported P values are adjusted in GraphPad prism to account for multiple comparisons. The protocols described in this study were replicated successfully more than five times in independent experiments, and they were also reproduced at least by three different operators. See also the Life Sciences Reporting Summary.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Code availability. The custom codes used to analyze the data in this study are available from the corresponding author upon request.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Reporting Summary

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Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

- **n/a** Confirmed
- [x] The exact sample size \((n)\) for each experimental group/condition, given as a discrete number and unit of measurement
- [x] An indication of whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
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- [x] A description of all covariates tested
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- [x] Clearly defined error bars
  
  *State explicitly what error bars represent (e.g. SD, SE, CI)*

Our web collection on statistics for biologists may be useful.

Software and code

Policy information about availability of computer code

- **Data collection**
  - The commercially available softwares used in this study to collect, stitch and visualize the data are: ImSpector (Version 5.295, LaVision BioTec GmbH), Vision4D (Version 2.12.6 x64, Arivis AG), Amira (Version 6.3.0, FEI Visualization Sciences Group), Imaris (Version 9.1, Bitplane AG), Zen 2 (Version 10.0.4.910, Carl Zeiss AG), Zen 2 (Version 2.0.0.0, Carl Zeiss AG).

- **Data analysis**
  - The commercially available softwares used in this study to analyze the data are: Amira (Version 6.3.0, FEI Visualization Sciences Group), Imaris (Version 9.1, Bitplane AG), GraphPad Prism (Version 6, GraphPad Software Inc.).
  - The already publicly or published codes/softwares used in this study to stitch and analyze the data are: Fiji (Version 1.51, https://fiji.sc/), TeraStitcher (Version 1.10, https://abria.github.io/TeraStitcher/), NeuroGPS-Tree (https://www.nature.com/articles/nmeth.3662), ClearMap (https://www.sciencedirect.com/science/article/pii/S0092867416305554).
  - All the custom-made codes: the code to pre-process the data for NeuroGPS-Tree in Fiji, the custom-made macros for Fiji to remove region of interests in images, the custom-script to calculate the volume of specific brain regions for the cellular density are available upon request.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.
Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The data of this study are available from the corresponding author upon reasonable request.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Data exclusions | Replication | Randomization | Blinding |
|-------------|-----------------|-------------|---------------|---------|
| Sample sizes were chosen based on prior experience with similar models. Sample sizes are specified in figure legends. | Exclusion criteria were pre-established. Animals that resulted negative for the expression of fluorescent proteins by genotyping were excluded from the study. For MCAO experiments, exclusion criteria were as follows: insufficient MCA occlusion (a reduction in blood flow to 15% of the baseline value) and blood flow recovery >80% within 10 min of reperfusion. | The indication of how many times each experiment was repeated independently showing similar results is written in the corresponding figure legend. Overall, the protocols described in this study were replicated successfully more than 5 times in independent experiments and they were also reproduced at least by 3 different operators. | The animals used in this study were selected for each experiment based on their genetic background (wt or fluorescent transgenes). Within each strain, animals were randomly selected. | Littermates were housed in the same cage. The investigator was not blinded to group allocation for data analysis as this was obvious during the experiments (e.g. stroke vs sham or TBI vs unlesioned). |

Reporting for specific materials, systems and methods

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
|     | Unique biological materials |
|     | Antibodies |
|     | Eukaryotic cell lines |
|     | Palaeontology |
|     | Animals and other organisms |
|     | Human research participants |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
|     | ChIP-seq |
|     | Flow cytometry |
|     | MRI-based neuroimaging |

Antibodies

Antibodies used

- Atto594 conjugated anti-RFP nanoboster (Chromotek, rba594-100)
- Atto647N conjugated anti-GFP nanoboster (Chromotek, gba647n-100)
- Atto488N conjugated anti-GFP nanoboster (Chromotek, gba488-100)
- Atto488 conjugated anti-Vimentin nanoboster (Vimentin-Label_Aatto488, Chromotek, vba488-100)
- Alexa647 conjugated anti-GFP antibody (Invitrogen, A31852)

All the nanoboosters were used in dilution 1:500 to label brain slices, 1:600 to label whole brains and 1:7000 to label whole bodies. The antibody was used in dilution 1:850 to label brain slices, 1:790 to label whole brains and 1:7000 to label whole bodies.
Validation

All the nanoboosters were coming from the same company which is the manufacturer: Chromotek https://www.chromotek.com/products/nano-boosters/. The manufacturer validated the nanoboosters and in its website (see link) they provide references, images of the validation from published papers and from validation experiments. We used nanobooster to label fluorescent proteins and vimentin expressed in mouse tissue, as indicated in example images from the manufacturer website. Before any deep tissue staining such as vDISCO whole body labeling, we further validated the compatibility of the nanoboosters with vDISCO in brain slices from transgenic animals as indicated in Method part of the manuscript.

The Alexa647 conjugated anti-GFP antibody (GFP Polyclonal Antibody Alexa Fluor 647, Invitrogen, A31852) has been validated by the manufacturer: Invitrogen https://www.thermofisher.com/antibody/product/GFP-Antibody-Polyclonal/A-31852. In the manufacturer website (see link) they provide references, images of the validation from published papers and from validation experiments. We used this antibody to label GFP expressed in mouse tissue, and this application has already been shown by references indicated in the manufacturer website. We further validated the compatibility of this antibody with iDISCO+ protocol in Supplementary Figure 3.

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

| Laboratory animals | We used the following mixed gender animals in the study: 6 weeks-12 months old CX3CR1GFP/+ (B6.129P-Cx3cr1tm1Litt/J, Jackson Laboratory strain code: 005582), 6 weeks-6 months old Thy1-GFP M, 3-7 months old Thy1-YFP, 4 weeks old Prox1-EGFP (Tg[Prox1-EGFP]K_SY221Gsat/Mmucd, MMRRC strain code: 031006-UCD), 6 months old VEGFR3-YFP, 2 months old CX3CR1GFP+/x CCR2RFP/+ (B6.129(Cg)-Ccr2tm2.1Ifc/J, Jackson Laboratory strain code: 017586), 6 months old LysM-EGFP (Ly2tm1.1Graf, MGI: 2654931), 3 months old CD68-EGFP (C57BL/6-Tg(CD68-EGFP)1Drg/j, Jackson Laboratory strain code: 026827), 3 weeks-4 months old C57BL/6J, 3-4 months old NMRI nu/nu mice. |
| Wild animals | This study does not involve wild animals |
| Field-collected samples | This study does not involve field-collected samples |