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Jensen, Kristian HR; Berg, Rune W.

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CLARITY-compatible lipophilic dyes for electrode marking and neuronal tracing

Kristian H. R. Jensen & Rune W. Berg

Fluorescent lipophilic dyes, such as DiI, stain cellular membranes and are used extensively for retrograde/anterograde labeling of neurons as well as for marking the position of extracellular electrodes after electrophysiology. Convenient histological clearing techniques, such as CLARITY, enable immunostaining and imaging of large volumes for 3D-reconstruction. However, such clearing works by removing lipids and, as an unintended consequence, also removes lipophilic dyes. To remedy this wash-out, the molecular structure of the dye can be altered to adhere to both membranes and proteins so the dye remains in the tissue after lipid–clearing. Nevertheless, the capacity of such modified dyes to remain in tissue has not yet been tested. Here, we test dyes with molecular modifications that make them aldehyde-fixable to proteins. We use three Dil–analogue dyes, CM-DiI, SP-DiI and FM 1–43FX that are modified to be CLARITY-compatible candidates. We use the challenging adult, myelin-rich spinal cord tissue, which requires prolonged lipid–clearing, of rats and mice. All three dyes remained in the tissue after lipid–clearing, but CM-DiI had the sharpest and FM 1–43FX the strongest fluorescent signal.

Tracing the position of an extracellular electrode after electrophysiological experiments in the nervous system is important to verify both the brain region and the distance between electrode and nearby neurons, which are often identified via either genetically expressed reporters or immunohistochemistry. The tracing is commonly accomplished using a non-toxic fluorescent and lipophilic dye, as the widely used carbocyanine dye DiI, which is painted on the electrode shank prior to insertion. The electrode then leaves remnants of the dye that can be easily be detected by fluorescent microscopy in histological preparations. An additional widespread application of lipophilic fluorescent dyes is for tracing neuronal connections and projections in the nervous system especially since these dyes, e.g. DiI, are well-suited for immunohistochemistry and have minimal photo–bleaching. DiI can be transported retrogradely and has been used for post-mortem axonal labelling in animals and humans. Nevertheless, this approach has an unfortunate limitation due to the need to do serial sectioning of the tissue, which is imprecise and labour intensive. The tracing can require several sections if the cutting angle is not parallel to the electrode track or neuronal trace and includes the risk of distorting the samples.

However, recent developed clearing techniques, such as CLARITY, PACT, CUBIC, IDISCO, and Sca/e6–16 conveniently allow immunohistochemistry, imaging and 3D-reconstruction in large volumes of unsectioned tissue. These clearing techniques work essentially by washing away the lipids with detergents and solvents. CLARITY works by polymerizing the fixed tissue into an acrylamide hydrogel prior to lipid–removal. This leaves molecules with amine ends, i.e. proteins, DNA, and RNA, into a protein skeleton structure. These lipid–clearing processes introduce a caveat when combined with the axonal tracing or electrode marking: The lipophilic dyes are also washed out since they adhere to the lipids. CLARITY and similar clearing techniques are thus currently incompatible with the employment of lipophilic dyes in tracing and marking. Clearing alternatives such as Clear and do not wash out the lipids and therefore leaves the lipophilic dyes secured, but they do not clear as well as CLARITY and therefore have low visualization depth. The clearing methods SeeDB3, 3DISCO4 and BABB5 have high visualization depth, but the penetration depth of immunohistochemistry is limited to 100–250 μm or lower. It would therefore be an advantage to find alternative dyes for tracing, which would be compatible with CLARITY and similar clearing techniques while also having a strong tie to the cellular membranes to limit the diffusion before fixing. Notably, a very recent improvement of the Sca/e clearing technique, called Sca/eS, is able to overcome these shortcomings and preserve DiI–labeling. This was demonstrated in axon tracks, which were...
labelled with DiI–crystals applied onto the tissue surface on post-mortem brain tissue. The tissue was fixed prior to the dye–application and then fixed again for a week. In spite of these improvements, fixability of lipophilic dyes, that remain in the tissue after lipid–clearing, continue to be a desirable quality. The simplest approach to obtain an alternative dye is to chemically change an existing dye, e.g. DiI, into a DiI–analogue, which possesses both the property of adhering to the cellular membranes and protein structures, such that it would remain in the tissue after lipid–clearing. Such dyes have already been developed, in particular the analogues of DiI: CM-DiI, SP-DiI and FM 1–43FX, with modifications that make them aldehyde-fixable to proteins (Fig. 1). Here, we tested the DiI–analogues on multi–electrodes placed in the spinal cord using CLARITY as clearing method. The spinal cord has an envelope of dense white matter and is therefore a difficult tissue to clear of lipids, though it has successfully been cleared previously.

Results
Fixable lipophilic dyes remain after lipid–clearing. We tested three dye–candidates, the dyes CM-DiI, FM 1–43FX, and SP-DiI (Fig. 1), in living tissue. Multi–electrodes, which had been dipped in the dye (diluted in ethanol), were inserted for 10 min into the spinal cord of an anesthetized mouse (Fig. 2a–c). Traces of the fixable dyes remained in the tissue as indicated by the fluorescent signals (Fig. 2d–e) after lipid–clearing of the spinal cord using CLARITY (Fig. 2f–g). In the widefield microscope, dye FM 1–43FX had the broadest and most vivid overall fluorescence, while CM-DiI had the sharpest traces. SP-DiI was barely visible with a poor overall fluorescent signal (Fig. 2d–e). Twelve traces were identifiable for CM-DiI, 13 for FM 1–43FX, and only 9 for SP-DiI, where 14 were expected for each dye. Electrode track marking using CM-DiI and FM 1–43FX were the most reliable (Fig. 3). They both enabled clear electrode localization in 3D (Fig. 3d,h), whereas the SP-DiI reconstruction required additional post-processing when imaged together with DAPI (subtraction of the spectrally overlapping DAPI signal, Fig. 3j–l) and was limited by the low fluorescence and few identifiable traces (Fig. 3i–l). For the electrodes with CM-DiI, 86% of the tracks were recovered, and 93% of the tracks for the electrodes applied FM 1–43FX (Fig. 4a).

Figure 1. Overview of chemical structures and properties of DiI and CLARITY–compatible analogues. DiIC18(3) (DiI in short) is a fluorescent lipophilic cationic indocarbocyanine dye used for single molecule imaging, fate mapping, and neuronal tracing, as it is retained in the lipid bilayers. CM-DiIC18(3) (CM-DiI in short) incorporates a mildly thiol–reactive chloromethylbenzamido (CM) substituent (green) that confers aldehyde fixability via conjugation to thiol-containing proteins. SP-DiIC18(3) (SP-DiI in short) has two sulfophenyl (SP) groups (magenta) that confers fixability and a greater ε than DiIC18(3). FM 1–43FX, which is less similar to DiIC18(3), is a lipophilic styryl dye with an aliphatic amine side chain (red) for aldehyde fixability. It has, notably the lowest ε, and the greatest Stokes shift, but also half the molar mass due to lacking the two 18-carbon-long alkyl tails (blue). *Spectral properties determined in methanol. ε: molar attenuation coefficient in cm−1 M−1.

| Dye               | Solvent        | Concentration | Excitation/Emission* | ε* | Molar mass |
|-------------------|----------------|---------------|----------------------|----|------------|
| DiIC18(3)         | Ethanol        | 2,5 mg/mL     | 549 / 565 nm         | 148,000 | 933,9 g/mol |
| SP-DiIC18(3)      | Ethanol        | 2,0 mg/mL     | 557 / 573 nm         | 164,000 | 1,145,7 g/mol |
| CM-DiIC18(3)      | Ethanol        | 1,0 mg/mL     | 553 / 570 nm         | 194,000 | 1,051,5 g/mol |
| FM 1–43FX         | Water or Ethanol | 1,0 mg/mL    | 510 / 626 nm         | 38,000  | 560,1 g/mol |
In euthanized rats, where the spinal cord had been PFA–fixed prior to insertion of electrodes for 30 min, there was a strong SP-DiI fluorescence after CLARITY–clearing (Supplementary Fig. 1).

As a control, we tested whether a non–fixable DiI variant, DID, the redshifted version of DiI, remained in the tissue following lipid–clearing using both CLARITY and Sca/eA2. DiD was applied on electrodes prior to insertion into the tissue. The tissue was then PFA–fixed and after lipid–clearing using both CLARITY (Supplementary Fig. 2) and Sca/eA2 (data not shown) there was no traceable remnants of the dye in the tissues.

**FM 1–43FX has the strongest fluorescence.** Fluorescence measurements demonstrated FM 1–43FX to have the strongest fluorescence signal, followed by CM-DiI (Fig. 3). The fluorescent signals were quantified as the intensity of signal in a rectangle place over each identifiable electrode vestige. The mean signal of these rectangles ± standard error of the mean show a clear difference between the intensities of the three dyes (Fig. 4b). SP-DiI had limited staining and required double the excitation laser power for all images (2.5 mW versus 1.25 mW at 514 nm). Both CM-DiI and FM 1–43FX had stronger fluorescent signals and the electrode traces were visible in imaging depths down to 866 μm (Supplementary Fig. 3). FM 1–43FX has a broader emission spectra than CM–DiI and SP–DiI, which led to crosstalk (bleed–through) when acquiring a multi–channel image with DAPI and GFP. This bleed–through required image post-processing to subtract the separate fluorescent signals (Supplementary Fig. 3).

In addition, in order to verify whether using water versus ethanol as the dye solvent changed the degree of dye binding to the electrode and tissue staining we tested the degree of staining using both these dye solvents for FM 1–43FX. The two solvents had no clear difference in the degree of staining (Supplementary Fig. 3). We also tested whether using organic solvents, such as glicerol and 2,2′-thiodiethanol (TDE) as the refractive index matching medium (RIMM) had an effect on the dyes. We compared glicerol and TDE with the aqueous 88% w/v Histodenz in PBS and found no noticeable difference in fluorescence and image quality (data not shown).
Dye diffusion and axon tracing. The width of the fluorescent electrode traces varied from 5 to 90 μm. CM-DiI and SP-DiI had thinner (<30 μm) traces, while FM 1–43FX had broader traces (>30 μm) (Fig. 5a–c). The broader staining using FM 1–43FX is likely due to the lateral diffusion along the membrane associated with labelling of cells and projections (Fig. 5c). FM 1–43FX extensively stained the white matter column in longitudinal direction at the electrode tip (right, f). Both CM-DiI and FM 1–43FX enable clear electrode localization in 3D (d,h). (i–l) SP-DiI had faint traces, that were difficult to distinguish from the background and localize in 3D (l). All images positioned with dorsal surface left, and images are from hemisection 2 (except a,e,i). Diffuse staining to the left is the superficial tissue at the point of entry. The multielectrode had seven shanks, each 50 μm thick and 200 μm apart. The laser power used to excite CM-DiI and FM 1–43FX was 1.25 mW and 2.5 mW for SP-DiI. Due to the high excitation laser power, the broad spectra of DAPI and the faint signal of SP-DiI, it was necessary to removed DAPI background fluorescence by subtracting 50% of the DAPI images from the SP-DiI images (j–l). The depths of the imaging plane from the surface are given as z-stack midpoints (hemisection 1 and 2): 915 μm and 805 μm (CM-DiI), 910 μm and 706 μm (SP-DiI), 667 μm and 675 μm (FM 1–43FX). ((a) Maximum Intensity Projection of 53 images, z = 338 μm, (b–d) 66 images, z = 455 μm, (e) 53 images, z = 335 μm, (f–h) 72 images, z = 497 μm, (i) 31 images, z = 200 μm, (j–l) 73 images, z = 504 μm).
Figure 4. Performance of the DiI–analogues quantified by the fluorescent residue from the electrode tracks after lipid–clearing. (a) Electrode tracks recovered by visual inspection from the two hemisections: twelve traces were identifiable for CM-DiI (86%), 13 for FM 1–43FX (93%), and only nine for SP-DiI (93%) out of 14. (b) Fluorescence intensity in rectangles placed on the identified electrode traces with background fluorescence subtracted. Values are the mean intensity of the identified tracks with the standard error of the mean. *The laser power used to excite SP-DiI was double (2.5 mW) that of CM-DiI and FM 1–43FX (1.25 mW) and therefore is not directly comparable, hence the broken line of separation.

Figure 5. CM-DiI gives sharp staining, while FM 1–43FX gives brighter traces of electrodes and axons. High resolution images of CM-DiI, FM 1–43FX and SP-DiI (as seen in Fig. 3b, f, j) reveal (a) CM-DiI has the sharpest traces while (b) FM 1–43FX has the strongest overall fluorescence (excitation laser power was 1.25 mW in both). (c) SP-DiI had a fainter signal even with the double laser power (2.5 mW). (d,e) Close-ups from (b) boxes show the lateral diffusion of FM 1–43FX along the axonal membranes in the ventral white–matter column at the electrode tip. (a) Maximum Intensity Projection of 65 images, z = 428 μm, (b) 38 images, z = 247 μm, (c) 53 images, z = 347 μm, (d) 11 images, z = 139 μm, (e) 10 images, z = 127 μm.)
Discussion

It has previously been possible to achieve only two out of three important qualities in histology: High visualization depth, compatibility with immunohistochemistry, or compatibility with lipophilic dyes. Fortunately, with fixable Dil–analogue based lipophilic dyes, such as CLARITY, it is now possible to get all three desired qualities. In the present study, we therefore tested the compatibility of three groups of lipophilic dyes with CLARITY, for the purpose of visualizing electrode placement and neuronal projections. Since other lipid–clearing methods work in similar yet gentler ways compared with CLARITY, these fixable Dil–analogue are likely to work with other lipid–clearing methods as well. We traced electrode tracks following electrophysiology and 3D reconstruction with the option of including immunostaining. All three dyes had preserved fluorescent signal even after prolonged lipid–clearing using CLARITY. CM-Dil had the sharpest and FM 1–43FX the strongest signal (Figs 2, 3, 4 and 5), which demonstrates a graceful compatibility with CLARITY. Together our results show that the Dil–analogue, including FM 1–43FX, are fixable, but show different staining patterns and fluorescent intensities (Figs 3, 4 and 5). We recommend using FM 1–43FX to achieve the strongest signal and CM-Dil for the sharpest signal.

There are alternative variants of the Dil and FM dyes, which have alterations in their chemical structures. Since the alterations are minor, these alternative dyes likely have similar properties regarding the fixation and lipid–extraction protection. The three lipophilic dyes that were tested in the present report, are representatives of the main types of lipophilic and fixable dyes: 1) the sulfonated Dil–variants, 2) Dil with a chloromethyl benzamide modification and 3) FM dyes with an attached aliphatic amine (Supplementary Fig. 4). These three dyes are suitable to compare since they have similar excitation and emission spectra and could thus be excited and imaged with the same microscope parameters, i.e. excitation laser and emission filter wavelengths. Other variants may have different excitation/emission spectra, but they are likely to have similar chemical properties regarding lipid–clearing, since their fixable parts are the same. CM-Dil and SP-Dil are variants of the commonly used long-chain lipophilic carbocyanine dye Dil. We selected SP-Dil as a representative for the group of anionic sulfonated Dil–variants with sulfo-, DiI-DS, DiD-DS, and sulfophenyl-modifications, SP-Dil, SP-DiO (Supplementary Fig. 4).

An example of a subtle difference between variants is the Dil with different length of the carbon bridge between the indoline rings, which gives different excitation and emission wavelengths e.g. DiD (DiIC1(5), ex/em 650/670 nm) a red-shifted version with a 5 rather than 3 carbon bridge (Supplementary Fig. 4). This neither effects fixability, nor does it make the dye less lipophilic. The lipophilic property is dependent on the length of the alkyl chain length (Fig. 1). Similarly, FM 1–43 has a red-shifted variant, FM 4–64, with a 6 rather than 2 carbon bridge between the aromatic rings. Nevertheless, the parts of the molecules, which confer the fixability of the dyes, e.g., the aliphatic amine, remain the same, and therefore the property of remaining in the tissue after lipid–clearing is likely to be conserved. The chloromethyl benzamide (CM) and sulfophenyl (SP) modifications (Fig. 1), keep the dyes retained throughout PFA–fixing and permeabilization procedures with detergents and solvents e.g. Triton-X and acetone. The mildly thiol–reactive CM substitution confers aldehyde fixability via conjugation to thiol–containing proteins. The mechanism of SP-Dil retention is currently unknown.

SP-Dil has the highest molar attenuation coefficient (Fig. 1) of the lipophilic fluorescent dyes that had tolerated fixing and permeabilization according to the manufacturer (Thermo Fisher Scientific). Nevertheless, SP-Dil only gave faint staining and the electrode tracks were difficult to identify (Figs 3, 4 and 5). In addition, SP-Dil required double the excitation laser power compared to CM-Dil and FM 1–43FX during imaging. When we inserted the electrode for 30 minutes in a pre-fixed rat spinal cord, a bright and sharp staining was found (Supplementary Fig. 1), which suggests that SP-Dil staining requires a longer exposure in the tissue. It may, therefore, work better for longer exposure such as during chronic electrode implantations. Dil is commonly used for tracing neuronal tracts in human post-mortem brain samples, and therefore SP-Dil would be a suitable analogue for similar neuronal tract tracing, where lipid–clearing and large volume microscopy are required.

CM-Dil gave a bright and sharp staining of electrode tracks after only 10 minutes in the tissue (Figs 2, 3 and 4). It had limited diffusion into surrounding cell membranes and gave narrow electrode traces. For electrode tracing, CM-Dil was bright, sharp and reliable (Fig. 3a–d) with 12 of the 14 electrode traces recovered (Fig. 3a). However, we observed an overall diffuse staining of CM-Dil when the insertion site was close to the incision of the pia mater (Supplementary Fig. 3). In conclusion, CM-Dil gave a bright and sharp staining, and is therefore well suited as a marker of acute electrode placement, and likely also chronic.

FM 1–43FX also gave a bright staining, despite its low molar attenuation coefficient (38,000 m2/ mol). The molar mass of FM dyes is half of Dil–variants (Fig. 1) and therefore the molar concentration is approximately double compared to CM-Dil in our experiments, which contributes to the stronger fluorescence. FM 1–43FX gave a broader and more diffuse staining of the electrode tracks, as well as lateral diffusion along the white matter fibers (Fig. 2f). Its small molecular weight gives FM 1–43FX the fastest diffusion of three dyes, which in parts could explain the broader staining of the electrode track. It should be noted that the diffusion in the extracellular space is rather limited and anomalous since the molecules are lipophilic and therefore have ‘sticky’ boundary conditions of the cellular membranes.

FM dyes may not be inert, but can bind to and block muscarinic acetylcholine receptors. Other FM dyes can block the nicotinic muscarinic acetylcholine receptors and pass through and block mechanotransduction channels in the hair cells of the cochlea. An important property of FM dyes is their rapid lateral diffusion and involvement in vesicle endocytosis. These dyes are absorbed into the outer lipid bilayer and then taken up by endocytosis at the pre–synaptic terminals, especially in active neurons. The low molar mass and the involvement in endocytosis likely contributes to the spread of staining compared to CM- and SP-Dil (Figs 1 and 2). Therefore, when chronically implanting...
electrodes marked with FM 1–43FX, the dye will likely rapidly stain the neuronal projections and synaptic terminals in addition to the cells at the electrode site. The rapid lateral membrane diffusion may also contribute to the high fluorescence yield by preventing dye aggregation, which is a common cause of fluorescence self-quenching36. Dif dyes are also effective neuronal tracers, but are known to have slow diffusion and therefore require longer incubation37. Their diffusion speed is not specifically dependent on alkyl chain length38, but it can be increased by modifying the alkyl chains to unsaturated alkenes as with EAST Di (DiβC12, C3)39. Alternatively, the application of an electrical field or the depletion of cholesterol using e.g. methyl-β-cycloexodrin in the tissue improved lateral diffusion speed40. However, fixable DiI variants are currently not commercially available with alkene chains. It is therefore unlikely that the CM and SP modifications affect diffusion speed and they are thus expected to diffuse and behave like regular DiI41,42. Therefore, SP-DiI and CM-DiI would be well suited for chronic electrode implantations.

FM and DiI dyes are considered to have limited cytotoxic and not to inhibit cell proliferation43. However, one study finds that carbocyanine dyes with long (≥C16) alkyl side chains, as DiI, are toxic to embryonic rat motoneurons. Another study that they inhibit mitochondrial electron transport chain activity44. Therefore, carbocyanine dyes with shorter (<C12) alkyl side chains may be more suitable for embryonic tissue or long-term labelling. CM-DiI is reportedly toxic to preadipocytes45.

There are other clearing methods similar to CLARITY that use organic solvents or detergents, which disrupts membranes and impairs reliable signal preservation of lipophilic dyes i.e. DISCO5, CUBIC4, Sca/e6 and TDE46-47. Sca/e uses a mixture of urea and glycerol, which disrupts the membranes and dissolves lipophilic dyes6,17. A promising novel type of sorbitol–based clearing method called Sca/e6 has been demonstrated to preserve membrane structure16. Remarkably, Sca/e also preserves DiI-labeling in axon tracks labelled with DiI–crystals applied onto the tissue, though it should be noted that it was performed on post-mortem brain tissue, which was fixed prior to the dye–application, after which the tissue again was fixed a week17. Other alternatives do not damage the membranes or remove lipophilic dyes. For instance, FocusClear TM and Clear7 are based on aqueous solutions that are compatible with lipophilic dyes and immunohistochemistry, but they do not clear as well as CLARITY and Sca/e17,22. SeeDB and FRUIT use fructose, which preserves membranes and lipophilic dyes, but the processed tissues has limited permeable for macromolecules and antibodies and are thus incompatible with large volume immunohistochemistry19,26,48.

We used the passive CLARITY technique, PACT42, which relies on incubation and agitation at 37–40 °C in 8% SDS. We consider the high detergent concentration and heating during PACT to one of the harshest clearing methods. Active CLARITY with electrophoresis produces oxygen radicals making it even harsher than passive CLARITY. These free radicals can chemically reduce the dye and quench fluorescence. However, fluorescent proteins are well preserved in active CLARITY and ACT-PRESTOP5. Since the DiI–analogues are covalently bound to primary amines on proteins and nucleic acids by methylene bridges following aldehyde fixation, they are not removed during lipid extraction by solvents or detergents. The fixable DiI–analogues were stable in CLARITY and the organic solvents i.e. 80% Glycerol and 65% TDE45 used as imaging mediums. When stored in 80% glycerol for 6 months the tissue and traces retained their image worthy fluorescent signal and did not require additional laser power to create identical images as presented here. We therefore predict that the fixable DiI–analogues tested in passive CLARITY are also compatible with other clearing methods based on organic solvents or lipid removal as Sca/e6, DISCO5, CUBIC4 and TDE46-47. Moreover, these dyes may also be useful in other animal or plant tissues; that require CLARITY clearing with electrophoresis or the addition of enzyme digestions50,51.

In general, dyes, as DiI, are used for antero- and retrograde tracing of neurons in vivo, in vitro, and post-mortem to better understand neuronal circuits5. Other common neuronal tracing methods rely on e.g. viral vectors, dextran-conjugates, Cholera Toxin Subunit B or biocytin52. To date, the only neuronal tracer studies in CLARITY have been done using viral vectors53,54. Viral vectors are powerful tools for studying neuronal circuits54. Fixable DiI–analogues could serve as a viable alternative to tracing using viral vectors (See Supplementary Table 1 for an overview of dyes).

Methods

All experiments and procedures were approved by the Department of Experimental Medicine, and according to procedures laid out by the Danish Ministry of Justice and the Danish National Committee for Ethics in Animal Research. Adult Sprague-Dawley rats of both sexes were used. Transgenic mice (PV-Cre:channelrhodopsin-2/ EYFP were provided by the Jackson lab via the Bartak lab36. Mice expressing enhanced green fluorescent protein (eGFP) under the promoter of glial fibrillary acidic protein (GFAP, Tgm/hgFAPEGEP) (GFEC 335) were provided by the Perrier lab37 who got the strain from the Kirchhoff lab38.

Tracer Dyes. The multi-electrode was submerged in 0.5 ml of a dye solution in a 1.5 ml microcentrifuge tube (Eppendorf) and the electrode air-dried out over at least 5 min before usage and protected from light under an aluminum cover. The dye solutions were either 0.25% w/v DiIC18(3), 0.25% w/v DiDC18(5), 0.2% w/v SP-DiI-C16(3), or 0.1% w/v CM-DiI-C16(3) (CellTracker™) in ethanol, or 0.1% w/v FM® 1–43FX (a fixable version of FM® 1–43) in ethanol or water (Thermo Fisher Scientific). The electrodes were washed by leaving them overnight in ethanol. When painting onto the electrodes, ethanol is preferable as solvent due to quicker drying of the dye after application. Chemical structures draw with ChemSketch (ACD/Labs).

Electrode insertion. The multi–electrode (A4x4-3 mm-50-125, Neuroexus Inc.) was attached to a micro-manipulator and gradually inserted dorsally 0.3–1 mm into the spinal cords in the sagittal plane slightly laterally from the hemicord midline. In the mouse brain, the multielectrode was inserted approximately 0.2 mm from the cortical surface. Measurements usually lasted an hour, but were not acquired here. Time in tissue was 8–30 minutes. The electrode painted with DiD was inserted in the mouse cortex and left for 2 hours. The multichannel
silicon electrodes (multi–electrodes) that were used had either 8 shanks, 40 μm wide, 15 μm thick, 200 μm apart (Neuronexus inc., part no. A8x8-5 mm-200-160, where one lateral shank broke off during a previous experiment), 4 shanks, 83 μm wide, 15 μm thick, 150 μm apart (Neuronexus inc., part no. A4x1-tet-3 mm-150-150-312-A16) or 4 shanks 125 μm apart (Neuronexus Inc., part no. A4x4-3 mm-50-125).

CLARITY. Mice and rats were perfused transcardially with 4% w/v paraformaldehyde (PFA) in phosphate buffered saline (PBS). The spinal cord and brain was then removed and stored in 4% w/v PFA overnight. All tissue was stored in PBS until further processing. Brain tissue was embedded in 3% w/v SeaPlaque® Agarose (Lonza) and cut with a vibratome (VT1000 S, Leica Biosystems) into 1000 μm slices. Spinal cord segments were manually cut with a scalpel along the midline or embedded in agar as brain tissue and cut into 600 μm transverse slices. The CLARITY clearing processing was performed via PACT: a passive CLARITY technique59,60 and accomplished according to Yang et al.13 by the following steps: The tissue was placed in a conical tube with 4% w/v acrylamide, 0.05% w/v bis-acrylamide, 0.25% w/v VA-044 (Wako Chemicals) in PBS, and incubated overnight at 5 °C. The following day, the solution was degassed with N₂ for 5 minutes, then incubated at 37 °C while gently shaking for 4 hours until the hydrogel had fully polymerized. The tissue was extracted from the gel with Kimwipes and washed in 8% w/v sodium dodecyl (SDS, Sigma-Aldrich) in PBS, pH 7.5, twice over a day at room temperature or 37 °C. The clearing process continued over several nights with replacement of SDS each day if possible to enhance the clearing speed. Coronal mouse brain sections (1 mm thick) and (1 cm long) adult mouse spinal cords segments were cleared over 4 days, while rat spinal cord segments were cleared over 6 days.

Staining and Mounting. After the CLARITY procedure, the sample was washed in PBS-T (PBS with 0.1% v/v Triton-X-100) at room temperature for 10 minutes during gentle shaking, which was repeated once. Nuclear DNA was stained with 2 μg/ml DAPI in PBS with 0.1% v/v TritonX-100, and 0.01% w/v sodium azide, at room temperature during gentle shaking over two nights. After staining, the tissue was washed twice with PBS for 10 minutes and gently dried with filter paper. Then tissue was transferred to a refractive index matching medium (RIMM) of either 80% v/v glycerol (Sigma-Aldrich Co), 88% w/v Histodenz (Sigma-Aldrich Co), or 63 v/v % 2,2′-thiodiethanol (TDE, VWR international) in PBS with 0.01% sodium azide and pH adjusted to 7.5 with NaOH. The tissue was incubated in RIMM at room temperature until transparent, usually overnight, prior to imaging. The tissue was placed in a chamber designed for live cell imaging (POC R2, PeCon) consisting of two 0.17 mm glass slides with silicon spacers to prevent compression. The tissue was sandwiched between the glass slides while being covered in RIMM and gently held in place without being compressed by the silicon spacers.

Imaging. We used confocal microscopes LSM 700 and LSM 710 (Zeiss) with EC Plan-Neofluar, air 10x/NA 0.3 with a working distance of 5.2 mm. Excitation sources were a 30 mW diode for 405 nm, a 25 mW Argon laser at 488 nm and 514 nm (CM-Dil and FM 1–43FX were excited at 5% and SP-Dil at 10% laser power) with an Airy unit of 0.98–1.5. Images were acquired with 2 or 4 times frame–averaging, tiles had 0–10% overlap and Z-stacks had 10% overlap. Overview images were acquired with a Zeiss SteREO Lunar V12 with a 1.5x Neolumar S. Post-processing (contrast adjustments, and subtraction in Fig. 3)–1 and Supplementary Fig. 3e: 50% of the DAPI, GFP and Dil images), maximum intensity projections (MIP) and 3D–reconstructions) was done in Zen Blue 2012 (Zeiss) and Vaa3D 3.05561. Macroscopic images where taken with a M320 Dental Microscope (Leica).

Fluorescent measurements. We measured the fluorescence of the electrode traces seen in the confocal images (Fig. 3) on unadjusted MIPs. The mean intensity in 80 μm × 600 μm rectangles placed on the identifiable electrode traces were subtracted the mean background intensity in a 160 μm × 600 μm rectangle placed elsewhere on the tissue in the images. This was done in Zen Blue 2012 (Zeiss). Measurements were handled in Prism 6 (GraphPad).

References
1. Watson, T. C., Becker, N., Apps, R. & Jones, M. W. Back to front: cerebellar connections and interactions with the prefrontal cortex. Front. Syst. Neurosci. 8, 4 (2014).
2. DiCarlo, J. J., Lane, J. W., Hsiao, S. S. & Johnson, K. O. Marking microelectrode penetrations with fluorescent dyes. J. Neurosci. Meth. 64, 75–81 (1996).
3. Blanche, T. J., Spacek, M. A., Herke, J. F. & Swindale, N. V. Polytoodes: high-density silicon electrode arrays for large-scale multunit recording. J. Neurophysiol. 93, 2987–3000 (2005).
4. Holmqvist, B. L., Ostholm, T. & Ekstrom, P. Dil tracing in combination with immunocytochemistry for analysis of connectivities and chemoarchitectonics of specific neural systems in a teleost, the Atlantic salmon. J. Neurosci. Meth. 42, 45–63 (1992).
5. Hono, M. G. & Hume, R. I. Dil and diO: versatile fluorescent dyes for neuronal labelling and pathway tracing. Trends Neurosci. 12, 333–335, 340–341 (1989).
6. Elberger, A. J. & Honig, M. G. Double-labeling of tissue containing the carbocyanine dye DiI for immunocytochemistry. J. Histochem. Cytochem. 38, 735–739 (1990).
7. Körbert, C. et al. Current concepts in neuroanatomical tracing. Prog. Neurobiol. 62, 327–351 (2000).
8. Vidal-Sanz, M., Villegas-Pérez, M. P., Bray, G. M. & Aguayo, A. J. Persistent retrograde labeling of adult rat retinal ganglion cells with the carbocyanine dye diI. Exp. Neurol. 102, 92–101 (1988).
9. Okabe, H. S. & Wahlsten, D. Prenatal formation of the normal mouse corpus callosum: a quantitative study with carbocyanine dyes. J. Comp. Neurol. 323, 81–90 (1992).
10. Mufson, E. J., Brady, D. R. & Kordower, J. H. Tracing neuronal connections in postmortem human hippocampal complex with the carbocyanine dye Dil. Neurobiol. Aging 11, 649–653 (1990).
11. Thal, D. R., Capetillo-Zarate, E. & Galuske, R. A. Tracing of temporo-entorhinal connections in the human brain: cognitively impaired argyrophilic grain disease cases show dementic alterations but no axonal disconnection of temporo-entorhinal association neurons. Acta Neuropathol. 115, 175–83 (2008).
12. Chung, K. et al. Structural and molecular interrogation of intact biological systems. Nature 497, 332–337 (2013).
13. Yang, B. et al. Single-Cell Phenotyping within Transparent Intact Tissue through Whole-Body Clearing. Cell 158, 945–958 (2014).
14. Susaki, E. A. et al. Whole-brain imaging with single-cell resolution using chemical cocktails and computational analysis. Cell 157, 726–739 (2014).
15. Renier, N. et al. iDISCO: A Simple, Rapid Method to Immunolabel Large Tissue Samples for Volume Imaging. Cell 159, 896–910 (2014).
16. Hama, H. et al. Scale: a chemical approach for fluorescence imaging and reconstruction of transparent mouse brain. Nat. Neurosci. 14, 1481–1488 (2011).
17. Richardson, D. S. & Lichtman, J. W. Clarifying Tissue Clearing. Cell 162, 246–257 (2015).
18. Hirashima, T. & Adachi, T. Procedures for the Quantification of Whole-Tissue Immunofluorescence Images Obtained at Single-Cell Resolution during Murine Tubular Organ Development. Plos One 10, e0133543 (2015).
19. Kim, S. Y., Chung, K. & Deisseroth, K. Light microscopy mapping of connections in the intact brain. Trends Cog. Sci. 17, 596–599 (2013).
20. Matsubayashi, Y., Iwai, L. & Kawasaki, H. Fluorescent double-labeling with carboxyfluorescein tracing neuronal and immunohistochemistry using a cholesterol-specific detergent digitonin. J. Neurosci. Meth. 174, 71–81 (2008).
21. Hama, H. et al. ScaleX: an optical clearing palette for biological imaging. Nat. Neurosci. 18, 1518–1529 (2015).
22. Kuwajima, T. et al. ClearT: a detergent- and solvent-free clearing method for neuronal and non-neuronal tissue. Development 140, 1364–8 (2013).
23. Ke, M.-T., Fujimoto, S. & Imai, T. SeeDB: a simple and morphology-preserving optical clearing agent for neuronal circuit reconstruction. Nat. Neurosci. 16, 1154–61 (2013).
24. Ėrtürk, A. et al. Three-dimensional imaging of solvent-cleared organs using 3DISCO. Nat. Prot. 7, 1983–1995 (2012).
25. Yokomizo, T. et al. Whole-mount three-dimensional imaging of internally localized immunostained cells within mouse embryos. Nat. Protoc. 7, 421–431 (2012).
26. Ke, M.-T. & Imai, T. Optical clearing of fixed brain samples using SeeDB. Curr. Prot. Neurosci. 66, (unit 2.22), 1–19 (2014).
27. Andrade, W., Seabrook, T. J., Johnston, M. G. & Hay, J. B. The use of the lipophilic fluorochrome CM-DiI for tracking the migration of lymphocytes. J. Imm. Meth. 194, 181–189 (1996).
28. Maurya, D. K. et al. Therapy with un-engineered nave rat umbilical cord matrix stem cells markedly inhibits growth of murine lung adenocarcinoma. BMC cancer 10, 590 (2010).
29. Verstreken, P., Ohya, T. & Bellén, H. J. FM 1–43 labeling of synaptic vesicle pools at the Drosophila neuromuscular junction. Meth. Mol. Biol. 440, 349–69 (2008).
30. Bertrand, C. A., Laboisse, C., Hopfer, U., Bridges, R. J. & Frizzell, R. A. Methods for detecting internalized, FM 1–43 stained particles in epithelial cells and monolayers. Biophys. J. 91, 3872–83 (2006).
31. Marra, V., Burden, J. J., Crawford, F. & Staras, K. Ultrastructural readout of functional synaptic vesicle pools in hippocampal slices based on Fluo-4 dye labeling and photoconversion. Nat. Prot. 9, 1337–47 (2014).
32. Berg, R. W., Chen, M.-T., Huang, H.-C., Hsiao, M.-C. & Cheng, H. A method for unit recording in the lumbar spinal cord during locomotion of the conscious adult rat. J. Neurosci. Meth. 182, 49–54 (2009).
33. Ėrtürk, A. et al. Three-dimensional imaging of the unsectioned adult spinal cord to assess axon regeneration and glial responses after injury. Nat. Med. 18, 166–171 (2011).
34. Bauer, M., Godec, A. & Metzler, R. Diffusion of finite-size particles in two-dimensional channels with random wall configurations. Phys. Chem. Chem. Phys. 16, 6118–24 (2014).
35. Gaffield, M. a. & Betz, W. J. Imaging synaptic vesicle exocytosis and endocytosis with FM dyes. Nat. Prot. 1, 2916–2921 (2006).
36. West, W. & Pearce, S. The dimeric state of cyanine dyes. J. Phys. Chem. 566, 1894–1903 (1965).
37. Swift, M. J., Crago, P. E. & Grill, W. M. Applied electric fields accelerate the diffusion rate and increase the diffusion distance of Dil in fixed tissue. J. Neurosci. Meth. 141, 153–163 (2005).
38. Derezo, Z. & Jacobson, K. Comparative lateral diffusion of fluorescent lipid analogues in phospholipid multilayers. Biochim. 19, 6050–6057 (1980).
39. Sparks, D. L., Lee, L. F., Martin, T. A. & Rogers, J. Neural tract tracing using Di-I: A review and a new method to make fast Di-I faster in human brain. J. Neurosci. Meth. 163, 3–10 (2000).
40. Pucadyil, T. J. & Chattopadhyay, A. Effect of cholesterol on lateral diffusion of fluorescent lipid probes in native hippocampal membranes. Chem. Phys. Lip. 143, 1–11 (2015).
41. Makowiecki, K., Garrett, A., Clark, V., Graham, S. L. & Rodger, J. Reliability of VEP Recordings Using Chronically Implanted Screw Electrodes in Mice. Trans. Vis. Sci. Tech. 4, 15 (2015).
42. Taylor, H. et al. Investigating local and long-range neuronal network dynamics by simultaneous optogenetics, reverse microdialysis and silicon probe recordings in vivo. J. Neurosci. Meth. 235, 83–91 (2014).
43. Anderson, W. M. & Trgovcich-Zacok, D. Carbocyanine dyes with long alkyl side-chains: Broad spectrum inhibitors of mitochondrial electron transport chain activity. Biochem. Pharmacol. 49, 1303–1311 (1995).
44. Hemmrich, K., Meersch, M., von Heimburg, D. & Pallua, N. Applicability of the dyes CFSE, CM-DiI and PKH26 for tracking of human predipocytes to evaluate adopse tissue engineering. Cells Tissues Organs 184, 117–27 (2006).
45. Costantini, L. et al. A versatile clearing agent for multi-modal brain imaging. Sci. Rep. 5, 9808 (2015).
46. Zheng, H. & Rinaman, L. Simplified CLARITY for visualizing immunofluorescence labeling in the developing rat brain. Brain Struct. Func. 14, 1–9 (2015).
47. Aoyagi, Y., Kawakami, R., Osanai, H., Hibi, T. & Nemoto, T. A rapid optical clearing protocol using 2,2’-thiodiethanol for microscopic observation of fixed mouse brain. Plos one 10, e0116280 (2015).
48. Rou, B. et al. Scalable and Dil-compatible optical clearance of the mammalian brain. Front. Neuroanat. 9, 19 (2015).
49. Lee, E. et al. ACT-PRESTO: Rapid and consistent tissue clearing and labeling method for 3-dimensional (3D) imaging. Sci. Rep. 6, 18631 (2016).
50. Lee, H., Park, J.-H., Seo, I., Park, S.-H. & Kim, S. Improved application of the electrophoretic tissue clearing technology, CLARITY, to intact solid organs including brain, pancreas, liver, kidney, lung, and intestine. BMC Devel. Biol. 14, 48 (2014).
51. Palmer, W. M. et al. PLA-CLARITY: 3D molecular imaging of whole plant organs. Sci. Rep. 5, 13492 (2015).
52. Lancia, J. L., Wouterlood, F. G. & Lancia, L. A half century of experimental neuroanatomical tracing. J. Chem. Neuroanat. 42, 157–83 (2011).
53. Menegas, W. et al. Dopamine neurons projecting to the posterior striatum form an anatomically distinct subclass. eLife 4, 1–30 (2015).
54. Lerner, T. et al. Intact-Brain Analyses Reveal Distinct Information Carried by SNC Dopamine Subcircuits. Cell 162, 635–647 (2015).
55. Nasi, J., Cepko, C. L., Born, R. T. & Beier, K. T. Neuroanatomy goes viral! Front. Neuroanat. 9, 80 (2015).
56. Barkat, T. R., Polley, D. B. & Hensch, T. K. A critical period for auditory thalamocortical connectivity. Nat. Neurosci. 14, 1189–94 (2011).
57. Costantini, L. et al. Optimization of CLARITY for Clearing Whole-Brain and Other Intact Organs. eNeuro 2, ENEURO.0022–15.2015 (2015).
60. Tomer, R., Ye, L., Hsueh, B. & Deisseroth, K. Advanced CLARITY for rapid and high-resolution imaging of intact tissues. *Nat. Prot.* **9**, 1682–97 (2014).

61. Peng, H., Ruan, Z., Long, F., Simpson, J. H. & Myers, E. W. V3D enables real-time 3D visualization and quantitative analysis of large-scale biological image data sets. *Nat. Biotech.* **28**, 348–53 (2010).

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**Author Contributions**

K.H.R.J. and R.W.B. conceived and conducted the experiment, and analysed the results. Both authors reviewed the manuscript.

**Additional Information**

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