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TISSUE CLEARING

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Competing interests

H.R.U. is co-founder of CUBICStars Inc and a co-inventor on the following patent applications covering the CUBIC reagents: PCT/JP2014/070618 (pending, patent applicant is RIKEN, other co-inventors are E. A. Susaki and K. Tainaka); PCT/JP2017/016410 (pending, patent applicant is RIKEN, other co-inventors are K. Tainaka and T. Murakami). K.C. is an inventor for patent applications covering some technologies described in this paper and co-founder of LifeCanvas Technologies. A.E. and C.P. have filed a patent on whole-body clearing and imaging related technologies.

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Abstract

Tissue clearing of gross anatomical samples was first described over a century ago and has only recently found widespread use in the field of microscopy. This renaissance has been driven by the application of modern knowledge of optical physics and chemical engineering to the development of robust and reproducible clearing techniques, the arrival of new microscopes that can image large samples at cellular resolution and computing infrastructure able to store and analyze large data volumes. Many biological relationships between structure and function require investigation in three dimensions and tissue clearing therefore has the potential to enable broad discoveries in the biological sciences. Unfortunately, the current literature is complex and could confuse researchers looking to begin a clearing project. The goal of this Primer is to outline a modular approach to tissue clearing that allows a novice researcher to develop a customized clearing pipeline tailored to their tissue of interest. Further, the Primer outlines the required imaging and computational infrastructure needed to perform tissue clearing at scale, gives an overview of current applications, discusses limitations and provides an outlook on future advances in the field.

INTRODUCTION

Tissue clearing refers to a collection of techniques that render biological samples transparent. These techniques enable the deep imaging of large tissue volumes using light microscopy approaches that are usually limited by the scattering of light by the tissue. Tissue clearing allows researchers to avoid time-consuming tissue sectioning approaches that can introduce artefacts and examine biological tissues in their native, three-dimensional state. Further, light microscopy enables the use of individual-molecule labelling techniques and imaging at sub-micrometre spatial resolutions, unlike traditional approaches for volume imaging such as magnetic resonance imaging (MRI), computed tomography (CT) and ultrasound imaging.

Tissue clearing protocols facilitate the passage of light through a biological sample by minimizing the refractive index (RI) differences between components of the tissue¹. In general, this requires the removal of lipids (RI \approx 1.47) and replacement of intracellular and extracellular fluids (RI \approx 1.35) with a solution of equivalent RI to the remaining protein constituents (RI $>$ 1.50). Despite this simple underlying principle, the design of a tissue clearing project can easily become complex and requires optimization of multiple variables. Factors that should be considered include the need for retention of molecules of

interest during processing (for example, proteins or mRNA), maintenance of the sample's structural integrity, conformational protection of endogenously expressed fluorescent proteins, enhanced porosity to allow for the diffusion of small molecule or immunoglobulin labels, and preservation of the structure of relevant epitopes if immunolabelling.

Previous reviews have highlighted the basic physics¹ and chemistry² behind tissue clearing and highlighted the promise of tissue clearing for the field of neuroscience³. This Primer will instead focus on the methodology for clearing various tissue types, the technologies for imaging cleared tissue and how to store and analyse image data. Specifically, the Primer will enable a researcher to understand the purpose of each step in a clearing protocol and assemble a clearing pipeline specifically designed for their sample. It will discuss the advantages and disadvantages of several imaging techniques and describe several common image analysis and quantitation routines. This comprehensive description of clearing, imaging, and data analysis aims to provide the necessary information for individual researchers to design simple clearing experiment and also guide institutional core facilities and commercial entities on how to clear samples at scale.

EXPERIMENTATION

Historically, clearing techniques were assigned to one of two categories based on the composition of their final clearing solution: solvent-based (hydrophobic) or aqueous-based (hydrophilic). Originally, solvent-based techniques were faster, provided a closer RI match to the delipidated, proteinaceous sample and shrunk the sample in size. Conversely, aqueous techniques better preserved the emission of fluorescent proteins (if present), retained biomolecules to a higher degree and expanded samples in size. However, many of the original limitations of each category have been addressed and solvent and aqueous techniques have become nearly equivalent⁴⁻⁶. In fact, it is now recognized that combining aqueous-based and solvent-based delipidation steps can be advantageous and aqueous RI-matching solutions can be combined with solvent-based delipidation (and vice-versa). Therefore, it no longer seems relevant to label techniques aqueous-based or solvent-based but rather to view tissue clearing as a pipeline composed of several modules that can be mixed-and-matched. These modules are sample fixation, pre-treatment, delipidation, fluorescent labelling, RI matching and image acquisition and analysis (Fig. 1). Each clearing protocol must be customized to the sample type and the labelling method used, and the imaging modalities (such as microscopes) or IT infrastructure available to a researcher can influence the choice of clearing technique and inform the scale of the project. Below is an in-depth discussion of the various modules that can be combined to create a complete tissue clearing workflow, with details summarized in Table 1.

Sample choice

Many commonly studied biological tissues from a host of organisms have been shown to be compatible with tissue clearing approaches. Entire rodents have been cleared and imaged while intact⁷⁻¹¹, as have several other model organisms including drosophila^{12,13} and zebrafish^{14,15}. Human tissues, including an entire human brain and kidney, have also been cleared successfully⁶. However, there are limitations to clearing and not all tissues

clear equally well. For example, successful clearing of whole rodents usually requires the removal of the skin from the animal. Additionally, tissues with pigments (such as skin), calcification (bone), or extensive extracellular matrix and/or structural proteins (tumours and cardiac tissues) do not obtain the same level of clearing as tissues without these features.

Specific and bright fluorescent labelling is essential for imaging, regardless of the tissue. Small molecule organic dyes, dye-conjugated antibodies and genetically encoded fluorescent proteins are commonly used labels. Small-molecule dyes can rapidly penetrate and label thick tissue, but are limited in their recognizable targets. Antibodies have the widest utility but suffer from long labelling times and reproducibility issues owing to batch-to-batch variation. Genetically encoded fluorescent proteins may label a higher percentage of target molecules and allow labelling of specific cell types¹⁶. Viral expression of fluorescent proteins avoids the complex molecular biology of stably introducing a transgene to a tissue and can achieve high levels of fluorescence emission depending on virus titre, serotype and choice of promoter^{17,18}. However, care must be taken as many fluorophores are susceptible to bleaching with hydrogen peroxide and fluorescent protein emission can be quenched by dehydration (see below).

Finally, it is important to match the final size of the sample to available imaging systems and data storage capacity. It is of note that cleared samples that are millimetres or centimetres in size can exceed the working distance of many microscope objectives and can produce TBs of data.

Fixation

Fixation of tissue is always required before clearing to avoid excessive loss of target biomolecules or tissue integrity. Fixing covalently crosslinks proteins and/or nucleic acids in the tissue to preserve the tissue's mechanical integrity and biomolecular architecture. Fixation is a balancing act; if fixation is too weak, the tissue will not withstand chemical treatments in later clearing steps. Conversely, over-fixation can result in fixative molecules blocking access to the epitopes required for fluorescent labelling.

The degree of fixation is determined by the concentration of the fixative and amount of time the tissue spends immersed in it. The most commonly used tissue fixative is paraformaldehyde (PFA), which crosslinks the tissue's biomolecules by reacting with amine groups on proteins and nucleic acids. PFA is usually used at a concentration of 3–4% w/v in phosphate-buffered saline (PBS). PFA diffuses rapidly into tissue, making it a good molecule for quickly and uniformly fixing samples greater than 500 µm in diameter. Glutaraldehyde (GA) can be used as an alternative to PFA, which — although chemically similar — reacts with biomolecular amine groups faster, resulting in more intermolecular covalent crosslinking and enhanced tissue fixation¹⁹. However, as GA is a larger molecule than PFA, diffusion into tissues is slower, leading to non-uniform tissue preservation. To address issues caused by diffusion, the SWITCH method can be used²⁰. In this method, inactivated GA molecules diffuse through the sample in a low-pH “OFF” buffer and once the GA molecules are uniformly distributed, the tissue is transferred to a neutral pH “ON” buffer to initiate fixation. Using this protocol, GA was demonstrated to provide improvements in biomolecular preservation (3–5% protein loss) over PFA tissues (30–40%

protein loss)²⁰. GA-preserved tissues exhibit high autofluorescence [G] in the 500–600 nm range²¹ and retain mRNA molecules less well than PFA, and therefore GA is most suitable for strengthening delicate tissues or retaining poorly expressed proteins if imaging proteins in downstream imaging pipelines.

Chemical crosslinks formed by aldehyde fixatives can block antibody access to epitopes and reduce the effectiveness of downstream immunostaining (see below). Enzymatic digestion and/or a high heat and pressure treatment is routinely used to permeabilize tissue, reduce aldehyde crosslinks and restore antibody accessibility in thin histology sections — a process referred to as antigen retrieval. However, applying these treatments to intact tissue is detrimental to its integrity. Interestingly, a combination of detergent, solvent and mild heat has been shown to have an antigen retrieval effect when included in a tissue clearing protocol (FLASH, see ref¹²).

Using polyepoxide molecules as fixatives, such as polyglycerol-3-polyglycidyl ether (P3PE), enables a high degree of intermolecular and intramolecular crosslinking for several biomolecules. Use of P3PE in the SHIELD protocol produces tissue that is more rigid and mechanically stable than tissue fixed with PFA or GA⁴. Additionally, this fixation method produces cleared tissue with significantly higher protein retention, mRNA retention and fluorescent protein emission than GA-fixed and PFA-fixed tissue. P3PE concentrations can be adjusted between 2–20% w/v, with epitope availability and chemical/label penetrability enhanced at low concentrations and protein preservation, mRNA preservation and tissue rigidity enhanced at high concentrations. In addition to the above fixation methods, 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) has shown utility for fixing genetic material in place⁴.

It is best to evaluate several fixation protocols, including various concentrations and durations of each fixative, to determine which best retains your targets of interest while still allowing for sufficient and timely clearing. For example, thicker or denser tissues may require use of the SWITCH method coupled with relatively low fixative concentrations to enable uniform fixation within a reasonable amount of time.

Optional pre-treatments

Preparing a cleared sample most often involves removing lipids from the sample and placing it in a solution matched to the RI of the remaining tissue components (usually protein, RI > 1.5). Refractive index matching alone may be adequate for clearing thin (< 0.2 mm), non-pigmented samples; however, most tissues require additional processing. The pre-treatment steps below are optional. Some are needed for specific tissue types, while others have a more general application. Experimentation on smaller samples is advised to identify which are necessary for a particular tissue of interest.

Decolorization—Many tissues contain natural endogenous pigments that can absorb light and prevent effective clearing and imaging, and in large tissue volumes even sparse pigments are obstacles to acquiring high-contrast images^{2,3,10,22}. Heme is the most abundant chromophore found throughout tissues and is associated with hemoglobin and myoglobin pigments found in the blood and muscles, respectively. Blood can be

removed after an animal is sacrificed by transcardial perfusion of large volumes of isotonic buffer; however, perfusion with PBS alone is insufficient to remove all heme from blood-rich and muscle-rich tissues such as the kidney, heart, muscle and liver. Extensive chemical screening has identified that amino alcohols effectively elute heme from both blood and muscle²³; these chemicals (specifically Quadrol) also assist in lipid extraction, are compatible with fluorescent proteins and are a key component in the CUBIC tissue clearing protocols^{11,24}. The amino alcohol N-methyldiethanolamine (NMDEA), in conjunction with 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS) detergent, can effectively remove heme while permeabilizing and delipidating tissue⁶. N-alkylimidazole also has high utility for heme elution²⁵. Sacrificed animals can be perfused with any of these heme solvating chemicals, or individual tissues — including human tissue — can be incubated with them after dissection.

Additional endogenous chromophores may be present depending on the tissue. Pigments of the compound eye such as pteridine and ommochrome and melanin in skin can be particularly difficult to remove. The amino alcohol 2,2',2'',2'''-(Ethylenedinitrilo)tetraethanol (THEED) has been shown to be effective at removing pigments of the compound fly eye while preserving GFP and mCherry emission (FlyClear¹³).

A more generalized method for decolorization is to break the chemical bonds of pigments using peroxides, a process often referred to as bleaching^{15,26–29}. Bleaching gives powerful decolorization performance, although it requires optimization of the peroxide concentration and treatment duration to avoid excessive damage to the tissue. Further, as peroxide treatment destroys the central chromophore of fluorescent proteins and bleaches some organic dye molecules, downstream immunostaining is required to visualize quenched fluorescent proteins and any fluorescent labelling must be carried out after the bleaching step.

Quenching autofluorescence—Autofluorescence can decrease the signal-to-noise ratio (SNR) and reduce contrast when imaging. To avoid this, samples can be photobleached to eliminate autofluorescence prior to staining. Photobleaching can be accomplished by exposing the tissue to broadband light for an extended period; for example, exposure using a custom device containing broadband visible LEDs in a cold room for up to three days can reduce autofluorescence in thick tissue samples^{30,31} (the cold room prevents tissue from overheating and degrading). A key disadvantage to this method is the need for immunostaining, as genetically encoded fluorescent proteins will be photobleached. A more common and efficient method to overcome autofluorescence is to select red and far-red dyes that fluoresce outside the 500–600 nm range where most autofluorescence occurs. If red and far-red dyes are selected, autofluorescence in the 500–550 nm range can be imaged to derive useful information, for example to identify anatomical structures or register the dataset against other samples or atlases (see Results section).

Hydrogel embedding—Hydrogel embedding techniques were originally developed to improve the mechanochemical properties of fixed tissue over standard PFA fixation. Hydrogel embedding technologies for tissue clearing employ the polymerization of

monomer solutions containing varying compositions of acrylamide (AA), sodium acrylate (SA), and bis-acrylamide (Bis) molecules to generate polyacrylamide (pAAm) meshes that physically or chemically fix biomolecules in place. In recent years, researchers have harnessed the unique properties of hydrogels — such as their expandability and elasticity — to further enhance the applications of hydrogel embedding technologies. The fundamental principles and novel methods in tissue-hydrogel transformation technologies are covered in-depth in a recent review paper on the subject³².

CLARITY was the first widely-used hydrogel embedding technique³³. CLARITY and its variants covalently anchor endogenous tissue biomolecules to a pAAm mesh that is generated from AA, PFA, and Bis (Fig. 2). Bis serves to chemically crosslink separate acrylamide chains to form the pAAm mesh, while PFA serves to crosslink tissue molecules to the mesh and decrease protein loss. Bis and PFA can be excluded from the monomer solution to increase pore size for enhanced clearing and immunostaining, with a reduction in mechanochemical preservation and stability³⁴.

More recently, the elastic properties of pAAm hydrogels have been harnessed to increase the toughness of hydrogel embedded tissue in a process called ELAST³¹. By significantly decreasing the Bis crosslinker concentration, more physical entanglement and less covalent crosslinking of long pAAm chains occurs. This decreases the rigidity and brittleness of the resulting tissue-gel, making it reversibly stretchable and more resistant to wear and tear when handled.

Certain hydrogels, such as the pAAm gels used in CLARITY, polymerize best when maintained in an oxygen-free environment and their polymerization is often activated by a temperature sensitive initiator. Custom (or commercial) equipment can be used to perform these steps; for example, a basic setup involves placing a conical tube or plate warmer inside a vacuum chamber. The samples are placed under vacuum prior to heating the sample to 37°C for approximately 3 hours.

Another interesting property of hydrogels is their ability to expand. These expansion properties have been harnessed to physically expand hydrogel-embedded tissue-gels and thus increase the achievable spatial resolution of conventional light microscopy³⁵. Although expansion occurs during the final RI-matching step, the fixed tissue must first be embedded in a polyacrylamide-polyacrylate gel that will later facilitate expansion in each dimension. Additionally, protein digestion or denaturation is required to permit final tissue expansion. This has been achieved using Proteinase K (ExM³⁵), LysC (Pro-ExM³⁶), SDS (MAP³⁷) or imidazole (CUBIC-X³⁸). LysC, SDS and Triton X-100 are now the preferred strategies as these reagents are the least destructive to tissue and better preserve biomolecules for post-expansion labelling^{36–38}. Recent protocols expand tissue up to 20x (iExM³⁹) and allow for fine tuning of the degree of expansion (ZOOM⁴⁰).

Decalcification—The clearing of limbs or whole vertebrates requires the decalcification of bone as calcified bone is a strong light scatterer^{2,3,10,22}. 50–70% of bone consists of carbonated hydroxyapatite (HAp) crystals distributed throughout a collagen matrix. Elution of calcium ions from HAp reduces light scattering and renders bony material transparent.

Decalcification can be performed by submerging the sample in an acidic solution (as illustrated by the classic classroom experiment in which immersing an egg in vinegar renders the shell transparent); however, this causes substantial soft tissue damage. Neutral buffered saline solutions that contain the calcium ion chelator EDTA are preferred⁴¹ and EDTA has been used in many tissue clearing methods, including CUBIC-B²⁵, PEGASOS⁴², PACT-deCAL⁴³, bone CLARITY⁴⁴ and vDISCO⁴⁵. Interestingly, imidazole may further enhance the decalcifying and clearing effects of EDTA on bone tissue²⁵.

Delipidation

Removing lipids (delipidation) is critical for successful tissue clearing as lipids act as an optical barrier to light and a physical barrier to fluorescent probe delivery. Delipidation can be performed using solvents, detergents or a combination of both, and achieved passively or with the aid of an electric field (Fig.3). Delipidation of hydrogel embedded samples is primarily performed with detergents in aqueous buffer to prevent dehydration of the gel.

Solvent-based delipidation—Solvent-based delipidation involves submersing the tissue in a series of solutions (Fig.3). First, an initial dehydration step serves to permeabilize cell membranes, initiate some lipid solvation, and enhance tissue bleaching if the final dehydrating agent is combined with peroxides^{46,47}. Dehydration is performed by moving the tissue through solutions with an increasing concentration of a water-miscible organic solvent. This process can cause tissue shrinkage, although increasing the number of steps in the gradient can minimize this effect and better preserve morphology⁴⁶. Preservation of nominal sample size and morphology is often essential for registration to an anatomical atlas (see below).

Historically, tetrahydrofuran (THF), ethanol or methanol have been used for dehydration, although these reagents have notable disadvantages. Rapid dehydration using THF can cause unwanted morphological changes⁴⁶. Further, tissues with a high lipid content do not clear well when dehydrated with ethanol as it does not have the same delipidating effect as methanol or THF (see below)⁴⁸. Similarly, dehydrating tissues with THF, ethanol or methanol will quench fluorescent proteins, as removing water molecules from the chromophore of fluorescent proteins causes a conformational change that eliminates fluorescence emission. Methanol and ethanol quench fluorescent proteins within hours^{46–48}. Fluorescence can be maintained in THF for a few days if the pH remains above 9.0 (as in fDISCO⁴⁹), dehydration and clearing is performed at 4°C (fDISCO⁴⁹) and/or by adding butylated hydroxytoluene (sDISCO⁵⁰). Because of this quenching effect, THF, ethanol and methanol are rarely used to clear tissues expressing fluorescent proteins and these tissues are instead best dehydrated using 1-propanol or tert-butanol (FluoClearBABB⁵¹). If tert-butanol dehydration is performed step-wise (uDISCO⁹), in alkaline conditions (a-uDISCO⁵²) or mixed with Quadrol and polyethylene glycol (PEGASOS⁴²), GFP fluorescence can be preserved for over a month. Methanol, ethanol and THF are still used for dehydration of tissues that do not express fluorescent proteins as synthetic dyes are usually stable in organic solvents. Methanol, ethanol and THF provide the strongest dehydration and delipidation effects, which is important for achieving large tissue transparency. Tissue dehydration can

affect the interactions of antibodies with their epitopes, either enhancing or preventing binding, and this effect must be independently tested for each novel antibody used⁴⁷.

All of the dehydrating agents described above perform some degree of delipidation. However, this is not sufficient to produce a well-cleared final sample. Therefore, an additional incubation in dichloromethane (DCM) — a water-immiscible solvent — is performed to complete lipid removal from the tissue^{53,54}. DCM induces the creation of inverse lipid micelles, which diffuse out of the tissue (Fig. 4). Owing to the high immiscibility of water in DCM, care must be taken to ensure that all water has been removed from the sample before submerging it in this solvent.

Detergent-based delipidation for aqueous-based clearing—As fixation is performed in aqueous buffers supplemented with fixative, these samples can be directly delipidated using detergents (Fig.3). The most commonly used detergents are SDS, Triton X-100 and CHAPS. Detergents act by disrupting lipid bilayers and forming micelles that can be transported out of the tissue (Fig.4). Alkaline pH, higher temperature, and longer incubations can all enhance the degree of detergent-based delipidation and result in more chemically and optically permeable tissue. Delipidation buffers generally contain between 4%–10% detergent and are incubated at or above room temperature to prevent the solution from becoming supersaturated. Incubations can take from hours to months depending on the tissue type, size, method of micelle removal and the desired level of clearing^{5,6,11,24,25,33,55,56}.

Detergents other than SDS and Triton X-100 have been used for delipidation, including sodium lauryl sulfate (SLS) in the CLARITY³³ workflow, sodium dodecylbenzenesulfonate in CUBIC-HL²⁵, Tween-20 in FocusClear¹ and 1,2-Hexandiol²⁵. CHAPS (used in the SHANEL workflow⁶) has also shown utility in very large samples as it generates smaller micelles than other detergents that can diffuse out of the tissue more quickly (Fig. 4).

Combined solvent and detergent delipidation—Physiological lipids have varying levels of polarity. Some lipids are better removed by detergents, others are better extracted through dehydration and delipidation by solvents. Lipid rich tissues can therefore benefit from the use of both detergents and solvents for delipidation (Fig.3). Adipose tissue, for example, can be delipidated by first incubating with 0.1% (v/v) Triton X-100 followed by a methanol dehydration and further lipid extraction with DCM, as shown in the Adipo-Clear⁵⁷ workflow. Likewise, insufficient delipidation by ethanol and DCM can be compensated for through an initial incubation with CHAPS, a strategy used in the SHANEL⁶ workflow.

Delipidation using electric fields—The speed of delipidation in large, thick tissue samples is limited by the rate that SDS and lipid micelles diffuse in and out of the sample, respectively. This rate can be increased using electric fields (Fig. 3). Unidirectional electrical fields were first used for this task³³ and are now implemented in many commercial electrophoretic tissue clearing (ETC) devices. ETC devices can speed delipidation dramatically, accomplishing in a day what can take weeks by passive diffusion; however, unidirectional electric fields can produce unequal lipid extraction as lipid micelles nearest the anode are forced to traverse through the entire sample. This differential

delipidation, along with the electrical force exerted on the tissue, can cause malformation of the tissue during the ETC process. A more efficient way to delipidate a sample is to use a rotating electric field or rotate the sample within a unidirectional field. This process, referred to as stochastic electrotransport, enhances the diffusion of highly electromobile species such as lipid micelles, while keeping species with low electromobility such as other tissue biomolecules relatively stationary⁵⁸. Hydrogel embedding, and crosslinking of biomolecules of interest to the gel is a requirement for ETC in order to maintain tissue stability and to retain electromobile molecules of interest. The original publications contain design suggestions for ETC³³ and stochastic electrotransport⁵⁸ hardware and both can now be purchased commercially.

Labelling

Extracting biologically meaningful information from transparent tissues requires specific and uniform fluorescent staining of target molecules such as nucleic acids or proteins. Small molecule dyes such as the nuclear dyes DAPI, propidium iodide, RedDot2 and SYTO^{3,24,33,42} and membrane integrating dyes such as fixable DiI can diffuse or be perfused through biological tissue. Other fluorescent dyes can be targeted to molecules of interest using short peptide tags⁵⁹ or antibodies. Passive diffusion of small molecules and antibodies is slow and the diffusion time scale correlates quadratically with the tissue thickness; when staining tissue at the organ scale, it can take days for the label to completely permeate to the tissue's core⁶⁰.

There are many well documented variables that need to be optimized for any immunostaining protocol including fixation conditions, antibody concentrations, temperature, and time⁶¹. These optimizations are best carried out on thin (~200 µm) tissue sections and imaged on a confocal microscope to ensure complete penetration through to the middle of the section. Canonical labelling with antibodies requires the use of two large molecules: a primary antibody that recognizes the molecular target and a secondary antibody that carries the fluorophore^{24,45,47}. Labelling with multiple primary antibodies creates further complication as the optimal conditions for each antibody may be different. Once uniform staining of a thin section has been established, the antibody concentrations can likely stay the same for thicker tissue and the volume of antibody solution is increased. Each antibody also has a unique rate of denaturation that increases with temperature. Highly unstable antibodies must be incubated at lower temperatures and replenished if staining protocols last multiple days.

The largest barrier to scaling up an immunolabelling protocol is that the combination of the slow movement and high affinity of the antibody label results in the aggregation of the antibody at the sample's surface⁵. Below, we outline several methods to prevent surface aggregation by reducing the distance the label must travel or temporarily manipulating its affinity for its target. Together, these methods permit more rapid and homogenous labelling of large, cleared tissues.

Reduce sample size—The simplest method to improve labelling speed is to reduce the size of the sample. Before initiating a clearing experiment, the tissue should be trimmed

as small as possible. Dehydration can decrease sample volume up to 60% if used, with the rate that the tissue is dehydrated influencing the level of shrinking. Some researchers have embraced this shrinking effect to allow for more rapid labelling and imaging⁹. It should be noted that shrinking the sample will also decrease the achievable resolution when imaging as some components of the tissue will have moved closer than the diffraction limit of a light microscope⁶². Reversible shrinkage can be exploited to speed up the labelling process; if a tissue is infiltrated with a tough and elastic hydrogel, the sample can be reversibly compressed to one-sixth its original size³¹. Labelling can be performed in the compressed state, decreasing the distance that molecular labels must diffuse to their targets. Similarly, high-ionic-strength solutions can be used to temporarily shrink non-elastic tissue during the staining process⁵. If reversible shrinkage is used for labelling, the sample is returned to its original size prior to imaging to avoid compromising the final image resolution.

Reduce label size—Smaller labels can diffuse faster through a tissue and the use of small molecule fluorescent dyes with affinity for specific molecular targets is advantageous over antibodies when possible. Nanobodies (~15 kDa) are one-tenth the size of a conventional antibody (~150 kDa) and show high tissue penetration and high thermal/chemical stability; these labelled single-variable-domain antibodies from Camelidae or shark species⁶³ have been shown to rapidly diffuse through fixed and cleared tissue, decreasing labelling time up to 5-fold^{45,60} over antibodies. Although labelling with nanobodies is fast, it should be noted it can also result in a relatively dim fluorescent signal. Standard immunolabelling protocols use polyclonal secondary antibodies, which can bind at multiple regions of a single primary antibody thereby amplifying the fluorescence signal. Nanobody labelling is a single step process where only one nanobody binds to each biomolecule of interest; therefore, no signal amplification occurs. Additionally, commercially available nanobodies are still limited to a small range of targets, although this range is rapidly increasing.

Increase porosity—Lipid membranes in biological tissue prevent the diffusion of imaging agents and immunolabelling is therefore often performed after delipidation. However, certain pre-treatment or delipidation steps may deplete or alter the target of interest, requiring immunolabelling to be performed first. In addition to delipidation by detergents^{6,23,33,34,64} or solvents⁶, weak fixation⁶⁵, partial protein digestion^{5,65-67} or increasing the pore size of hydrogels can improve label diffusion³⁴. Further, it is possible to harness the natural porosity of tissue for label delivery; for example, transcatheter perfusion of fluorescent labels during pre-treatment is an excellent method for labelling whole animals^{11,34,45}. Unfortunately, this perfusion technique is technically demanding and requires large amounts of label.

Manipulate label affinity—The binding affinity of fluorescent labels can be manipulated to allow them to diffuse freely throughout the tissue before their binding properties are reactivated in a way analogous to the SWITCH method described above. For ionic bonding stains such as nuclear stains, high salt concentrations can be used to inhibit binding until homogenous tissue distribution is achieved^{5,23}. Alternatively, antibody binding can be temporarily inhibited (“turned off”) by introducing urea, Quadrol or SDS to a physiological buffer solution, or increasing the temperature or pH^{5,20,23,55}. Once the

antibody has distributed itself throughout the tissue, the tissue is returned to the standard buffer solution and binding will be activated (“turned on”). Careful optimization of solute concentration, temperature and pH is required if using this technique as antibodies can undergo irreversible denaturation and lose their binding capability when exposed to the above conditions. Changes in temperature and antibody concentration and/or the addition of Quadrol or urea appear to effect antibody penetration. Conversely, changes to salt and detergent concentrations tend to influence binding efficiency^{5,23,55}. Finally, use of primary and secondary antibody complexes – pre-formed by co-incubating these antibodies prior to applying them to the sample – only requires a single diffusion event and results in a two-fold decrease in labelling time.

Apply an outside force—Certain fluorescent dyes and antibodies are electromobile and stochastic electrotransport can be used to increase the speed that these labels move through a cleared tissue sample^{20,58}. When a SWITCH method is combined with stochastic electrotransport, multiple, rapid rounds of labelling can be performed. This methodology (eFLASH²⁰) enables a highly multiplexed approach to investigating cleared tissues, where tens of target molecules can be investigated in a single sample rather than the standard 3–4. A custom (now commercially available) active immunolabelling device is needed to speed labelling; the device consists of a sample chamber across which an electric field can be applied and the sample or the electric field rotated. Active cooling is required to prevent heat damage to the sample. Another option to further increase the labelling speed is to focus the electrical field by adding magnets to the sides of the sample chamber. This technique, referred to as electromagnetic focused immunohistochemistry (EFIC) or magnetohydrodynamic force (MDH), drives the labels into the tissue and prevents them from flowing around the outside of the tissue^{68,69}.

Refractive index matching

The last step of any clearing protocol involves immersing the sample in a final clearing solution. This solution must match the RI of the remaining biological components — primarily protein — and be miscible with the solutions used prior to it. The optimal RI for final clearing should be 1.50–1.60 to best match the remaining protein in the sample after delipidation (with an exception for expand samples, see below). Care must be taken to ensure fluorescent labels are compatible with the chosen solution. Post-fixation in 4% PFA may be required to better retain antibody labels, although this will prevent de-staining and is therefore incompatible with multi-round labelling. A wide variety of solutions have been used and are categorized as aqueous-based or solvent-based; formerly, aqueous solutions better preserved fluorescent protein emissions, whereas solvents better cleared tissue due to their higher RI. These differences are much less pronounced owing to recent advances.

Aqueous-based solutions—Many hydrophilic tissue-clearing reagents have been reported, from simple sugars such as sucrose and fructose to advanced chemistries such as the medical grade GI imaging reagent iohexol^{2,3,10,22}. RI-matching solutions must achieve an RI of 1.50–1.60 in non-expanded samples; however, due to solubility limitations in water, many aqueous solutions of these reagents can only reach an RI of ~1.45. A new generation of aqueous solutions are now available that can match the clearing potential of solvents;

these use high-RI chemicals that have high solubility in water, including antipyrine/nicotinamide (CUBIC-R⁷⁰, RI = 1.52–1.54), N-methylacetamide/Histodenz (Ce3D⁷¹, RI = 1.50) and 2,2-thiodiethanol (TDE)/dimethylsulfoxide (DMSO)/iohexol (dPROTOS³¹, RI = 1.52).

It should be noted that the expansion or contraction of a sample will affect its RI. Aqueously cleared samples do not contract and therefore do not require as high an RI for their final clearing solution (RI~1.52) compared to dehydrated solvent cleared samples (RI~1.56). In the most extreme examples, where hydrogel embedded tissues are expanded 4–20 times, the sample is primarily composed of water and has an RI near 1.33; no specialized clearing solution is needed for these types of expansion methods.

Solvent-based solutions—If samples have been dehydrated and delipidated in solvents, the remaining tissue can be rendered optically transparent by impregnating it with an additional high RI solvent. Formerly, this has included benzyl alcohol benzyl benzoate (BABB) or dibenzyl ether (DBE)^{53,54}. Like the dehydrating solutions described above, BABB and DBE can quench fluorophores contained in the sample. Several approaches can improve fluorescent protein preservation during BABB or DBE incubation, including maintaining a basic pH by supplementing the final clearing solution with triethylamine⁵¹ or Quadrol⁴² and preventing peroxide formation by adding DL-alpha-tocopherol⁹ or propyl gallate⁵⁰. Additional fluorescent protein preservation has been achieved through the incorporation of diphenyl ether (DPE⁹) or polyethylene glycol (PEGMMA⁴²). Supplementing BABB and DBE with the fluorescent protein protectants described above can preserve fluorescence emission for weeks to months in the cleared tissue^{8,42,50,51}.

The toxicity of solvents, particularly BABB, remains a concern for some researchers and multi-user core facilities where cleared samples are often imaged. Ethyl cinnamate and 2,2-thiodiethanol (TDE) are less toxic alternatives that can be used to clear tissue. Ethyl cinnamate has shown equivalent clearing effects and prolonged fluorescent protein emission relative to BABB and DBE when clearing mouse organs⁷². A 97% TDE water solution results in a similar refractive index to BABB and DBE; however, many fluorescent dyes become unstable in TDE at this concentration^{73–75}. Therefore, TDE is better suited to increase the RI of aqueous clearing solutions when used at concentrations below 70% (v/v) as described above.

Imaging

Cleared tissue samples require imaging techniques capable of optical sectioning [**G**] over scales of millimetres to centimetres. Some technologies, such as confocal microscopy and 2-photon microscopy, are widely accessible in individual labs and institutional core facilities whereas others, such as light sheet microscopy and optical projection tomography, are more limited. Additional components such as special objectives or sample holders may also be required (see ‘Results – Complications’ below). Table 2 displays a comparison of several microscope modalities commonly used for imaging cleared tissue samples.

Point-scanning and spinning disk confocal microscopes are the most widely available optical sectioning microscopes. Upright point-scanning confocal microscopes equipped with high

numerical aperture (NA), long working distance dipping objectives [G] produce the highest quality images of cleared tissue. Inverted point-scanning confocal microscopes are limited in imaging depth and resolution (Table 2) owing to the need to use long working distance air objectives that cause sample-induced spherical aberrations when imaging thick, cleared tissues⁷⁶. Spinning-disk microscopes allow fast acquisition of 3D volumes, but quickly experience a loss of confocality as they image deeper into tissue and are often optimized for use with high-NA objectives with working distances less than 200 μm . 2-photon excitation is advantageous for imaging cleared tissue as it can extend the imaging depth over standard confocal microscopy for less optimally cleared samples and eliminates out-of-plane photobleaching when imaging large axial volumes. Unfortunately, both point-scanning confocal and 2-photon microscopes are slow and can require tens of seconds to image a single field of view, and imaging samples greater than 1–3 mm^3 in size requires a faster approach.

Light sheet microscopes have quickly become the preferred system for imaging cleared tissue. This is primarily owing to their speed, which arises from the perpendicular orientation of the excitation and detection objectives. Cleared tissues are well-suited to light sheet microscopy as the lack of scattering inside the tissue allows light sheets to be stably projected through the tissue and the emitted fluorescence can traverse large spans of tissue to the detection objective without scattering. Light sheet systems can be optimized for high resolution imaging using short, thin excitation sheets or lattice light sheets [G], although these configurations have steric hinderances [G] that prevent the imaging of cleared tissue greater than 100–200 μm in size. Systems can be optimized for larger samples using long, thick excitation sheets and low-resolution (NA < 0.5), large-field-of-view detection objectives (for example, the ultramicroscope⁴⁸), where the light sheet (5–10 μm) determines the thickness of the optical section. Whenever we refer to light sheet microscopy in this Primer, we refer specifically to these high-capacity systems, which can image a cm^3 volume in minutes-hours depending on the desired resolution and sample properties.

Optical projection tomography (OPT) can also be used for imaging cleared samples. By rotating the sample during imaging, OPT can achieve isotropic spatial resolution [G]. OPT achieves a lower lateral and higher axial resolution in comparison to light sheet or confocal microscopes using similar optics. Imaging can be accomplished in minutes to hours if the sample fits in a single field of view. A limitation of OPT is that the sample cannot be larger than a few millimetres in each dimension. OPT has primarily been used for imaging small vertebrate embryos and intact rodent organs that fit these dimensions and only require low-resolution imaging (cellular resolution at best)^{77,78}.

Objectives

Two categories of detection objectives are used for imaging cleared tissue: clearing-specific, high-NA dipping objectives and low-NA air objectives. The decision of which objective to select is sample independent and solely based on the imaging system used and the level of resolution required. Dipping objectives are primarily used on upright point scanning confocal or multi-photon microscopes and are also be found in certain light sheet designs. Dipping objectives for cleared tissue imaging optimally have a high NA (> 0.9), long

working distance (> 5 mm), low magnification ($< 25\times$) and correction collars to allow imaging in solutions of varying refractive indexes. These objectives enable sub-cellular resolution imaging. Low-NA air objectives are more commonly used on light sheet systems and are not recommended for use with confocal or multi-photon microscopes as they have poor axial resolution and suffer from sample-induced spherical aberration. Preventing sample-induced spherical aberration requires methods that ensure the path length light travels through air is constant at all axial positions, for example by fixing objectives in place and translating the sample or placing a dipping cap over the objective⁷⁶.

Data handling

Tissue clearing experiments, especially when performed at scale, produce large amounts of data. For simple experiments such as imaging single fields of view in tissues < 500 μm thick, the microscope's acquisition computer or a standard desktop will suffice. However, imaging intact organs or animals is data intensive and can require extensive IT investments. Whereas confocal and multi-photon microscopes produce between 5–10 GB of data per hour per colour channel, light sheet microscopes can produce hundreds of GB of data per hour; indeed, a single sample labelled with 2–3 fluorophores and imaged using light sheet microscopy can produce over a TB of data. Researchers aiming to perform large-scale or high-throughput imaging of cleared tissue must carefully consider their IT infrastructure to account for these high rates of data production and it should be noted that the investment in IT infrastructure could equal or exceed the costs associated with purchasing the imaging system itself. The first consideration is whether to procure local or cloud storage. If a local data centre exists, expanding its storage capacity may be more cost effective than renting cloud space. Network upgrades may also be needed as the transfer of datasets 100 GBs to TBs in size is prohibitively slow over a standard 1 Gbit/s network. 10 Gbit/s or higher speeds are preferred. Installing microscopes, data storage and workstations in close physical proximity — preferable on the same network switch [G] — will increase transfer speeds and aid in network stability. If renting cloud storage, researchers may need to purchase dedicated bandwidth between their location and the cloud storage server to ensure uploads occur at practical speeds. It is essential to consider the data output of tissue clearing experiments from the early planning stages and ensure adequate IT infrastructure exists.

RESULTS

At the completion of a tissue clearing experiment, a researcher can expect to obtain a sample through which a fluorescence microscope can image beyond the standard limit of 50–100 μm imposed by light scatter. How much deeper will be a function of the tissue composition and the clearing protocol. It is common for large cleared tissues to retain a yellow tint owing to the continued absorption of blue and near-UV light by the remaining protein in the sample (Fig. 5) and a lack of this yellow colour in large samples may even indicate a loss of protein during the clearing process. The most common qualitative method for determining the effectiveness of clearing is to place the tissue over printed text and attempt to read the words below. A transilluminator with text or a grid printed on a transparent film can allow for better viewing. A more quantitative approach is to remove a > 1 mm section of cleared tissue and view it under a dark-field microscope; when compared to uncleared tissue, the

cleared tissue should not be visible under dark-field illumination owing to a lack of light scatter. The most quantitative method is to analyse proton concentration across a sample using MRI; again, the less contrast that is visible, the better cleared the tissue is⁷⁹.

The integration of tissue clearing and high-resolution light-sheet microscopy has enabled the volumetric visualization of large biological specimens while simultaneously creating challenges for the subsequent assessment of cellular content. Analysis of cleared tissue imaging data can be intensive and time consuming by eye and requires image analysis routines, algorithms and software. Many open-source and commercial tools now exist in this space. Some general software tools capable of handling large data files include the BigDataBrowser for Fiji/ImageJ⁸⁰, Napari, Arivis Vision4D, Imaris, and Aivia. Data analysis routines for cleared tissue follow five basic steps: correction of imaging induced artefacts; filtering to improve contrast; segmentation of objects and assessment of segmentation accuracy; registration to anatomical atlases (optional); and quantitative measurements of segmented objects. Quantitative measurements can take several forms, including object counting, volume measurements, distance measurements, determination of colocalization and tracing of filamentous objects such as neurons and blood vessels. We discuss these steps in more detail below, along with details of specialized software tools for each.

Image stitching

Cleared samples will exceed the field of view of most imaging systems. Therefore, imaging of the entire tissue requires the acquisition of multiple Z-stack “tiles” that must be fused together by image stitching. During stitching, neighbouring tiles are translated horizontally and vertically until a best fit is found within overlapping regions of the tiles; it is therefore important to acquire each tile with an overlap of 5–10% with all neighbouring views. This approach serves to correct two common artefacts: the misalignment of objects that cross tile boundaries (caused by sample movement or non-perfect stage translation) and the inhomogeneity of signal intensity across the field of view^{81,82}. Several software tools are available for stitching including the BigStitcher for Fiji/ImageJ⁸³, Arivis Vision4D and Imaris.

Correction of sample-induced artefacts

The two most common imaging artefacts associated with imaging cleared tissue are spherical and chromatic aberration. Spherical aberration occurs when light rays striking the outer portion of a spherical lens do not focus to the same point on the optical axis as the rays that enter at the centre of the lens. Spherical aberration in fluorescence microscopes manifest as a dim, blurry, and axially compressed image. Chromatic aberration occurs if different wavelengths of light fail to focus at the same point of the optical axis. This is the result of light passing through a material with high dispersion [**G**] and in a fluorescent image manifests as a shift of each colour channel —most obvious axially— relative to one another. Microscope objectives contain several lens elements that together reduce spherical and chromatic aberration; however, these lens elements are selected assuming that only immersion and mounting media with specified RI and dispersion values will be used (usually those of water, 80% glycerol or oil). If a clearing solution does not exactly match

the RI and dispersion values specified in the objective's design, sample-induced spherical and chromatic aberrations are produced⁷⁶. Spherical aberration can be avoided by using an objective specifically calibrated to the RI and dispersion of the clearing solution (see Objectives, above), fixing the objectives in place and translating the sample through the clearing solution, installing a dipping cap over the front lens of the detection objective or using software (available from certain microscope manufacturers or a recent tutorial for correcting spherical aberration⁷⁶) to correct the axial distortion after acquisition. Chromatic aberration in the axial dimension can be corrected by identifying a fiducial marker (such as a broadband autofluorescent structure or a position where all fluorophores colocalize) and obtaining a short Z-stack from which the axial shift of this singular object can be calculated for each colour channel. Z-stacks for each colour channel can then be offset (confocal) or the beam of a lightsheet microscope can be shifted axially for each colour channel according to this calculation. Alternatively, channel registration software can be used to correct chromatic aberration post-acquisition using a similar approach. If several fiducial markers can be identified, the program can automatically calculate the required axial shift and realign the dataset. Post-processing has the advantage of correcting for lateral chromatic aberrations as well. If the dataset does not contain easily identifiable objects, a tissue phantom comprised of agarose and a sparse concentration (1–30 per field of view) of 200 nm broad-band fluorescent or gold beads submerged in the final RI-matching medium of choice can be used. The required channel offset is identified by imaging the phantom and then applied to the dataset set through the hardware or software methods described above. Fluorescent beads (such as TetraSpeck from ThermoFisher) are preferred; however, they are unstable in most solvents, necessitating the use of gold beads.

Filtering to improve contrast

Microscopy data is inherently noisy and this noise determines how successfully objects of interest can be segmented and analysed in automated routines. Therefore, some degree of filtering is often required prior to segmentation. Because cleared tissue datasets are large, it is important to select filtering techniques with the least computational burden. A simple median filter with a diameter (in pixels) set to the same size as the objects of interest in the data set is often sufficient. Where a median filter is insufficient or the data was acquired with low-NA objectives via confocal microscopy or thick light sheets, deconvolution may be required to eliminate out-of-focus light and improve optical sectioning. Nearly all open source and commercial image analysis software packages offer these methods.

Segmentation of objects

Segmentation refers to the grouping of pixels or voxels from a data set into objects. Once defined, these objects can be 3D rendered free of background noise or measured in numerous ways to obtain quantitative data. The simplest method for segmentation is to choose a single intensity threshold and count all pixels above that value as being part of one or more objects. Any pixels below this threshold are considered background. When applied to an entire image, this method is referred to as global thresholding and can be performed using the open source^{9,46} and commercial software^{45,84} packages described above to segment objects in cleared tissue volumes. The most common object segmented in cleared tissue is nuclei. In most tissues, nuclei are spaced far enough apart to permit highly

accurate segmentation even when general nuclear stains that label all cells are used^{7,22}. Fig.6 displays an example of single-cell counting in several different mouse organs, which were cleared with the CUBIC-R+ protocol and imaged by light-sheet microscopy; here, over 75 million cells are identified in each organ by using a 3D Hessian-based difference of Gaussian algorithm²⁴. The accuracy of nuclear segmentation can be further improved by only labelling a subset of cells, which can also produce biologically relevant insights when linked to a physiological readout such as identification of active neurons (via c-fos staining) after providing a stimulus to an animal⁴⁶. Segmentation of individual cell bodies labelled with cytoplasmic or membrane expressed fluorescent proteins is also possible if the fluorophore is sparsely expressed¹⁷.

Global thresholding is difficult to perform in large cleared tissue as the signal-to-noise ratio can vary dramatically from region to region within the sample, making selecting a single intensity volume that can accurately separate signal from noise impossible. In recent years, the application of machine learning-based algorithms to segment objects of interest in microscopy images has provided robust methodologies for cellular profiling⁸⁵. At least two studies have shown that machine learning algorithms are superior to global thresholding when used to detect, segment and quantify cells in large cleared tissues^{6,86}.

Quality control for automated segmentation—It is impossible for a human to assess the accuracy of segmentation by eye throughout a whole dataset. However, it is possible to perform quality control on portions of a dataset to infer the overall accuracy. For example, a researcher could count sparse objects by eye within multiple, small, representative regions throughout the sample. The automated segmentation algorithm can then be run on the same regions and the results compared. Counting by eye quickly becomes impossible as objects become denser and “by hand” segmentation can be used, where a researcher makes annotations on each object on sequential 2D planes by circling or painting with a mouse or pen that are then combined into 3D objects. Again, a series of smaller regions are annotated by hand, counted, and the results are compared to those obtained by automated segmentation. All open source and commercial software programs identified above have annotation functionality. Automated segmentation should be > 90% accurate and preferably > 95% accurate when compared to by-eye or by-hand methods.

When approaching machine learning for the first time, a researcher should determine if pre-trained algorithms already exist for their structure of interest (for example, StarDist⁸⁷ for detecting nuclei). Due to the unique labelling patterns of individual experiments, it is likely that such resources will not be available and in these cases, multiple user-friendly options exist for training machine learning algorithms using the annotate-by-hand approach described above, including Trainable Weka Segmentation (ImageJ/Fiji), ilastik, Intellis (Zeiss), Vision4D (Arivis), and Aivia. When using these types of painting approaches for training, it is important to paint the edges of objects so that the algorithm is trained using pixels at object boundaries. At least two objects, or classes, must be trained: “Object1” and background. It should be noted that many of these software packages are not compatible with the large data sets obtained by imaging cleared tissue and custom coding is often required. First, training data must be established by manual annotation, as described above,⁸⁶ or by creating synthetic training data that mimics the structural features to be

imaged⁸⁸. Enough training data must be produced so that a portion can be reserved to test the algorithm's ability to achieve satisfactory results with data it has not seen previously. Secondly, a machine learning approach must be chosen. Random forests [G] are the simplest approach and require the least computing power, although they are less flexible than complicated neural networks⁸⁹. In this approach, the only parameter to set is the number of decision trees, with more decision trees improving the performance of the model but requiring additional computing power and processing time. Neural networks require in-depth understanding to correctly design their architecture including the number of layers, number of neurons in each layer, activation functions, and initialization rates; therefore, an equal or greater number of hyperparameters must also be set for the training algorithm^{85,90}. Therefore, customized neural networks should only be attempted by those with expertise in deep learning and proper testing is essential. A good algorithm can correctly identify objects chosen at random from the training data with few false positives and few false negatives; that is, its precision (correct annotation rate among all the annotations it made) and its recall (correct annotation rate among all the elements in the dataset that should have discovered) are high. Although the manual annotation step can be time-consuming, it is a worthwhile investment. A well-established algorithm facilitates the 3D analysis of large specimens, leading to a better understanding of various biological and biomedical questions^{86,88}.

Quantitative measurements of segmented objects—Quantitative measurements performed in cleared tissue are similar to those performed in volumetric images of non-cleared tissue, although all measurements must be performed in three dimensions and the size of the dataset can cause an analysis routine to exceed the maximum capacity of many current image analysis software programs.

The least computationally complex and most-performed quantitation of 3D imaging data from cleared tissue is object counting. Once objects have been segmented (although this is not a requirement), they can easily be counted by scanning the dataset for areas of intensity maxima. Objects that are easily counted include nuclei^{23,46} and sparsely labelled cell bodies¹⁷. Volume measurements are easily calculated if the X, Y and Z dimensions of a voxel are known and this information is often contained in the metadata of the image file and automatically read by most image analysis software; the volume of an object can be calculated as the number of voxels in the object multiplied by the volume of the voxel. Several distance measurements can also be performed; for example, the centre of mass for each object can be identified and a region-growing technique — in which a virtual sphere is extended pixel-by-pixel until the sphere contacts the next nearest centre of mass — can be used to measure the distance between objects in a sample⁹¹. If two or more fluorescent labels are used, the region-growing technique can be specified to only detect objects in another channel, providing a readout of the distance between different molecules of interest. In cleared tissue, co-localization measurements usually require segmentation of objects of interest in two or more channels, locating their centres of mass and setting a colocalization limit – the maximum distance two centres of mass can be from one another and still be considered colocalized. This is usually set to 0.5–1x the diameter of the object⁹². Finally, filament tracing of blood vessels and neurons are popular quantitative measures. Filament tracing often remains semi-automated and requires substantial user input

to direct the propagation of these structures through poorly labelled or noisy regions of the image⁹³. Once filaments have been segmented, several measurements can be performed, including length, volume, orientation, tortuosity [**G**] and number of branch points. See the Applications section below for more detailed descriptions of common applications for each of these quantitative measures.

Registration to tissue atlases

Atlases are three dimensional maps that are used to delineate different anatomical or functional regions of an organ. In complex organs, the boundaries between these regions require hundreds of labels to be visualized⁹⁴. As incorporating these labels into an experiment is currently not achievable, determining the exact region of an organ in which a fluorescent signal resides requires registering the 3D dataset to an existing atlas^{24,38,95–104}. Atlases are used extensively in the analysis of cleared tissue 3D datasets; for example, several studies have mapped data from multiple cleared brains onto the Franklin and Paxinos Atlas¹⁷, the Waxholm Space Atlas^{23,100} and the Allen Brain Atlas⁴⁶. Open-source software for performing these types of registration are available including ClearMap⁴⁶ and large deformation diffeomorphic metric mapping (LDDMM)^{105–107}. When mapping data to an atlas there are two key considerations. First, the dataset must contain a fluorescence channel with a similar fluorescence intensity across the entire organ, most often a channel containing tissue autofluorescence or a general nuclear dye. Second, the resolution of the sample image must be similar to the resolution of the images in the atlas. A limitation of many established atlases is that their regional-level resolution, often developed using MRI or CT^{108,109}, is not sufficient for registering the cellular resolution of microscopy data. To address this, a single-cell resolution atlas was developed for the mouse brain (CUBIC-Atlas) along with a cloud-based framework to perform 3D analysis at single-cell resolution (CUBIC-Cloud)¹⁰⁴. High-resolution reference atlases of entire organisms are needed to assist whole-body clearing methods. Currently, manual annotation⁸⁶ of individual organs is being replaced with deep-learning-based organ segmentation¹¹⁰; this approach has potential to transform the analysis of whole-body scans, eventually leading to the generation of more accurate mouse body atlases.

Required computing specifications

High-end workstations are needed to perform the above analyses. For optimal performance, data must be rapidly accessible, ideally in a solid state drive (SSD); if SSDs are cost prohibitive, multiple hard disk drives (HDDs) in a RAID array can suffice. Large amounts of RAM (> 200 GB) can reduce the need for disc access, which is usually an image analysis bottleneck. Finally, the amount of GPU memory and number of GPU processing cores are also important. For image analysis routines that can be GPU accelerated such as CLIJ¹¹¹, DeepMACT⁸⁶, TubeMap¹¹² and others, these specifications will determine how rapidly the processing can be completed. Additionally, GPU memory will determine the resolution limits when displaying 3D data. For example, when running a deep learning-based algorithm such as DeepMACT, a workstation with 64 GB RAM and a 12 GB Nvidia Titan XP GPU is capable of processing a series of whole-mouse images in 45 min⁸⁶. Workstations equipped with 512 GB RAM and a 24 GB NVIDIA Quadro P6000 GPU have been used to process the TubeMap algorithm¹¹².

APPLICATIONS

Tissue clearing now provides a viable alternative to serial sectioning of large biological samples. In comparison to sectioning, tissue clearing is often a lengthier process owing to the requirement for incubations in various solutions and the additional time required for diffusion of labels; however, these incubations require very little hands-on time and no specialized skills. Clearing protocols that do not require specialized equipment allow for a high degree of parallelization that cannot be achieved with a single vibratome or cryostat. Further, tissue clearing avoids mechanical damage that can occur within the tissue during sectioning and the heavy computational burden of aligning, de-warping and assembling hundreds or even thousands of 2D images into a 3D rendering. Analysis of intact, cleared samples shows improved accuracy compared to quantitative stereology¹¹³ and traditional 2D histopathology¹¹⁴. Finally, clearing allows for studies across entire organs (such as long-distance nerve tracing¹⁷) or entire organisms (such as identification of cancer metastasis^{7,86}) that cannot be accomplished through serial sectioning.

Biologically, the applications of tissue clearing methods are wide-ranging. Tissue clearing has primarily shown use in the fields of neuroscience, oncology and cardiology, although nearly every organ and organ system in rodents has now been cleared, including the brain, intestine, spleen, lymph node, heart, kidney, lung, eye, and bone^{7,10,11,27,34,43–45,70,71,110}. Other model and non-model systems such as fish and salamanders have also been used^{15,115,116}. The ever-increasing size of *in vitro* models such as spheroids and organoids — now many millimetres in size¹¹⁷ — necessitates the use of tissue clearing to image them in their entirety. Finally, methods to clear living samples and enhance imaging depth for *in vivo* studies now show promise.

Applications in neuroscience

In many organisms including mice and humans, neurons are the largest cells in the body with processes up to several centimetres long. Their large size requires that they be investigated in complete volumes spanning a whole organ or even organism, and for most animals tissue clearing or whole animal sectioning¹¹⁸ is required (aside from in the most translucent aquatic creatures, as demonstrated in a recent pre-print article¹¹⁹). Nervous system tissues are abundant in lipids and relatively lacking in pigment and extracellular matrix proteins such as collagen and elastin, making them ideally suited to clearing as once the lipid is removed, only a sparse protein matrix is left behind. Solvent-based and aqueous-based techniques have both^{6,57} proven successful in clearing tissues of the nervous system; however, techniques that combine both may be best suited for regions of high lipid density such as the spinal cord and myelinated axon bundles in the brain.

A major goal in neuroscience is to dissect the complete wiring diagram of the brain, a field known as connectomics. Even after tissue clearing, the density of neurons remains beyond the resolution limit of fluorescence microscopes¹²⁰ and current practice is to create maps of inter-areal projections by sparsely labelling collections of neurons^{121,122} (Fig.7). Several techniques exist to sparsely label subsets of neurons throughout the brain, including genetically encoded fluorescent proteins under the control of a cell-type specific promoter and immunolabelling with known markers of specific cell types. Expression of fluorescent

proteins may involve a single fluorescent protein per cell type or use combinatorial expression of multiple fluorescent proteins to enable the identification of hundreds of individual cells (for example Brainbow^{123,124} and other similar approaches). Transgenic animals can be produced expressing fluorescent proteins or they can be expressed by infecting cells with a virus carrying DNA encoding a fluorescent protein. Rabies and AAV viruses are commonly used for transducing cells of the nervous system; for example, a combination of AAV viruses were directly injected into the brains of mice in order to determine patterns of dopamine neuron innervation in the midbrain¹⁷. A less invasive approach is to use AAV viruses that can be injected into the blood stream and infect specific cell types of the brain¹⁸. To aid in the analysis of sparsely labelled neurons in cleared tissue volumetric, neuronal tracing algorithms have been developed that can perform manual or semi-automated segmentation⁹³. Most promising are new, fully automated algorithms such as NeuroGPS-Tree, which has traced over 1000 neurons in thick, cleared tissue sections¹²⁵, and a software package called BIRDS that uses deep learning to perform long-distance neuronal projection tracing and parallel counting of cell populations in different brain regions¹²⁶.

Tissue clearing has also been used to map areas of brain activity after an organism performs a behaviour or is administered a stimulus. Brain activity can be inferred by immunostaining for the protein products of immediate-early genes such as *c-fos*. If *c-fos* is present in a neuron, this indicates it was recently activated. Using this method, brain region activity mapping was accomplished in cleared mouse brain, harvested after sensory whisker stimulation or observed parenting behaviours in mice⁴⁶.

Finally, tissue clearing shows potential to aid in the investigation of neuronal pathologies. The SWITCH clearing and immunolabelling protocol was recently used to show that amyloid-beta peptides (A β) that contribute to the onset and progression of Alzheimer's dementia can be labelled and imaged in a brain-wide manner¹²⁷. This methodology allowed for microscopic mapping of A β plaques and identified several novel sub-cortical hubs where A β accumulates.

To date, tissue clearing of brains has primarily focused on rodent tissue as human tissue presents several barriers to tissue clearing, including enhanced autofluorescence, over-fixation from long-term preservation and large physical size. Recently, the SHANEL and ELAST tissue clearing protocols have been adapted to overcome many of these barriers and permit investigation of large brain slabs or even whole brain^{6,31}. Numerous exciting applications exist in this domain and we refer readers to a recent review for further applications³.

Applications in oncology

Whole animal clearing using aqueous-based (CUBIC) and solvent-based (vDISCO) techniques has enabled the visualization of micro-metastases throughout an entire rodent at single-cell resolution^{7,86}. Additionally, by fluorescently labelling a therapeutic antibody, it is possible to detect the percentage of metastatic sites receiving treatment across the entire animal⁸⁶ (Fig.8).

In the field of cancer pathology, there is interest in determining whether three-dimensional samples can provide a more accurate diagnosis or prognosis than traditional two-dimensional surgical sampling techniques. A form of expansion microscopy, termed ExPATH, suggested that imaging of cleared biopsies from breast cancer patients was able to discriminate between early breast neoplastic lesions better than surgical sampling¹¹⁴. Large (30 × 20 × 5 mm) surgically resected breast tumours were cleared in less than 48 hours using the PathoDISCO solvent-based clearing technique¹²⁸; in PathoDISCO the key step is using 2,2-dimethoxypropane to react with water to create acetone and methanol, which rapidly dehydrates the tissue.

The microenvironment that surrounds a tissue is highly heterogeneous and influences its growth and development¹²⁹. Hydrogel embedding protocols have been shown to preserve important elements of the microenvironment surrounding developing breast tissue and allow for its investigation in 3D¹³⁰. In this manner, the CUBIC tissue clearing method revealed the role of TGF- β in the tumour microenvironment by profiling multicellular interactions in a mouse model of experimental lung metastasis.

Interestingly, studies investigating the administration of novel, non-lipid-based nanoparticles — which are being investigated as potential delivery vehicles for therapeutics — showed both solvent-based¹³¹ and aqueous-based^{132,133} techniques allow for the retention of these nanoparticles in the tissue. These studies demonstrated that tissue clearing can be used for measuring the biodistribution of novel cancer therapeutics.

Applications in cardiovascular research

Blood vessels are easily labelled prior to clearing through the perfusion of labelling reagents. Multiple labels have been used, including antibodies targeting molecules that line the vasculature¹³⁴, wheat germ agglutinin and Evans blue⁸⁸, or lipophilic dyes¹³⁵. It is important that any label contain a moiety that can react with fixatives such as PFA or else it will be washed away during subsequent clearing steps. Fig.9 displays several cleared tissues in which blood vessels have been stained. Tissue clearing allows for the study of vessel repair and remodelling during development or after injury in any organ throughout the body; of particular interest are studies of blood vessel rupture in the brain in the case of aneurysm and stroke, and infarctions of the heart¹³⁶. Cardiovascular studies investigating blood vessel structure and function are now aided by several semi-automated vasculature segmentation, and quantification methods have been developed in commercial¹³⁷ and custom¹³⁸ software to analyse small regions of interest (ROI) in cleared mouse brains. These methods are not sufficient to reliably detect capillaries with variable signal to noise ratio (SNR) in different brain regions or to systematically analyse the vascular alterations at the whole brain or organ scale^{88,112}. Machine learning-based strategies have overcome these hurdles and currently two machine learning-based methods (VesSAP⁸⁸ and TubeMap¹¹²) are available to reliably detect, segment and register images of whole-brain vasculature to the Allen brain atlas. Recently, tissue clearing has been used to map the parasympathetic and sympathetic circuits that control heart rate in mice¹³⁹, produce a single-neuron-scale map of the intrinsic cardiac nervous system of the rat¹⁴⁰, and quantify sympathetic hyperinnervation of murine hearts after infarction¹⁴¹.

Non-murine applications

Aquatic organisms—Genome modification technologies such as CRISPR have enabled biological investigation in species beyond traditional model systems. Clearing protocols must be adapted to each species as many express unique pigments throughout their tissues that aid in protecting them from dangerous UV radiation (such as protection of the haematopoietic niche by melanin in fish)¹⁴². Many pigments such as melanin, ommochromes and pterins are resistant to hydrogen peroxide bleaching and do not elute with heme in amino alcohols. The DEEP-Clear method has shown that combining hydrogen peroxide, THEED and acetone can bleach or remove most of the pigments from annelids, molluscs, bony fish and tetrapods¹⁵. Others have shown that the addition of decalcification and bleaching steps render crustaceans clear and may enable the creation of developmental atlases with single-cell resolution¹¹⁶. Finally, an enzymatic treatment with collagenase was added to a customized tissue clearing protocol for flatfish to show that thyroid hormone is involved in their eye migration during metamorphosis¹¹⁵.

Plants—The plant wall imposes an additional RI mismatch that is not present in mammalian tissue and must be addressed. Additionally, chloroplasts acts as a pigment absorbing blue and red light and generates fluorescence during imaging, which may or may not be desired depending on the experiment. Plant tissue has historically been cleared using a single solution of sodium hydroxide and chloral hydrate that breaks down cell walls and equilibrates the RI throughout the sample¹⁴³. Urea¹⁴⁴, xylitol powder and sodium deoxycholate¹⁴⁵ and NaClO¹⁴⁶ have been used as less-toxic alternatives to chloral hydrate. Interestingly, the general principle of RI matching has even been used to create transparent wood with the potential to replace glass as a building material¹⁴⁷.

Humans—Three-dimensional imaging of clinical samples could have additional diagnostic potential over two-dimensional histopathology investigations¹⁴⁸; however, imaging these samples is difficult owing to over fixation, high levels of autofluorescence, the need for exogenous fluorescent labels and the overall high density of tissue from older patients. For very dense tissue, images of routine, thin, pathology tissue sections can be improved via expansion microscopy¹¹⁴. For larger samples — including intact human organs — the use of relatively small detergent (CHAPS) and solvent molecules for delipidation are needed overcome the density of human tissue⁶. Elasticizing human tissue has been shown to improve probe delivery when the tissue is stretched, while simultaneously providing better preservation over time to allow for multiple rounds of labelling³¹. Conversely, human embryonic and foetal tissue is less dense and far easier to clear. Solvent-based clearing methods have been used to clear human embryos and foetuses from gestational weeks 6–14¹⁴⁹.

***In vitro* applications**—Tissue clearing protocols have become essential for imaging spheroids and organoids in their entirety as traditional light microscopy approaches cannot image more than 100–200 µm beyond their surface. Even the smallest organoids (on the scale of 50–100 µm) can benefit from simple RI matching. Human airway, colon, kidney, liver and breast tumour, and mouse mammary gland organoids of this size have been cleared with simple RI matching in high-concentration sugar solutions¹⁵⁰. More complex

tissue clearing methods are needed for larger and denser organoids such as 3D neuronal spheres (200–300 μm diameter)¹⁵¹ and retinal organoids¹⁵². Finally, for organoids many mm in diameter, the SCOUT protocol has been used to clear, label, image and correlate hundreds of phenotypes between control and experimentally treated organoids¹¹⁷. SCOUT was able to identify several effects of Zika infection on cerebral organoids approximately two millimetres in diameter at the single-cell level¹¹⁷.

***In vivo* and label-free applications**—The modular tissue clearing methodologies described in this Primer are targeted toward fixed, non-living tissues stained with fluorescent labels. Although not the focus of this article, researchers have demonstrated clearing strategies that increase the imaging depth for several optical technologies in *in vivo* samples, including single-photon and multi-photon fluorescence, second harmonic generation (SHG) and optical coherence tomography (OCT) (reviewed in ¹⁵³). To date, *in vivo* methods have primarily focused on the topical application of biocompatible high-RI solutions to the skin^{154–158}, skull¹⁵⁹ and superficial tumours^{160,161} to provide increased imaging depth in both animal and human systems. As fluorescent labelling of live human tissue can be difficult owing to the inability to use genetic modification, human samples are often imaged by label-free techniques such as SHG and OCT. The ability to perform deep, label-free imaging *in vivo* is intriguing as it may hold clinical potential for the diagnosis and understanding of certain pathologies such as tumour biology.

Applications for systems biology

Systems biology generally entails analysing hundreds to thousands of targets within a tissue. Often, labelling is the limiting step as it requires substantial time and, although studies exist where tens of organs¹⁷ have been imaged, it is more practical to perform systems-level investigations on smaller samples. Tissue clearing can be used on this smaller scale; for example, spatial transcriptomic techniques like MER-FISH and seqFISH use imaging to detect thousands of mRNA species in a sample^{162,163} and tissue clearing has been adopted with these techniques to enable three-dimensional spatial transcriptomics. Incorporation of tissue clearing with the MER-FISH technique allowed a read out of the positions of thousands of mRNA molecules in each cell of a tissue sample in 3D¹⁶⁴.

REPRODUCIBILITY AND DATA DEPOSITION

The reliability and reproducibility of tissue clearing protocols has dramatically improved in recent years. Solvent-based clearing methods now generally use chemical components that are affordable and effective. Aqueous-based techniques have been aided by commercially available equipment for hydrogel polymerization and ETC. Further, the availability of commercial pre-formulated clearing solutions has allowed for standardization of clearing protocols between research groups.

Fluorescent labelling in tissue clearing experiments still primarily relies on antibodies. Commercially available antibodies are prone to reproducibility issues caused by batch-to-batch variation, which are more apparent (specifically background caused by off-target binding) when labelling large tissues. Antibodies should be carefully selected based on

past results and validated on thin, uncleared tissue sections. An online repository for tissue clearing-validated antibodies can also be used.

Many journals and funding bodies encourage the deposition of raw data in publicly accessible searchable databases. Recently, imaging data repositories such as the Image Data Resource and the Cell Image Library have been made available¹⁶⁵ and authors are encouraged to deposit their raw data in a publicly accessible database if possible. In addition to raw images, essential metadata should be reported to ensure reproducibility and set expectations for a researcher who would like to replicate a tissue clearing experiment (Box 1).

Studies that involve clearing tens or hundreds of samples may produce hundreds of TBs or even PBs of data, and public image repositories are limited in the amount of data they can store. Additionally, available bandwidth and a researcher's proximity to the repository's server can prevent uploads from occurring in a reasonable amount of time. Powerful lossy and non-lossy compression algorithms such as JPEG XS, AVIF, and WebP can reduce data size with little reduction in image quality; these provide better quality and compression than using .jpeg or .png files.

LIMITATIONS AND OPTIMIZATIONS

There are some outstanding limitations of clearing techniques. First, the methods described in this Primer are most often applied to fixed tissue as *in vivo* applications of tissue clearing are still limited¹⁵³. Second, several factors restrict the throughput of tissue clearing experiments such as the limited sample capacity in tissue clearing equipment, high cost of antibodies and clearing solutions, the long time required for immunolabelling, lack of common conditions for antibody binding, sample capacity limitations in imaging devices, and limitations in the IT infrastructure for data storage and analysis. Although one could imagine developing an automated pipeline for clearing, labelling and imaging, this would require an extensive investment in parallel equipment for sample preparation and imaging. Third, imaging of large samples requires microscope objectives with equally large working distances. Most imaging systems designed for cleared samples use moderate-NA, low-magnification, long-working-distance objectives from zoom microscopes. In general, increasing the working distance of an objective decreases its NA and therefore its achievable resolution⁸¹. The goal of tissue clearing experiments is primarily to survey a large tissue or organ; however, if subcellular resolution is also required, secondary imaging of select areas may be better suited to a second, higher resolution imaging device. Finally, the maximum size of samples that can be cleared now exceeds the maximum sample size able to fit under a microscope^{6,113}. Before investigation of human organs can be fully realized, imaging devices able to accommodate samples of this size must be developed. It is important to note that every tissue is different and researchers should not automatically follow a previously published method; for example, protocols that clear high lipid content tissue such as brain are not optimal for pigmented, protein-rich kidney tissue. By viewing tissue clearing protocols as modular, a researcher can select and assemble an optimal pipeline for clearing their tissue of interest.

OUTLOOK

The past decade has seen dramatic innovations in tissue clearing methods. Clearing solutions are no longer discovered serendipitously; researchers with extensive knowledge of optical physics, chemistry and tissue engineering are designing tailored solutions that improve on past methods. A decade ago, tissue clearing techniques were complex, unreliable and each had obvious limitations. Now, through incremental improvements, both aqueous and solvent-based techniques are able to preserve emission from fluorescent proteins and dyes and their final clearing solutions have coalesced around an ideal RI of 1.52–1.56. Commercially manufactured systems for steps such as hydrogel polymerization and electrophoretic tissue clearing and labelling are now available from multiple manufacturers, as are microscopes and objectives that have been designed specifically for cleared tissue.

Barriers do remain despite the positive outlook. The use of genetically encoded fluorescent proteins is not always possible and immunolabelling techniques must still be used in many systems. No matter how quickly antibodies are delivered to their target, if the antibody is not specific or stable, the result will be a failed experiment. Further, the current variability in ideal labelling conditions between individual antibodies makes it unlikely that a researcher's unique panel of probes will all function under identical conditions. We hope that nanobodies will be able to solve the many variability and stability issues of antibodies, although they are not without limitations. First, the random attachment of a fluorophore to a nanobody can block its binding site, necessitating directed attachment of the fluorophore with genetically encoded click chemistry^{166,167}. Second, fewer fluorophores can be attached to a nanobody than an antibody owing to its small size, providing a weaker signal. Third, nanobodies are primarily monoclonal and can be directly conjugated to a fluorescent dye. Therefore, polyclonal secondary antibodies that normally produce substantial fluorescent signal amplification are not used. Amplification strategies that are widely used in the single-molecule-RNA FISH field such as SABER¹⁶⁸ and HCR¹⁶⁹ could be adopted for nanobodies (as they have been for antibodies¹⁷⁰) to alleviate the above limitations. Single chain variable fragments (scFvs) - small immunolabels known to penetrate faster through biological tissue than nanobodies - have seen little application in tissue clearing to date but may be a promising technology for labelling large samples¹⁷¹.

As tissue clearing experiments become commonplace, deep biological insight will follow. It is difficult to imagine a field of biology that will not benefit from the ability to investigate the 3D relationships between all components of a tissue. The recent clearing of entire, rhesus monkey¹⁷² and human brains⁶ has shown that clearing methods are outpacing imaging technology. Labs looking to image on this scale will need to partner with industry to design and build objectives and optical systems with working distances of tens of centimetres that can still achieve reasonable lateral and axial resolution. There can be no doubt that the democratization of tissue clearing has begun and exciting biological insights are sure to follow.

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Glossary

Autofluorescence

Fluorescence arising from endogenous fluorescent molecules contained within a biological specimen. Can also be introduced exogenously (i.e. some hydrogels autofluorescence).

Dipping objectives

Objective lenses that are designed to be submerged into a liquid. Dipping objectives are found on upright microscopes and samples are mounted without a coverslip.

Optical section

An image of a 2D plane within a 3D object that is derived by optical, rather than mechanical, means.

Lattice light sheet

A light sheet that is formed using a specialized interference pattern that results in the projection of thin beams of excitation light into a sample, which are rapidly dithered to form the light sheet.

Steric hindrances

Refers to the inability to mount a sample on a microscope when the objective's working distance is shorter than the thinnest dimension of the sample.

Isotropic spatial resolution

Refers to specialized light microscopy techniques that produce an identical lateral and axial resolution

Network switch

A computer network hardware device that allows multiple computers to communicate.

Dispersion

A measure of the change in refractive index relative to the wavelength of light passing through a substance. If a substance has high dispersion, it means blue light and red light will refract differently when passing through it.

Random forest

A machine learning algorithm that is comprised of many “estimators” that each make a prediction as to which segmentation group a pixel should belong. When many estimators are combined into a “forest” the final prediction is highly accurate.

Tortuosity

Describes the degree of curvature/twist in a blood vessel.

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Box 1:**Suggested metadata to report when preparing and imaging cleared tissue**

1. Approximate dimensions of the sample being cleared.
2. Composition of all clearing solutions used, duration of incubation in each solution and the purpose/function of each solution.
3. Degree of expansion or contraction of the sample.
4. Antibody manufacturer, lot number, concentration, incubation temperature and incubation duration.
5. Imaging modality used.
6. Numerical Aperture (NA) of imaging objective.
7. Light sheet thickness (if used).
8. Voxel size of raw images.
9. Total raw data size.
10. A summary of all data processing steps (SA correction, chromatic correction, filtering).
11. Description of data quantitation methods.
12. Deposition of any custom software in an open source depository.
13. A subset of manually annotated ground-truth data to assess accuracy of automated data quantitation.

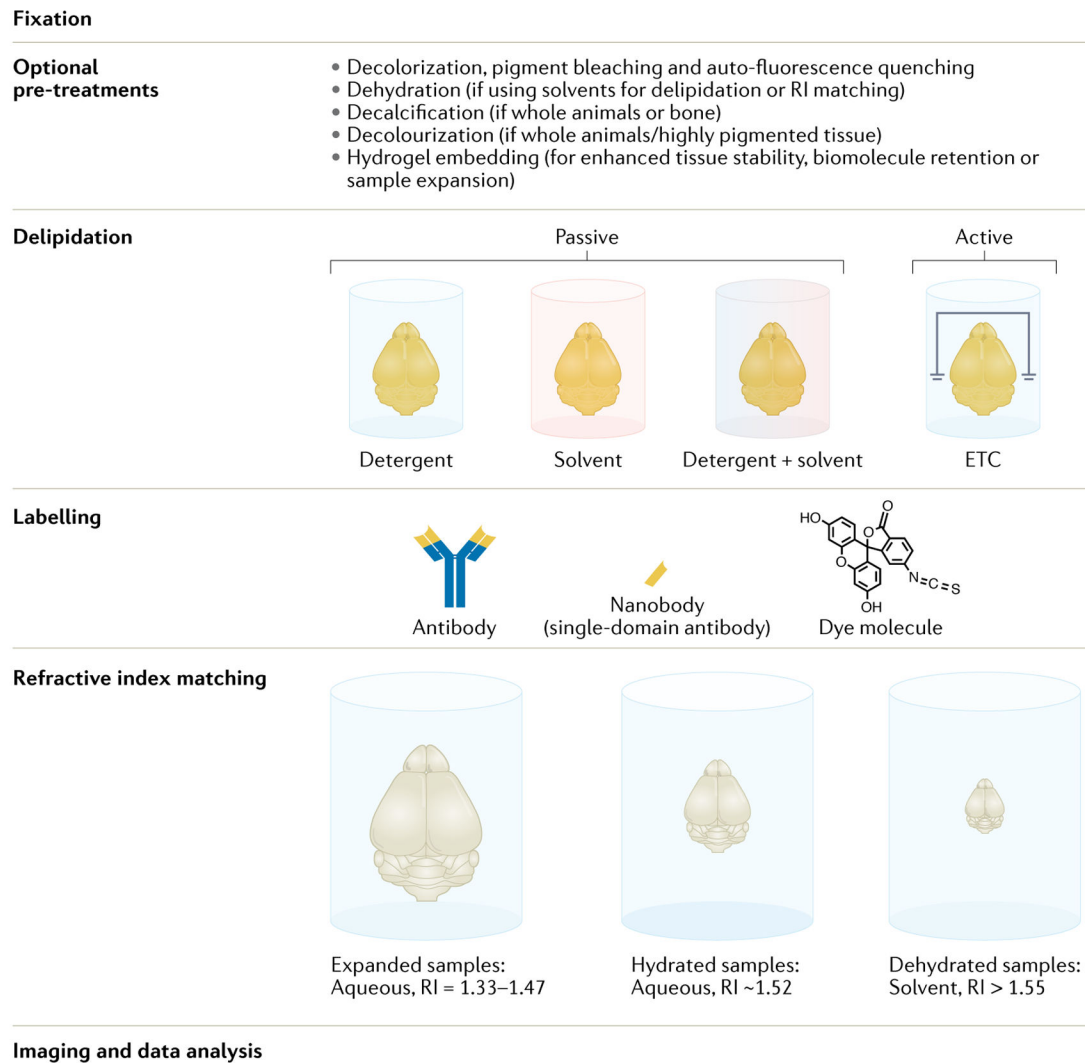


Figure 1: An overview of the components of a tissue clearing experiment.

Tissue clearing workflows are numerous, making them difficult to summarize. However, workflows can be defined as a series of modules (fixation, pre-treatment, delipidation, labelling and refractive index matching), where each module can be customized to suit the tissue that is to be cleared. Multiple modules must be combined to clear large organoids, organs and whole animals, whereas small samples (< 0.5 mm) may only require a refractive index matching step. Blue shading indicates that samples are immersed in polar, water-based, aqueous solutions whereas red shading indicates samples are in less/non-polar solvents. Expanded samples are hyperhydrated and grow in size, whereas dehydrated samples commonly shrink. ETC, electrophoretic tissue clearing; RI, refractive index.

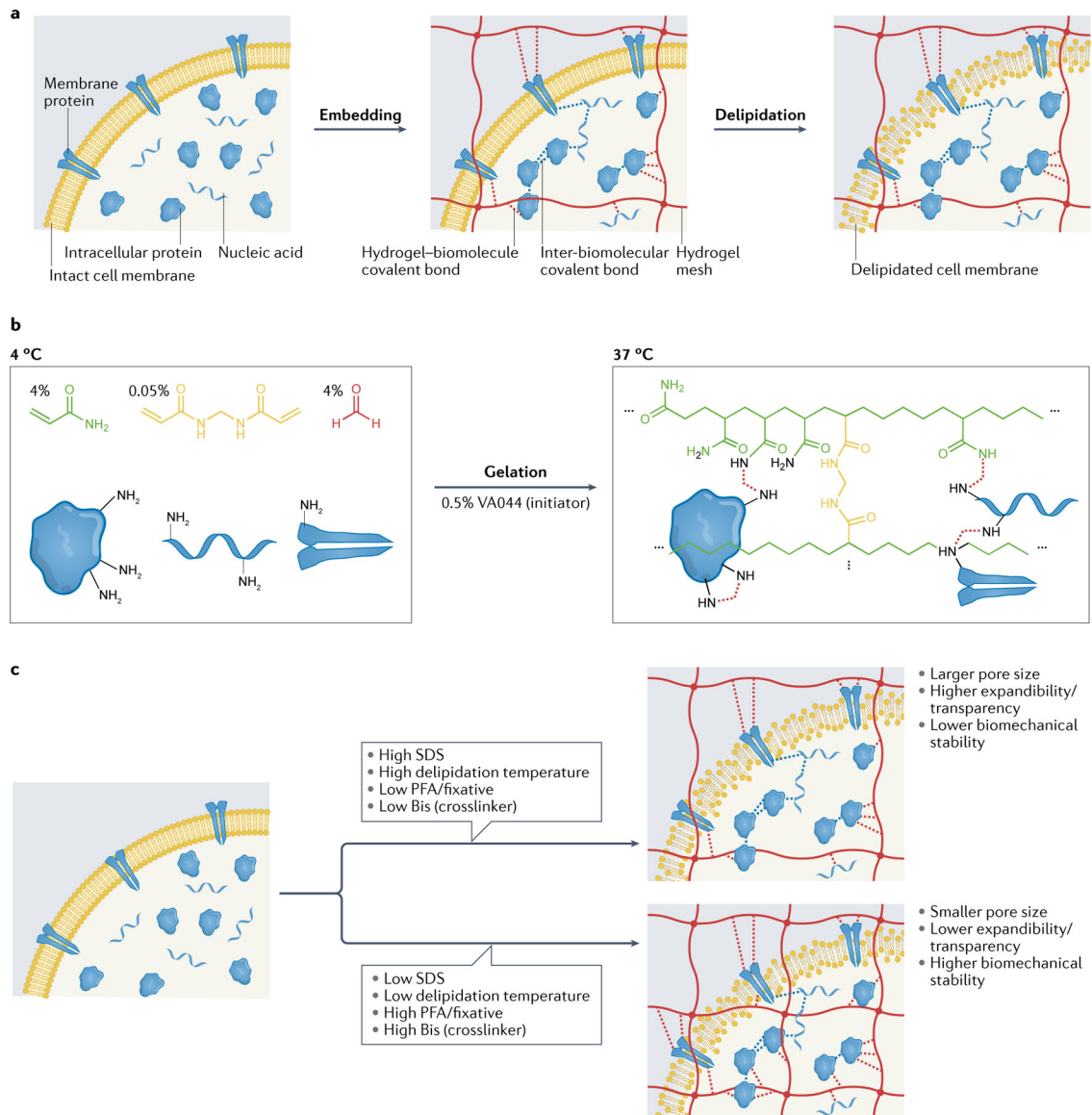


Figure 2: Concept of hydrogel embedding.

(a) Tissue biomolecules are fixed chemically or physically to a hydrogel mesh generated *in situ*, and then cell membranes are removed during delipidation to enable chemical transport and optical transparency. (b) In CLARITY hydrogel embedding, paraformaldehyde (PFA), bis-acrylamide (Bis) and acrylic acid (AA) are used to covalently link the primary amines of proteins and nucleic acids to a polyacrylamide (pAAm) hydrogel mesh generated through free radical polymerization. Free radical generation and subsequent polymerization is initiated by the chemical VA044 when it is warmed to a temperature of 37°C. (c) A tissue-hydrogel's pore size, expandability, transparency, and biomechanical stability can be modulated based on the delipidation temperature and the concentrations of the sodium dodecyl sulfate (SDS) delipidation detergent, the PFA fixative and the Bis crosslinker.

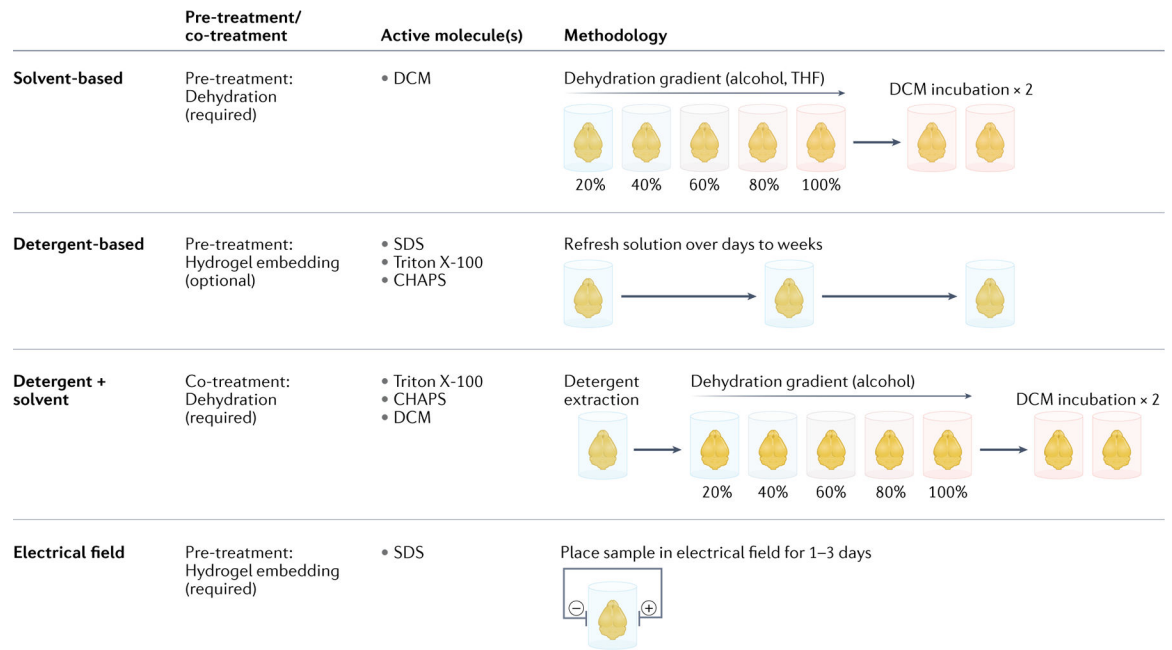


Figure 3: Protocols for delipidation.

A summary of methods for delipidating tissue using the most common detergents and solvents. Methodology example represents clearing of a murine brain. CHAPS, (3-((3-cholamidopropyl) dimethylammonio)-1-propanesulfonate); DCM, dichloromethane; SDS, sodium dodecyl sulfate; THF, tetrahydrofuran.

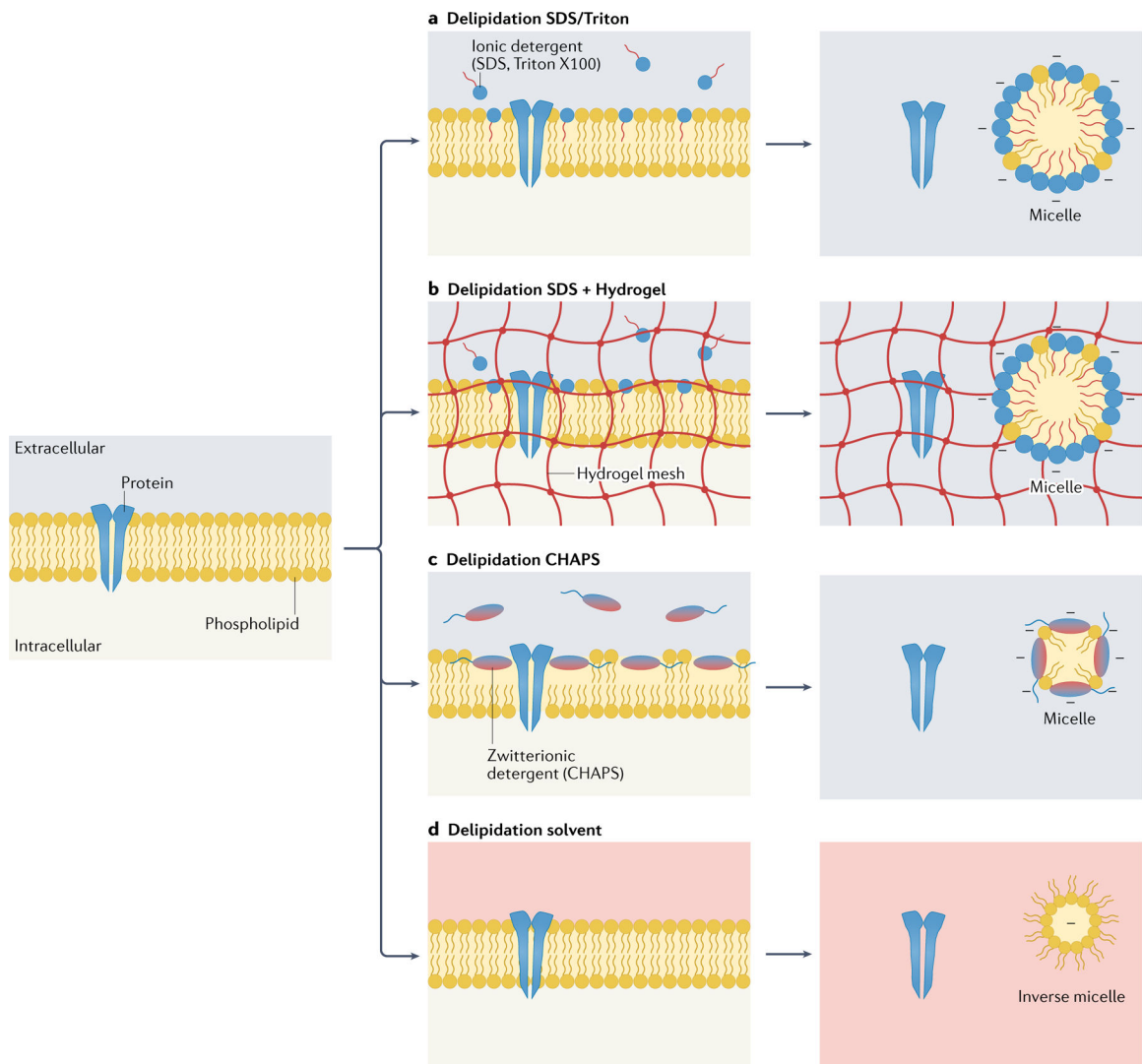


Figure 4: Mechanisms of delipidation.

(a) The long non-polar tails of ionic detergents such as sodium dodecyl sulfate (SDS) or Triton X100 intercalate with membrane lipids. If the detergent concentration is high enough, large micelles will form consisting of detergent and membrane lipids. **(b)** Intercalation of membrane lipids by ionic detergents can occur in the presence of a hydrogel mesh. **(c)** The hydrophobic moiety of zwitterionic detergents such as CHAPS embed into the top of the plasma membrane. If the detergent concentration is high enough, small micelles will form. **(d)** When tissues are transitioned from an aqueous liquid (polar) to a solvent (non-polar), membrane lipids form reverse micelles in which their non-polar tails are oriented toward the surrounding solvent environment. In all situations, micelles migrate out of the tissue and are washed away. CHAPS, (3-((3-cholamidopropyl) dimethylammonio)-1-propanesulfonate).

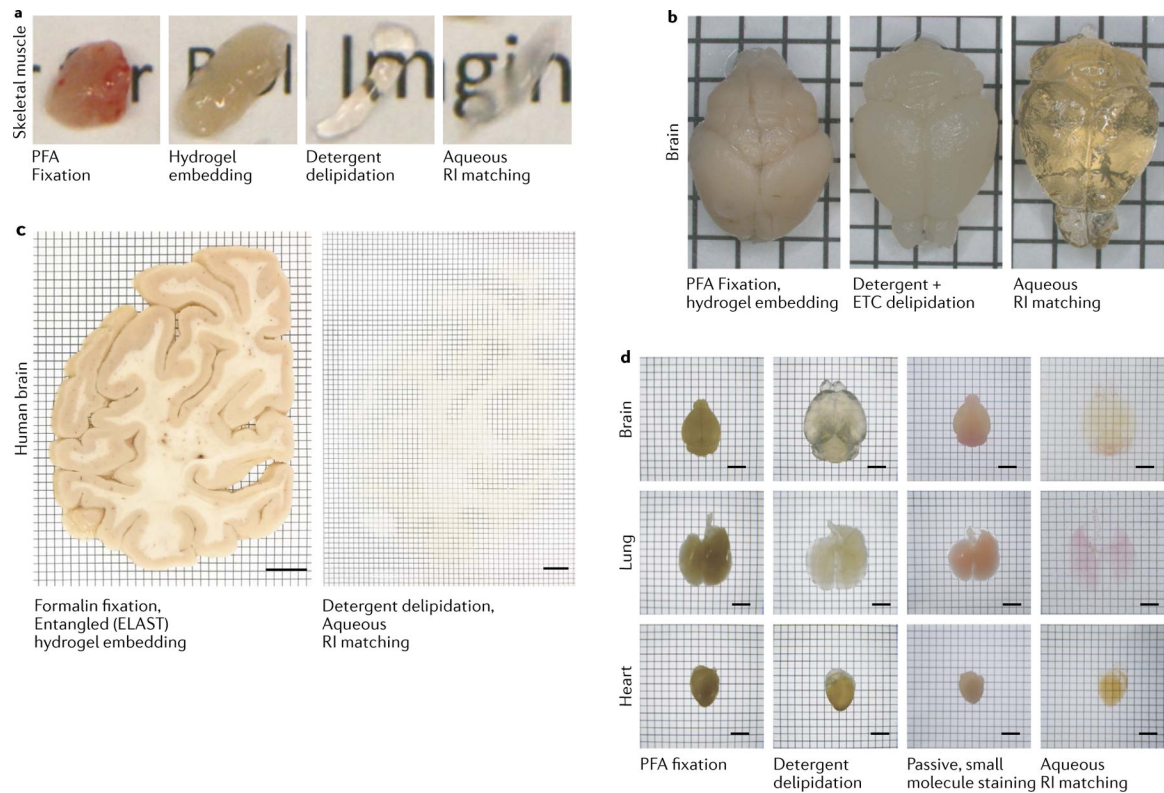


Figure 5: Examples of the clearing process.

(a) Four pieces of mouse skeletal muscle at various stages of the clearing process. From left to right: muscle extracted from the mouse and fixed in paraformaldehyde (PFA); muscle tissue extracted from a mouse that was transcardially perfused with phosphate buffered saline (PBS) and hydrogel embedded; muscle tissue passively delipidated for 4 days in SDS/boric acid; muscle tissue that has been refractive index matched in an RI = 1.47 aqueous solution. (b) Mouse brains cleared using the CLARITY process. Left: hydrogel embedded sample. Centre: sample that has been delipidated using active electrophoretic tissue clearing (ETC) for 24 hours. Right: Sample after refractive index matching in a 1.47 RI solution for 48 hours. (c) 2mm-thick coronal human brain hemisphere slabs after formalin banking. Scale bars, 1cm. Left: slab before clearing. Right: the same sample after ELAST tissue transformation and RI matching. (d) Mouse whole-organ clearing using the CUBIC process and nuclear staining with propidium iodide. RI matching was performed using the CUBIC R+(N) protocol (RI= 1.522). Scale bars, 5 mm.

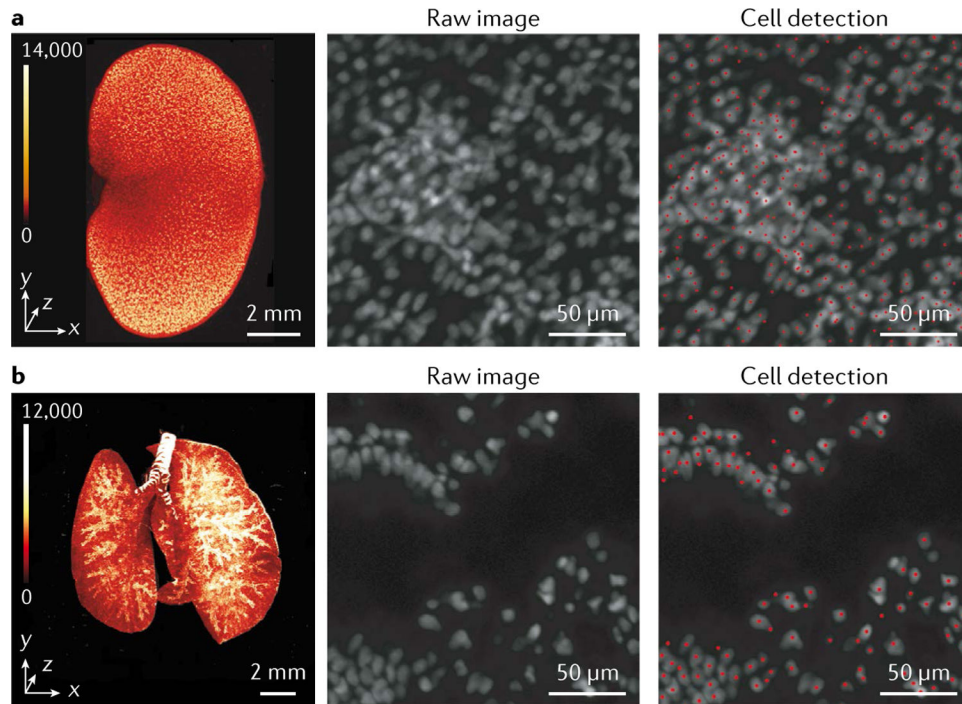


Figure 6: Whole-organ cell profiling using the latest CUBIC-L/R+ protocol.

Volume-rendered and single-plane images of mouse organs of 8-week-old C57BL/6N male mice that were cleared using CUBIC-L/R+. **(a,b)** The organs were stained with propidium iodide (PI) and individual cells were detected using a custom, machine-learning, GPU-based cell detection algorithm with over 90 % accuracy (see ref. ²⁴). **(a)** Imaging of the mouse kidney identified 79–83 million cells. **(b)** Imaging of the mouse lung identified 99.3 million cells.

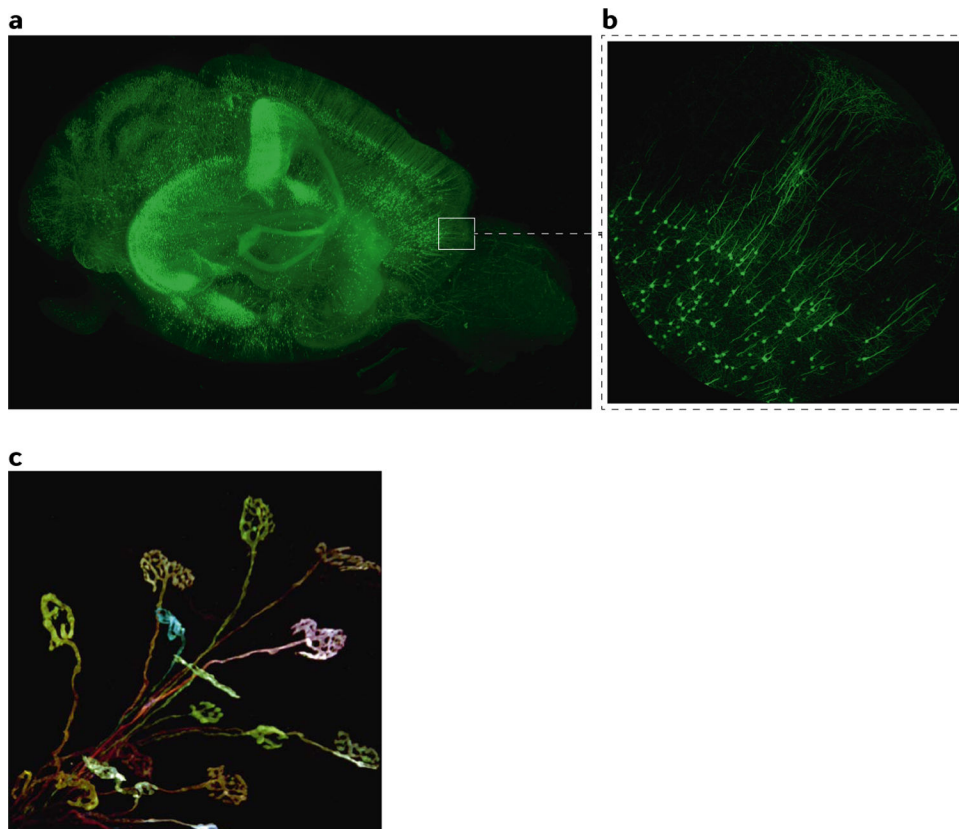


Figure 7: Example of neuronal staining in the mouse brain.

(a) 3D rendering of an entire mouse expressing GFP downstream of the *thy1* promoter¹²² in a subset of cells. The brain was cleared using a CLARITY protocol. Major fibre tracks are clearly visible. (b) Zoomed cortical region of the brain shown in panel a. Individual neuronal processes can be visualized. (c) Brainbow-expressing neuromuscular junctions in mouse skeletal muscle. Muscle was cleared using a passive hydrogel-embedding method. Each neuron expresses a unique ratio of blue, green and red fluorophore, allowing for the observation of individual neurons.

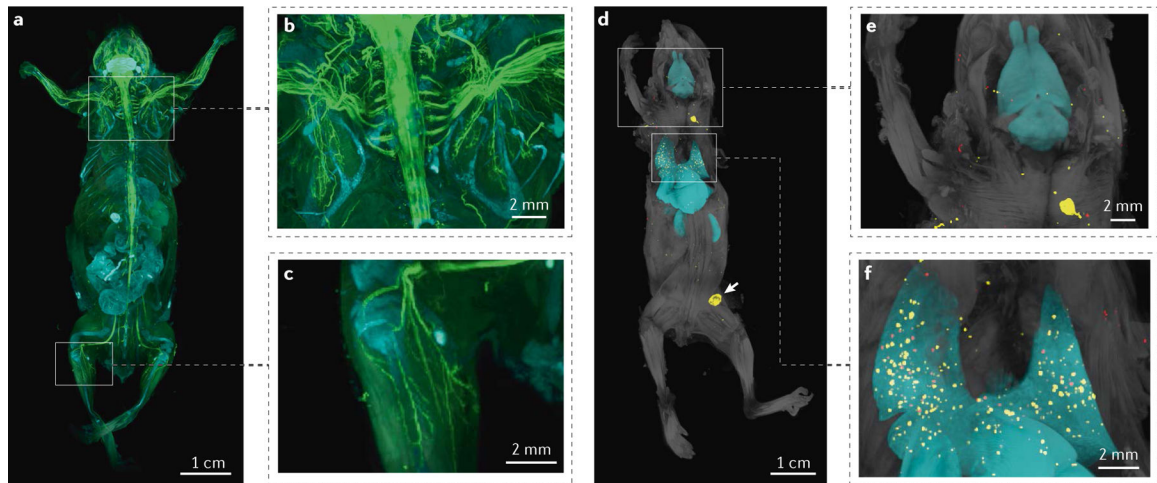


Figure 8: Representative results of vDISCO panoptic imaging.

a, 3D rendering of a Thy1-GFPM mouse after vDISCO whole-body immunolabelling/clearing and imaging by light-sheet microscopy. Neuronal fibres expressing GFP are enhanced by anti-GFP nanobodies conjugated with Atto647 dye and shown in green, bones and internal organs highlighted by propidium iodide are in cyan, and the background signal — mainly from autofluorescence of the muscle tissue — is in white. **b** and **c**, Zoomed-in images of the boxed regions in panel **a**, showing detailed neuronal structures with sub-cellular level resolution. **d**, 3D reconstruction of a vDISCO-processed NSG mouse implanted with MDA-231 human breast cancer cells in the mammary fat pad for approximately 2 months, followed by intravenous injection of an anti-human CA12 therapeutic antibody named 6A10 which is conjugated with Alexa-568 dye. The cancer cells were previously transduced by mCherry viral vectors and further enhanced by anti-mCherry nanobodies conjugated with Atto647 dye. The 6A10-targeted primary tumour (pointed by the white arrowhead) and metastases are shown in yellow, untargeted metastases are in red, and selected organs including the brain, lungs, liver, kidneys are manually segmented and shown in cyan. **e** and **f**, enlarged view of the boxed regions in **d** showing greater details of the distribution of micrometastases and targeting efficacy of the 6A10 therapeutic antibody.

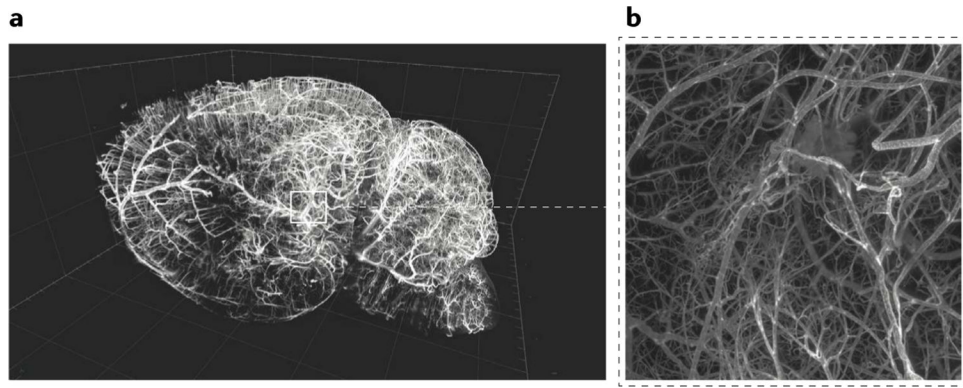


Figure 9: Example of vascular staining in the mouse brain.

(a) 3D render of an entire mouse brain that was perfused with CM-DiI lipophilic dye prior to fixation. After fixation, the brain was cleared using a modified iDISCO protocol and imaged via light sheet microscopy. (b) Zoomed region of brain on left showing the level of detail in vasculature staining.

Table 1:

Tissue Clearing Modules

Modules and reagents	Indications for use	Required equipment	Additional considerations	Originating protocol
Fixation (1–12 hours)				
Paraformaldehyde (PFA)	• General fixation protocol	None	• Concentration and timing require optimization	Scafe ¹⁷³ , CLARITY ³³
Glutaldehyde (GA)	• Used to retain low-abundance biomolecules • Used to stabilize fragile tissues	None	• Concentration and timing require optimization • Often combined with PFA	SWITCH ²⁰
Polyglycerol-3-polyglycidyl ether (P3PE)	• Used to retain low-abundance biomolecules • Used to stabilize fragile tissues • Used to preserve fluorescent protein emission • Used to retain mRNA	None	• High protein and mRNA fixation efficiency	SHIELD ⁴
1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC)	• Used to retain mRNA	None	• High mRNA fixation efficiency • Poor protein fixation efficiency	STARmap ¹⁷⁴
Fixation with sodium dodecyl sulfate (SDS), urea, borate and heat	• Used for antigen retrieval	Oven	• May enhance antibody staining • Concentration of reagents and incubation length should be carefully determined	CLARITY ³³ , FLASH ¹²
Decolorization (5–24 hours)				
Amino alcohols (Quadrol, THEED)	• Used when tissues contain pigments such as kidney, liver, spleen and muscle (heme); skin (melanin); brain (lipofuscin)	None	• Used to remove heme from blood (hemoglobin) and muscle (myoglobin)	CUBIC ²⁴ , FlyClear ¹³
N-alkylimidazole		None	• Effective at heme removal	CUBIC ²⁵
N-methyldiethanolamine (NMDEA)		None	• Effective at heme removal	SHANEL ⁶
Bleaching by hydrogen peroxide		None	• Widely effective across many pigments • Can cause tissue damage if left too long or used at too high a concentration	iDISCO ⁴⁷
Quenching/autofluorescence (12–24 hours)				
Photobleaching with high-intensity light	• Used when tissues have high autofluorescence (human brain)	LED array inside refrigeration unit	• Broadband white light is most effective • Will bleach genetically encoded fluorescent proteins	ELAST ³¹ , Multispectral LED array ³⁰
Use of red and far-red fluorophores		Antibody conjugated fluorescent dyes	• Use dyes with emission at wavelengths > 600 nm	iDISCO ⁴⁷
Hydrogel embedding (12–24 hours)				
Acrylamide, bis-acrylamide and PFA	• Used to retain low expression biomolecules • Used to stabilize fragile tissues • Used to retain mRNA	Heat block under vacuum	• Best for retention of mRNA and low expressing proteins • Gel density can be adjusted to promote clearing and labelling (less	CLARITY ³³ , PACT/PARS ³⁴

Modules and reagents	Indications for use	Required equipment	Additional considerations	Originating protocol
	• Required when performing tissue expansion		dense) or biomolecule retention (more dense)	
Acrylamide		Heat block under vacuum	• Acrylamide is polymerized in the absence of cross-linking molecules (bis-acrylamide), creating a stretchable gel	ELAST ³¹
Acrylamide, acrylate, protein enzymes, detergents and imidazole		Incubator	• Hydrogels can expand many fold when placed in pure water (or an imidazole solution in the case of CUBIC-X)	ExM ³⁵ , ProExM ³⁶ , MAP ³⁷ , CUBIC-X ³⁸ , iExM ³⁹
Decalcification (12 hours–3 weeks)				
EDTA and imidazole	• Required when clearing tissue containing calcified bone		• Used to remove calcification in tissue	CUBIC-B ²⁵ , PEGASOS ⁴² , PACT-deCAL ⁴³ , bone CLARITY ⁴⁴ , vDISCO ⁴⁵
Delipidation (1 hour–3 days)				
By solvent	• Required to clear any tissue > 0.2 mm thick	None	• Tissue must first be dehydrated with tetrahydrofuran (THF) or alcohol • If tissue contains fluorescent proteins, dehydrating solutions must be maintained at 4°C and > pH 9.0. Consider supplementing with: butylated hydroxytoluene or Quadrol and polyethylene glycol (PEG) • Complete delipidation requires additional incubation in dichloromethane (DCM)	fDISCO ⁴⁹ , sDISCO ⁵⁰ , uDISCO ⁹ , a-uDISCO ⁵² , FluoClearBABB ⁵¹ , PEGASOS ⁴²
By detergent		Optional: electrophoretic tissue clearing (ETC) device	• Most common detergents: SDS, TritonX-100, CHAPS • Degree of clearing increased by alkaline pH, increased temperature and increased incubation time	CLARITY ³³ , PACT-PARS ³⁴ , CUBIC ²³
By solvent and detergent		None	• Must be performed sequentially	Adipo-Clear ⁵⁷ , SHANEL ⁶
Refractive index matching (1 hour–2 days)				
In solvent	• Required for all tissue clearing experiments	None	• Commonly used solvents: BABB, dibenzyl ether (DBE) and ethyl cinnamate • If tissue contains fluorescent proteins, a pH > 9.0 should be maintained. Consider supplementing with triethylamine, Quadrol, DL-alpha-tocopherol, propyl gallate, DPE or PEG. • RI should be ~ 1.56	sDISCO ⁵⁰ , uDISCO ⁹ , PEGASOS ⁴²
In aqueous solution		None	• Commonly used reagents are antipyrine and nicotinamide; N-methylacetamide and Histodenz; or TDE, DMSO	CUBIC ⁷⁰ , Ce3D ⁷¹ , ELAST ³¹

Modules and reagents	Indications for use	Required equipment	Additional considerations	Originating protocol
			and iohexol • RI should be ~ 1.52	
With expansion		None	• Use pure water with RI = 1.33 (ExM, ProExM, iExM, MAP) or 5% imidazole/55% antipyrine with RI = 1.467 (CUBIC-X) solutions for RI matching	ExM ³⁵ , ProExM ³⁶ , MAP ³⁷ , CUBIC-X ³⁸ , iExM ³⁹

BABB, benzyl alcohol benzyl benzoate; CHAPS, 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate; EDTA, ethylenediaminetetraacetic acid; RI, refractive index; THEED, 2,2',2'',2'''-(Ethylenedinitrilo)tetraethanol;

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Table 2:

Microscopy options for imaging cleared tissue

Microscopy technique	Best for	Not for use with	Speed	Numerical aperture (NA) range ^a	Approximate resolution (lateral, axial) ^b	Availability
Spinning disk confocal	<ul style="list-style-type: none"> • Samples < 100 μm thick • Fast imaging 	<ul style="list-style-type: none"> • Samples > 200 μm (when disk optimized for high NA) 	Fast	Up to 1.4 (for < 200 μm working distance oil immersion objectives)	250 nm, 400 nm	High
Point-scanning confocal (inverted)	<ul style="list-style-type: none"> • Samples < 2 mm thick • Subcellular resolution 	<ul style="list-style-type: none"> • Samples prone to bleaching 	Slow	Up to 0.5 (for long-distance air objectives)	700 nm, 4 μm	High
Point-scanning confocal (upright)	<ul style="list-style-type: none"> • Samples < 6 mm thick • Subcellular resolution 	<ul style="list-style-type: none"> • Samples prone to bleaching 	Slow	Up to 1.1 with dipping objectives	300 nm, 750, nm	High
2-photon (upright)	<ul style="list-style-type: none"> • Samples < 6 mm thick • Subcellular resolution 	<ul style="list-style-type: none"> • Multicolour (>2) imaging 	Slowest	Up to 1.1 with dipping objectives	300 nm, 750, nm	High
Light sheet	<ul style="list-style-type: none"> • Samples < 6 cm thick • Cellular resolution 	<ul style="list-style-type: none"> • High resolution/ subcellular imaging 	Fastest	Below 0.5 (for low magnification air objectives)	700 nm, 5 μm	Lower
Optical projection tomography	<ul style="list-style-type: none"> • Samples < 5 mm thick • Isotropic imaging 	<ul style="list-style-type: none"> • High resolution/ subcellular imaging 	Fast	Below 0.1 (for low magnification air objectives)	3 μm , 3 μm	Lowest

^a higher NAs give better resolution^b assumes emission light at a wavelength of 550 nm; calculated at highest NA for each modality