

REVIEW

Optical clearing methods: An overview of the techniques used for the imaging of 3D spheroids

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Abstract

Spheroids have emerged as *in vitro* models that reproduce in a great extent the architectural microenvironment found in human tissues. However, the imaging of 3D cell cultures is highly challenging due to its high thickness, which results in a light-scattering phenomenon that limits light penetration. Therefore, several optical clearing methods, widely used in the imaging of animal tissues, have been recently explored to render spheroids with enhanced transparency. These methods are aimed to homogenize the microtissue refractive index (RI) and can be grouped into four different categories, namely (a) simple immersion in an aqueous solution with high RI; (b) delipidation and dehydration followed by RI matching; (c) delipidation and hyperhydration followed by RI matching; and (d) hydrogel embedding followed by delipidation and RI matching. In this review, the main optical clearing methods, their mechanism of action, advantages, and disadvantages are described. Furthermore, the practical examples of the optical clearing methods application for the imaging of 3D spheroids are highlighted.

KEYWORDS

imaging, light scattering, optical clearing, optical microscopy, spheroids.

1 | INTRODUCTION

In the early years of the 20th century, cell culture techniques (i.e., cell proliferation in environmentally controlled conditions) emerged as a powerful methodology to study and understand the behavior of the cells within the human body (Souza, Ferreira, Marangoni, Bastos, & Goulart, 2016). Such approach allowed the characterization of the cellular differentiation, migration, proliferation, and mechanics, as well as their response to external stimuli (e.g., drugs, environment changes, and mechanical stress; Souza et al., 2016). In this way, researchers could decipher the background of several *in vivo* processes, such as the tissue development and the pathophysiology of the diseases (Hudu, Alshrari, Syahida, & Sekawi, 2016; Souza et al., 2016). Therefore, *in vitro* culture of cells is now broadly used in several fields, such as cancer research, tissue regeneration, gene

therapy, and drug screening, among others (Duval et al., 2017; Kapałczyńska et al., 2018).

In literature, 2D cell cultures (i.e., the culture of cells as monolayers attached to a plastic or glass surface) are the most attractive for research purposes, due to its simplicity, reproducibility, and low cost (Kapałczyńska et al., 2018). Nevertheless, different studies have been demonstrating that 2D cellular systems display biochemical profiles and bioactivities divergent from those observed *in vivo*, which is correlated with the lack of capacity of these cell culture models to reproduce the 3D cellular organization (Edmondson, Broglie, Adcock, & Yang, 2014; Kapałczyńska et al., 2018). For instance, Sandberg & Ernberg (2005) reported that ≈30% of the 7,000 genes analyzed in different cell lines showed a statistically significant differential gene expression when compared to their original tissues. Thus, 3D cell cultures, such as spheroids, have received great attention from the research community, as these *in vitro* models can represent more accurately the different properties of human tissues, such as liver (Yoon, Lee, Lee, & Lee, 2015), thyroid

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(Cirello et al., 2017), cartilage (Jukes et al., 2008), pancreatic tissue (Lumelsky et al., 2001), cardiac muscle (Kehat et al., 2001) or of solid tumors (e.g., breast, colon, pancreas, prostate, ovary, among others (Eiraku et al., 2008; Eiraku et al., 2011; Ham et al., 2018; Ham, Joshi, Luker, & Tavana, 2016; Hamilton, 1998; Khawar et al., 2018; Lazzari et al., 2018; Suga et al., 2011)). Spheroids are microtissues with a diameter within hundreds of micrometers to few millimeters that present a spatial architecture, cellular organization, cell-cell, and cell-extracellular matrix interactions quite similar to those found in the human tissues (as reviewed in detail elsewhere (Costa et al., 2016; Duval et al., 2017)).

On the other hand, the microscopy techniques (e.g., optical and fluorescence microscopy) arise as one of the easiest, low cost, and safe methodologies for the analysis of *in vitro* samples (Genina, Bashkatov, & Tuchin, 2010). Particularly, the fluorescence microscopy assumes great importance since it allows the observation of fluorescent markers that stain a specific area or molecule of interest within the cells (Graf & Boppart, 2010). Furthermore, the fluorescence microscopy also supports the acquisition of images with high resolution of the subcellular structures when the cells are cultured as monolayers. However, the imaging of large 3D spheroids by fluorescence microscopy is challenging due to their thickness and to the light-scattering phenomenon. The light scatter results from the mismatches between the cellular constituents refractive index (RI)

that prompt the dispersion of the excitation light through the sample and thus limiting its penetration, or by decreasing the amount of emitted light that reaches the detector (Richardson & Lichtman, 2015). For this reason, it is difficult to obtain high-resolution images of intact spheroids, especially from its interior (Achilli, Meyer, & Morgan, 2012). Therefore, researchers often use thin slices of spheroids (5–7 μm) to obtain images at a submicron resolution by fluorescence microscopy (Costa et al., 2016; Graf & Boppart, 2010). Notwithstanding, the spheroids sectioning is a laborious and time-consuming process that can permanently alter the spheroid initial structure (e.g., distort, disrupt, fold, compress or stretch; Langenbach et al., 2011; Richardson & Lichtman, 2015). In addition, the post-acquisition 3D reconstruction of the slices performed using computer software may introduce artifacts in the images (Richardson & Lichtman, 2015). Another approach for the imaging of the deeper regions of intact spheroids is the utilization of optical sectioning microscopies, such as confocal laser scanning microscopy (CLSM), multiphoton microscopy (MPM; e.g., two-photon microscopy [2PM]), and light-sheet fluorescence microscopy (LSFM; e.g., single [or selective] plane illumination microscopy [SPIM]; Table 1; reviewed in detail in (Graf & Boppart, 2010; Santi, 2011)). These microscopic techniques are capable of acquiring images of a thin single plane of the sample (so-called “stack”) with minimal interference of the remaining parts of the sample. Then, the 3D reconstruction of

TABLE 1 Comparison of the optical sectioning microscopy techniques that can be used for the imaging of 3D spheroids

Technique	Confocal laser scanning microscopy	Multiphoton microscopy (e.g., two-photon microscopy)	Light-sheet-based fluorescence microscopy (e.g., single [or selective] plane illumination microscopy)
Concept	<ul style="list-style-type: none"> Optical sections are produced by scanning the sample point-by-point with a laser beam focused on the sample; Uses a pinhole to exclude out-of-focus background fluorescence from detection. 	<ul style="list-style-type: none"> Uses localized nonlinear excitation to excite fluorescence only within a thin plane; More than one photon is emitted by a pulsed infrared laser source at a time to excite a fluorophore. 	<ul style="list-style-type: none"> A sheet plane of light illuminates a plane within a sample; The illumination is done perpendicularly to the direction of the imaging axis.
Optical sectioning	Yes	Yes	Yes
Light scatter influence	+++	+	+++
Penetration depth	$\approx 100 \mu\text{m}$	up to 1 mm	>1 cm
Resolution	<micron	<micron	micron
Photodamage	+++	++	+
Handling	+	+	+++
Time	+++	++	++
Equipment cost	++	+++	+
References	Benninger and Piston (2013); Combs and Shroff (2017); Costa et al. (2016); Genina et al. (2010); Marques et al. (2015)	Benninger and Piston (2013); Combs and Shroff (2017); Costa et al. (2016); Genina et al. (2010); Marques et al. (2015); Zipfel, Williams, and Webb (2003)	Benninger and Piston (2013); Combs and Shroff (2017); Costa et al. (2016); Feuchtinger, Walch, and Dobosz (2016); Genina et al. (2010); Marques et al. (2015); Reynaud et al. (2008); Santi (2011)

+: Low; ++: moderate; +++: high.

TABLE 2 Refractive index of several cellular components

Cellular component	Refractive index	Ref(s)
Cell membrane	1.46	Bigio and Bown (2004)
Cytoplasm	1.36–1.39	Bassnett (2009); Choi et al. (2007); Zhang et al. (2017)
DNA	1.44	Bigio and Bown (2004)
Lipids	1.39–1.48	Bigio and Bown (2004); Feuchtinger, Walch, and Dobosz (2016); Zhang et al. (2017)
Lysosome	1.6	Wilson, Cottrell, and Foster (2007)
Mitochondria	1.36–1.42	Haseda et al. (2015); Zhang et al. (2017)
Nucleus	1.36–1.39	Bassnett (2009); Choi et al. (2007)
Proteins	1.38–1.43	Bigio and Bown (2004); Genina et al. (2010)
Water	1.33	Bigio and Bown (2004)
Organelles	1.33–1.35	Feuchtinger et al. (2016)

different stacks allows the observation of a great part or even the totality of the sample (Richardson & Lichtman, 2015). In particular, LSFM is the fluorescence-based microscopy technique with the highest penetration depth (>1 cm), and therefore has been broadly used for the imaging of intact spheroids (Achilli et al., 2012; Benninger & Piston, 2013; Combs & Shroff, 2017; Costa et al., 2016; Desmaison et al., 2018; Feuchtinger, Walch, & Dobosz, 2016; Genina et al., 2010; Gualda, Simão, Pinto, Alves, & Brito, 2014; Marques, Oliveira, Chang, Paula-Neto, & Menezes, 2015; Masson et al., 2015; Paiè, Bragheri, Bassi, & Osellame, 2016; Pampaloni, Ansari, & Stelzer, 2013; Reynaud, Kržič, Greger, & Stelzer, 2008; Santi, 2011; Schmitz, Fischer, Mattheyer, Pampaloni, & Stelzer, 2017; Smyrek & Stelzer, 2017; Weber et al., 2015; Wenzel et al., 2014).

Alternatively, to perform the imaging of intact spheroids researchers can employ optical clearing methods before samples are analyzed by microscopy. These methods aim to reduce the light-scattering phenomenon in the biological samples, rendering samples with increased transparency that promotes deeper imaging, as well as the cells structural and functional analysis (discussed hereafter).

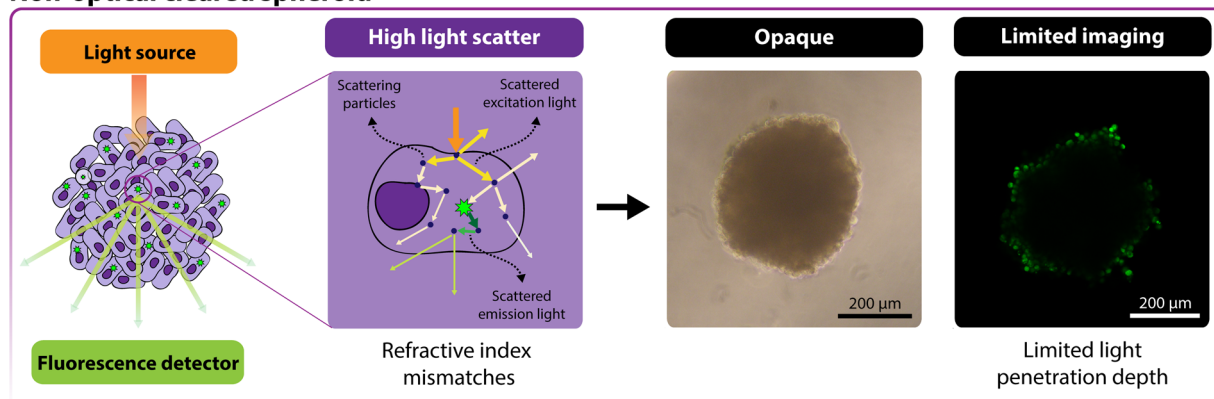
2 | LIGHT SCATTERING AND TACTICS OF OPTICAL CLEARING

Fluorescence imaging of biological samples is quite challenging due to the limited light penetration, which is influenced by the optical properties of the acellular and cellular sample's constituents. In this way, the light absorption by molecules, such as hemoglobin, myoglobin, and melanin, present in some tissues will reduce the light intensity that reaches both the deeper regions of the tissue (excitation light) and the detector (emitted light; Richardson & Lichtman, 2015). Therefore, the sample bleaching allows to minimize or even prevent the light absorption by the endogenous pigments. In fact, different methodologies can be explored to reduce the light absorption by heme (major light absorber and the most abundant chromophore in the body): (a) sample can be

treated with hydrogen peroxide that oxidizes the pigment (Azaripour et al., 2016); (b) samples washing with highly acidic or basic solutions that lead to the heme dissociation from hemoglobin (Susaki & Ueda, 2016); and (c) sample incubation with aminoalcohols (e.g., N,N,N,N-tetrakis(2-hydroxypropyl) ethylenediamine) that elute the heme chromophore from hemoglobin (Susaki & Ueda, 2016).

On the other hand, the limited light penetration in tissues and spheroids can also be attributed to the nonhomogeneous RI (i.e., how light propagates through a given substance compared to vacuum conditions) through the sample and consequent light refraction (Table 2; Richardson & Lichtman, 2015). For instance, the cells' lipids and proteins present high RI values (≈ 1.44 and ≈ 1.43 , respectively), while cells' cytoplasm presents the lower RI (≈ 1.35 ; Feuchtinger et al., 2016; Seo, Choe, & Kim, 2016; Susaki & Ueda, 2016). These RI mismatches induce changes on the speed and angle of the light (visible and near-infrared region) propagation, that is, light refraction from molecules, membranes, organelles, and cells (Figure 1; Richardson & Lichtman, 2015). This phenomenon promotes light scattering and it is the main cause of tissues opacity. The light dispersion through the sample difficult its penetration into deeper regions and decreases its intensity (Figure 1; Keereweer et al., 2013). In this way, the light penetration is usually limited to a range of 100–150 μm , which is insufficient for the imaging of the spheroids whole volume (Grist, Nasser, Poon, Roskelley, & Cheung, 2016; Richardson & Lichtman, 2015). On the other hand, the fluorescence light resulting from the excited fluorophore can also be scattered and therefore it may not reach the detector with high intensity or indicate inaccurately its origin, which results in images with poor resolution and contrast (Keereweer et al., 2013). Hence, several optical clearing approaches have been investigated to homogenize the RI across the sample and consequently reduce the light-scattering phenomenon. This process renders 3D large biological samples with higher transparency and therefore improves the light penetration, imaging depth, and contrast, that is, the majority of the light will attain the focus point and then the majority of the fluorescence light will be detected without suffering deviations in its path (Figure 1).

Non-optical cleared spheroid



Optical cleared spheroid

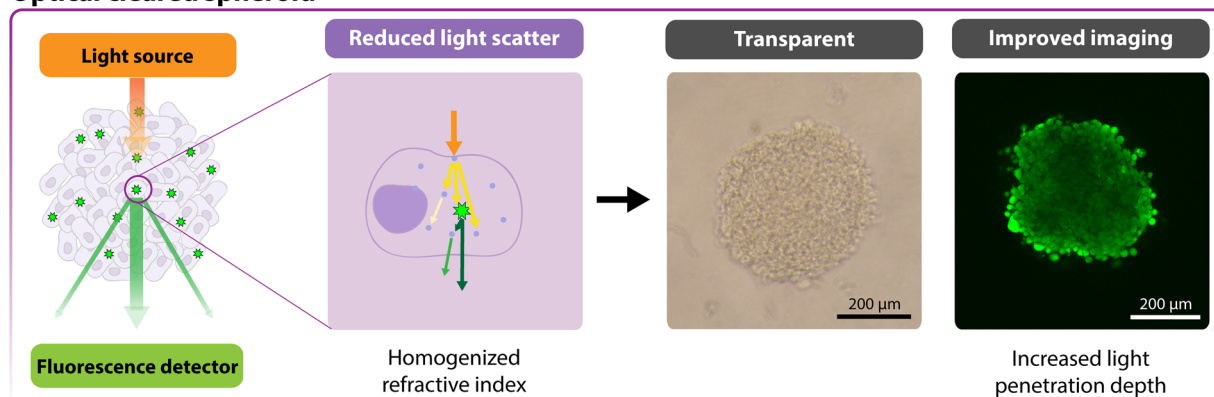


FIGURE 1 Representation of the light propagation in a noncleared and cleared spheroid. A noncleared spheroid has several mismatches in the refractive indexes (RI) of its constituents inducing the light scattering and consequently light dispersion through the spheroids, which results in opaque samples that do not allow deep imaging. A cleared spheroid has a transparent appearance due to the homogenization of the RI of its cellular constituents and consequently, light penetrates deeper in the spheroid making possible the imaging of the inner regions [Color figure can be viewed at wileyonlinelibrary.com]

3 | CLASSIFICATION OF THE OPTICAL CLEARING METHODS

Optical clearing methods are used to modify the optical properties of biological samples by removing, replacing, and modifying some of its components, to match the overall RI. These methods can be divided into four major groups according to their basic procedures, as previously reported (Seo et al., 2016): (a) simple immersion in an aqueous solution with high RI; (b) delipidation and dehydration followed by RI matching; (c) delipidation and hyperhydration followed by RI matching; and (d) hydrogel embedding followed by delipidation and RI matching (Figure 2 and Table 3).

The clearing methods were originally developed for processing large samples obtained from animals (e.g., bone, brain, embryo, heart, intestine, kidney, lung, muscle, pancreas, spinal cord, spleen, testis, among others), as already reviewed in detail in (Feuchtinger et al., 2016; Genina et al., 2010; Lee, Kim, & Sun, 2016; Richardson & Lichtman, 2015; Seo et al., 2016; Silvestri, Costantini, Sacconi, & Pavone, 2016; Susaki & Ueda, 2016; Tainaka, Kuno, Kubota, Murakami, & Ueda, 2016; Tuchin, 2015; Yu, Qi, Gong, Luo, & Zhu, 2018; Zhu, Larin, Luo, & Tuchin, 2013). Herein, the basic concepts and mechanisms of the different

clearing methods are presented, highlighting their application in the handling of spheroids samples. For sake of brevity, only the 3D imaging of solvent-cleared organs (3DISCO) and its derivatives, BABB, CLARITY and its derivatives, *Clear^T*, *Clear^{T2}*, CUBIC, FocusClear™, Scale, See Deep Brain (SeeDB), Spalteholz's technique, and TDE (2,2'-thiodiethanol) will be described.

3.1 | Simple immersion in an aqueous solution with high RI

This group includes the SeeDB, TDE, *Clear^T*, *Clear^{T2}*, and FocusClear™ clearing methods. In these methods, biological samples are gradually cleared by immersing them in aqueous solutions that have a high RI (1.4–1.5; Figure 2; Ariel, 2017; Seo et al., 2016). Due to the osmotic pressure, the water content in the sample (that has low RI \approx 1.33) will be passively replaced by the clearing solution. Therefore, the average RI of the sample will be homogenized to 1.4–1.5 since the clearing solution RI value match that exhibited by proteins (RI \approx 1.43) and lipids (RI \approx 1.44; Feuchtinger et al., 2016).

These immersion techniques are simple, easy to perform, and low cost (Genina et al., 2010). Additionally, the absence of detergents

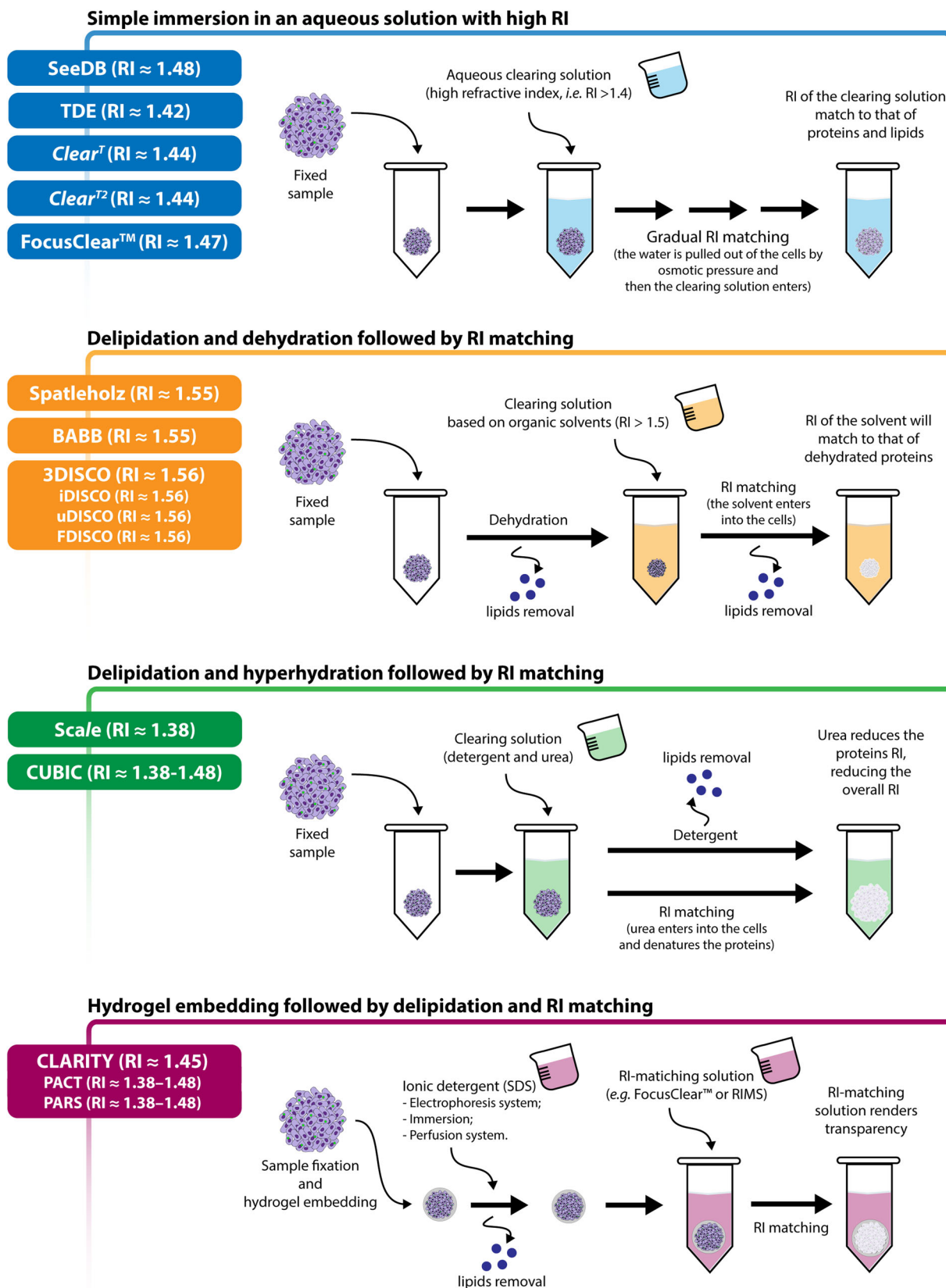


FIGURE 2 Overview of the three major groups of optical clearing methods: simple immersion in an aqueous solution with high refractive index (RI); delipidation and dehydration followed by RI matching; delipidation and hyperhydration followed by RI matching; and hydrogel embedding followed by delipidation and RI matching. The final sample RI generally obtained with each of the clearing methods is also displayed [Color figure can be viewed at wileyonlinelibrary.com]

TABLE 3 Overview of the optical clearing methods applied for the imaging of spheroids

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (μm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
Simple immersion in an aqueous solution with high RI									
SeeDB	<ul style="list-style-type: none"> • Fructose; • α-thioglycerol. 	2% formaldehyde	<1 hr	<ul style="list-style-type: none"> • MCF-7 human breast cancer. 	370 \pm 90	<ul style="list-style-type: none"> • Fluorescent Ubiquitination-based Cell Cycle Indicator (FUCCI). 	MPM (2PM)	<ul style="list-style-type: none"> • Clearing was performed with the assistance of a microfluidic device; • Spheroids size is minimally reduced; • Transparency is slightly improved; • Allow more than 250 μm of image depth; • Nuclei blur is less extent; • Fluorescence intensity (green autofluorescence) is increased. 	Grist et al. (2016)
		4% paraformaldehyde	72 hr	<ul style="list-style-type: none"> • Rat adult neural stem cells (NSC). 	104.64 \pm 22.01	<ul style="list-style-type: none"> • DAPI; • Cy3. 	CLSM	<ul style="list-style-type: none"> • Leads to spheroids shrinkage; • Moderate transparency; • Allows the imaging of the whole spheroid; • Difficult handling of the viscous clearing solutions. 	Boutin and Hoffman-Kim (2015)
TDE	<ul style="list-style-type: none"> • 2,2'-thiodiethanol (TDE). 	n.a.	30 min	<ul style="list-style-type: none"> • Tissue spheroids 	300	<ul style="list-style-type: none"> • H2B-mCherry (histone H2B fluorescent nuclear reporter protein). 	LSFM (SPIM)	<ul style="list-style-type: none"> • High-throughput imaging at the subcellular level is performed; • Detailed imaging of the whole spheroid. 	Paiè et al. (2016)
Clear ^T	<ul style="list-style-type: none"> • Formamide. 	4% paraformaldehyde	35 min	<ul style="list-style-type: none"> • Normal human dermal fibroblasts (NHDF). 	378 \pm 47	<ul style="list-style-type: none"> • Propidium iodide (PI). 	CLSM	<ul style="list-style-type: none"> • Spheroids size is not affected; • Fluorescence intensity may be reduced; 	(Costa et al., 2018a)

(Continues)

TABLE 3 (Continued)

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (μm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
Clear ^{T2}	<ul style="list-style-type: none"> Formamide; Polyethylene glycol (PEG; 8,000 Da). 	4% paraformaldehyde	75 min	<ul style="list-style-type: none"> C6 rat glioma; Primary cortical neurons; Rat adult neural stem cells (NSC). 	104.64 \pm 22.01	<ul style="list-style-type: none"> DAPI; Cy3; Dylight@488. 	CLSM	<ul style="list-style-type: none"> Allows up to \approx207.72 μm of image depth; Imaging in the interior of the spheroids is improved; Reversible. Longer incubation times does not improve the clearing; Spheroids size was not affected; Allows up to \approx100 μm of image depth; Imaging in the spheroids interior is improved. 	Boutin and Hoffmann-Kim (2015)
		10% neutral buffered formalin	75 min	<ul style="list-style-type: none"> PLHC-1 human liver cancer; LNcaP human prostate cancer; BEAS-2B human lung epithelium. 	n.a.	<ul style="list-style-type: none"> Hoechst 33342; Alexa FluorTM 647; CellROX[®] green. 	CLSM	<ul style="list-style-type: none"> Imaging depth is augmented; Imaging in the spheroids interior is improved. 	Kabadi et al. (2015)
		<ul style="list-style-type: none"> Formamide; Polyethylene glycol (PEG; 4,000 or 8,000 or 10,000 Da). 	4% paraformaldehyde	75 min	<ul style="list-style-type: none"> Normal human dermal fibroblasts (NHDF). 	396.17 \pm 28.72	<ul style="list-style-type: none"> Propidium iodide (PI). 	CLSM	<ul style="list-style-type: none"> Spheroids size was not affected independently of the PEG molecular weight; Fluorescence intensity is preserved independently of the PEG molecular weight;

(Continues)

TABLE 3 (Continued)

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (μm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
FocusClear™	<ul style="list-style-type: none"> FocusClear™ solution (derivative of polyoxyalkalene, diatrizoic acid, dimethyl sulfoxide (DMSO), ethylenediaminetetraacetic acid, glucamine, sodium diatrizoate, β-nicotinamide adenine dinucleotide phosphate). 	4% paraformaldehyde	2 hr	<ul style="list-style-type: none"> Intestinal stem cells isolated from mice. 	≈ 300	<ul style="list-style-type: none"> Propidium iodide (PI). 	CLSM	<ul style="list-style-type: none"> The PEG 4,000 Da allowed the higher imaging penetration depth ($\approx 211.67 \mu\text{m}$); PEG 4,000 enhanced the imaging of the spheroids interior; Reversible. Transparency is improved; Imaging performed with high resolution. 	Chen et al. (2013)
Delipidation and dehydration followed by RI matching									
BABB	<ul style="list-style-type: none"> Benzyl alcohol; Benzyl benzoate; Ethanol. 	4% paraformaldehyde	>20 min	<ul style="list-style-type: none"> T47D human breast cancer; DLD1 human colorectal cancer; DU145 human prostate cancer; Primary human colon cancer cells. 	≈ 400	<ul style="list-style-type: none"> SYTOX™ Green; Fluorescein isothiocyanate (FITC); Alexa Fluor™ 555. 	<ul style="list-style-type: none"> monolithic digital LSFM (mDSLIM) 	<ul style="list-style-type: none"> Clear imaging of spheroids hypoxic, dead, and proliferative cells 	Wenzel et al. (2014)
		Formalin	n.a.	<ul style="list-style-type: none"> HCT116 human colorectal cancer. 	650	<ul style="list-style-type: none"> Propidium iodide (PI). 	LSFM	<ul style="list-style-type: none"> Imaging depth is improved; Imaging performed with high resolution 	Desmaison et al. (2018)
		n.a.	n.a.	<ul style="list-style-type: none"> T47D human breast cancer. 	150–500	<ul style="list-style-type: none"> Drag5. 	<ul style="list-style-type: none"> Monolithic Digital Scanned 	<ul style="list-style-type: none"> Allows the acquisition of high-quality images of 	Schmitz et al. (2017)

(Continues)

TABLE 3 (Continued)

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (µm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
		70% ethanol or methanol or 4% paraformaldehyde	>14 min	<ul style="list-style-type: none"> U343 human glioblastoma. 	≈600	<ul style="list-style-type: none"> Alexa Fluor™ 488; Alexa Fluor™ 568; DAPI. 	LSFM (mDSLML) Monolithic Digital Scanned LSFM (mDSLML)	intact spheroids at cellular resolution; • Transparency is improved; • Occurs spheroids shrinkage.	Smyrek and Stelzer (2017)
Delipidation and hyperhydration followed by RI matching									
Scale	<ul style="list-style-type: none"> Glycerol; Triton X-100; Urea 	4% paraformaldehyde	72 hr	<ul style="list-style-type: none"> Rat adult neural stem cells (NSC). 	104.64 ± 22.01	<ul style="list-style-type: none"> DAPI; Cy3. 	CLSM	<ul style="list-style-type: none"> Double concentration of Triton X-100 improves the clearing; Longer incubations times does not improve the clearing; Transparency is improved; Occurs spheroids expansion; Spheroids become mechanically fragile. 	Boutin and Hoffmann-Kim O (2015)
	<ul style="list-style-type: none"> Sorbitol; Urea 	<ul style="list-style-type: none"> 2% formaldehyde 	<1 hr	<ul style="list-style-type: none"> MCF-7 human breast cancer cells 	370 ± 90	<ul style="list-style-type: none"> Fluorescent Ubiquitination-based Cell Cycle Indicator (FUCCI). 	MPM (2PM)	<ul style="list-style-type: none"> Transparency is improved; Occurs spheroids expansion; Nuclei blur is less extent; Allows more than 250 µm of image depth; Fluorescence intensity is increased; Occurs fluorescence shift. 	Grist et al. (2016)

(Continues)

TABLE 3 (Continued)

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (µm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
CUBIC	<ul style="list-style-type: none"> • 2,2',2''-nitriiotriethanol; • Glycerol; • N,N,N',N'-tetrakis(2-hydroxypropyl) ethylenediamine; • Triton X-100; • Sucrose; • Urea 	10% neutral buffered formalin	≈6–10 days	<ul style="list-style-type: none"> • HCT116 human colorectal cancer 	400	<ul style="list-style-type: none"> • Alexa Fluor™ 594; • H2B-mCherry (histone H2B fluorescent nuclear reporter protein). 	LSFM (adaptive SPIM)	<ul style="list-style-type: none"> • Transparency is improved; • Allows the acquisition of high-resolution in-depth images; • Improved image quality in terms of peak intensities and contrast. 	Masson et al. (2015)
Hydrogel embedding followed by delipidation and RI matching									
CLARITY	<ul style="list-style-type: none"> • Acrylamide; • Bisacrylamide • VA-044 • SDS • X-Clarity™ 	4% paraformaldehyde	2 days	<ul style="list-style-type: none"> • Human-derived adipose stem cells (hASC). 	200 ± 50	<ul style="list-style-type: none"> • Alexa Fluor™ 647; • Atto488™; • DAPI. 	CLSM	<ul style="list-style-type: none"> • Clearing was performed with the assistance of a microfluidic device; • Transparency is improved; • Leads to spheroids shrinkage; • Nuclear structure and the positions of the proteins are not disturbed; • Fluorescence intensity is acquired in the entire spheroid; • Clearing time is reduced to 5 hr by alternating the pH in the surrounding hydrogel-embedded spheroids. 	Silva Santibañ et al. (2018)
	<ul style="list-style-type: none"> • Acrylamide • VA-044 • RIMS (lohexol, 1,4-diazabicyclo[2.2.2]octane, sodium borate, and sodium azide). 	4% formaldehyde	<5 hr	<ul style="list-style-type: none"> • MCF-7 human breast cancer cells; • MDA-MB-435 human melanoma cells; 	250 and 450	<ul style="list-style-type: none"> • CellTracker™ Red; • DAPI; • DiO; • GFP; 		<ul style="list-style-type: none"> • Clearing was performed with the assistance of a microfluidic device; • Transparency is improved; 	Chen et al. (2016)

(Continues)

TABLE 3 (Continued)

Clearing method	Reagent (s)	Fixative	Clearing process time ^a	Spheroids cells	Spheroids diameter (µm)	Fluorescent probes	Microscopy technique	Observation(s) ^b	Reference
				<ul style="list-style-type: none"> B16F10 murine melanoma cells (endogenously expressing tdTomato); U87MG human glioblastoma cells (endogenously expressing GFP); HUVEC human umbilical vein endothelial cells. 	<ul style="list-style-type: none"> LIVE/DEAD® Fixable Near-IR Dead Cell Stain; Phalloidin-488; tdTomato; Transferrin-647. 		<ul style="list-style-type: none"> Spheroids not embedded in the gel are deformed by the fluidic forces; Higher concentrations of acrylamide allow better preservation of spheroids structure; DIO fluorescence is lost during spheroids treatment with SDS, while this process does not affect the spheroids stain with Transferrin-647 and facilitated the spheroids stain with DAPI, and Phalloidin-488; Imaging depth is improved by 150% for MCF-7 spheroids, and 250% for U87MG-GFP spheroids; Method is compatible with live/dead imaging of drug-treated spheroids. 		

n.a.: not available.

Abbreviations: BABB: benzyl alcohol/benzyl benzoate; CLSM: confocal laser scanning microscopy; DAPI: 4',6-diamidino-2-phenylindole; LSM: light-sheet fluorescence microscopy; M2M: multiphoton microscopy; SeeDB: See Deep Brain; SPIM: single (or selective) plane illumination microscopy; 2PM: two-photon microscopy.

^aTime spent in the clearing process, excluding the spheroids fixation time.

^bInformation available about the clearing technique.

preserves the lipidic content of each sample, thus allowing the imaging of these structures using lipophilic tracer dyes (e.g., Dil and FM 1-43FX; Yu et al., 2018). Nevertheless, the preservation of the lipid content (which are one of the main factors responsible for light scattering in biological tissues) restricts the sample transparency (Tuchin, 2015). Furthermore, the incorporation of the aqueous clearing solution in the sample may induce an increase in the sample volume and consequently change its initial structure (Richardson & Lichtman, 2015). Such also impacts on the imaging of the whole spheroid due to the increased image acquisition time and post-processing issues (Lee et al., 2016). Lastly, since these methods rely on the passive diffusion of the solutions into the sample, the clearing process may need extensive incubation times depending on the sample size (Lee et al., 2016; Seo et al., 2016).

3.1.1 | See Deep Brain

The RI of the biological samples' constituents can be homogenized by immersing them in sugar-based aqueous solutions that have a high RI. In this way, Ke, Fujimoto, and Imai (2013) used fructose to develop the See Deep Brain (SeeDB) method that was applied for the clearing of large mouse samples, like whole embryos and neonatal brains. In SeeDB method, samples are successively immersed in 20, 40, and 60% (w/v) fructose aqueous solutions during 4–8 hr, followed by their incubation in 80% fructose during 12 hr and lastly in the SeeDB solution (RI \approx 1.49 at 25°C; 115% (w/v) fructose and 0.5% (v/v) α -thioglycerol) during 24 hr (Ke & Imai, 2014; Ke et al., 2013). The α -thioglycerol is used to avoid the Maillard browning discoloration derived from the reactive ketone group of fructose, which can lead to autofluorescence (Dills, 1993). Grist et al. (2016) applied SeeDB to clear MCF-7 breast cancer spheroids with $370 \pm 90 \mu\text{m}$ of the diameter that were transfected with fluorescent ubiquitination-based cell cycle indicator (FUCCI) for imaging by using 2PM. The clearing process was performed with the assistance of a microfluidic system, where spheroids were trapped. Then, clearing solutions (20, 40, 60, 80, 100, and 115% (w/v) fructose solutions with 0.5% (v/v) α -thioglycerol) were perfused through the microchannels using a syringe pump at a rate of approximately $20 \mu\text{l}/\text{min}$. The results demonstrated that the method induces a minimal reduction of the size of the spheroids, that is, the ratio of spheroids area before and after the clearing was 1.02 (Grist et al., 2016). This reduction is dependent on the flow rate used in the microfluidic channel, that is, higher flow rates lead to higher shrinkage, which may be related to the fluidic compressive forces or to the osmotic pressure. The transparency was slightly improved since the light transmittance increased to about 5.5% in the cleared spheroids (Grist et al., 2016). An analysis of the average FUCCI fluorescence intensity as a function of penetration depth into the sample demonstrated that the SeeDB method allowed to image the spheroids at depths greater than $250 \mu\text{m}$, while the imaging of noncleared spheroid was limited to about $150 \mu\text{m}$ (Grist et al., 2016). Nevertheless, the authors also observed that SeeDB solution increased the green

fluorescence intensity of the spheroids (Grist et al., 2016), which was attributed to the formation of Maillard reaction products (Berke, Miola, David, Smith, & Price, 2016).

Boutin & Hoffman-Kim (2015) also applied the SeeDB method to clear rat adult neural stem cells (NSC) spheroids (mean diameter of $104.64 \pm 22.01 \mu\text{m}$). Authors embedded the spheroids in 20, 40, and 60% (w/v) fructose solutions (8 hr each). Then, spheroids were immersed in 80% (w/v) and in 100% (w/v) fructose solutions (12 hr each) and lastly during 24 hr in 115% (w/v) fructose, with all solutions containing 0.5% (v/v) α -thioglycerol. The spheroids after the clearing process presented moderate transparency in comparison to the noncleared spheroids (maintained in phosphate buffered saline [PBS]; Boutin & Hoffman-Kim, 2015). Interestingly, despite being described that the SeeDB does not significantly influence the samples size (Grist et al., 2016; Ke et al., 2013), these authors observed that the spheroids shrink (104.64 ± 22.01 vs $76.2 \pm 13.77 \mu\text{m}$ in diameter) occurred as a consequence of the clearing process (Boutin & Hoffman-Kim, 2015). Authors also observed that 4',6-diamidino-2-phenylindole (DAPI)-labeled nuclei were difficult to distinguish. Furthermore, during the execution of the experimental protocol, the use of highly viscous solutions made the spheroids manipulation difficult, which resulted in the loss of many samples during the assay (Boutin & Hoffman-Kim, 2015). Moreover, the viscosity of highly concentrated solutions of fructose also made challenging the acquisition of in-focus images containing both the spheroid and the background (Boutin & Hoffman-Kim, 2015).

Alternatively, poor viscous clearing aqueous solutions with high RI can be obtained by using TDE, formamide (used in *Clear^T* and *Clear^{T2}* methods) or diatrizoic acid (used in *FocusClearTM* method), as discussed hereafter.

3.1.2 | TDE

2,2'-Thiodiethanol (TDE) is a nontoxic, cheap, and water-soluble liquid displaying a low viscosity that has been widely used as a mounting media, since it has a RI (for 100% TDE the RI is 1.52) similar to that of typical immersion oils used in fluorescence microscopy (Smyrek & Stelzer, 2017; Staudt, Lang, Medda, Engelhardt, & Hell, 2007). Moreover, TDE has also been used as a clearing agent, since it allows the control of the RI value by diluting it with water, that is, 30, 60, and 97% (v/v) TDE:water solutions have RI \approx 1.39, 1.45, and 1.52, respectively (Aoyagi, Kawakami, Osanai, Hibi, & Nemoto, 2015; Costantini et al., 2015). Aoyagi et al. (2015) applied TDE solutions (30, 60, and 97% (v/v) in PBS) to clear mice brains and brain slices after their incubation for 6 hr to 7 days, for improving the light penetration depth in both CLSM and 2PM. TDE has also been applied for the clearing of spheroids. Paiè et al. (2016) applied TDE to clear H2B-mCherry (histone H2B fluorescent nuclear reporter protein) expressing spheroids with an average diameter of $300 \mu\text{m}$. For that purpose, authors dipped spheroids in different TDE solutions (25, 50, and 68% (v/v) in PBS) for 10 min each. Then, the high-throughput imaging of the spheroids was performed in an optofluidic lab-on-a-chip that integrates SPIM illumination and

continuous sample delivery in a microfluidic channel. Briefly, after the clearing, the spheroids circulated at a constant speed (20 $\mu\text{m/s}$) in a microchannel containing a 0.1% (v/v) TDE solution and their imaging was performed quickly (50 ms acquisition time) while passing through the light sheet. Combining the use of TDE with SPIM, it was possible to acquire images of the entire spheroid with a subcellular resolution that allowed the clear identification of the cells' nuclei, the cell counting and the observation of the cell mitosis (Paiè et al., 2016). Furthermore, it was possible to perform the 3D rendering of the acquired images and consequently obtain a 360° detailed view of the spheroids (Paiè et al., 2016).

3.1.3 | Clear^T

Formamide is a water-miscible polar solvent that was used for the first time by Kuwajima et al. (2013) in a method, termed *Clear^T*, for clearing mice tissues (whole brains, brain sections, and embryos). In this method, samples are dipped in gradient solutions of formamide in water (20 up to 95% (v/v) formamide) to homogenize the RI of the tissue to about 1.44 (Richardson & Lichtman, 2015). The sample incubation time in each solution varies according to the sample size, for example, the clearing of whole brains took more than 2 days, while 20–1,000 μm brain sections took less than 1 hr. Our research group also applied this method to allow the analysis of spheroids composed of normal human dermal fibroblasts (NHDF), with a diameter of $378 \pm 47 \mu\text{m}$ (Costa, Moreira, de Melo-Diogo, & Correia, 2018a). In this method, spheroids were incubated for 5 min in each formamide/water solution (20, 40, 80, and 95% (v/v)), followed by a final incubation step of 15 min in formamide 95% (v/v), and then imaged by CLSM. The method improved the spheroids transparency and can be reversed by immersing the samples in PBS, which assures that the samples are not chemically modified by the clearing method Ke et al., 2013. The propidium iodide (PI) signal depth on the cleared spheroids was about 43% higher than that on the noncleared spheroids (207.72 ± 19.54 vs. $145.71 \pm 15.66 \mu\text{m}$; Costa et al., 2018a). Moreover, the modification of spheroid size was not significant, after the clearing process the spheroids presented a diameter of $407 \pm 27 \mu\text{m}$, while the noncleared ones displayed a diameter of $378 \pm 47 \mu\text{m}$ (Costa et al., 2018a). Nevertheless, although *Clear^T* allowed a successful clearing of the samples, the formamide-mediated protein denaturation can result in the quenching of the fluorescent proteins signal, as observed for green fluorescent protein (GFP) in mouse embryos (Kuwajima et al., 2013; Richardson & Lichtman, 2015). Furthermore, *Clear^T* may also reduce the PI fluorescence in spheroids (Costa et al., 2018a).

3.1.4 | Clear^{T2}

The *Clear^{T2}* method is an adaptation of the *Clear^T* with the objective of avoiding the formamide-mediated quenching of the fluorescent protein dyes. In this way, the *Clear^{T2}* method described by Kuwajima et al. (2013) aimed to preserve the fluorescence of the samples by promoting the protein stabilization using polyethylene glycol (PEG)

8,000 Da in combination with formamide (Rawat, Raman Suri, & Sahoo, 2010). In brief, the clearing of the biological samples (mice whole brains, brain sections, and embryos) was achieved through their immersion in graded series of formamide/PEG solutions (once in 25/10% (v/w) and then twice in 50/20% (v/w); Kuwajima et al., 2013). Similarly, the time of incubation in each solution depends on the sample size (clearing of the whole mouse brains takes ≈ 18 hr, while 20–1,000 μm sections require less than 2 hr).

The *Clear^{T2}* method has been widely described for spheroids clearing (Boutin & Hoffman-Kim, 2015; Boutin et al., 2018; Kabadi et al., 2015). Boutin and Hoffman-Kim (2015) used *Clear^{T2}* to process C6 glioma, primary cortical neuron, and NSC spheroids by embedding them during 10 min in 25/10% (v/w) formamide/PEG 8,000, followed by 5 min in 50/20% (v/w) formamide/PEG 8,000 and lastly 60 min in 50/20% (v/w) formamide/PEG 8,000. The authors observed that no significant changes occurred in the volume of the NSC spheroids, after their clearing through *Clear^{T2}*, that is, the diameter of cleared spheroids was $99.77 \pm 22.3 \mu\text{m}$ and $104.64 \pm 22.01 \mu\text{m}$ for the noncleared ones. Furthermore, the fluorescence signals of tracers (nucleus [DAPI]) and antibody immunostaining (Cadherin (Cy3), laminin (Dylight@488), nestin (Cy3), β -III tubulin (Cy3), S100 (Cy3)) were detectable by CLSM at all depths within spheroids as well as the fine features within the center of the spheroids (Boutin & Hoffman-Kim, 2015). The authors also described that prolonging the spheroid incubation time did not improve the clearing efficacy. Recently, the same research group also showed the applicability of the *Clear^{T2}* method for the imaging of immunostained spheroids composed of primary-derived postnatal rodent cortex cells by CLSM and MPM, to study in vitro the formation of endothelial cell capillary-like structures (Boutin et al., 2018).

In another study, *Clear^{T2}* was used in conjunction with immunostaining and biochemical staining for the imaging of spheroids of PLHC-1, LNCaP, and BEAS-2B cells by CLSM (Kabadi et al., 2015). *Clear^{T2}* allowed the sharp visualization of the nuclear structure (Hoechst 33342) even in the center of the microtissue at depths superior to 75 μm along the z-axis, contrasting with the maximum depth of 30 μm in the noncleared spheroids (Kabadi et al., 2015). The method also allowed improved imaging of the spheroids' E-cadherin cell-cell interactions (Alexa Fluor™ 647), as well as the reactive oxygen species by using the CellROX® green assay (Kabadi et al., 2015).

In our research group, the *Clear^{T2}* method was used for the clearing of PI-stained NHDF spheroids (diameter of $396.17 \pm 28.72 \mu\text{m}$; Costa, Moreira, de Melo-Diogo, & Correia, 2018b). In this study, the influence of PEG molecular weight (4,000, 8,000, and 10,000 Da) on the clearing efficacy of the *Clear^{T2}* method described by Boutin and Hoffman-Kim (2015) was evaluated. Independently of the PEG molecular weight, the *Clear^{T2}* clearing method contributed to increasing the spheroids transparency and for the preservation of the PI fluorescence intensity (Costa et al., 2018b). Furthermore, no significant changes in the size of the spheroids were observed for all the conditions tested. Nevertheless, the *Clear^{T2}* method performed using PEG 4,000 allowed a better PI signal depth, that is, the use of PEG 4,000 Da allowed the detection of the PI signal up to $211.67 \pm 16.81 \mu\text{m}$, while for the PEG 8,000 and

10,000 Da the fluorescence was only detected up to 183.89 ± 14.13 and $176.43 \pm 11.44 \mu\text{m}$, respectively (Costa et al., 2018b). Moreover, spheroids fluorescence signal in the cross-section penetration depth was also improved by the use of PEG 4,000 Da (more than 24% compared to that of noncleared spheroids, at a penetration depth of 100 μm in the z-axis; Costa et al., 2018b). These results may be related with the fact that smaller PEG chains can penetrate and distribute more easily through the spheroids and consequently promote the stabilization of the fluorescence probe in deeper regions of spheroids, thus allowing better imaging.

3.1.5 | FocusClear™

FocusClear™ is a nontoxic, ready to use clearing solution that was originally developed for the processing of fruit fly brains and now is commercialized by CelExplorer Labs (U.S. Pat. No. 6472216B1 and China Taiwan Patent No. 206390). FocusClear™ solution comprises a mixture of diatrizoic acid, the clearing agent, with dimethyl sulfoxide (DMSO), ethylenediaminetetraacetic acid, glucamine, β -nicotinamide adenine dinucleotide phosphate, sodium diatrizoate, and a derivative of polyoxyalkalene (Genina et al., 2010). In the literature, it was already demonstrated that FocusClear™ can enhance the transparency of several types of biological tissues (e.g., mouse intestine (Fu et al., 2009) and mouse brain tissues (Moy et al., 2015)), as well as in spheroid samples (Chen et al., 2013). Chen et al. (2013) reported the clearing of crypt spheroids composed of intestinal stem cells isolated from mice, with approximately 300 μm of thickness, by performing their immersion during 2 hr in the FocusClear™ solution. Optical light microscopic observation revealed that it was possible to observe that spheroids maintained in PBS were opaque and dark in their central regions, while the cleared spheroids were almost transparent (Chen et al., 2013). In addition, the clearing process also allowed the visualization of the PI fluorescence signal across the spheroid by CLSM as well as the clear identification of the crypt spheroid shape and the protruded villi-crypt domains (Chen et al., 2013).

3.2 | Delipidation and dehydration followed by RI matching

To expedite the clearing process, researchers developed methods based on the lipids removal through the use organic solvents, that is, delipidation and dehydration, followed by RI matching methods (Spalteholz's technique, BABB, 3DISCO, and its derivatives). These approaches usually require two major steps: (a) initial dehydration of the sample, and (b) immersion of the samples in organic solvents (Figure 2). The initial dehydration step is usually performed using ethanol or methanol, since the solvents used in the following steps are not miscible with water (Seo et al., 2016). During the dehydration, step can also occur the removal of some lipids. After the water removal, the overall RI of the sample will increase since dehydrated proteins have a RI value >1.5 , which is higher than that of water and lipids (1.33 and 1.44, respectively; Richardson & Lichtman, 2015). Therefore, in the second step, samples are immersed in an

organic solvent with higher RI (>1.5) to match the value of the dehydrated sample. Ideally, the organic solvents should also promote the removal of the remaining lipids since this will ensure a better homogeneous RI matching (Richardson & Lichtman, 2015).

Although these methods result in improved transparencies in large samples in shorter periods of time, these approaches are well known for disrupting the fluorescence of some probes (e.g., GFP), due to the use of fixatives like ethanol, methanol, and acetone as well as organic solvents that cause structural changes in the proteins (Lee et al., 2016). Moreover, the samples dehydration can lead to their shrinkage, which can affect the fluorescence of some proteins. Fluorescent protein chromophores need water molecules to sustain their fluorescence emission (Richardson & Lichtman, 2015). This sample shrinkage occurs due to the loss of the 3D hydrogen-bonding networks between the water (70–80% of the samples) and the remaining intracellular constituents, an interaction that the dehydrating agents cannot fully replicate (Tainaka et al., 2016). Lastly, the organic solvents are toxic and can destroy the glues, as well as the equipment used in samples imaging.

3.2.1 | Spalteholz's technique

The first optical clearing method based on organic solvents was described a century ago by Spalteholz and it was developed to allow the histological analysis of large biological samples (e.g., entire organs and organ systems; Spalteholz, 1914). This method included an initial sample fixation with formalin, followed by its bleaching with hydrogen peroxide (used to perform hemoglobin discoloration which is responsible for the biological tissues visible light absorption). Then samples were dehydrated, using a series of alcohol concentrations, and immersed in a mixture of methyl salicylate and benzyl benzoate. Nevertheless, this method leads to samples shrinkage, browning, and also to the damage of the biological samples outer layers (2–3 cm), due to the formation of bubbles and cell death prompted by hydrogen peroxide (Alnuami, Zeedi, Qadri, & Ashraf, 2008; Cumley, Crow, & Griffen, 1939; Steinke & Wolff, 2001). Additionally, the Spalteholz method can also affect the fluorescence emission by protein dyes (e.g., GFP loses its fluorescence upon exposure to oxyradicals that are produced by the hydrogen peroxide; Alnuami et al., 2008). Therefore, this method has not been used for the clearing of 3D cell cultures but is the basis of other clearing methods aimed for the spheroids processing (described hereafter).

3.2.2 | Benzyl alcohol/benzyl benzoate

Inspired by the Spalteholz method, Dent, Polson, and Klymkowsky (1989) and Dodt et al. (2007) reported the application of benzyl alcohol/benzyl benzoate (BABB)-based optical clearing protocol (also known as Murray's clear). The BABB method was initially used to clear mice embryos and whole brains for allowing their imaging by ultramicroscopy. The method includes an initial dehydration step based on the samples' incubation for 1 day in each ethanol solution (i.e., 30, 50, 70, 80, 96, and twice in 100% (v/v) ethanol) with an

additional 1 hr in 100% (v/v) hexane to achieve the maximum dehydration. Then, the samples are incubated with the clearing solution of benzylalcohol/benzyl benzoate (1:2 (v/v)) during 2 days.

Wenzel et al. (2014) adapted the BABB method to clear spheroids of T47D, DLD1, DU145, and primary colon cancer cells with a mean diameter of $\approx 400 \mu\text{m}$. For that purpose, spheroids were initially dehydrated in ascending ethanol concentrations (50, 70, 85, and 99% (v/v); 5 min each) and then transferred to a BABB (1:1 (v/v)) solution. The authors demonstrated that the spheroids clearing allowed the visualization of the hypoxic (antipimonidazole antibody conjugated with FITC, a marker for hypoxia) and the dead cells (SYTOX™ Green stain for dead cells) in the inner regions of the spheroid, as well as the proliferative cells (Click-iT™ EdU Alexa Fluor™ 555) in their periphery using a custom build monolithic digital LSFM (mDSLIM; Wenzel et al., 2014). Furthermore, the authors could evaluate the presence of dead cells in the spheroids core region after the administration of different compounds that target dormant cells (e.g., Antimycin, Cisplatin, Metformin, Paclitaxel, Rotenone, and Staurosporine). These results allowed the authors to confirm that the spheroids mimic the *in vivo* tumor microenvironment and additionally that this technique can be used in the high-throughput and high-content microscopy analysis for identifying substances that specifically target dormant cells in spheroids core regions (Wenzel et al., 2014).

Desmaison et al. (2018) also applied the BABB clearing and LSFM microscopy for the structural analysis of large HCT116 spheroids (average diameter of $650 \mu\text{m}$). The clearing was performed through the sequential incubation of spheroids in 25, 50, 75, and 95% (v/v) ethanol solutions and lastly in BABB solution (benzyl alcohol:benzyl benzoate, 1:2 (v/v)). After the clearing, the authors demonstrated that the imaging of the PI-stained cell nuclei in the deep interior regions of the spheroids could be performed with improved resolution and detail. The obtained results revealed that the shape of the cell nuclei and the cell division is influenced by the spheroid growth conditions, that is, free growth over a nonadhesive surface versus physically confined growth conditions (spheroids embedded in agarose; Desmaison et al., 2018). Similarly, Schmitz et al. (2017) applied the BABB clearing method and mDSLIM to image Draq5 stained T47D spheroids (displaying diameters ranging from $150 \mu\text{m}$ to more than $500 \mu\text{m}$). After the clearing procedure, the authors could perform the study of several parameters in spheroids, such as spheroids volume, number of cells, and cell nucleus volume by using the automated nuclei segmentation and image computational analysis.

Smyrek and Stelzer (2017) applied the BABB method to clear U343 spheroids (average diameter of $600 \mu\text{m}$) with the aim to optimize an immunofluorescence staining protocol for 3D cell biology. The spheroids GM130 (cis-Golgi matrix protein), α -tubulin, and β -catenin proteins were immunolabeled and the cell nucleus was stained with DAPI. For the clearing, spheroids were dehydrated with increasing concentrations of ethanol (30, 50, 70, 90, 96, and twice in 100% (v/v), 2 min each) and then transferred to the BABB solution (benzyl alcohol:benzyl benzoate, 1:2 (v/v)) until the transparency was

attained (Smyrek & Stelzer, 2017). After, the authors quantitatively analyzed the specificity and homogeneity of the stain, as well as the Alexa Fluor™ 568, Alexa Fluor™ 488, and DAPI signal intensity. Overall, the results demonstrated that the best protocol for the spheroids immunofluorescence labeling included: (a) fixation of the spheroids during 15 min with 4% paraformaldehyde at room temperature (RT); (b) detergent-based permeabilization by Triton X-100, during 15 min, at RT; (c) blockage of unspecific binding sites with a solution of 0.1% bovine serum albumin, 0.2% Triton X-100, 0.05% Tween-20% and 10% goat serum for 1 hr at RT; (d) incubation of the first antibody at 37°C during 18–24 hr; (e) incubation of the second antibody at 37°C during 4 hr; (f) dehydration and clearing by BABB during 15 min; and (g) imaging by mDSLIM (Smyrek & Stelzer, 2017).

Although the utilization of the BABB clearing method can result in samples with high tissue transparency, the use of ethanol/methanol during the dehydration step still results in a reduction of the size of the spheroids, which can impact the analysis of the data extracted from these samples (Smyrek & Stelzer, 2017).

3.2.3 | 3D imaging of solvent-cleared organs

Since the ethanol dehydration step used in the BABB method results in the quenching of the proteins fluorescence (Becker, Jährling, Saghafi, Weiler, & Dodt, 2012; Feuchtinger et al., 2016), Becker et al. (2012) investigated alternative chemicals (RI between 1.5 and 1.7) that could be used for samples dehydration. The results obtained demonstrated that the utilization of tetrahydrofuran (THF) in brain hemispheres has a similar dehydration effect to that of ethanol, with the advantage of the background fluorescence intensity be considerably reduced and the intensity of the fluorescence signal be enhanced (Becker et al., 2012). This solvent adaptation was termed the 3DISCO clearing method. In this procedure, the samples were initially dehydrated by using grading THF solutions (50, 70, 80% (v/v)) followed by their incubation in THF 100% (v/v), 1–3 times, dichloromethane, and then in dibenzylether (Erturk et al., 2012). The incubation time of the samples depends on its size, that is, few hours for small organs (few millimeters in size) such as spinal cords, mammary glands or lymph nodes and a day for large organs such as the brain.

3DISCO protocol variations have been also developed, such as iDISCO (immunolabeling-enabled 3D imaging of solvent-cleared organs), uDISCO (ultimate 3D imaging of solvent-cleared organs), and FDISCO (DISCO with superior fluorescence preserving capability), that were describe by Renier et al. (2014), Pan et al. (2016), and Qi et al. (2019), respectively. In the iDISCO, samples are pretreated with methanol, Triton X-100, and dimethyl sulfoxide (DMSO) to facilitate the antibody penetration and consequently improve the whole-organ immunolabeling (Renier et al., 2014). On the other hand, the uDISCO was developed to significantly reduce the size of the samples (up to 65% of its original size) by using tert-butanol as a dehydration reagent, which can make possible the observation of large samples, with sizes up to the whole mouse body

(Pan et al., 2016). The FDISCO method allows overcoming the quenching of the fluorescent proteins that happens in the 3DISCO method by controlling the pH and temperature conditions during the clearing procedure (Qi et al., 2019).

Despite the 3DISCO and its modified methods being widely used for the clearing of several animal samples, its application in spheroids processing is yet to be described.

3.3 | Delipidation and hyperhydration followed by RI matching

The delipidation and hyperhydration followed by RI matching methods (Scale and CUBIC) use solutions of urea (denaturing agent) and Triton X-100 (nonionic detergents) for promoting the clearing of the samples (Figure 2). During this process, the detergents will remove the lipids that have high RI (≈ 1.44) and therefore reduce the sample overall RI. Furthermore, the urea will penetrate the cells denaturing the folded proteins (RI ≈ 1.43), which also reduces the sample RI (Richardson & Lichtman, 2015). Moreover, the urea also creates an osmotic gradient that will favor the water diffusion into the sample (hydration) and therefore homogenizes the overall RI value to ≈ 1.38 (Richardson & Lichtman, 2015).

These methods are easy to perform and have a high clearing capacity, due to the removal of lipids from the sample, without leading to fluorescence quenching and toxicity problems associated with the solvent-based clearing methods (Seo et al., 2016). Furthermore, this approach is also compatible with immunoblotting techniques (e.g., immunofluorescent labeling with antibodies) since the cellular membrane is permeable to the antibodies after its treatment with the detergent (Mattei, Lira, Perez, & Riske, 2017). On the other hand, the samples can undergo hydration-induced expansion, which is undesirable during the analysis of the spheroids (Seo et al., 2016).

3.3.1 | Scale

Hama et al. (2015) developed the Scale clearing method for the imaging of brain samples obtained from mice and patients. These researchers initially observed that the use of 8 M urea solutions can improve the tissue transparency, but also results in great tissue expansion. In this way, these authors tested different Scale solutions composed of urea, as well as Triton X-100 and glycerol with the objective to reduce the expansion of the samples and improve the tissue transparency. The Triton X-100 detergent was used to promote the lipids removal from the sample and therefore improve the sample transparency. Glycerol was added to counterbalance the urea-induced tissue expansion, assist the dehydration, and to target the lipophilic tissue regions. Among the different solutions investigated, the better clearing results were obtained when ScaleA2 (4 M urea, 10% (w/v) glycerol and 0.1% (w/v) Triton X-100; RI ≈ 1.378) solution were used. However, the clearing took several weeks (e.g., more than 2 weeks for mouse embryos) and the samples became highly fragile, difficult to handle, and tissue swelling was still

observed. Having this in mind, in a subsequent work this study group developed the ScaleS clearing method, where solutions composed of urea (4 M), Triton X-100 ($\leq 0.2\%$ (v/v)) and sorbitol (20, 27, 36.4, and 40% (w/v)) were used (Hama et al., 2015). The sorbitol replaced the glycerol due to its clearing properties and superior potential to reduce the sample volume expansion caused by urea.

Scale clearing solutions have also been employed for processing spheroid samples (Boutin & Hoffman-Kim, 2015; Hama et al., 2015). ScaleA2 solution (4 M urea, 0.2% (w/v) Triton X-100% and 10% (w/w) glycerol in water) was used to clear NSC spheroids (with an average diameter of $104.64 \pm 22.01 \mu\text{m}$) for imaging by CLSM (Boutin & Hoffman-Kim, 2015). In this work, a two times higher concentration of Triton X-100, in comparison to that described by Hama et al. (2015), was used for improving the samples clearing. The spheroids clearing was achieved by performing 3 cycles of 24 hr incubation with fresh ScaleA2 solutions. Further, the authors observed that the realization of the ScaleA2 protocol during longer periods did not improve the samples clearing. Moreover, the spheroids processing with the ScaleA2 resulted in a size expansion to an average diameter of $139.2 \pm 32.64 \mu\text{m}$ (Boutin & Hoffman-Kim, 2015). Lastly, the authors reported that the spheroids, in comparison to those incubated with PBS, become mechanically fragile after the clearing process.

In another study, the ScaleSQ (one of the Scale solutions investigated by Hama et al. (2015), composed of 9.1 M urea and 22.5% (w/v) sorbitol; RI ≈ 1.439) was used to clear MCF-7 breast cancer spheroids (diameter = $370 \pm 90 \mu\text{m}$; Grist et al., 2016). The method increased the average transmitted light intensity through the spheroids, that is, the transmittance after the clearing increased by 7.5%. Furthermore, the authors observed that the ScaleSQ significantly increased the FUCCI fluorescence intensity in the deeper regions of the spheroid and also the depth until each spheroid could be imaged (more than $250 \mu\text{m}$ deeper in the z-axis) by 2PM (Grist et al., 2016). Nevertheless, the clearing solution induced the sample swelling, about 1.7-times higher spheroids area after the clearing and appear to mediate a shift in the color, that is, the FUCCI orange fluorescence (characteristic of the cells' nucleus in interphase stages of their cell cycle) appear more yellow (Grist et al., 2016).

3.3.2 | Clear, unobstructed brain/body imaging cocktails, and computational analysis

Another clearing method that uses urea and Triton X-100 (like the Scale solutions) is the clear, unobstructed brain/body imaging cocktails and computational analysis (CUBIC). This method was initially developed for the clearing of mouse whole-brain imaging with LSFM (Susaki et al., 2014), and it involves the samples treatment with two clearing solutions, termed ScaleCUBIC-1 (also known as Reagent 1; 25% (w/w) urea, 25% (w/w) N,N,N',N'-tetrakis(2-hydroxypropyl) ethylenediamin and 15% (w/w) Triton X-100) and ScaleCUBIC-2 (also known as Reagent 2; 50% (w/v) sucrose, 25% (w/v) urea, 10% (w/v) 2,2',2''-nitrioltriethanol and 0.1% (v/v) Triton

X-100; RI = 1.48). The amino alcohol N,N,N,N-tetrakis(2-hydroxypropyl) ethylenediamine in the Reagent 1 solution was used as a tissue solubilizing agent which decolorizes blood and makes the organs much more transparent (Tainaka et al., 2014). In brief, the CUBIC protocol was performed by immersing the whole-brain samples in the Reagent 1 for 6–7 days (renewing the media after the first 3 days of incubation), then washing them several times with PBS followed by their incubation with 20% (w/v) sucrose in PBS and then a final immersion in the Reagent 2 solution for 2–7 days.

Masson et al. (2015) applied the CUBIC clearing method to clear HCT116 spheroids (average diameter of 400 μm) to develop a high-resolution in-depth imaging approach of optically cleared thick spheroids using an adaptive SPIM. For this purpose, spheroids were immersed with the Reagent 1 solution for either 2 or 4 days (temperature 37°C), then samples were transferred into a fresh Reagent 1 solution for additional 3–4 days. Then, spheroids were washed several times with PBS and incubated with 20% (w/v) sucrose and 50% (w/v) glycerol solution for a minimum of 2 hr. Lastly, spheroids were immersed in the Reagent 2 solution with gentle shaking for 1–2 days, before the imaging. The optical clearing guaranteed high transparency and reduced the aberration patterns of the spheroids, contributing to the high-resolution in-depth imaging of the Alexa Fluor™ 594 (Click-iT™ EdU) and H2B-mCherry signals (Masson et al., 2015). Moreover, the authors demonstrated that, after the clearing procedure, the spheroids' images improved in quality, namely the fluorescence intensity and contrast.

3.4 | Hydrogel embedding followed by delipidation and RI matching

The hydrogel embedding followed by delipidation and RI matching optical clearing methods comprehends the CLARITY and its derived approaches (e.g., PACT and PARS). These methods involve three major steps. The first step comprises the perfusion of the sample with a mixture of paraformaldehyde (or formaldehyde), acrylamide, and bisacrylamide monomers (that form a polyacrylamide gel in the sample after its polymerization). Then, the lipids are removed from the sample-gel hybrid by using a solution of an ionic detergent, namely sodium dodecyl sulfate (SDS). Lastly, the hydrogel-embedded samples are immersed on a RI-matching solution (e.g., FocusClear™) to render it transparency.

The gel will provide support for the biological structures to sustain their initial integrity, but it will also stabilize the proteins, nucleic acids, and other small molecules by cross-linking them with the mesh of the gel (Silvestri et al., 2016). Despite the sample being immersed in a hydrogel, its porous nature makes possible the penetration and diffusion of exogenous macromolecules, such as immunolabeling dyes and fluorochromes (Feuchtinger et al., 2016). Since CLARITY-based methods include the lipids removal, it is possible to achieve a great clearing in a similar way to other delipidation-based methods. However, due to the sample protection in a gel, the proteins concentration and folding is not affected in the same extent like when the methods containing harsh solvents or high

concentrations of detergents. Additionally, since the removal of the lipid is performed by using SDS instead of organic solvents (like those used in the BABB and 3DISCO methods), the fluorescence is preserved (e.g., GFP fluorescence; Chung et al., 2013).

3.4.1 | CLARITY and derived methods

The original clear lipid-exchanged acrylamide-hybridized rigid imaging/immunostaining/in situ hybridization-compatible tissue-hydrogel (CLARITY) method (also known as active CLARITY) was proposed in 2013 by Chung et al. (2013) to clear adult mouse whole brains and human brain samples of 500 μm in thickness. Authors, initially infused a mixture of 4% (w/v) paraformaldehyde, 4% (w/v) acrylamide, 0.05% (w/v) bisacrylamide and 0.25% (w/v) 2,2'-azobis [2-(2-imidazolyl)propane] dihydrochloride (VA-044; thermal radical initiator) into the tissues. Then, after the hydrogel polymerization (3 hr at 37°C), samples were subjected to an electric field potential (created by a custom electrophoresis system) and a 4% (w/v) SDS solution (1 or 2 days) to actively remove the lipidic content. Lastly, the samples were incubated in FocusClear™ (2 days for mouse brains and 1 day for human brain samples).

Nevertheless, the active CLARITY method requires specific equipment, as well as an expensive commercial RI-matching clearing solution (FocusClear™). Additionally, the strong electric field can change the molecular and structural integrity of the sample (Seo et al., 2016). Therefore, several CLARITY-derived methods emerged, such as PACT (passive clarity technique), PARS (Perfusion-assisted agent release in situ), and others (reviewed in detail by Jensen and Berg (2017) and Du, Hou, Zhang, and Li (2018)). The PACT and PARS methods use lower concentrations of fixatives and hydrogel monomers in conjugation with higher SDS concentrations (8% (w/v)), to facilitate and increase the passive diffusion of SDS into the sample and thus obtain a greater clearing without using the electrophoresis system. Additionally, both methods usually use RIMS (Refractive Index Matching Solution composed mainly of Histodenz or Sorbitol), which is a more economical RI-matching solution than FocusClear™ (Yang et al., 2014b). In the PACT method, sample-hydrogel hybrid is simply merged with an SDS solution, while in the PARS method the removal of the lipids is performed by perfusing continually the sample with the SDS solution (Yang et al., 2014b).

CLARITY-based methods were also used to clear spheroids. Silva Santisteban, Rabajania, Kalinina, Robinson, and Meier (2018) used CLARITY and a microfluidic system to clear an image human-derived adipose stem cells spheroids (200 \pm 50 μm in diameter) by CLSM. In brief, after spheroids trapping in the chambers of the microfluidic chip, these were perfused with a solution of paraformaldehyde, acrylamide, and bisacrylamide (4:4:0.25% (v/v)) for 1 hr (replaced every 10 min). After 2 hr of gel polymerization (VA-044 at 37°C), the lipids were removed by using 0.14 M SDS solution (pH = 8.5) that was renewed every 30 min during a total of 3 days. All solutions (except the thermal radical initiator) were flushed with a flow rate of 2 $\mu\text{l}/\text{min}$ using a flush time per chamber of 20 s. At last, X-Clarity™ (Logos Biosystems) was introduced as mounting media to match the RI. The

(Ariel, 2017; Zhu et al., 2013). In fact, optical clearing methods (e.g., SWITCH and PRESTO) were recently designed to actively improve the penetration of macromolecules into deep regions of tissues by altering electrical, physical and chemical factors (Lee et al., 2016).

On the other hand, researchers are still searching for optical clearing methods that permit the imaging of the cells without damaging or killing them, which will impact on the realization of time-course experiments (Calve, Ready, Huppenbauer, Main, & Neu, 2015; Neu, Novak, Gilliland, Marshall, & Calve, 2015). In fact, all the clearing methods described so far damage the cells (e.g., cells fixation and organic solvents utilization; Calve et al., 2015). Therefore, it is crucial to identify a RI-matching agent that does not harm the spheroids (Boothe, Hilbert, Heide, Berninger, & Huttner, 2017). Recently, Iodixanol has been investigated as a nontoxic medium supplement and RI-matching agent that improves image quality in live-imaging experiments involving primary cell cultures, planarians, zebrafish and human cerebral organoids (Boothe et al., 2017).

The development of a unarmful optical clearing method will also open several windows in terms of the therapeutic applications. In other words, the increased light penetration in cleared tissues may be useful for photo-based therapies, as well as drug-delivery monitoring.

5 | CONCLUSION

For more than 100 years, optical clearing methods have been used for the clearing of large biological samples obtained from animals. These methods have been developed to fulfill the following two major premises: (a) provide high transparency without affecting the initial structure and size of the sample; and (b) preserve endogenous and exogenous fluorescence (e.g., immunofluorescence), which is fundamental for the analysis of the proteins of interest.

More recently, optical clearing methods started to be explored for allowing the whole imaging of large 3D spheroids, as reviewed in this article. Till nowadays, SeeDB, TDE, *Clear^T*, *Clear^{T2}*, FocusClear™, BABB, Scale, CUBIC, and CLARITY optical clearing methods have been used for spheroids imaging under different types of microscopes (2PM, adaptive SPIM, CLSM, LSM, and mDSLIM). Gathering the information available in the literature it is possible to organize the four major types of optical clearing methods accordingly to their simplicity of execution, duration, clearing capacity, as well as their influence on samples integrity and preservation of the fluorescence of the proteins (Figure 3). Nevertheless, there are several other optical clearing methods (e.g., 3DISCO, FRUIT, PRESTO, and SWITCH) that are yet to be investigated in spheroids.

With future efforts, it will be possible to obtain a high-throughput-compatible approach to combine clearing, high-content imaging, and analysis of spheroids. Thus, optical clearing methods will contribute significantly to the widespread use of spheroids in the evaluation of different cellular events in normal and diseased human tissues.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

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