3D-Non-destructive Imaging through Heavy-Metal Eosin Salt Contrast Agents

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Abstract: Conventional histology is a destructive technique based on the evaluation of 2D slices of a 3D biopsy. By using 3D X-ray histology these obstacles can be overcome, but their application is still restricted due to the inherently low attenuation properties of soft tissue. In order to solve this problem, the tissue can be stained before X-ray computed tomography imaging (CT) to enhance the soft tissue X-ray contrast. Evaluation of brominated fluorescein salts revealed a mutual influence of the number of bromine atoms and the cations applied on the achieved contrast enhancement. The dibromo fluorescein barium salt turned out to be the ideal X-ray contrast agent, allowing for 3D imaging and subsequent complementing counterstaining applying standard histological techniques.

Medical diagnosis providing microscopic information on (sub-)cellular level is currently realized through conventional 2D histopathology, which is mainly based on light microscopy techniques. In combination with a large diversity of dyes and staining methods that have been developed over time various specific biological structures can be individually targeted. However, the current techniques are limited to the staining of only thin (2–20 μm) 2D microscopic slides that originate from a 3D biopsy sample. As the demand to extend biological and medical investigations to three dimensions has grown significantly over the last decade, several approaches for 3D imaging have been developed. Beyond serial-sectioning based approaches, these imaging methods include confocal and light sheet laser-scanning microscopy and block-face imaging (episcopic microscopy). Even though these methods have improved considerably over the years, they still require salt correction and registration steps, and the resulting reconstructions are often incomplete or do not represent the 3D structure of the imaged sample in a reliable way.

To overcome these limitations, X-ray microscopic computed tomography (μCT) and nanoscopic CT (nanoCT) imaging have proven to provide valuable 3D information of biological samples in a fast, convenient, and non-destructive way (Figure 1A). Here, non-destructive refers to the ability of X-ray CT to investigate biological samples using a multiscale approach ranging from whole organisms over whole organs to pieces of organs of animals without the need to embed or destruct the biological material. As a result, a 3D data set is received, which allows for virtually extracting any desired slice under any arbitrary angle throughout the entire volume (as an example see the video in the Supporting Information).

X-ray CT devices can be found at large synchrotron facilities but also in a laboratory environment capable of providing resolutions comparable to conventional 2D histology. Because of the intrinsically low attenuation properties of soft tissue at hard X-ray energies, the use of stains bearing high atomic number elements is inevitable in order to reach sufficient contrast (Figure 1B, C). Nonetheless, there is currently very limited availability of easy-to-handle X-ray staining agents that are (i) speedily penetrating the tissue without creating artefacts, (ii) targeting a specific biological morphology by staining the probe homogenously and completely, and (iii) at the same time suitable for large and dense tissue samples. In addition, staining procedures need to be introduced that are fully compatible with standard histology and thus allow further investigations of the region of interest (ROI) by the histologist, if necessary. First attempts to develop such dyes as X-ray contrast agents included modifications of elemental iodine I₂ for whole organ morphology and eosin Y disodium salt (2a) to target specifically the cell cytoplasm.

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The contrast within the soft tissue by using heavy-metal eosin Y salts 2b–2f. Because of the high Z value of heavy metals (Figure 1C), their implementation should allow to lower the concentration of contrast agent needed to reach a suitable contrast enhancement comparable to that of 2a. As water solubility might be impaired by the involvement of heavy metal atoms, we furthermore investigated the influence of the bromine atoms at the fluorescein core in 2 in response to solubility in aqueous media, tissue penetration, staining properties, and contrast enhancement, in order to implement a new “gold standard” for μCT of biopsy samples.

The heavy metal-eosin Y salts 2b–2f were synthesized by treating the lactone form of eosin Y (1) with the corresponding metal hydroxide and metal acetate, respectively (Scheme 1A and Supporting Information, chapter 2). The salts 2b–2f were obtained as homogenous solids after recrystallization in 12–51% yield.

In a preliminary solubility study, all eosin Y salts 2b–2f showed a significantly lowered maximum concentration in water when compared to the disodium compound 2a (see the Supporting Information, chapter 4). The Ag+ salt 2b as well as the Gd3+ salt 2f did not meet the sensitivity threshold needed for μCT measurements (<10 mg mL−1) and were thus not further investigated. The other compounds 2c–2e were then studied with regard to their staining properties.

The use of X-ray contrast agents to stain ex vivo biological samples often suffers from a spatially and temporally anisotropic stain penetration into soft tissue, leading to an artificial contrast gradient between the surface and the sample core. We thus chose turkey liver samples as soft tissue test pieces for staining as liver tissue is one of the densest soft tissues present in both birds and mammals. If the liver tissue can be completely and homogeneously stained within a reasonable time frame the dye should be capable of successfully staining other tissue types as well. Cuboidal turkey liver pieces of approximately 3 mm edge length were therefore treated in triplicates with 31.9 mm aqueous solutions of each dye 2c–2e, whereby the maximum water solubility of the barium-eosin Y salt (2c) was used as the concentration of the staining agents. Macroscopic investigations showed the best results using the Ba2+ eosin Y salt 2c with respect to uniform staining. In addition, no physical deformation of the tissue cubes, such as shrinking, was visible. An inhomogeneous and incomplete staining was observed for the samples treated with the Cu2+ 2d and Pb2+ salts 2e[13] even after an incubation time of 144 h (see Scheme 1B and Supporting Information, chapter 8). To allow for an X-ray contrast agent to enhance the soft-tissue contrast, the standard histological staining protocol had to be tailored towards X-ray μCT by means of pH adjustment of the tissue prior to staining, increasing the incubation times and the concentration of the contrast agent. As such acidification of the tissue sample prior to staining turned out to be crucial for 2a and 2c. The enhanced accumulation of both 2a and 2c in the cell cytoplasm can be explained by increased ionic interactions between the contrast agent and basic protein residues that are protonated at acidic pH values.[14] Under these conditions, a macroscopically complete and homogeneous staining for the

**Non-toxic compound**[10, 11] and readily available as the disodium salt 2a (Scheme 1), but its contrast enhancement is limited despite the four covalently bound bromine atoms (Z = 35). Best results for soft-tissue visualization employing 2a are achieved only by using very high concentrations of 2a (30% w/v) and cannot be further enhanced because of its limited water solubility (300 mg mL−1). Based on recent reports on 2a as contrast agent for μCT[10, 11, 6d–e] we investigated the possibility to further improve

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**Figure 1.** (A) Schematic representation of a μCT setup showing the X-ray source on the left and the detector on the right. In the middle is a rotation stage holding the soft-tissue sample. The Lambert–Beer law describes the reduction of the X-ray intensity by passing through a material. Here the linear absorption coefficient μ, which is unique for each material, is directly proportional to the atomic number Z to the power of 4.[10, 11] (B) Unstained soft tissue is mainly made of hydrogen (H), carbon (C), oxygen (O), and nitrogen atoms (N), which consist of low atomic numbers resulting in an inherent low contrast for soft tissue seen on the CT slice. (C) Stained tissue on the other hand has accumulated contrast agent that holds elements of high atomic number Z (see stained CT slice). The contrast agent used here was dibromo fluorescein 9c (see chemical structure) holding two bromine atoms (Z = 35, highlighted in apricot) and a barium atom (Z = 56, highlighted in green).

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**Scheme 1.** A) Synthesis of eosin Y salts 2a and 2c) macroscopic investigations of stained tissue. The tissue samples were pre-treated with a 4% formaldehyde solution and 0.5 mL glacial acetic acid. Staining was performed using a 31.9 mm aqueous solution of eosin Y salt 2a or 2c–2e. For each condition, an individual number of three turkey liver pieces (≈27 mm3) was stained to ensure reproducibility; one of these pieces was investigated macroscopically.
barium-eosin Y salt (2c) was achieved after incubation of the tissue for 72 h, while the tissue samples treated with 2a suffered from inhomogeneities (see the Supporting Information, chapter 8).

In order to study the contrast enhancement of the barium-eosin Y salt (2c) in comparison to the eosin Y disodium salt (2a) in more detail, CT measurements at a μCT system were performed for the tissue samples incubated for 72 h under different conditions. The obtained μCT data were analyzed and evaluated for (i) completeness of staining, (ii) presence of diffusion rings, (iii) contrast enhancement, (iv) appearance of CT artifacts as streaks, and (v) homogeneity of the staining.

The barium-eosin Y-treated samples (1 and 3), which have been acidified during fixation, offered a complete and homogeneous staining of the soft-tissue samples, as indicated by the CT slices (Figure 2A) and the constant plateau seen in the line plot (Figure 2B). This result clearly confirms the macroscopic observations (see Scheme 1B and Supporting Information, chapter 8) and shows the reliability and reproducibility of the staining method using 2c as the dye. In addition, these probes (2c, acidified) displayed a significantly higher gray value (Figure 2B) when compared to the equimolar eosin Y disodium-stained samples 2 and 6 and the non-acidified barium-eosin Y samples 5 and 7. As the gray values represent a measure for contrast enhancement (the higher the gray value the better the attenuation contrast), this refers to a significantly increased contrast of the acidified liver samples stained with 2c compared to 2a and thus demonstrates the impact of the heavy metal cation. This effect, however, was only observed for the barium-eosin Y salt 2c while other Ba salts containing low atomic number elements, such as Ba(OH)$_2$, or Ba(OAc)$_2$, failed as specific staining agents. These negative control experiments show nicely the mutual influence of the organic anion eosinate Y and the heavy-metal barium cation for cytoplasm interaction and contrast enhancement. Besides this, the variations in the line plots of the acidified eosin Y disodium samples (2 and 6) as well as the non-acidified barium-eosin Y samples (5 and 7) seen at the beginning or end of the plateau, clearly indicate the formation of a diffusion ring, indicative of an incomplete and inhomogeneous staining of the soft tissue sample.

Encouraged by the results obtained above using the barium eosin Y staining agent (2c), we further explored a possible contrast enhancement by raising the water solubility of the barium eosinates. We thus developed a directed synthesis to selectively brominate the xanthenecore in 3 to obtain the mono-, di-, and tri-brominated fluorescein derivatives 4–6 (Scheme 2). Treatment of 3–6 with NaOH or Ba(OH)$_2$ afforded after recrystallization the corresponding sodium and barium salts 2 and 7–10. To get a first hint at the hydrophilicity of these compounds, we determined the logP values of the lactones 1 and 3–6 as well as of the acid salts 2, 7, and 9 (Supporting Information, chapter 3). Here, an explicit correlation of the number of Br atoms and the partition coefficient was revealed, in particular for the barium fluoresceines 2c and 7c–10c.

Because of the significantly lowered lipophilicity of the dibromo barium fluoresceinate 9c in comparison to the eosin Y salt 2c in combination with its straightforward synthesis, the dibromo fluorescein 9c was evaluated in more detail. In a second staining experiment the influence of the halogenation degree on the fluorescein core was studied with respect to contrast enhancement using μCT. Since the first staining experiment proved already a better performance of the barium-eosin Y salt (2c) compared to the eosin Y disodium salt (2a), we decided to exclude 2a from the current study. In line with the results described for 2c before (see Figure 2), the soft-tissue samples being acidified during fixation or prior to staining with 2c and 9c, respectively, displayed a better contrast enhancement compared to the non-acidified samples (for details see the Supporting Information, chapter 10). The dibromo barium fluorescein derivative (9c) performed better than 2c when the maximum concentration of each compound in water was applied (c = 45 mg mL$^{-1}$ [c = 72.0 mM] for 9c; c = 25 mg mL$^{-1}$ [c = 31.9 mM] for 2c) offering a homogeneous staining of the
whole tissue sample (see the Supporting Information, chapter 9, Figure S3). The increased contrast for 9c can be explained by its enhanced water solubility of the dibromo compound 9c. Overall, the barium xanthene derivatives 2c and 9c showed much better contrast enhancement compared to the sodium analogues 2a and 9a at such low concentrations.

To allow for comparison with the histological results and to showcase the compatibility of the developed barium fluorescein staining protocol with standard histological methods, the cell nucleus-specific staining with the counter stain Mayr’s hematoxylin was applied to the histological microscopic slide after staining with 2c and μCT measurements. As expected, the cell cytoplasm still appeared pinkish (see the Supporting Information, chapter 12, Figure S4A) and the cell nuclei were highlighted purple in color (see the Supporting Information, chapter 12, Figure S4B). The barium dibromo fluoresceinate salt 9c, staining of the cell cytoplasm, as well as the additional Mayr’s hematoxylin counterstain (H-stain) were therefore not disturbed by each other and thus standard histological slides (see the Supporting Information, chapter 12, Figure S4A, B) were obtained following standard histological H-staining procedures. The resulting images were of high quality and suitable to undergo further histological analysis, which contributes to the practicality of our method and significantly enhances the sample availability.

After the in-depth evaluation on the ability of the barium salts 2c and 9c to act as X-ray CT tracers, we wanted to compare the staining quality of our newly developed dye 9c to the recently introduced X-ray stain 2a. Therefore, we applied the dibromo fluorescein barium salt (9c) to mouse kidney-tissue pieces (∼5 mm edge length). The mouse kidney with its different tissue types offers various biological structures at multiple scales. After the acquisition of an overview scan, a volume of interest (VOI) was chosen and measured with high resolution as a local tomography (see the Supporting Information, video). The ability to obtain high-resolution 3D data with isotropic resolutions of desired VOIs while keeping the soft-tissue sample intact is a unique feature of μCT that cannot be found among other microscopy techniques such as most light or electron microscopy techniques.

Our results are displayed in Figure 3 and confirm homogenous staining throughout the sample without diffusion arti-

![Scheme 2](Image)

**Scheme 2.** Selective bromination of fluorescein (3) and preparation of the corresponding sodium 7a–10a and barium salts 7c–10c.

![Figure 3](Image)

**Figure 3.** High-resolution μCT data of the cortex and outer medulla region of a mouse kidney after application of the dibromo fluorescein barium salt (9c). (A) Representative μCT volume of 1.20 mm × 1.20 mm × 1.20 mm (effective voxel size = 3.3 μm). (B–D) Individual μCT slices derived from the orthogonal planes through the volume shown in (A). (E) Representative X-ray barium eosin Y-stained histological microscopic slide. The cell nuclei were counter stained with Mayr’s sour hematoxylin. Legend: I proximal convoluted tubules, II: artery, III: vein, IV: capillary, V: medullary rays, and VI: loop of Henle.
facts. The perspective view of a representative volume in Figure 3A highlights the 3D arrangement of inner structures and connectivity between different tissue types. The provided contrast enabled the visualization of relevant anatomical structures within this volume of interest, the cortex and outer medulla region such as convoluted tubules, vessels, or medulla rays. Representative individual 2D µCT slices derived from the orthogonal planes of the volume in Figure 3A are displayed in Figure 3B–D. The validation of the µCT results was performed through histological analysis. The representative histological light microscopy image shown in Figure 3E was obtained from the very same dibromo fluorescein barium salt (9c) stained mouse kidney tissue piece seen in Figure 3A. Here, the obtained histological section was counterstained with the standard histological procedure according to Mayr’s sour hematoxylin to stain the cell nuclei, which were not targeted by the toplasm selective 9c. This resulted in a typical H&E-stained histological section, which corresponds very well with the µCT slice shown in Figure 3C.

Different gray values within the sample display varying concentrations of contrast agent accumulated in the soft tissue, that is, the higher the protonated protein/peptide content in the cytoplasm the more contrast agent can interact with these structures and the higher the contrast. This can be seen in the perspective view of a representative virtual volume in Figure 3A and the individual CT slices in Figure 3B–D, for example, labelled structures such as V appear very bright and III being black meaning no accumulation of contrast agent. The CT results are very well reflected in the histological slide where the structures V appear very pink indicating a high concentration of the contrast agent 9c, while II and III being white highlight the absence of contrast agent. Thus, the staining results and image quality compare very well with the respective results obtained with the eosin Y sodium salt (2a). A clear advantage is seen here in the enormous reduction of the contrast agent used to stain the biopsy samples from 300 mg·mL⁻¹ for 2a to 45 mg·mL⁻¹ for 9c. Even though the color of the dibromo fluorescein barium salt (9c) shifted to orange-pink when compared to the eosin Y disodium salt (2a), the pathologists observed no problems to perform their histological analysis. Thus, the new X-ray stain dibromo fluorescein barium salt (9c) proved superior and is thus suitable for µCT even at low concentrations.

To conclude, the improved solubility of 9c in water was crucial to obtain this new X-ray staining agent that allows to non-destructively and selectively visualize the cell cytoplasm of biological and medical soft-tissue samples in three dimensions. The application of the staining protocol to turkey liver and mouse kidney tissue pieces underlines the reliability of the protocol and emphasizes the use for different tissue types. The ability to counterstain the biopsy samples using standard histological methods paves the way for establishing a convenient 3D X-ray histology approach as a complementary tool for future histological analysis. This will offer access to additional information and support histologists where 2D imaging is facing its boundaries and thus meeting the demands to provide answers to advanced medical questions, which will benefit from targeted staining of specific biological structures as well as non-destructive 3D imaging techniques.

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Conflict of interest

The authors declare no conflict of interest.

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