Excitation-resolved cone-beam x-ray luminescence tomography

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Abstract. Cone-beam x-ray luminescence computed tomography (CB-XLCT), as an emerging imaging technique, plays an important role in in vivo small animal imaging studies. However, CB-XLCT suffers from low-spatial resolution due to the ill-posed nature of reconstruction. We improve the imaging performance of CB-XLCT by using a multiband excitation-resolved imaging scheme combined with principal component analysis. To evaluate the performance of the proposed method, the physical phantom experiment is performed with a custom-made XLCT/XCT imaging system. The experimental results validate the feasibility of the method, where two adjacent nanophosphors (with an edge-to-edge distance of 2.4 mm) can be located.

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X-ray luminescence computed tomography (XLCT) has been proposed as a new molecular imaging modality.1 In principle, when irradiated with x-rays, the x-ray-excitable nanophosphors (or other similar materials) emit light (luminescence) when they are too close to each other (see Fig. 3). In this work, we propose a new method to improve the imaging resolution of CB-XLCT, which is achieved by using the excitation-resolved imaging scheme combined with the principal component analysis (PCA) method. Note that in the proposed excitation-resolved imaging scheme, the nanophosphors within an imaged object are irradiated by multiband x-ray voltages. The imaging scheme is different from the conventional XLCT imaging mode where nanophosphors are only irradiated by a defined x-ray tube voltage.1,5-8 Considering that nanophosphors have a variety of excitation efficiency versus x-ray tube voltage,2,9 in the dimension of energy spectrum, the regions containing different nanophosphors will show different excitation behaviors. As a result, by using a multivariate image analysis method, it is possible to resolve and identify these regions even if they are close to each other.9,10 In this letter, PCA, as a common multivariate image analysis method, is used to resolve the adjacent nanophosphors, which is achieved by using XLCT tomographic images from the multiband x-ray voltages as the input data.

In detail, in x-ray luminescence imaging, when x-rays are transported to nanophosphors in biological tissues, the nanophosphors will emit visible or near-infrared light, $S(r)$. Based on Lambert–Beer law, $S(r)$ can be expressed as follows:

$$S(r) = I(r)\xi \rho(r),$$  \hspace{1cm} (1)

$$I(r) = I_0 \exp \left[-\int_{r_0}^{r} \mu_a(t)dt\right],$$  \hspace{1cm} (2)

where $\rho$ is the nanophosphor density, $\xi$ is the light yield, $I_0(r)$ is the x-ray source intensity at the initial position $r_0$, and $\mu_a$ is the x-ray attenuation coefficient. Furthermore, the emitted light will propagate in biological tissues, which can be modeled by the following diffusion equation:11

$$-\nabla \cdot [D(r)\nabla \Phi(r)] + \mu_a(r)\Phi(r) = S(r),$$  \hspace{1cm} (3)

where $\Phi$ is the photon density, $\mu_a$ is the absorption coefficient, and $D$ is the diffusion coefficient.

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Based on the finite-element method (FEM), the above imaging model can be converted into the following linear relationship:

\[ \Phi_{\text{meas}} = WP, \]

(4)

where \( W \) is the weighted matrix, used to map the unknown nanophosphor distribution \( \rho \) to the known measurements \( \Phi_{\text{meas}} \). In this work, the unknown \( \rho \) is obtained by solving Eq. (4) using algebraic reconstruction technique (ART) with non-negative constraints.

To clearly resolve the adjacent nanophosphors within imaged object, in this work, PCA transformation is used. In detail, here, the input data of PCA are \( N \) frames 3-D XLCT tomographic images, where \( N \) is the number of the used multiband x-ray voltages. If the reconstructed image has \( M \) pixels, then we can represent one frame data as an \( M \) column vector. By transforming all frames, the input data are arranged as a \( M \times N \) matrix \( \mathbf{P} \).

Referring to our previous studies Ref. 10, in this work, the principal component (PC) of the input data can be obtained by solving the following equation:

\[ \mathbf{Z} = \tilde{\mathbf{P}} \times \mathbf{R}, \]

(5)

where \( \mathbf{R} \) is the eigenvector matrix and the column of \( \mathbf{R} \) is eigenvector of the matrix \( \mathbf{R} = [1/(M-1)]\mathbf{P}^T \mathbf{P} \) with \( \mathbf{P} \) being a dataset obtained from the original data \( \mathbf{P} \) by subtracting its mean value. \( \mathbf{Z} \) is a PC matrix and the first column of \( \mathbf{Z} \) is the PC1, the next column is the PC2, and so on. Further, the absolute values of positive and negative elements of the PCs are, respectively, utilized as a weight factor for creating images, termed as PC-XLCT image, which can be used to illustrate the distributions of nanophosphors within imaged object.

The physical phantom experiment was performed with a custom-made XLCT/XCT imaging system, similar to that described in Liu et al. As shown in Fig. 1, in the hybrid system, the conebeam XLCT system was placed vertically on the optical bench, while the micro-XCT system was placed horizontally. These two subsystems used a communal rotational stage. For the XLCT imaging, the x-ray source used was a microfocus x-ray source with a maximal power of 80 W (Oxford Instrument, United Kingdom). The luminescent photons emitted from nanophosphors were recorded by an electron-multiplying CCD (EMCCD) camera (iXon DU-897, Andor, United Kingdom) coupled with a Nikkor 50-mm f/1.8D lens (Nikon, Melville, New York). To protect the EMCCD chip from x-ray irradiation, a 4-mm depth of lead shield was used. The micro-XCT imaging was performed by the aforementioned x-ray tube combined with a CMOS x-ray flat-panel detector (2923, Dextela, United Kingdom).

Figure 2(a) shows the schematic diagram of the phantom geometry, where a transparent glass cylinder (outer diameter 3.0 cm) was used as the phantom. The phantom was filled with 1% intralipid and water with \( \mu_s = 0.02 \text{ cm}^{-1} \) and \( \mu_t = 10 \text{ cm}^{-1} \). Two transparent glass tubes (outer diameter 4.2 mm) filled with the red nanophosphor \( \text{Y}_2\text{O}_3: \text{Eu}^{3+} \) and the blue nanophosphor \( \text{BaMgAl}_{16}\text{O}_{27}: \text{Eu}^{2+} \) were used as the x-ray luminescent targets. To evaluate the imaging performance of CB-XLCT, the two tubes were separated with an edge-to-edge distance of about 4.2 mm along X-axis. In this work, the nanophosphors \( \text{Y}_2\text{O}_3: \text{Eu}^{3+} \) and \( \text{BaMgAl}_{16}\text{O}_{27}: \text{Eu}^{2+} \) were purchased from Jiangxi Illuma Fluorescent Materials Co., Ltd., China.

To implement the proposed excitation-resolved imaging scheme, in XLCT imaging processes, the nanophosphors were irradiated by multiband x-ray source voltages \((55, 60, 65, 70, \text{ and } 75 \text{ kV})\). Note that the x-ray source current was fixed at 1 mA in all the aforementioned cases. For each specific voltage, 360 deg x-ray luminescence tomography imaging was performed using 24 projections in a 15 deg step. When collecting these luminescence projections, the integrating time of EMCCD was set to 3 s, the EM gain was set to 260, and the EMCCD binning was set to \( 1 \times 1 \).

Figure 3 shows the reconstructed XLCT tomographic images, which are obtained by using different x-ray voltages as excitation. For XLCT reconstruction, the imaged object (phantom) was discretized into 2787 nodes and 11,476 tetrahedral elements. The reconstruction region was a 3.0 cm \( \times \) 3.0 cm \( \times \) 5.5 cm 3-D region. The experimental results indicate that it is difficult to resolve the distributions of nanophosphors in the two adjacent tubes based on these reconstructed tomographic images. This is mainly because of the diffusive nature of photon migration in tissues.

Figure 4 shows the generated PC-XLCT images. As expected, when PCA was applied to the above reconstructed XLCT images (see Fig. 3), we generated the corresponding PC-XLCT images which could be used to illustrate the distributions of nanophosphors within the imaged object even if they were close to each other. As shown in Fig. 4(a), the distribution of nanophosphors \( \text{Y}_2\text{O}_3: \text{Eu}^{3+} \) in the phantom is illustrated by the positive PC2-XLCT image. Similarly, the distribution of nanophosphors \( \text{BaMgAl}_{16}\text{O}_{27}: \text{Eu}^{2+} \) in the phantom is illustrated by the negative PC2-XLCT image [see Fig. 4(b)]. Furthermore,
Figs. 4(d) and 4(e) show the 3-D visualization results of the positive and the negative PC2-XLCT images using different views, respectively. In conclusion, by using a multiband excitation-resolved imaging scheme combined with PCA method, we resolve two nanophosphors with edge-to-edge distance 2.4 mm. This method has potential for improving the imaging performance of CB-XLCT. However, when using the multiband excitation-resolved imaging scheme, the nanophosphors must be irradiated by different x-ray voltages in the imaging processing. As a result, it increases the data acquisition time. In addition, the generated PC images only reflect the structural information of the nanophosphors. To obtain the concentration of nanophosphors, independent component analysis can be considered as an effective method. Further, we must also point out that the x-ray energy information may also be used in the XLCT reconstruction, which is helpful for improving the imaging performance. Finally, more complex phantom experiments (e.g., resolving the nanophosphors of the same color with different or same concentrations) and in vivo experiments should also be considered, which will extend the application of the proposed method. Systematic studies will be investigated in our future work.

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References