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TITLE: High Resolution X-ray Phase Contrast Imaging with Acoustic Tissue-Selective Contrast Enhancement

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14. ABSTRACT We show that laser-driven thermal modification can be used to selectively enhance regions of a flowing carbon suspension. The preliminary proof-of-principle experiments illustrate the ability to measure thermal modifications in a time-resolved manner utilizing x-ray phase-contrast imaging paving the way for implementation in a biological model. Additional results of biological tissue sample measurements are presented. X-ray phase-contrast images of murine hepatic and pulmonary samples reveal fine structure usually visible only by utilizing histological or microscopy techniques. The method holds promise for future applications to study murine models of pulmonary and hepatic disease.					
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Introduction

Thermally Modulated X-ray Phase Contrast Imaging

The experimental setup for modifying a phase contrast x-ray image by using laser-driven thermal gradients is illustrated in Figure 1. The output of a 527nm Nd:YLF laser is focused onto a liquid water jet of carbon suspension and imaged using a microfocus x-ray source coupled in-line with a synchronously gated intensified optically coupled CCD camera.

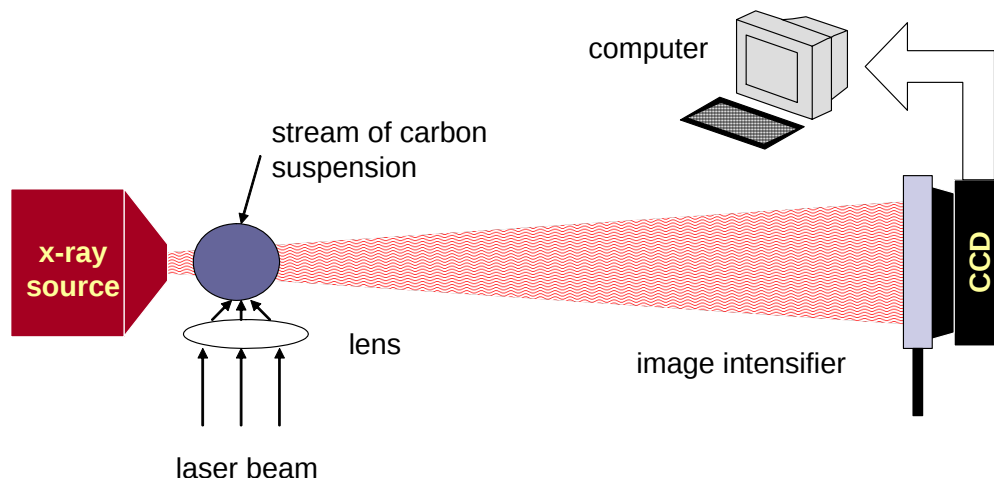


Figure 1 : Diagram of the experimental apparatus. X-radiation generated by a microfocus tube penetrates a sample irradiated by a the focused output of 527nm Nd:YAG laser. A gated intensified optically coupled CCD camera detects the transmitted x-rays and is read by the computer that stores the images.

In principle, the sensitivity of x-ray phase-contrast imaging allows the visualization of weak phase effects imparted to x-ray photons as they traverse a thermally modulated sample. Biologically, these experiments hold promise to take advantage of the rather large optical contrast of blood in the infrared to modulate x-ray images. By using a liquid jet of carbon suspension we experimentally investigate the basic principles of laser modulated x-ray phase-contrast imaging.

In addition to image modulation, the need to develop biological sample preparation methods designed to take advantage the proliferation of high-resolution x-ray sources. The goal of the current research is to explore the uses of phase-contrast x-ray imaging for investigating both static and laser-modulated biological samples.

Body

The right-hand image in Figure 2 shows the results of a set of experiments involving the interaction of a focused laser with a flowing carbon suspension in water. The figure on the left illustrates an x-ray image of the jet without laser modification. The features labeled a, b and c in Figure 2 shows the modification of the jet induced by the laser-driven thermal heating. Each of the three features arises from thermal jet expansion

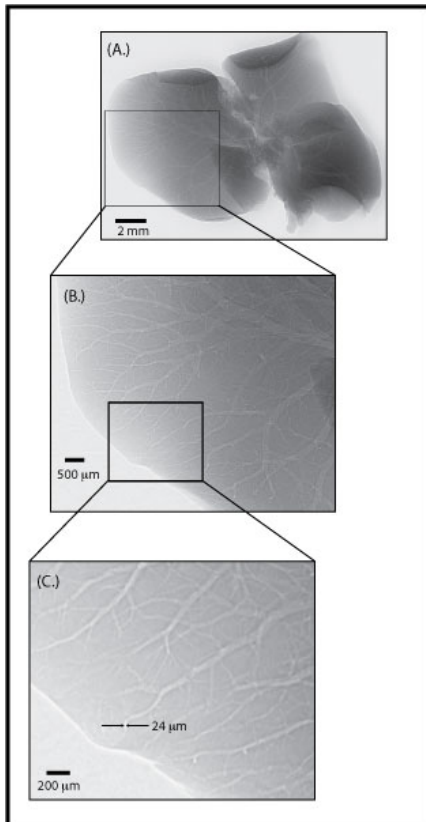


Figure 3. X-ray phase contrast image of a formaldehyde-fixed and dried mouse liver sample. Microvessels are resolved to their finest levels.

Given the inherent advantages of having air and soft-tissue interfaces for x-ray phase-contrast imaging, murine pulmonary samples were also imaged. Figure 3 displays a typical result of an excised rat lung, which has been clamped and partially inflated with air. Alveolar features down to $10\ \mu\text{m}$ are visible. The speckled pattern is indicative of air and soft-tissue interfaces expected with the inflation of the alveolar sacs. Also visible are darker web-like features, attributed to pulmonary venous vasculature. The ability to image pulmonary samples with this type of detail naturally lends itself to the study of pulmonary pathologies. For example, utilizing the imaging technique to image murine models of pulmonary fibrosis, emphysema, and cancer could provide valuable access to fine structure usually attained only by histological methods.

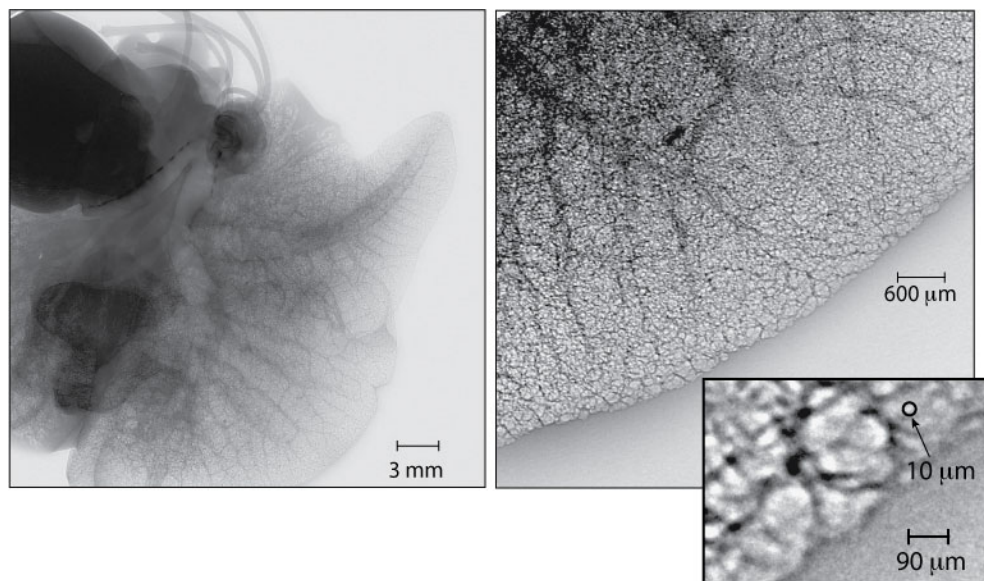


Figure 4 Three magnified views of a phase contrast image of an air-filled excised rat lung.

Reportable Outcomes

“High-resolution angiography: Computed tomography coupled X-ray phase contrast imaging of biological tissue samples” with C. M. Laperle, T. J. Hamilton, P. Wintermeyer, D. Shi, M. Anastasio, C. Rose-Petruck, G. and J. R. Wands *in preparation*

C. M. Laperle, G. Cao, T. J. Hamilton, C. Rose-Petruck, and G. J. Diebold, “Photothermal Modification of X-ray Phase Contrast Images”, *Progress in Biomedical Optics and Imaging, Photons Plus Ultrasound: Imaging and Sensing 2007*, 8 64371N, (2007).

“X-ray Phase Contrast Imaging: Transmission Functions Separable in Cartesian Coordinates”, with Guohua Cao, Theron Hamilton, Christopher Laperle and Christoph Rose-Petruck *submitted to Journal of the Optical Society of America A*.

Conclusions

We have proven the concept of laser-driven thermally modified x-ray phase contrast imaging. We have shown the method has the ability to visualize thermally modified regions of objects. Furthermore, we have shown the sensitivity of phase-contrast x-ray imaging on excised hepatic and pulmonary murine tissue samples.

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