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Nonlinear coherent four-wave-mixing in optical microscopy

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08.30-10.30 CWC - Lasers in Medicine I Presider: T.G. Papazoglou, F.O.R.T.H. - I.E.S.L., Crete, GREECE

08.30 CWC1

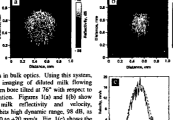
METHOD FOR REAL TIME COLOUR DOPPLER OPTICAL COHERENCE TOMOGRAPHY

A. V. Zvyagin, J. B. FitzGerald, K. K. M. B. Silva, and D. D. Sampson Optical + Biomedical Engineering Laboratory. Department of Electrical and Electronic Engineering, The University of Western Australia, Nedlands W. A. 6907. Tcl.: +61-8-9380-1522, Fax. +61-8-9380-1065

Tel: +61-8-9380-1522, Fax.:+61-8-9380-1065 Colour Doppler optical coherence tomography (CDOCT) has recently shown great promise in two-dimensional, high-spatial-resolution (on the order of tens of microns) tomographic velocity mapping of blood in living tissue.¹ CDOCT is based on optical interference in a scanning Michelson interferometer under broadband illumination. When a scattering object is in motion, it generates a signal in the detection circuit, an interferogram, the envelope of which gives the spatial reflectivity, and the frequency of which is proportional to the sum of the object's velocity and that of the reference are scanner. Conventionally, post-processing of the entire interferogram is performed, requiring a large amount of computation, which procludes the real time operation of CDOCT necessary to provide micro-artefact-free images in vice. Another major problem with the conventional approach concerns the bandwidth of the detection electronics.² For high dynamic range (sensitivity), the detection bandwidth bould be as narrow as possible. However, the need to measure rapid and variable flows present in the vast majority of blood vessels requires a wider detection bandwidth to accommodate the variable Doppler frequency. Thus, either the dynamic range or the velocity range must be compromised.

present in the two series of the second seco

interferogram envelope (ref detection bandwidth. Furthermore, for a given sample period, only the values of the frequency and reflectivity must be stored on computer, representing a massive reduction in stored data compared to the conventional approach. In order to test our order to test our ection scheme, we detection



detection scheme, we performed ross-sectional imaging of diluted milk flowing through a glass pipe of 0.58 mm bore tilted at 76° with respect to the incident near-infrared radiation. Figure 21(a) and 1(b) with respect to the incident near-infrared radiation. Figure 21(a) and 1(b) with respect to the system schliss high dynamic range, 98 diluted wilk low we las large velocity range, -20 to +20 mm/s. Fig. 1(c) shows the velocity profile across the sample's centroid (solid line). The good fit to a parabola (dashed line) is velocited that the velocity map correctly reproduces the laminar flow of the milk solution.

0.2

The application of a PLL detection technique to CDOCT provides the best prospects reported to date for real-time, *in vivo* tomographic imaging of blood flow velocity and, simultaneously, tissue morphology in the absence of motion artefact.

¹Z. Chen et al., "Noninvasive imaging of *in vivo* blood flow velocity using optical Doppler tomography", Opt. Lett. 22, 1119 (1997)

²M. Kulkami et al., "Velocity-estimation accuracy and frame-rate limitations in color Doppler optical cohe tomography", Opt. Lett. 23, 1057 (1998).

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Nonlinear coherent four-wave-mixing in optical microscopy

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The use of nonlinear optical techniques in microscopy has intensified the versatility of optical The use of nonlinear optical examples at microscopir ans meaning the versatility of optical methods to explore biological samples at microscopir resolution. In addition to multiphoton excited fluorescence, coherent four-wave-mixing (FWM) optical methods like third harmonic generation (THG)⁴ and coherent anti-Stokes Raman scattering (CARS)², have been adopted in microscope configurations. The application of coherent FWM techniques with high spatial resolution has paved the way for the possible implementation of a broad spectrum of

resolution has paved the way for the possible implementation of a broad spectrum of spectroscopic tools in microscopy. Opposed to multi-photon excited fluorescence, which is an incoherent process, FWM methods yield coherent signals. It is therefore expected that the spatial distribution of coherent emission near the focal volume may significantly differ from its incoherent counterpart. Consequently, in order to define resolution criteria for FWM microscopes, insight into the spatial organization of the coherent emission field is desirable. In this contribution we present a detailed theoretical analysis of the imaging properties of coherent nonlinear microscopes. We have developed a model that allows calculation of the generation and propagation of coherent signals under high numerical aperture (NA) conditions without invoking the slowly varying envelope approximation. Based on calculations of coherent anti-Stokes Raman scattering (CARS) signals it is shown that diffraction effects play a prominent role in the spatial distribution of the coherent signal networks of a point spread function (PSF) and the object but is shaped by the complex interplay of object size and coherent build-up dynamics.

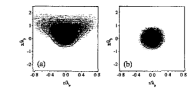


Fig .1. Intensity plot of CARS emission field near the focal volume of a NA = 0.9 air objective (a) and the corresponding illumination volume (b)

Y. Barad, H. Eisenberg, M. Horowitz and Y. Silberberg, Appl. Phys Lett. 70 (1997) 922
A. Zumbusch, G.R. Holtom and X.S. Xie, Phys. Rev. Lett 82 (1999) 4142