Compressed single pixel imaging in the spatial frequency domain

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Abstract. We have developed compressed sensing single pixel spatial frequency domain imaging (cs-SFDI) to characterize tissue optical properties over a wide field of view (35 mm x 35 mm) using multiple near-infrared (NIR) wavelengths simultaneously. Our approach takes advantage of the relatively sparse spatial content required for mapping tissue optical properties at length scales comparable to the transport scattering length in tissue (l_tr ~ 1 mm) and the high bandwidth available for spectral encoding using a single-element detector. cs-SFDI recovered absorption (μ_a) and reduced scattering (μ_s^r) coefficients of a tissue phantom at three NIR wavelengths (660, 850, and 940 nm) within 7.6% and 4.3% of absolute values determined using camera-based SFDI, respectively. These results suggest that cs-SFDI can be developed as a multi- and hyperspectral imaging modality for quantitative, dynamic imaging of tissue optical and physiological properties. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.22.3.030501]

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1 Introduction

Spatial frequency domain imaging (SFDI) is a noncontact wide-field imaging technique that uses sinusoidal patterns of intensity-modulated light to characterize multiply scattering media, such as biological tissue. SFDI separates the contributions of light scattering from absorption by measuring the frequency-dependent modulation transfer function of diffusively reflected structured light. This information is used to map and form images of tissue optical parameters μ'_a and μ_s, respectively. Optical properties at multiple wavelengths are used to derive quantitative images of biochemical composition, such as the tissue concentration of oxy- and deoxyhemoglobin and water. SFDI techniques have been utilized widely to characterize tissue composition and function in humans, animal models, and ex vivo tissue specimens.

There is generally a tradeoff between spectral and spatial information content in SFDI instruments. This is due to the fact that wide-field imaging is typically performed serially at several discrete wavelengths using selective illumination and/or optical filters. Spatial frequency domain spectroscopy (SFDS) was introduced to enhance spectral content using a broadband light source, fiber-based collection, and a spectrometer. However, this approach has a field-of-view (FOV) that is limited to the collection fiber (~2 mm). Hyperspectral imaging cameras can expand the FOV; however, these systems are generally relatively expensive and can have limitations in speed, dynamic range, and sensitivity. In this letter, we describe an alternative approach that incorporates compressed sensing (CS) and a single pixel detector for wide-field multispectral imaging. This strategy is compatible with both broadband light sources imaged with fiber-coupled spectrometers and single element detectors using several temporally encoded wavelengths/sources.

Imaging applications of CS were first introduced in a “single pixel camera” form factor that transferred workload from hardware to postacquisition computation. CS addresses image reconstruction by relying on random samples from a scene under view instead of capturing individual pixels. This approach can be particularly effective in near-infrared (NIR) and SWIR spectroscopic imaging by replacing expensive cameras with a single photodetector. In addition, resolution requirements for SFDI mapping of μ'_a and μ_s are determined by the relatively low frequency of variations in the tissue transport scattering length (l_tr ~ 1 mm). This allows typical SFDI sinusoidal projection patterns to sample tissue sparsely at f ≈ 1/(3l_tr). Since CS methods utilize principles of sparsity, the number of random samples imposed by Nyquist Shannon boundary can be reduced and CS is well-matched with diffuse optical imaging and tomography.

2 Materials and Methods

2.1 Spatial Frequency Domain Imaging

The SFDI instrumentation, data acquisition, and postprocessing have been previously enumerated elsewhere. Briefly, spatially modulated light is projected onto a sample via a digital light processing (DLP) unit. At each spatial frequency, three evenly spaced modulation phases of 0 deg, 120 deg, and 240 deg are cycled through. The diffusively reflected light is then captured with a camera. The phase-offset images are next fed in a three-phase demodulation formula, Eq. (1), to extract spatial frequency-dependent reflectance. The demodulated reflectance is also calibrated to account for instrument function. As it was shown in Ref. 1, the reflectance sensitivity to optical absorption and scattering varies as a function of spatial frequency. This can be modeled with diffusion theory or Monte Carlo simulations. Using a minimum of two spatial frequencies, the calibrated reflectance at every pixel is then applied to the simulations resulting in μ'_a and μ_s maps. Once the μ'_a and μ_s are decoupled at several wavelengths, μ_s is used to calculate hemodynamic parameters by assuming that oxy/deoxyhemoglobin and water are the dominant absorbing chromophores in the medium.
\[ AC(x, y) = \frac{2^{1/2}}{3} \left[ \left| I_{0 \text{ deg}}(x, y) - I_{120 \text{ deg}}(x, y) \right|^2 \\
+ \left| I_{120 \text{ deg}}(x, y) - I_{240 \text{ deg}}(x, y) \right|^2 \\
+ \left| I_{240 \text{ deg}}(x, y) - I_{0 \text{ deg}}(x, y) \right|^2 \right]^{1/2}. \] (1)

### 2.2 Compressed Sensing Framework

Nyquist–Shannon is a fundamental sampling theorem in digital signal processing that states that a signal can exactly be recovered if the sampling rate is at least twice its highest frequency. Considering a two-dimensional array with \( N \) pixels on a digital camera sensor, the raw image can be considered as vector \( \mathbf{x} \), where \( \mathbf{x}[i], i = 1, 2, \ldots, N \). This vector can be expanded as a combination of its orthonormal basis such as discrete cosine transform, wavelet, etc.,

\[ x = \sum_{i=1}^{N} \Psi_i C_i, \] (2)

where \( C_{i=1:N} \) and \( \Psi_{i=(1:N)\times(1:N)} \) are the coefficient vector and transform operator, respectively. In the case of \( k \)-sparse expansion of \( \mathbf{x} \) vector, only \( k \) number of \( C_i \) coefficients are nonzero. The CS technique\(^{15}\) suggests using only \( M \ll N \) (\( M \approx k \)) samples of the signal \( x \) with sensing matrix \( \Phi_{i=(1:M)\times(1:N)} \) which gives the measurement vector

\[ y = \Phi \mathbf{x}. \] (3)

This rewrites Eq. (2) as

\[ y = \Phi \Psi \mathbf{C}. \] (4)

The sensing matrix \( \Phi \) is independent of the raw image, \( \mathbf{x} \). As part of the CS algorithm, a convex optimization problem is solved using \( I_1 \)-minimization to recover image \( \mathbf{x} \) based on measurement vector \( y \),

\[ \hat{C} = \text{arg min} \| C_1 \|_1 \text{ in a way that } \Phi \Psi \mathbf{C} = y. \] (5)

The raw image can then be reconstructed with \( \hat{\mathbf{x}} = \Psi \hat{\mathbf{C}} \). In this paper, we have utilized the denoising-based AMP (D-AMP) CS algorithm and source code developed in Ref. 17 on the postprocessing side.

### 2.3 cs-SFDI

On the detection side of our cs-SFDI instrument, a sample, e.g., tissue phantom, is imaged via a collection lens onto a digital micromirror device (DMD) (CEL5500, Digital Light Innovation, Austin, Texas) with \( N \) mirror bins. The sensing matrix \( \Phi \) is also uploaded to the DMD memory as a stack of random patterns from a Bernoulli distribution.\(^{18}\) This binary sensing matrix matches well with the ON/OFF orientation of the DMD. The reflected light from the DMD is then focused on the photodetector (DET100A, Thorlabs, Newton, New Jersey) and finally builds the measurement vector \( y \). The CS algorithm allows for reducing the number of sampling patterns stored on a DMD to effectively reconstruct an image. This approach reduces both the duration of data acquisition and the cost of the DMD. Figure 1 shows a schematic of our cs-SFDI setup. On the excitation side, the output of three coaligned LEDs (M660L4, M850L3, and M940L3, Thorlabs, Newton, New Jersey) is condensed onto a sinusoidal-patterned mask (Applied Image Inc., Rochester, New York), and then the mask is imaged onto the tissue simulating phantom. The LEDs are simultaneously intensity-modulated in the frequency range of 3 to 5 kHz, which is higher than the sampler DMD’s refresh rate. The reflected power of the frequency-encoded sources is determined for each random pattern via temporal frequency domain analysis. This feature is important because the low dynamic range of the photodetector enables adding more frequency-encoded light sources to the setup without affecting the total acquisition speed.

A LABVIEW platform (National Instruments, Austin, Texas) controls the linear stage (T-LS28M, Zaber Technologies Inc, Vancouver, BC), the sampler DMD, and the photodiode, thus accelerating the data acquisition. In this configuration, we used a sinusoidal mask on a linear stage instead of a second DMD to project structured light. This was due to the fact that each of the DMD array elements flickers on and off to generate grayscale intensities. In a camera-based SFDI system, the camera’s exposure time can be adjusted to match the pulse width modulation (PWM) on the DMD. However, this can be problematic in a single pixel imager because of the photodiode’s high bandwidth and lack of access to the PWM sequence of every single micromirror element. An additional advantage to transmission-based masks for SFDI is their low-cost and simplicity.

### 3 Results

We constructed tissue phantoms to investigate the ability of the cs-SFDI instrument to estimate bulk optical properties.\(^{19}\) The “sample” phantom consists of two regions: a central lesion, which is made of Naphthal Green B dye as the main absorber, and TiO\(_2\) as the main scatterer both dispersed in a cured polydimethylsiloxane (PDMS) base. Naphthal Green is an NIR dye with an absorption peak at 714 nm. The surrounding area of the lesion is made of India Ink as the main absorber, TiO\(_2\), and PDMS.\(^{19}\) The central lesion has a lower concentration of scattering agent than the surrounding region. Figure 2 shows the work flow of the SFDI technique based on the images...
reconstructed with the cs-SFDI instrument. In Fig. 2(a), the reflectance from phase-shifted sinusoidal projections is first captured by the compressed single pixel camera. These raw images are then demodulated to AC and DC components, Fig. 2(b), calibrated against another tissue phantom with known optical properties, Fig. 2(c), and finally interpolated pixel-by-pixel to generate the $\mu_s^\prime$ and $\mu_a$ maps, Fig. 2(d).

We compared cs-SFDI derived optical properties to those obtained with a conventional camera-based SFDI measurement. In the latter case, we replaced the cs-SFDI photodiode with a CCD camera (Flea3 1394B, Point Grey, Richmond, BC) and set all of the DLP mirrors to ON status. The three LEDs were serially controlled for each sinusoidal pattern projection to achieve spectral and spatial content. Figure 3(a) shows good agreement of the extracted bulk optical properties with the single pixel detector and the CCD camera on two contrasting regions of the phantom, the central lesion and the surrounding area. The average percent difference between these two imaging configurations were 7.6% and 4.3% for $\mu_s^\prime$ and $\mu_a$, respectively. Each point in Fig. 3(a) corresponds to a mean optical property from a $10 \times 10$ mm region of interest (ROI) on a reconstructed map at a specific wavelength. The error bars are calculated from the standard deviation of the values on the same ROIs. According to Fig. 3(b), the absorption coefficient of the central lesion follows the absorption spectra trend of Naphthol Green B while it remains almost unchanged on the surrounding PDMS background. The reduced scattering coefficients also follow the expected spatial contrast and power law dependence.

One of our primary goals in SFDI instrument development is to increase the data acquisition rate, which mainly relies on three parameters: image resolution, number of phase shifted patterns, and number of wavelengths. From the perspective of the single pixel camera, higher image resolution requires increasing the number of sampling patterns on the DMD to achieve a certain accuracy of reconstruction. We have chosen our optimum image resolution based on the transport scattering length in tissue ($l_{tr}$), which is typically $\sim 1$ mm in the NIR. Thus, for $64 \times 64$ pixels in a $35 \times 35$ mm FOV, the pixel size is about 0.55 mm, which is adequate and comparable to $l_{tr}$. A sampling rate of about 10%, i.e., 400 samples, is adequate for a tissue phantom illuminated with sinusoidal patterns at a spatial frequency of 0.1 mm$^{-1}$. This sampling rate results in a maximum acquisition rate of 12.5 frames per second (FPS) on our DMD. However, other DMD chips, such as DLP7000 (Texas Instruments, Dallas, Texas) provide refresh rates of 32,550 Hz and acquisition rates of up to 80 FPS. Because cs-SFDI uses high-bandwidth single pixel detectors, spectral content can be increased using frequency encoding without reductions in image

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**Fig. 2** (a) Raw reflectances from the tissue phantom reconstructed with the single pixel detector. (b) Demodulated AC and DC reflectance, (c) calibrated demodulated reflectance, and (d) absorption and reduced scattering coefficient maps at three wavelengths.

**Fig. 3** Bulk optical property measurements. (a) Comparison of bulk $\mu_a$ (top left) and $\mu_s^\prime$ (top right) using standard camera-based SFDI and cs-SFDI calculated at multiple wavelengths at two distinct regions of a tissue phantom: background and lesion (tagged in a white light image of the phantom). (b) Spectral and spatial contrast in phantom’s bulk $\mu_a$ (bottom left) and $\mu_s^\prime$ (bottom right) extracted using the cs-SFDI instrument.
acquisition speed. In the current cs-SFDI configuration, the linear translational stage used for shifting the sinusoidal phase pattern is a limiting factor in achieving higher frame rates and switching between multiple spatial frequencies. As a result, it takes 5 s in total to switch among three phase-shifted frames and capture images. Recently, signal processing techniques in the SFD have been developed to reduce the number of phases required to achieve the demodulated reflectance and achieve “single snapshot” performance. These methods, however, require a high number of cycles in the modulation pattern to avoid ringing artifacts. This imposes faster sampling rates to preserve higher frequency content in the image, thus increasing acquisition time. Therefore, further studies are required to evaluate the fidelity and performance of the cs-SFDI system when using single snapshot demodulation techniques.

4 Conclusion
In conclusion, we have developed a single-pixel CS method for quantitative imaging of optical properties in the spatial frequency domain. A CS algorithm was used to decrease the size of the sensing matrix required to accurately reconstruct and map optical properties. Performance was validated using tissue-simulating phantoms and demonstrating agreement in absorption and reduced scattering coefficients between cs-SFDI and a conventional camera-based system. When combined with temporal encoding of multiple sources, cs-SFDI takes advantage of the high-bandwidth of single pixel detectors and provides a fixed frame rate strategy for simultaneous multispectral imaging. Future work will emphasize improving imaging speed and spectral content by incorporating single-snapshot demodulation strategies, broadband light-labeling, and single-pixel, fiber-coupled spectrometer detection. These results provide a new framework for broadband multi- and hyperspectral functional tissue imaging with important implications for accurately and dynamically characterizing tissue composition, structure, and metabolism.

Disclosures
No conflicts of interest, financial or otherwise, are declared by the authors.

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