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Mueller matrix polarimetry for improved liver fibrosis diagnosis

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An experimental Mueller matrix polarimeter is used to quantify human liver fibrosis by measuring retardance and depolarization of thin biopsies. The former parameter is sensitive to fibrillar collagen, the latter is specifically sensitive to fibrillar collagen around blood vessels, which is not significant for liver fibrosis diagnosis. By using depolarization like a filter, retardance distribution enables to distinguish between disease stages and limit the high degree of observer discrepancy. © 2012 Optical Society of America
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Human liver fibrosis is characterized by a modification of hepatic tissue with the deposition of fibrillar collagen (type I and III) in excess [1]. The assessment of this disease is mainly established by practiced pathologists from liver biopsies whose extra-cellular matrix (including fibrillar collagen) is stained with a specific dye, usually Sirius red. A five-stage grading system (F0-F4) was thus developed based on a Fibrosis-Metavir scoring system [2]. Grade F0 corresponds to healthy livers, grade F4 to the most aggressive stage (cirrhosis). However, it is currently difficult to obtain highly reproducible results from scores due to the high degree of intra- and inter-observer discrepancy [3]. An effort into quantifying automatically the liver disease has been made but fibrillar collagen that is also abundant around blood vessels in healthy tissues, can distort diagnosis. Thus, image processing methods are necessary to mask fibrotic areas but these processes are complex, generally semi-automatic and time consuming [4]. This Letter demonstrates the potentialities of Mueller polarimetry to first discriminate collagen around vessels from the one associated to fibrosis thanks to depolarizing properties, and then quantify the relevant fibrillar collagen by measuring retardance properties. This study relies on stained samples with Sirius red as it is known that stained fibrillar collagen enhances its natural birefringence [5,6]. A subsequent statistical method is developed to quantify fibrillar collagen with a smaller incidence of human subjectivity and variations in staining.

Mueller matrix (MM) polarimetry is used to obtain the full polarimetric response of samples (depolarization, birefringence and dichroism). The studies have been carried out with a Snapshot Mueller Matrix Polarimeter (SMMP) developed in our lab [7]. Its principle is to encode polarization states in the spectral domain by means of a broadband source and high-order retarders. Thanks to a specific retarder-thickness configuration, the full Mueller matrix of a sample is available in a single spectrum, \( I(\lambda) \), measured with a dispersive detection system (spectrometer 1200 grooves/mm).

Fig. 1. Diagram of the Snapshot Mueller Matrix Polarimeter setup. O1, O2: 10x objectives, SM: scattering medium, L1, L2: lenses, OF: optical fiber, other abbreviations defined in text.

The SMMP in Fig. 1 is composed of a 15-nm-broadband source (SLD from B&W Tek, Inc) emitting around \( \lambda_0 = 830 \, \text{nm} \), a linear polarizer (P1) oriented at 0°, two calcite retarders \((\text{Ret}_1, \text{Ret}_2)\) of thickness \( e = 2.08 \, \text{mm} \) respectively oriented at 45° and 0°, two calcite retarders \((\text{Ret}_3, \text{Ret}_4)\) of thickness \( 5e = 10.4 \, \text{mm} \) respectively oriented at 0° and 45°, a linear polarizer (P2) oriented at 90°, and a spectrometer. The sample under study is imaged on a scattering medium (Bfi Optilas light diffuser, 10° FWHW) by L1 in order to avoid coherence effects. The signal \( I(\lambda) \) is periodic and composed of several frequencies. With this retarder-thickness configuration, 13 frequencies are generated on the 10 nm-analysis window. The coefficients of a Mueller matrix \((m_i)\) are retrieved through application of a Fourier transform (real part and imaginary part) to \( I(\lambda) \). By application of the calibration procedures described in [8], the accuracy on the \( m_i \) coefficients (normalized by \( m_0 \)) is below 0.03 for measurements of well-known media (polarizer, waveplate). The interest of this polarimeter lies essentially in its short acquisition time since it depends only on the detector (aperture time and repetition rate, respectively 10 μs and 1 kHz in this work).

The parameters issued from experimental Mueller matrices \((M)\) were extracted by the Lu and Chipman decomposition [9], which is based on the assumption that \( M \) is the product of three matrices (diattenuation, \( M_D \), retardance, \( M_R \), depolarization, \( M_A \) matrices),
corresponding to each polarimetric effect (dichroism, birefringence, depolarization) so that $M = M_A \cdot M_\lambda \cdot M_p$. In this study, only the depolarization index, $P_D$, calculated directly from $M$ [10], and the retardance, $R$, from $M_R$ are analyzed. One should note that the $P_D$ index is equal to 1 for a non depolarizing medium and theoretically less than 1 otherwise (although the noise may rise it to above 1). The diattenuation signal, attributed to the Sirius red dye, is weak at the wavelength 830 nm due to the low absorption of the dye [6]. Moreover, the measurement of the fast axis orientation of the retarder, which gives the direction of collagen fibers, is not considered with here.

We carried out experiments with five 16-μm thick acute surgical human liver biopsies that were fixed in formalin and embedded in paraffin. They are then deparaffinized and stained by Sirius red. Samples were mounted between a microscope slide and a 0.17mm-thick coverslide. The liver histological status was assessed by two trained pathologists, which accounts for the different scores for some biopsies studied in this Letter. Liver cryosections were set in an afocal system built with two 10x 0.25NA objectives. Slices were mounted on motorized translation stages in order to scan the samples (x,y directions) and change the cross-section of the beam, i.e. the resolution of the image (by moving samples in z direction). All of the samples were set at normal incidence by auto-collimation and thus illuminated in the same direction by the laser source. For a 50-μm estimated resolution (knife-edge technique), retardance $R$ and depolarization index $P_D$ of two biopsies (for example F0 and F4) have been measured and represented pixel by pixel in Fig. 2 for 2.5x2.5 mm² regions of interest (ROI's). These ROI's were selected from full-field Non-Polarized (NP) images taken with a CMOS camera (not represented on the setup of Fig.1). Second Harmonic Generation (SHG) diffraction-limited images (2.5-μm estimated resolution) of the same ROI’s were obtained on the scanning nonlinear microscope of our lab (Olympus BX51WI-FV300 confocal system equipped with a Coherent Mira femtosecond laser tuned at 830nm and a home-built SHG photodetector at 415nm settled in transmission) using a low-NA objective (Olympus UPLSAPO 4x 0.16NA).

In the classical Non-Polarized image of the F4 biopsy, fibrotic areas appear darker (due to Sirius dye) than the normal tissue. Similar structures are observed on the retardance image, which is also closely correlated to the corresponding SHG image. As SHG imaging is highly specific to fibrillar collagen [11], we can conclude that retardance is mainly due to fibrillar collagen. On the other hand, depolarization index slightly changes on the ROI and is close to 1 in the F4 biopsy. However if $R$ and $P_D$ are analyzed in the area of blood vessels (for example in the F0 biopsy which possess a lot of vessels), results are quite different. As expected, retardance appears in the vessel region due to fibrillar collagen (confirmed by the SHG image), but there is also a stronger depolarization effect (i.e. depolarization index less than 1) when the laser beam focuses on the vessel walls. In this case, there is an important spatial variation of retardance within the laser beam cross section (one part of the beam sees birefringence and the other part sees no polarization effects). The polarimetric properties distributed in the beam are incoherently added, leading to depolarization. This effect has been observed on every vessel regions, whatever the degree of the disease. This could be explained by the fact that on vessels walls, the spatial variation of retardance is the strongest. Thus fibrotic collagen could be distinguished from collagen around vessels thanks to depolarization index. One should note that given the thickness of the biopsies (16 μm), depolarization due to forward scattering is negligible (scattering mean free path = 100 μm).

Then, after using images to assign depolarization to the collagen of vessels, a statistical method has been developed in order to quantify the grade of the disease without building any image. Retardance and depolarization index were measured by scanning, with a 50-μm estimated resolution, a surface of 1.5x1.5 cm² area with a 250-μm step size (i.e. a sampling of about 3600 measurements on the whole liver biopsies). The depolarization index $P_D$ is used to filter retardance values due to collagen around blood vessels and select only those resulting from the modification of hepatic tissue by keeping measurements with experimental $P_D$ values ranging from 0.98 to 1.02, due to Gaussian noise. Every time the laser beam probes a diseased area, the retardance value is superior to the baseline (Fig. 3)

The probability distribution of retardance is assessed for the samples after $P_D$ filtering and is different from a Gaussian distribution according to the modification of the hepatic tissues. The retardance distribution is
characterized by the mean, $R_0$, and a retardance value $R_{\text{lim}}$ for which the frequency of the theoretical Gaussian distribution is equal to 1% of the maximum. It can be noted that $R_0$ and $R_{\text{lim}}$ depend on: 1) the stability of the room temperature which slightly modifies retardance measurements, 2) the beam focusing conditions and 3) the staining variability.

The choice of the beam focus can be a relevant element in order to enhance retardance values and obtain a more sensitive measurement. Indeed due to the spatial inhomogeneity of the birefringence properties (discrepancy of optical axis directions), only the mean retardance is measured and it increases with the focusing as Fig. 5 shows. Nevertheless, if the spot size drops to below 50 $\mu$m when the sample is moved between the two objectives, the image background increases. The origin of this modification is not clear, and could come from the residual paraffin in which liver specimens were first embedded [12].

In conclusion, the present paper illustrates and discusses a novel automated quantitative method based on a scanning system that measures the polarimetric properties of hepatic tissues. This tool is able to quantify the fibrillar collagen of liver fibrosis without scoring the fibrillar collagen around vessels thanks to depolarization. The study of retardance distribution after filtering enables to distinguish between disease stages and should limit the high degree of observer discrepancy. Thus, our method represents a complement to semi-quantitative indexes of fibrosis to define the evolution stage of several hepatic diseases more accurately.

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References

Full references


