Quantitative correlation between light depolarization and transport albedo of various porcine tissues

Sanaz Alali
Manzoor Ahmad
Anthony Kim
Nasit Vurgun
Michael F. G. Wood
I. Alex Vitkin
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Abstract. We present a quantitative study of depolarization in biological tissues and correlate it with measured optical properties (reduced scattering and absorption coefficients). Polarized light imaging was used to examine optically thick samples of both isotropic (liver, kidney cortex, and brain) and anisotropic (cardiac muscle, loin muscle, and tendon) pig tissues in transmission and reflection geometries. Depolarization (total, linear, and circular), as derived from polar decomposition of the measured tissue Mueller matrix, was shown to be related to the measured optical properties. We observed that depolarization increases with the transport albedo for isotropic and anisotropic tissues, independent of measurement geometry. For anisotropic tissues, depolarization was higher compared to isotropic tissues of similar transport albedo, indicating birefringence-caused depolarization effects. For tissues with large transport albedos (greater than ~0.97), backscattering geometry was preferred over transmission due to its greater retention of light polarization; this was not the case for tissues with lower transport albedo. Preferential preservation of linearly polarized light over circularly polarized light was seen in all tissue types and all measurement geometries, implying the dominance of Rayleigh-like scattering. The tabulated polarization properties of different tissue types and their links to bulk optical properties should prove useful in future polarimetric tissue characterization and imaging studies.

Keywords: optical properties; polarimetry; tissues.

1 Introduction

In an effort to better visualize biological tissue structure, polarized light imaging has been used extensively to enhance spatial resolution and to provide contrast that cannot be achieved using ordinary light alone.1–7 In addition, polarized light-based techniques (polarimetry) have allowed tissue characterization via intrinsic properties such as birefringence (arising, for example, from fibrous connective tissue networks containing collagen and elastin), orientation (arising from structural organization), and optical activity (arising from the presence of chiral molecules such as glucose).8–10 Polarimetry-based metrics have demonstrated sensitivity to changes in tissue structure and composition and may be closely associated with underlying pathology. For instance, scar formation in the myocardium has been linked to a decrease in the birefringence signal.8–10 One of the limitations to polarized light-based techniques in optically thick turbid media (such as bulk tissue) is the effect of multiple scattering, which effectively randomizes the photon direction, phase coherence, and polarization. Consequently, only a fraction of the light emerging from a sample in particular detection geometry will remain polarized. Since polarimetry relies on measuring, quantifying, and interpreting the surviving polarized light fraction, examining the depolarization rates in different tissue types and relating these to tissue structure and pathology are of great importance.

Previously, several studies have concentrated on light depolarization in biological tissues. Demos et al. have suggested using the degree of polarization for discriminating between different types of tissues.13 Jacques et al. have used polarized light imaging to distinguish superficial backscattered light from the collagen fibers versus highly depolarized light from deep skin layers.12 The same group also has compared chicken liver and breast and porcine muscle and skin with a transmission mode polarimetry system, and it reported that linearly polarized light depolarizes faster in (anisotropic) tissues with birefringence, compared to isotropic tissues.13 Vitkin et al. have demonstrated in vivo that light that is diffusively scattered by human skin retains significant polarization in exact backscattering directions, and the degree of polarization increases with increased absorption.14 Sankaran et al. have investigated the degree of polarization in five porcine tissues (fat, tendon, blood, myocardium muscle, and artery) of different thicknesses (0.1 to 2 mm) in transmission geometry. They have shown that the Rayleigh-scattering regime is dominant for all the tissues except blood, with the degree of linear polarization exceeding the degree of circular polarization.15 More recently, Antonelli et al. compared the Mueller matrix in the backscattering direction from healthy and cancerous human colon biopsy samples, indicating that the cancerous samples were less depolarizing.16 A common theme in these studies is to quantify and understand the depolarization mechanisms in tissues and to use depolarization as a diagnostic tool. The current article thus describes our efforts in measuring, quantifying, and understanding...
depolarization behavior in a variety of bulk porcine tissues in the context of their measured optical properties.

A commonly used approach to characterize tissues is by measuring their optical properties, specifically the reduced scattering coefficient $\mu_s'$ and the absorption coefficient $\mu_a$. The reduced scattering coefficient is the probability of the photons scattering in the forward direction per infinitesimal path-length. The absorption coefficient is the probability of the photons absorption per infinitesimal path-length. Several suitable methods have been developed, with their relative advantages, disadvantages, and ranges of validity closely examined.15 In the current study, we performed polarimetric and optical property measurements and analysis on optically thick samples from six different porcine tissue types. The isotropic (low birefringence) tissues examined were the liver, kidney cortex, and brain. The anisotropic (high birefringence) tissues were the myocardium muscle, loin muscle, and tendon. To quantify the dependence of the depolarization rates on source-sample-separations was used. The optical properties

$\mu_a$, $\mu_s'$, of 0.8, 1.5, 2.4, 4.0, and 5.3 mm was connected to an optical control system that consisted of a diode laser (Thorlabs, Newton, NJ) at 635 nm, an optical multiplexer (MPM-2000, Ocean Optics, Dunedin, FL), and a visible-light spectrometer (S2000, Ocean Optics).

The collector fiber was connected to the spectrometer. The multiplexer was used to index the laser beam from fiber-to-fiber so as to achieve the various source-collector distances required for the spatially resolved diffuse reflectance algorithm. The well-known diffuse reflectance equations were employed to model the reflectance at each $r$.23,24 The probe was calibrated against a diluted Intralipid (Fresenius Kabi, Sweden) solution with known optical properties and validated against a series of Intralipid-dye phantoms with $\mu_a = 0.5–5$ cm$^{-1}$ and $\mu_s' = 3.8–11.5$ cm$^{-1}$. The two optical property coefficients, $\mu_a$ and $\mu_s'$, were computed from the five reflectance measurements by solving an inverse problem. It is worth noting that the algorithm did not utilize some of the reflectance measurements in the case of low $\mu_s'$ because the diffusion theory reflectance equation breaks down at low $\mu_s'$ for low $r$, as outlined in Kim et al.25 Hence, the algorithm would omit the reflectance value at, say, $r = 0.8$ for $\mu_s' < 5.75$ cm$^{-1}$, and use only four reflectance measurements to solve in the inverse problem in this scattering range. With this probe, we measured the optical properties of the six different tissues, periodically examining a well-characterized scattering solution of Intralipid as a calibration/system validation control phantom.

For polarimetric measurements, two slabs of each type of tissue, one with 2 mm of thickness and one with 1 cm of thickness, were sectioned. No formalin fixation was used to avoid possible crosslinking-induced polarization property changes, so the results apply to freshly excised tissues. Cutting the fresh tissue with a blade to a uniform thickness is challenging; therefore, several sections were made and the three most uniform slabs with the desired thicknesses were chosen for the experiment. The resultant tissue slabs were placed between two glass slides. The 2-mm samples were imaged in both transmission and backscattering geometry, while the 1-cm samples were imaged only in backscattering mode (not surprisingly, complete polarization loss was observed in transmission for all 1-cm tissues).

The polarimetric imaging setup shown in Fig. 1 was used. A diode laser (Thorlabs) with a wavelength of 635 nm was used to illuminate the sample. The polarizer, P1, and the removable quarter waveplates, WP1, were used to generate different linear and circular input polarization states. The first lens, L1, was used to project a 1-cm-diameter spot size onto the sample. Light exiting the sample in a particular detection direction then was channelled through an output arm consisting of a polarizer, P2, a removable waveplate, WP2, and a collecting lens, L2, that focused the light onto a charge-coupled device (CCD).

Fig. 1 Schematic of the polarimetric imaging setup. P1 and P2 are polarizers; WP1 and WP2 are removable quarter waveplates; L1 and L2 are lenses. In transmission mode, the detection arm is collinear with the incident beam. In the backscattering geometry, the analyzer arm is positioned $\Theta = 25$ deg off the exact retro-reflection direction.
camera (CoolSNAPK4, Photometrics, Tuscon, AZ). The output polarization was selected by P2 and WP2. Imaging was performed in two different geometries. In transmission mode, the output arm was placed collinear to the incident beam. In backscattering mode, light was collected by the output arm at an angle of 155 deg off-axis relative to the incident beam (25 deg off-axis from the exact backscattering direction). The camera field of view was set to a 1.5-cm-wide square, with the collected light spot in its center.

To characterize the interaction of polarized light with a sample of tissue for a specific measurement geometry, Mueller algebra was employed. In Mueller formalism, a polarized light beam is described by the Stokes vector \( S = [J Q U V]^T \), and its interactions are described by multiplication by a 16-element square \((4 \times 4)\) matrix. This matrix, called the Mueller matrix, represents a particular interacting entity (polarizer, wave plate, tissue sample, etc.). If the incident light beam is represented by the Stokes vector \( S_i \), and the output light beam vector is \( S_o \), then the Mueller matrix \( M \) acts as the polarization transfer function, such that \( S_o = M \cdot S_i \). To calculate the Mueller matrix for a particular tissue sample at particular detection geometry, we performed 24 sequential measurements. For each of four input polarizations \([H \text{ (linear at 0 deg)}, V \text{ (linear at 90 deg)}, P \text{ (linear at 45 deg)}, R \text{ (right circular)}]\), intensity profiles at six output polarizations \([H \text{ (linear at 0 deg)}, V \text{ (linear at 90 deg)}, P \text{ (linear at 45 deg)}, B \text{ (linear at }-45\text{ deg)}, R \text{ (right circular)}, L \text{ (left circular)}]\) were measured. From these 24 measurements, Mueller matrices for each sample for that specific geometry (transmission/backscattering) were constructed, as described previously in several studies (e.g., Wallenburg et al.\textsuperscript{26}).

A complicating issue in turbid polarimetry is the simultaneous occurrence of several polarized light-tissue interactions. This results in complicated cross-talk within and between the Mueller matrix elements and obscures the contribution of the individual sample polarization properties. Several mitigating strategies have been developed recently to deal with this problem.\textsuperscript{28-30} One approach is termed polar decomposition, originally proposed by Lu-Chipman and adapted by our group for tissue polarimetry.\textsuperscript{31-34} This decomposition method allows for extraction of individual properties of interest (birefringence, optical activity, depolarization, and diattenuation) from a single measured Mueller matrix.\textsuperscript{31-34} We have validated this approach using polarized-light Monte-Carlo simulations and phantoms with controlled polarization properties, as well as ex vivo and in vivo tissues.\textsuperscript{8-10,35} In this method, the tissue Mueller matrix \( M \) is decomposed into three constituent basis matrices—the depolarization matrix \( M_\Delta \), the retardance matrix \( M_R \), and the diattenuation matrix \( M_D \):\textsuperscript{35}

\[
M = M_\Delta M_R M_D. \quad (1)
\]

Ignoring for the moment the other two derived basis matrices and the information that they contain, the depolarization matrix can be written in the following form:

\[
M_\Delta = \begin{bmatrix} 1 & \vec{P}_\Delta & m_\Delta \end{bmatrix}, \quad (2)
\]

where \( \vec{P}_\Delta \) is the polarization vector and \( m_\Delta \) is a symmetric submatrix. The diagonal elements of \( m_\Delta \) describe the preserved portion of each of the polarization states \((Q, U, V)\) in the Stokes vector of the light beam. The net depolarization factor, also called the depolarization power, can be defined as\textsuperscript{27,33}

\[
\Delta_T = 1 - \frac{|m_{11}| + |m_{22}| + |m_{33}|}{3}, \quad (3)
\]

where \( m_{11}, m_{22}, \) and \( m_{33} \) are the diagonal elements of \( m_\Delta \). \( \Delta_T \), here termed the total depolarization, yields a quantitative average measure of how a material with this particular Mueller matrix depolarizes light. One can go further and define the linear depolarization, \( \Delta_L \), and circular depolarization, \( \Delta_C \), as follows:

\[
\Delta_L = 1 - \frac{|m_{11}| + |m_{22}|}{2}, \quad (4)
\]

\[
\Delta_C = 1 - |m_{33}|. \quad (5)
\]

Linear depolarization \( \Delta_L \) is a measure of the average depolarization of the \( Q \) and \( U \) components (linear polarization at 0 to 90 deg and 45 to 135 deg) of the Stokes vector interacting with the sample. Correspondingly, circular depolarization \( \Delta_C \) is a measure of the depolarization of the \( V \) component (circular polarization) of the Stokes vector. Using the resultant linear, circular, and total depolarizations allows quantitative comparison of the depolarizing behaviors of different biological tissues at a specific geometry, independent of the illumination polarization state (incident Stokes vector).

It is important to distinguish between depolarization metrics derived from the Mueller matrix of the sample and the degree of polarization calculated from the Stokes vector of the light. The former are intrinsic properties of the transfer function of the sample in a particular geometry (as embodied in the tissue Mueller matrix), whereas the latter depend on the sample and on the Stokes vector of the incident light. The derived Mueller metrics thus are more specific to the tissue being studied in that they factor out the confounding effects of the measurement system (incident light polarization). Further, the polar decomposition methods discussed previously permit one to isolate the depolarization effects from other polarizing interactions, something that is not possible with the Stokes descriptors alone. Mathematically, the total degree of polarization in the Stokes description is always higher than the linear and circular degrees of polarizations; an analogous relationship does not hold true for the Mueller matrix-derived depolarizations [Eq. (3) through Eq. (5)]. For the remainder of this paper, we discuss depolarization (\( \Delta_T, \Delta_L, \Delta_C \)) as derived from the decomposed Mueller matrices.

Following the decomposition procedure, we quantify the intrinsic depolarization parameters of the tissues in each geometry, as derived from Mueller algebra (independent of the incident light polarization state and other tissue polarizing interactions, as per the previous discussion). To gauge the uncertainty in the entire polarimetry methodology, a given Mueller matrix (each
with 24 measurements) was recorded three times for all samples in both geometries. Total, linear, and circular depolarization metrics reported in this paper are the average results of these measurements.

3 Results and Discussion

The measured optical properties \((\mu_a, \mu'_a)\) of the swine tissues at 635 nm are shown in Table 1. The reduced scattering coefficient contributes to the extent of photon diffusion in tissue. It is related to the scattering coefficient \(\mu_s\) and anisotropy factor \(g\)—the mean cosine of the scattering angle—through the relation \([\mu'_a = \mu_s(1 - g)]\). The transport albedo \(a'\) is a dimensionless quantity defined as \([\mu_s/(\mu'_a + \mu_a)]\). In addition to their role in interpreting tissue polarimetry results, these measured optical properties may contribute to recent tissue optics compilations.\(^{37-40}\)

Polarized light imaging allows the construction of spatial maps in which the Mueller matrix derived parameters may be plotted. In Fig. 2, we illustrate the calculated total, linear, and circular depolarization of kidney cortex samples in the transmission and backscattering geometries. The data shown in each map is the average of three independent measurements. The average standard deviation over all the pixels in the image is 1.5%. This number is representative of the average uncertainty in the derived polarization metrics, characteristic of the system noise over the measurement period.

Several prominent features are visually evident from Fig. 2. First, linear depolarization is lower than circular depolarization for both transmission and backscattering geometries, as well as for both 2-mm and 1-cm kidney cortex samples. In other words, linear polarization is better preserved than circular polarization in this tissue type. Second, the total depolarization is observed to be lowest in transmission mode for 2-mm samples and greatest in the backscattering mode for the 1-cm sample. Such images have been obtained for all the examined tissues; to enable quantitative analysis and inter-tissue comparisons, we averaged the data along a 0.1-cm central strip in the \(y\) direction, and plotted the total depolarization along the \(x\) direction, as shown in Fig. 3.

Comparing panels (a) and (b) of Fig. 3, one can conclude that all the tissues except tendon and brain are less depolarizing in transmission geometry rather than in backscattering. For brain and tendon, the transmission mode is not as useful as depolarization becomes too severe (greater than 98.5% even for this relatively thin, 2-mm slab) and essentially all polarized light information is lost. Measurements with such large depolarization are not reliable and lie in the range of the polarimetry system’s noise level. The transmission geometry thus becomes untenable for tissue polarimetry as either physical thickness or optical properties increase sufficiently. Hence, the brain and tendon results are not included in Fig. 3(a). Such extensive depolarization is likely because brain and tendon both have transport albedos higher than 0.97 (Table 1). A large transport albedo signifies the predominance of multiple scattering, with its variety of long, zigzag paths that effectively depolarize the light.\(^{41,42}\)

Furthermore, scattering in biological tissues is generally forward-peaked, with the anisotropy factor \(g\) varying between 0.9 and 0.99.\(^{19}\) A large transport albedo and forward-peaked

\[ \text{Table 1 Measured optical properties of several swine tissues at 635 nm.} \]

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Reduced scattering coefficient (cm(^{-1}))</th>
<th>Absorption coefficient (cm(^{-1}))</th>
<th>Transport albedo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td>6.90 ± 0.10</td>
<td>4.10 ± 0.11</td>
<td>0.62 ± 0.01</td>
</tr>
<tr>
<td>Kidney cortex</td>
<td>6.00 ± 0.10</td>
<td>1.20 ± 0.20</td>
<td>0.83 ± 0.02</td>
</tr>
<tr>
<td>Myocardium muscle</td>
<td>8.22 ± 0.40</td>
<td>1.62 ± 0.35</td>
<td>0.83 ± 0.07</td>
</tr>
<tr>
<td>Loin muscle</td>
<td>4.00 ± 0.07</td>
<td>0.37 ± 0.04</td>
<td>0.91 ± 0.02</td>
</tr>
<tr>
<td>Brain</td>
<td>11.27 ± 0.48</td>
<td>0.26 ± 0.10</td>
<td>0.97 ± 0.05</td>
</tr>
<tr>
<td>Tendon</td>
<td>13.50 ± 0.54</td>
<td>0.35 ± 0.18</td>
<td>0.97 ± 0.06</td>
</tr>
</tbody>
</table>

Fig. 2 Depolarization maps of the kidney cortex derived from the polar decomposition of the measured Mueller matrices. The columns indicate the total, linear, and circular depolarizations, and the rows indicate the different detection geometries and sample thickness. The \(x\) and \(y\) axes indicate the length and width, respectively. The color bar indicates depolarization levels as specified on the percent scale, where dark blue signifies regions where light remains most polarized and deep red indicates regions where light has lost most of its polarization. The error bar size is equal to the line thickness.
scattering implies that photons detected in backscattering geometry have experienced fewer scattering events and on average have traveled shorter path-lengths compared to the photons detected in transmission geometry. In fact, most photons detected in the reflection mode in these types of scattering media have sampled mostly near-surface layers, as we have recently demonstrated by Monte Carlo modeling studies.42–44 As a result, less depolarization is observed for high transport albedo tissues in backscattering mode.

A general trend in the total depolarization values of the tissues emerges from a closer examination of Fig. 3. Liver has the minimum total depolarization in all measurement geometries, whereas tendon appears to have the highest total depolarization among all tissue types. The tissues arranged in the order of increasing depolarization are liver, kidney, loin muscle, brain, myocardium muscle, and tendon. Comparing Fig. 3 and Table 1, one notices a correlation between total depolarization rates and the transport albedos of the corresponding tissue type.

To explore this further, the minimums of total, linear, and circular depolarization (Fig. 3) are plotted against the tissues’ transport albedos (Table 1) in Fig. 4. As previously mentioned, brain and tendon data in transmission mode are not shown because they exhibit total depolarization greater than 98.5% in transmission mode, which is in the range of the system noise level. As a result, less depolarization is observed for high transport albedo tissues in backscattering mode.

Another interesting trend evident from Fig. 4 is the preferential retention of linear polarization states compared to circular polarization states, in all tissues, for both geometries. Interestingly, the correlation coefficient values are also higher in isotropic tissues; as seen, anisotropic tissues shows higher depolarization compared to the isotropic one with the same transport albedo. For example, myocardium muscle, whose \( a^l \) is not exceptionally high (close to the value for kidney cortex), exhibits very high total depolarization. In addition, when comparing tendon tissue depolarization to that of the brain, tendon exhibits a higher depolarization despite having a comparable \( a^l \). One possible reason for both of these observations could arise from the intrinsic anisotropy exhibited by the different tissue types. Myocardium muscle and tendon (and loin muscle, to a lesser extent) are known to be highly anisotropic relative to the other tissues.17 The underlying birefringence that causes this tissue anisotropy thus may contribute to depolarization over and above the transport albedo effect invoked previously. Specifically in tissues, birefringence magnitude and orientation may be spatially inhomogeneous, changing in different regions of tissues/microdomains;34,35,45,46 in these regions, polarized light undergoes additional randomization, and therefore the total depolarization increases.15,45 Note that we specifically concentrate on depolarization phenomena here and do not analyze the derived birefringence magnitude and orientation (derived from the retardance matrix \( M_R \)), which are important in anistropic tissues such as skeletal muscle;33,46 an ongoing study examining birefringence phenomena in greater detail (e.g., the effects of variable spatial domains of different magnitude/orientation of birefringence) will be reported elsewhere.
cases of linear depolarization. Better preservation of linear polarization is usually associated with Rayleigh-like scattering, whereas tissues are often considered Mie-like in their scattering behavior. Thus, identifying the appropriate dominant scattering regime in biological tissues is tricky. Conventional wisdom is that tissues are composed of large scatterers and thus exhibit Mie-like scattering behavior, consistent with the predominantly forward-peaked nature of the observed scattering phase function (high values of $g$, as discussed previously). While cells and subcellular organelles such as the nucleus and mitochondria are within the Mie scattering regime, extracellular matrix elements such as collagen fibrils fall in the Rayleigh scattering regime. In suspensions containing both small and large scatterers, Ghosh et al. have shown that scattering behavior is dominated by smaller particles. This was also observed in simulations whereby small organelles were shown to contribute heavily to light scattering in cells. On the basis of size alone, one can therefore expect that biological tissues may be described better by the Rayleigh regime, with an alternate explanation for the high observed $g$-values.

The other important determining factor is the relative refractive index. Based on another study by Ghosh et al., if a phantom has large scatterers and a high anisotropy factor but a small relative refractive index contrast, it does not fall in the Mie scattering regime (e.g., circular polarization is no longer preserved better than linear, as it should in the “classical” Mie regime). Relative refractive index contrast and scatterer size thus are both important in determining the dominant scattering behavior. Interestingly, Sankaran et al. also reported better retention of linear over circular polarization states in transmission, measured though ~1-mm-thick slabs of tissues and through dense suspensions of high-contrast tissue simulating phantoms (compared to dilute phantoms); for the latter, they invoked the mechanisms of correlated (dependent) scattering engendered by dense packing to explain their findings. The preferential retention of linear polarization states observed in our study is also indicative of Rayleigh scattering, although the exact mechanism remains unclear. Given the complicated nature of tissue scattering, likely both small scatterers and large scatterers (with small relative refractive index and large anisotropy factor $g$), augmented by

![Fig. 4 Depolarization of different tissues related to their transport albedos.](image-url)

The correlation coefficient ($r$) indicates how well data fits a straight line, and it is calculated as $r = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2}(y_i - \bar{y})^2}$ for data points, with $x_i$ and $y_i$ being their mean values. In this case, $x_i = \text{transport albedo}$ and $y_i = \text{depolarization}$. Note that brain and tendon data is not included in (a)–(c) owing to their high depolarization (>98%).
compactness, are contributing. It is also worth mentioning that in all these studies, linear and circular degrees of polarization were based on the Stokes parameters of the light beam and thus depend somewhat on the input light parameters and are unable to account for other tissue polarizing effects; in contrast, our reported results are derived from the tissue Mueller matrix via polar decomposition and thus are free of these two confounding effects. Despite these differences, both approaches suggest the preferred preservation of linear polarization states over circular polarization states in biological tissues. We are currently conducting a controlled phantom study to explore further the linear and circular depolarization behavior as a method to ascertain the dominant scattering regime and the essential medium characteristics determining it.\(^{51}\)

4 Conclusions

In summary, we have measured depolarization and optical properties in a variety of biological tissues, and we have quantified the correlations between them. We used polarized light imaging and Mueller matrix polar decomposition to extract linear, circular, and total depolarizations in several freshly excised bulk swine tissues (liver, tendon, kidney cortex, brain, myocardium muscle, and loin muscle). Both transmission and reflection geometries were used to examine 2-mm and 1-cm tissue slabs. Bulk optical properties of the different tissues were measured and related to the derived depolarization behavior. A linear correlation between transport albedo and depolarization was observed and quantified. Furthermore, in highly anisotropic tissues like cardiac muscle and tendon, birefringence increases depolarization over and above the multiplescattering effects, as determined by the high transport albedo (i.e., in spite of the relatively low \(a\) of these birefringent tissues). Also observed was the preferential preservation of linear over circular polarized light in all tissue types and for all detection geometries. A tentative explanation invoking effective scatterer size and relative refractive index contrast was provided. Finally, for high-transport albedo tissues such as the brain and tendon, light experiences more depolarization through propagation in the forward direction (transmission mode) compared to the backscattering direction. In fact, transmission detection geometry becomes untenable for polarimetry applications for even 2-mm samples of brain and tendon because of extensive polarization loss. Detailed studies of depolarization behavior in various bulk animal tissues should help researchers understand the underlying biophysics of polarized light-tissue interactions, provide polarization properties of various tissue types that are generally unavailable in the literature, and assist in experimental design for polarized light imaging and for polarimetry-based tissue characterization studies.

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