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Abstract. We present an automated and systematic two-dimensional discrete Fourier transform (2D-FFT) approach to analyze collagen fiber organization through the use of second harmonic generation (SHG) microscopy. Average orientations of individual domains and Ising-like order parameters introduced to characterize the correlation between orientations of adjacent domains may be used to quantitatively characterize fibrous tissues. Our approach was applied to analyze tissues including rat tail tendon, mouse skin, bovine corneas, and human corneas. We also show that collagen fiber organization in normal and keratokonus human corneas may be distinguished. The current approach may be used for the quantitative differentiation of SHG collagen fiber morphology in different tissues and may be applied for diagnostic purposes. © 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.

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1 Introduction

Second harmonic generation (SHG) imaging is a minimally invasive optical imaging technique with intrinsic optical sectioning capability. The coherent nature of SHG leads to an intensity that is sensitive to the noncentrosymmetric arrangements of molecules and in biomedical applications, SHG has been shown to be effective in imaging fibrous collagen-rich tissues such as tendons, cartilage, and cornea. Specifically, SHG microscopy has been successfully used in the label-free identification of structural changes in connective tissues under conditions such as skin photoaging, corneal abnormalities (keratokonus), liver fibrosis, and Osteogenesis imperfecta. In tissue engineering applications, SHG imaging has been useful in the quantification and monitoring of collagen growths in engineered cartilage tissues. However, most efforts in using SHG imaging have focused on morphological characterization rather than quantification of the structural motifs of the fibrous structures. Therefore, combining SHG imaging with the appropriate methodology to quantify fiber structure can lead to objective evaluation in their structural organization. Recently, two-dimensional discrete Fourier transform (2D-FFT) has been shown to be an effective tool in analyzing fibrillar organization. In particular, the average orientation and anisotropy of fiber tissue within any given region of interest (ROI) can be determined using 2D-FFT transformation. Both of these parameters vary with the domain size of analysis and the variation can be viewed as a quantitative standard to monitor changes in the fibrous tissues. However, in previous studies, the sizes of the imaging area used for 2D-FFT analysis are usually subjectively chosen. Therefore, the major purpose of the present study was to introduce a systematic approach to investigate the effect of the size of the analyzed region on the Fourier transformation analysis. In addition to local fiber orientation and aspect ratio, we introduce a quantitative parameter evaluating the correlation between the orientations of neighboring domains. For this purpose, the Ising-like order parameter was calculated for the fiber angular matrices. In addition to normal connective tissues, we show that our approach can be used to differentiate between normal and pathological fiber orientations in the cornea.

2 Materials and Methods

2.1 Sample Preparation

Excised pieces of rat tail tendon (RTT), mouse skin (MS), bovine cornea (BC), normal human cornea (NHC), and human keratoconic cornea (KHC) were used in this study. RTT and MS were prepared from unfrozen tissues, and BC was freshly obtained from a local market and prepared within an hour prior to experiment. Keratoconic corneas were obtained from surgeries of penetrating keratoplasty from a local hospital. Two samples of NHs were obtained postmortem (Minnesota Lions Eye Bank, MN), transferred frozen, prepared, and imaged within 3 h after thawing. Prior to multiphoton examination, corneal specimens were stored in 20% dextran/PBS solution for about an hour (dextran T500, Pharmacosmos, Denmark) to avoid swelling. All specimens were sealed in glass well slides to avoid dehydration.
2.2 Second Harmonic Generation Microscope

The SHG microscope used in this study has been described previously.10,11 In short, an 80 MHz, 100-fs laser pulse from a Ti:sapphire oscillator (Tsunami, Spectral Physics, Mountain View, CA) tuned to 780 nm was used as the excitation source. The laser beam, directed by scanning mirrors, was expanded and reflected onto the back aperture of the focusing objective by a short pass dichroic mirror (700dcpxruv-2p, Chroma Technology, Rochester, VT). The oil-immersion objective (40x, S Fluor, NA1.3, Nikon, Japan) was used for imaging the RTT and MS, while for the corneas the water-immersion objective was used (40x, S Fluor, NA 0.8, Nikon, Japan). For RTT and MS, the SHG signal from the sample was collected in the backward direction by the focusing objective and further guided by a dichroic mirror (435DCXR, Chroma technology), whereas SHG from corneas was detected in forward geometry using collection lens (f = 5 cm, Chroma). In both cases, the SHG signals were isolated by a narrow band pass filter (HQ390/22m-2p, Chroma technology), and detected by photomultiplier tubes (R7400P, Hamamatsu, Japan).

2.3 SHG Image Acquisition

SHG images were acquired at the depths of approximately 15 μm from the surface of sample for RTT and MS. An automatic translation stage was used to translate the specimen following the acquisition of small-area images (512 × 512 pixel, 105 × 105 μm²). These adjacent images were subsequently stitched to form large-area images. The large-area images were 210 × 210 μm² (2 × 2 tiles) for RTT, 525 × 525 μm² (5 × 5 tiles) for MS, and 630 × 630 μm² (6 × 6 tiles) for NHC. All imaging experiments were performed at the ambient temperature of 19°C.

2.4 Fiber Orientation Analysis

To perform fiber organization analysis, the images were first divided into equally sized domains. Next, 2D-FFT was performed on each domain. To demonstrate our approach, we first applied our method on an artificially created fibrous pattern (192 × 192 pixels) composed of adjacent domains, each 64 × 64 pixels in size, with mutually orthogonal fibrous patterns in adjacent domains [Fig. 1(a) to 1(d)]. Next, the pattern was arbitrarily subdivided into domains of various sizes of 32 × 32, 48 × 48, 64 × 64, and 96 × 96 pixels in Fig. 1(a) to 1(d), respectively. The 2D-FFT analysis was applied on each subdivided domain, and the corresponding transformed images in the Fourier domain are shown in Fig. 1(a*) to 1(d*) in green. This procedure then was followed by a single pixel erosion-dilation process in order to smooth and delineate the contours of the transformed image. Quantification of the fiber geometry was then performed by fitting by an ellipse [shown in red in Fig. 1(a) to 1(d)] which allows the characterization of each domain by two parameters: (1) the angle between horizontal axis and the short axis of the ellipse, and (2) the aspect ratio (AR) between the short and long axes of the ellipse. The orientation of the short axis of the ellipse for each domain is shown by the white lines in Fig. 1(a) to 1(d). Since it is known that the short axis of the fitted ellipse is parallel to the average fiber orientation within each domain and that the AR describes the directional anisotropy, orientation properties associated with the fibers in each subdivided domain can be determined.

In this example, when fibers within each domain lie along the same direction, the 2D-FFT analysis accurately reflects the direction of the fiber orientation [Fig. 1(a) and 1(c)]. However, when the domain contains fibers along two orthogonal directions, the FFT analysis then reflects the average orientation of the fibers within the domain. Note, that in the latter case AR values are comparatively bigger [Fig. 1(b) and 1(d)]. According to our estimate, for a good correlation between the pattern line directions on images and those estimated from 2D-FFT the average AR value should be less than 0.5. Note that this criterion, as well as the optimal domain sizes, would depend on the characteristic extent of correlation in organization and needs to be derived for each case. In the case of collagen fiber factors, such as tissue type, image quality, resolution, and contrast would matter. Once these parameters are fixed, the corresponding estimate can be used unambiguously henceforth.

2.5 Correlation Parameter

In addition to performing FFT analysis, we determined the collinearity of directions between adjacent domains through the use of two-dimensional the Ising-like order parameter.
\[
H = \frac{1}{(N-1)^2} \sum_{i,j=1}^{N-1} \left[ (\vec{S}_{i,j} \cdot \vec{S}_{i,j+1})^2 + (\vec{S}_{i,j} \cdot \vec{S}_{i+1,j})^2 \right] - 1
\]

(1)

where \(N\) is number of domains, \((\vec{S}_{i,j} \cdot \vec{S}_{i,j+1})\) is scalar product of unit vectors \(\vec{S}_{i,j}\) and \(\vec{S}_{i,j+1}\), which represent orientations of \((i, j)\) and \((i, j + 1)\) domains with tilt angles \((\theta_{i,j}, \theta_{i,j+1}) \in (-\pi/2, \pi/2)\). Specifically, \(H\) is an analog of configuration energy in the XY model of the nearest neighbor case without external field. \(H\) can be used to quantify overall directional dispersion, hence it provides a numerical value of image turbidity. Since

\[
(\vec{S}_{i,j} \cdot \vec{S}_{i,j+1}) = |\vec{S}_{i,j}| |\vec{S}_{i,j+1}| \cos(\theta_{i,j} - \theta_{i,j+1})
\]

\[= \cos(\theta_{i,j} - \theta_{i,j+1}),\]

(2)

for our case, \(H\) may be written as

\[
H = \frac{1}{(N-1)^2} \sum_{i,j=1}^{N-1} \cos^2(\theta_{i,j} - \theta_{i,j+1})
\]

\[+ \cos^2(\theta_{i,j} - \theta_{i+1,j}) - 1.
\]

(3)

Therefore, \(H\) represents the nearest neighbor correlation between domains, allowing us to describe the whole image with a single parameter. Theoretically, \(H\) takes on values between \(-1\) and \(1\), where \(H = 0\) stands for a completely random orientation of domains without any correlation, \(H = 1\) represents the case of an image with parallel domains and, finally, \(H = -1\) corresponds to the case of perfectly orthogonal neighbors, i.e., we have completely anti-correlated alignment. Calculation of the order parameter \(H\) was performed using custom IDL (IDL 6.4, ITT Visual Information Solutions, CO), Wolfram Mathematica (WM 8, Wolfram Research, Inc., IL), and MS VBA (MS Office Plus 2010, Microsoft Corporation, WA) software.

### 3 Results and Discussion

Following a demonstration of our method with an arbitrarily generated pattern in Fig. 1, we applied our approach to more complicated fibrous tissues acquired through SHG imaging. Shown in Fig. 1(a) to 1(d) are the SHG images of RTT, MS, BC, and NHC, respectively. Next, the corresponding images were subdivided into different domains (each \(32 \times 32\) pixels, \(6.6 \times 6.6 \mu m^2\)) in which 2D-FFT was performed [Fig. 2(a*) to 2(d*)]. In these examples, since the fibers are not perfectly aligned along a given direction within each domain, the AR values derived from the fitted ellipses of the FFT image would increase compared to the case of perfectly aligned fibers. Also shown in Fig. 2(a*) to 2(d*) are red lines corresponding to the derived average fiber orientation within each domain.

As in the case of the arbitrarily generated fibrous pattern, the result of the 2D-FFT analysis is highly dependent on the size of the domain chosen. With a reasonably small domain size, the local fiber orientation can be better determined. However, a domain size too small would contain an insufficient amount of information of fiber morphology for accurate determination of fiber direction. With a large domain size, the 2D-FFT analysis would yield an average orientation within the domain and details of local fiber orientation would be lost.

To demonstrate these features, 2D-FFT analysis was performed using a range of domain sizes from \(16 \times 16\) to \(512 \times 512\) pixels which corresponded to actual image size of \(3 \times 3 \mu m^2\) to \(100 \times 100 \mu m^2\), respectively. Following the analysis, the average AR values were obtained from all domain sizes and the results are shown in Fig. 3. In the cases of MS and RTT, the AR values are the result of averaging 10 images taken from 2 specimens; 5 each. In the case of BC, ARs from images obtained at 10 different depths at \(100 \mu m\) apart are averaged. NHC data represent values from forward SHG images obtained at 6 depths, \(100 \mu m\) apart obtained from 2 specimens. A two-sided student’s t-test was used to compare parameters obtained for different specimens and the level of statistical significance was calculated using IBM SPSS software (Chicago, IL).

As Fig. 3 shows, the minima of the averaged AR values occur when the domain sizes are between \(19 \times 19\) and \(36 \times 36\) pixels (corresponding to \(3.7 \times 3.7\) and \(8.2 \times 8.2 \mu m^2\)). Furthermore, the ARs of different fibrous architectures show similar trends at domain sizes below that at minimum AR. This observation suggests that the spatial variations of fiber orientation would be lost when the domain is too small to capture a statistical sample of fiber orientation within each domain, while the large domain size would not be able to provide a detailed information of fiber orientation.
is indicative of the fact that when domain size is decreased, the determination of general fiber orientation becomes erroneous. On the other hand, increasing the domain sizes will result in the inclusion of too many details of local fiber orientations, hence the averaging and loss of information on fine structure. However, as domain size increases beyond the point of minimum AR, our analysis starts to reveal differences between the different tissue morphologies. First, note that for the case of RTT, a relatively low AR value (<0.4) was obtained for domain size 96 x 96 pixels (19.8 x 19.8 μm²). This is indicative of the fact that collagen fibers in RTT specimens are well aligned. Note that at large domain sizes, the ARs for the four tissue types approached different terminal values of 0.40/C6 0.14 (RTT), 0.76/C6 0.12 (MS), 0.52/C6 0.08 (BC), and 0.59/C6 0.03 (NHC). Our results indicate the different collagen-containing tissues possess different degrees of short- and long-term order in organization.

Based on derived directional maps, we determined the Ising-like model correlation parameter according to Eq. (3). The average correlation parameter, calculated for different tissue types using the domain size at maximal AR value for the different tissues are presented in Table 1. Our results show that the calculated correlation parameters are consistent with the overall image morphology ranging from as low as 0.29 ± 0.08 for the more random organization in MS dermis up to 0.75 ± 0.03 for the highly organized RTT (p < 0.005).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Correlation parameter</th>
<th>Domain size</th>
<th>Average AR</th>
</tr>
</thead>
<tbody>
<tr>
<td>RTT</td>
<td>0.75 ± 0.03</td>
<td>19</td>
<td>0.36 ± 0.11</td>
</tr>
<tr>
<td>MS</td>
<td>0.29 ± 0.08</td>
<td>26</td>
<td>0.47 ± 0.21</td>
</tr>
<tr>
<td>BC</td>
<td>0.64 ± 0.12</td>
<td>36</td>
<td>0.43 ± 0.07</td>
</tr>
<tr>
<td>NHC</td>
<td>0.53 ± 0.02</td>
<td>36</td>
<td>0.44 ± 0.03</td>
</tr>
</tbody>
</table>

As a final example of the potential applicability of our method in distinguishing normal and pathological tissues, we applied the 2D-FFT approach to analyze collagen fiber morphology in normal and pathological human corneas. Specifically, both forward SHG images from normal and keratoconical human corneas were obtained and analyzed. Representative large area images of normal and keratoconus cornea obtained at the depth of 100 μm are shown in Fig. 4(a) and 4(b). Scleral examination of the keratoconus sample revealed a scar, which introduced additional turbidity into the orientation of collagen fibers of this sample [Fig. 4(b)]. The correlation parameter calculated using domain sizes of 26 to 48 pixel (5.3 to 9.8 μm) for NHC and KHC acquired by forward SHG at 100 μm depth. Arrows indicate the correlation parameter values corresponding to minimal AR values for both cases.
parameter was calculated for entire images obtained at 6 depths for each specimen. AR values were calculated for different domain sizes and the minimum was determined to be $32 \times 32$ ($6.6 \times 6.6 \, \mu m^2$) for KHC and $36 \times 36$ ($7.4 \times 7.4 \, \mu m^2$) for NHC. Figure 4(a*) and 4(b*) show 2D-FFT analysis results of arbitrarily chosen ROIs from images 4(a) and 4(b) using the domain sizes corresponding to minimum average AR.

In addition, the calculated average correlation parameters for the entire area of normal and keratoconus corneal images are shown in Fig. 6. The average correlation parameters corresponding to domain size $36 \times 36$ pixels of minimum AR values are $0.53 \pm 0.02$ and $0.43 \pm 0.03$ for normal and keratoconus corneas, respectively ($p < 0.005$). This serves as an example of the quantification of image morphology and shows that quantitative comparison of differences of fiber order can be performed.

4 Conclusion

In this study, 2D-fast Fourier transform analysis was developed and applied for morphological analysis of SHG images of different connective tissues. An approach, based on average AR estimation was developed, which allows for the evaluation of actual fiber directions. We found that both the aspect ratios and correlation parameter can be used to quantify differences in tissue morphology. While the aspect ratio analysis characterizes local fiber orientation within the ROI domains, the correlation parameter is effective in revealing the orientation order among the ROIs. To avoid ambiguity, our analysis was performed on systematically selected ROI sizes and we found that the dependence of AR and correlation parameters on ROI sizes can be used as additional metrics for characterization fiber orientation. In addition, to distinguish fiber morphologies in normal tissues such as RTT, MS, or BC, we found that organizational differences between normal and pathological cornea (keratoconus) can be achieved. The demonstrated approach can be used for automated morphological analysis of tissue fibers and may be extended for diagnostic purposes.

Acknowledgments

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