Automated cardiac sarcomere analysis from second harmonic generation images

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Abstract. Automatic quantification of cardiac muscle properties in tissue sections might provide important information related to different types of diseases. Second harmonic generation (SHG) imaging provides a stain-free microscopy approach to image cardiac fibers that, combined with our methodology of the automated measurement of the ultrastructure of muscle fibers, computes a reliable set of quantitative image features (sarcomere length, A-band length, thick–thin interaction length, and fiber orientation). We evaluated the performance of our methodology in computer-generated muscle fibers modeling some artifacts that are present during the image acquisition. Then, we also evaluated it by comparing it to manual measurements in SHG images from cardiac tissue of fetal and adult rabbits. The results showed a good performance of our methodology at high signal-to-noise ratio of 20 dB. We conclude that our automated measurements enable reliable characterization of cardiac fiber tissues to systematically study cardiac tissue in a wide range of conditions. © 2014 Society of Photo-Optical Instrumentation Engineers (SPIE) [Doi: 10.1117/1.JBO.19.5.056010]

Keywords: second harmonic generation; sarcomere length; autocorrelation; fiber orientation.

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1 Introduction

Sarcomeres are the elementary contractile units in cardiac and skeletal muscles and are composed of three basic structural elements as shown in Fig. 1. Two transverse structures: the Z-discs, anchoring the (actin) thin filaments and the M-band that crosslinks the (myosin) thick filaments in the center. The elastic thin filaments connect both structures in the longitudinal plane. The interaction of sarcomere thin and thick filaments generates the force necessary for contractile function. The expression of sarcomeric proteins is well documented to respond to physiological activity or disease, making the sarcomere a prime candidate for a visible indicator of myofiber integrity.1-3 The study of sarcomere morphometry provides information about the sarcomere’s structure and function and thus about cardiac contractility.4 For this, end-diastolic sarcomere length (SL) has been widely used as a feature of sarcomere morphometry;5,6 however, reliable and automated techniques for measuring these sarcomeric features in living intact tissue are limited.

The second harmonic generation (SHG) microscopy has shown an enormous potential for providing information with high resolution and contrast of certain unstained structures, such as collagen, myosin, microtubulin, starch, and cellulose, which are of interest for many biomedical applications.7-13 This unique contrast mechanism combined with recent technological advances in multiphoton microendoscopes/fibers14-16 and ultrashort pulse laser sources17 suggests that SHG might find clinical applications as an optical biopsy tool.9 SHG has been studied in many applications related with collagen,10,18,19 as it is a highly efficient frequency doubling biomaterial that can be detected by standard multiphoton microscopes. Myosin, a molecule present in all types of muscle, shows a smaller harmonic signal9,10,20 and has been studied less.

The SHG signal along a muscle fiber presents a bipериodical pattern4,17 as shown in Figs. 1(d) and 1(e); the two periods are associated with characteristic sarcomeric lengths that provide information of the ultrastructure of the cardiac tissue.3,21,22 Particularly, the shortest period corresponds to the A-band length (ABL) and the longest period to SL (distance between Z-discs). Recently, the Fourier transform (FT) has been used to compute the periodicity and the preferred orientation from SHG in biological tissues,18,19 and it was demonstrated that image-processing techniques can extract quantitative information from this imaging modality. However, the generalization of this approach is limited, and there is a need to develop more automated and accurate methodologies. Recent papers have shown that the pattern analysis of striated muscle correlates with physiological functions, and this might be useful for developing tools that assist diagnosis and possibly to monitor chronic diseases.23-25 However, the pioneering work of Plotnikov et al. was related to abstract image features, providing no supportive information to clinicians and biomedical researchers. More recently, these image features have evolved toward describing more comprehensive concepts of biology that allow a multidisciplinary exchange of ideas among experts. Our prior research reported that the automated quantification of the average fiber orientation (FO), SL, and ABL in cardiac tissue was more reliably computed by means of a method based on the analysis of the autocorrelation rather than a Fourier approach.24 Another

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recent paper, focused on skeletal muscle tissue, developed a technique to quantify local disruption of the regular sarcomere lattice (verniers) that was associated with diseased muscle.23 These papers show that the SHG is, indeed, a promising stain-free image modality for assessing muscle and cardiac tissues because it is highly specific for myosin thick filaments and targets one of the major factors of the contractile machinery. Under some circumstances, SHG from collagen can also be found (i.e., in fibrotic or adult hearts) but this can be filtered out by different means, like postprocessing in combination with optical methods10,26 by taking advantage of the differences from the SHG generated by these two structures. Apart from that, there are no other major interference effects when detecting SHG signals from myosin in most tissues.

This paper extends the prior study of our model to understand the use and the limitations of the proposed algorithm and set some of the imaging conditions required to achieve a given level of accuracy. Our goal is to evaluate the robustness of our algorithm for measuring morphometric features of cardiac tissue in SHG images. For this, we first performed numerical simulations to evaluate the performance of our methodology in one- and two-dimensional computer-generated muscle fibers, by adding different artifacts that can influence the measurements, such as noise, depth of modulation, and fiber bending and twisting. Second, automatic measurements of sarcomere ultrastructure and fiber orientation of cardiac tissue samples were compared with manual measurements. Finally, we applied our algorithm to analyze the morphometric features of cardiac tissue in a set of SHG images from fetal and adult rabbit hearts, acquired at different signal-to-noise ratios (SNRs) to evaluate the practical applications of our study in biomedical research.

2 Materials and Methods

2.1 Image Processing

The proposed image processing algorithm is explained elsewhere in detail.24 In summary, it consists of: (1) estimation of local fiber orientation, (2) tracking of muscle fibers, (3) muscle fiber profiling, (4) correlation of muscle fibers SHG signal, and (5) model fitting and parameter extraction. Briefly, local fiber orientation is estimated by computing the local autocorrelation of small blocks of pixels, followed by direct binarization of the autocorrelation and principal component analysis. The major axis orientation of the muscle fiber is computed as the first principal component. Once the local fiber angle is estimated, the muscle fiber tracking is performed. After obtaining the intensity profile for every tracked muscle fiber, the autocorrelation is calculated and fitted with a parametric model. After the model fitting, the mean of the ultrastructure sarcomere measurements within a muscle fiber is obtained for each fiber in the image. Therefore, distributions of FO, SL, ABL, and thick–thin interaction length (TTIL) for a SHG image of cardiac tissue were the quantitative descriptors obtained as output parameters of our algorithm. Finally, the mean ± standard deviation (SD) of the four features were calculated from the distributions, rejecting all the values outside the range ± 2.7 · std, since they were considered as outliers.

2.2 Simulation of Computer-Generated Muscle Fibers

The SHG intensity signal from a sarcomere can be modeled by two Gaussian functions.5,24 In this manner, the whole cardiac muscle fiber \( f(z) \) can be defined as

\[
 f(z) = \frac{1}{\sigma \sqrt{2\pi}} e^{-\frac{(z-z_0)^2}{2\sigma^2}}
\]
\[ f(z) = \sum_{n=1}^{M} \alpha_k \exp \left[ -\left( \frac{z - T_1/2 - nT_2}{\sqrt{2\sigma_k}} \right)^2 \right] + \alpha_k \exp \left[ -\left( \frac{z + T_1/2 - nT_2}{\sqrt{2\sigma_k}} \right)^2 \right], \]

where \( T_1 \) corresponds to the ABL, \( T_2 \) to the SL, and \( \sigma_k \) to the TTIL. \( \alpha_k \) accounts for the illumination variation when imaging the \( k \)th sarcomere [see Fig. 1(c)]. \( M \) denotes the number of sarcomeres in the muscle fiber.

Artificial cardiac muscle fibers were numerically generated, according to Eq. (1), varying the different ultrastructural morphometric features: (1) TTIL, (2) ABL, and (3) SL, in one dimension (1-D). However, for this calculation skips many steps of our methodology that add error. For this, we included the two-dimensional (2-D) generation of muscle fibers, as shown in Fig. 2(b). Similarly to 1-D, the 2-D SHG signal from a sarcomere can be modeled by two 2-D Gaussians. Therefore, the whole 2-D muscle fiber \( f(x,y) \) can be defined as

\[
f(x,y) = \sum_{n=1}^{M} \alpha_k \exp \left[ -\left( \frac{y - T_1/2 - nT_2}{\sqrt{2\sigma_k}} \right)^2 \right] - \left( \frac{x}{\sqrt{2\sigma_k}} \right)^2 \] + \alpha_k \exp \left[ -\left( \frac{y + T_1/2 - nT_2}{\sqrt{2\sigma_k}} \right)^2 \right] - (x/\sqrt{2\sigma_k})^2,
\]

where the \( y \) and \( x \) axes are aligned to the long and short axes of the muscle fiber, respectively [see Fig. 2(b)]. \( \sigma_{x,y} \) corresponds to the TTIL and \( \sigma_{x,y} \) to the thickness of the muscle fiber, assumed to be \( 2.5 \cdot \sigma_{x,y} \), according to the estimated fiber thickness measured manually in a SHG image from cardiac tissue Fig 5(a).

Also, we considered the addition of white Gaussian noise (WGN) in both 1-D and 2-D computer-generated muscle fibers, as shown in Figs. 2(a) and 2(b). Besides noise, many illumination changes occur during acquisition as a consequence of many factors like nonuniform illumination and tissue bending in and out of the focal plane. This has been modeled as a slowly varying image that modulates the image, as shown in Figs. 2(c) and 2(d). The proposed methodology is finally challenged by simulating the natural fiber bending within the 2-D plane (or fiber twisting) as shown in Fig. 2(e). Therefore, the artifacts that were modeled are: (1) addition of WGN of zero mean and a variance \( \sigma^2_{x,y} \) in 1-D and 2-D muscle fibers, (2) amplitude modulation of the SHG signal and 3-D fiber bending for 2-D data, and (3) fiber twisting: in plane fiber bending for 2-D data. The WGN power was varied to estimate an estimated SNR ranging from 0 to 50 dB. Amplitude modulation was simulated by generating a modulating image \( (I_{mod}) \) composed of two different cosines of unitary frequency \( f_x \) and \( f_y = 1 \) and two cosines of frequencies \( 2f_x-f_y \) and \( 2f_y-f_x \), with random amplitude and phase. With this combination of cosines, a random slow intensity variation modulating image, with a longitudinal component in the same direction as the muscle fibers, was generated. An example of a modulating image is displayed in Fig. 2(c). The modulated image \( (I_m) \) was obtained as \( I_m = I \cdot (1 + (I_{mod} - 0.5) \cdot m_x) \), where \( m_x \) denotes the depth of the modulation factor, which varies from 0 to 1. Fiber twisting was modeled by varying the local orientation of each sarcomere within the muscle fiber randomly, but limiting this angular change to a maximum of 15 deg. The maximum angular change within a fiber was chosen after estimating the maximum local angle variation in SHG images from cardiac tissue [see Figs. 5(a) and 5(b)]. Muscle fibers of five sarcomeres were considered in all the simulations.

The different simulated scenarios were: (1) 1-D and WGN with SNR from 0 to 50 dB; (2) 2-D and WGN with SNR from 0 to 50 dB; (3) 2-D, WGN with SNR from 0 to 50 dB, and depth of modulation with \( m_x = 0.2, 0.4, 0.6, \) and 0.8; (4) 2-D, WGN with SNR from 0 to 50 dB, fiber twisting and depth of modulation with \( m_x = 0.2, 0.4, 0.6, \) and 0.8. In total, 100 computer-generated muscle fibers with different morphometric dimensions \( (T_1, T_2, \sigma) \) were generated for each simulated scenario. The performance of the method was evaluated by computing the success rate, calculated as the ratio between the number of computer-generated fibers measured successfully, the total number of computer-generated muscle fibers, and the relationship between the experimentally estimated SNR of the signal and the ground truth using the root-mean-square error (RMSE) of the \( T_1, T_2, \) and \( \sigma \) values from the model of muscle fiber based on Eqs. (1) and (2). The SNR was estimated as \( SNR = E(I_w')^2 / E((I_s - I_w')^2) \) where \( I_w \) corresponds to the Wiener filtered image, with a window’s size of five pixels, and \( I_s \) corresponds to the noisy image \( (I_s = I + \text{WGN}) \).

2.3 Analysis of Biological Samples

Fetal and adult hearts from New Zealand white rabbits were analyzed to illustrate the practical implications of our study in biomedical research. Animals were provided by a certified animal source.
breeder. Animal handling and all procedures were performed in accordance to applicable regulations and guidelines and with the approval of the Animal Experimental Ethics Committee of the University of Barcelona. Hearts from these animals were obtained one day prior to birth (fetal) and seventy postnatal days (adult) following previously reported protocols. Briefly, hearts were fixed by immersion in 4% paraformaldehyde in phosphate buffered saline, for 24 h at 4°C. The tissue was then dehydrated and embedded in paraffin, and 30-μm thick transversal heart sections were cut in a microtome. Finally, sections were mounted onto silane (Sigma-Aldrich) coated thin slides. Tissue sections were deparaffinized with xylene and decreasing concentrations of ethanol (100°/96°/70°) and finally covered with Mowiol 4–88 mounting medium (Sigma-Aldrich).

A Leica TCS-SP5 laser scanning spectral confocal multiphoton microscope (Leica Microsystems Heidelberg GmbH, Manheim, Germany) equipped with a near-infrared laser (Mai Tai Broad Band 710–990 nm, 120 femtosecond pulse) and a DMI6000 inverted microscope from the Advanced Optical Microscopy Unit from Scientific and Technological Centres from the University of Barcelona was used to acquire SHG images from unlabeled heart tissues. A 25× water immersion objective (HCX IR APO L), with a numerical aperture (NA) of 0.95 and an oil immersion condenser (NA 1.40 OilS1), both from Leica Microsystems were used to acquire the SHG images. The images were sampled at a pixel size of 40 nm.

Samples were identified to contain mostly cardiomyocytes and no collagen. Cardiac longitudinal fibers were oriented at 45° to maximize SNR of SHG from muscle fibers. The image size was 2048 × 2048 pixels.

First, manual measurements of sarcomere morphometry and local muscle fiber orientations were performed in 30 different regions within the SHG image of cardiac tissue acquired at high SNR. Moreover, the automatic measurement of the muscle fiber morphometry was performed in the same image, and both manual and automatic measurements were compared by calculating the mean ± SD. Finally, we evaluated the automatic measurement of muscle fiber morphometric features (SL, ABL, TTIL, and FO) in cardiac tissue, in two sets of 10 images each, acquired at two different SNRs by calculating for each image the mean ± SD of each of the four features.

## 3 Results

### 3.1 Robustness Analysis with Numerical Simulations

The computed success rate in retrieving measurements from computer-generated 1-D muscle fibers is above 90% at 15 dB and RMSE of SL, ABL, and TTIL is below one pixel for all three measurements at SNR of 7 dB [see Figs. 3(a) and 3(b)]. In the case of computer-generated 2-D muscle fibers, the computed success rate is above 90% also at 15 dB. Additionally, the computed RMSE of SL, ABL, and TTIL is below one pixel for all three measurements at a SNR of 15 dB [see Figs. 3(c) and 3(d)].

Results with a depth of modulation of 40% show already an effect at SNR below 15 dBs, and in this context the RMSE is almost doubled [see Figs. 3(e) and 3(f)]. However, at a SNR of 15 dBs the RMSE is still below one pixel. Higher depth of modulation shows increased errors but beyond 17 dB the RMSE is always maintained below 1 pixel and the success rate is above 90%.

The results of the natural fiber bending within the 2-D plane simulation in combination with all the prior artifacts show that, with a depth of modulation of 40%, a SNR of 20 dB should produce an error below one pixel [see Figs. 3(g) and 3(h)]. Finally, in order to illustrate the effect of WGN together with the other artifacts on the average and variability of the
Fig. 4 Real values of the SL of the simulated computer-generated muscle fibers plotted against the corresponding estimated values, and for the four different simulated scenarios and at three different SNRs; (a)–(c): 1-D at (a) SNR = 12 dB, (b) 17 dB and (c) 20 dB; (d)–(f): 2-D at (d) SNR = 12 dB, (e) 17 dB and (f) 20 dB; (g)–(i): 2-D + depth of modulation (DOM) of 40% at (g) SNR = 12 dB, (h) 17 dB and (i) 20 dB; and (j)–(l): 2-D + fiber twisting + DOM of 40% at (j) SNR = 12 dB, (k) 17 dB and (l) 20 dB. CI denotes confidence interval. The blue line corresponds to the linear fitting with the function $y = ax + b$. 

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measurements, real SLs were plotted against the corresponding estimated value in Fig. 4, at three different SNRs: 12, 17, and 20 dB and for each of the four simulated scenarios: 1-D, 2-D, 2-D with a depth of modulation of 40% and 2-D with twisted fibers and a depth of modulation of 40%.

3.2 Manual Versus Automatic Quantification of Muscle Fibers

The automatic measurement of sarcomere ultrastructure shows excellent agreement with 30 manual delineations in different regions of the image shown in Fig. 5(a), and it offers a similar uncertainty as shown in Table 1. The absolute difference between manual and automated measurements was less than one pixel for two of the morphometric features (ABL and SL). TTIL was not measurable manually. The measurement of local fiber orientation also shows excellent agreement with the manual delineations of different image regions as shown in Table 1. An example of a local fiber orientation estimation is shown in Figs. 5(b), 5(c) and 5(d), which also illustrates the good performance of the algorithm in computing the local fiber angle. Finally, the distributions of FO, SL, ABL, and TTIL for the same SHG cardiac tissue sample shown in Fig. 5(a) are plotted in Fig. 6 to illustrate the output of our algorithm.

Table 1 Comparison of muscle fibers morphometric measurements (mean ± standard deviation) calculated with our algorithm against manual measurements in thirty different regions within the same image.

<table>
<thead>
<tr>
<th>Morphometric feature</th>
<th>Manual</th>
<th>Automated</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-band length ($T_1$) (pixels)</td>
<td>20.57 ± 2.97</td>
<td>21.39 ± 1.63</td>
</tr>
<tr>
<td>Sarcomere length ($T_2$) (pixels)</td>
<td>45.51 ± 2.67</td>
<td>44.60 ± 2.66</td>
</tr>
<tr>
<td>Thick-thin interaction length (a) (pixels)</td>
<td>,</td>
<td>2.66 ± 0.37</td>
</tr>
<tr>
<td>Fiber orientation (degrees)</td>
<td>38.92 ± 9.22</td>
<td>36.46 ± 5.57</td>
</tr>
</tbody>
</table>

*Thick-thin interaction length was not measurable manually.

3.3 Quantification of Sarcomeres at Different Signal-to-Noise Ratios and Status

We show, in this section, the results comparing images from an animal model at two different SNRs. SNR was estimated after the image acquisition. The estimated SNR of the first set of images corresponding to young adult rabbit cardiac tissue was 23.21 ± 0.95 dB. The estimated SNR of the second set of images corresponding to fetal rabbit cardiac tissue was 16.72 ± 0.21 dB. The mean and SD of the four morphometric features measured automatically are shown in Table 2. The estimated depth of modulation for the set of images from fetal rabbit hearts was 0.88 and 0.59 for the set of images from young adult rabbit hearts. Two representative images from both sets of SHG images are displayed in Fig. 7. It can be appreciated that the fetal cardiac muscle fibers are more twisted compared to the young adults. Also, it can be observed the higher depth of modulation in the image corresponding to fetal cardiac tissue. The higher influence of both artifacts and also the lower SNR in the set of fetal cardiac tissue images account for the higher variability obtained in the measurements of the muscle fibers morphometry.

4 Discussion

In the present paper, we have shown the good performance of our methodology to assess morphometric features of cardiac fibers in SHG images of cardiac tissue.

Many times, sample preparation, storage, microscope alignment, maintenance, and other issues related to image acquisition limit the quality of the tissue and the delivered images for the analysis. For this reason, we analyzed the performance of our method by including numerical simulations in 1-D and 2-D computer-generated muscle fibers. We modeled different artifacts that are present during the image acquisition process, including noise, depth of modulation, fiber bending, or twisting. The best performance of the system is found at 20 dB as it obtains a high success rate (above 90%) and high accuracy (RMSE below one pixel of 40 nm).

The manual validation performed on a subset of images enables us to ensure the good performance of our method at high SNR values (above 20 dB). In this context, we propose our methodology to extract quantitative image features from SHG microscopy to characterize cardiac tissue in the form of distributions of FO, SL, ABL, and TTIL as shown in Fig. 6. These
features are highly reproducible because they are related to ultrastructural morphology of the tissue and are not purely based on local signal ratios. The average value of FO is certainly arbitrary, but its distribution or standard deviation is related to fiber disruption, and this is related to muscle atrophy. End-diastolic SL is widely used as a feature of sarcomere morphometry, however, reliable and automated techniques for measuring these sarcomeric distances in living intact tissue are limited. Similarly, ABL and TTIL should also provide valuable information about the cardiac status, but they are submicron structures, which are highly difficult and tedious measurements to be performed manually, and thus there are no studies using them. Systematic analysis with our methodology provides an objective manner to quantify numerous unstained histological sections of cardiac tissue (or muscle tissue) that is not available by other means. The representation of the cardiac tissue in the form of four distributions offers a way to compare many samples and look for statistical differences within large datasets. Furthermore, statistical means can be used to reject spurious results that might occur at the tails of the distributions.

The condition of a minimum SNR of 20 dB is very strict and is recommended when studying small groups of subjects to obtain reliable conclusions. Automated analysis of real images of animals at different SNRs provided evidence that for a particular experiment, with similar conditions, concerns on the SNR can be reduced, as the averaging among many samples will converge to the right measurement. Also, the variability obtained in the automated analysis of both sets of real images is consistent with the expected variability obtained in the numerical simulations of computer-generated fibers. For example, according to the simulation results, the 95% confidence interval (CI) for the SL measurement in twisted fibers with a depth of modulation of 60% and with a SNR of 20 dB is ±3.68 pixels (data not shown), and the SD for the SL measurements in real SHG images from young adult cardiac tissue was ±1.40 pixels. On the other hand, the 95% CI for the SL measurement in twisted fibers with a depth of modulation of 80% and with a SNR of 17 dB is ±5.68 pixels (data not shown), and the SD for the SL measurements in real images from fetal rabbit cardiac tissue was ±5.82 pixels, which means a 95% CI of ±3.61 pixels. These results reflect the good performance of our algorithm in measuring the morphology of cardiac muscle fibers in SHG images and also how the numerical simulations can help us to improve the design of our experiments and also the posterior analysis by addressing some types of corrections like image exclusion, bias correction, or image re-acquisition when possible. Developments in this direction aim to bring together recent advances in biophotonics to clinical applications as this makes it possible to reproduce experiments objectively, many times and in different locations, as well as setting the new advances to develop robust image processing tools that extract relevant information for biomedical researchers and clinically driven applications. Additionally, our methodology was used to evaluate sarcomere ultrastructural differences in a control/case study of human fetuses, and reporting shorter cardiac sarcomere length in fetuses with intraterine growth restriction.

To conclude, our approach based on automated autocorrelation analysis shows similar results to manual delineations and is a robust technique providing a sufficient SNR in the images. We have evaluated the impact of noise, amplitude modulation, and fiber twisting, and minimal errors are found for images beyond 20 dB, which is a realistic amount for automated histology analysis. We further supported our numerical results of computer-generated data with real experiments based on two different acquisition conditions. Finally, the results showed that our technique for measuring the sarcomere distances is accurate and

### Table 2 Results of the automatic measurement of muscle fibers morphometry in two sets of ten images acquired at different SNRs. For each image, the mean ± standard deviation (SD) of the four morphometric features and of the SNR was computed, and it is reported here the average of the mean and of the SD for all the images.

<table>
<thead>
<tr>
<th>SNR (dB)</th>
<th>A-band length (pixels)</th>
<th>Sarcomere length (pixels)</th>
<th>Thick-thin interaction length (pixels)</th>
<th>Fiber angle (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mean</td>
<td>SD</td>
<td>mean</td>
<td>SD</td>
</tr>
<tr>
<td>23.21 ± 0.95</td>
<td>22.53</td>
<td>1.26</td>
<td>46.34</td>
<td>1.40</td>
</tr>
<tr>
<td>16.72 ± 1.66</td>
<td>16.20</td>
<td>3.29</td>
<td>34.38</td>
<td>5.82</td>
</tr>
</tbody>
</table>
allows us to reproduce changes previously described, concerning changes in the SL under pathological conditions.29–31

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Biographies of the authors are not available.