Histogram-based standardization of intravascular optical coherence tomography images acquired from different imaging systems

Shengnan Liu, Oleh Dzyubachyk, and Jeroen Eggermont

Division of Imaging Processing, Department of Radiology, Leiden University Medical Center, Leiden, 2300 RC, The Netherlands

Shimpei Nakatani

Division of Cardiology, Sakurabashi Watanabe Hospital, Osaka 530-0001, Japan

Boudewijn P. F. Lelieveldt

Division of Imaging Processing, Department of Radiology, Leiden University Medical Center, Leiden, 2300 RC, The Netherlands Intelligent Systems Department, Delft University of Technology, Delft, 2628 CD, The Netherlands

Jouke Dijkstra^{a)}

Division of Imaging Processing, Department of Radiology, Leiden University Medical Center, Leiden, 2300 RC, The Netherlands

(Received 4 December 2017; revised 11 June 2018; accepted for publication 5 July 2018; published 14 August 2018)

Purpose: Intravascular optical coherence tomography (OCT) is widely used for analysis of the coronary artery disease. Its high spatial resolution allows for visualization of arterial tissue components in detail. There are different OCT systems on the market, each of which produces data characterized by its own intensity range and distribution. These differences should be taken into account for the development of image processing algorithms. In order to overcome this difference in the intensity range and distribution, we developed a framework for matching intensities based on the exact histogram matching technique.

Methods: In our method, the key step for using the exact histogram matching is to determine the target histogram. For this, we proposed two schemes: a global scheme that uses a single histogram as the target histogram for all the pullbacks, and a local scheme that selects for each single image a target histogram from a predefined database. These two schemes are compared on a unique dataset containing pairs of pullbacks that were acquired shortly after each other with systems from two vendors, St. Jude and Terumo. Pullbacks were aligned according to anatomical landmarks, and a database of matched histogram pairs was created. A leave-one-out cross validation was used to compare performance of the two schemes. The matching accuracy was evaluated by comparing: (a) histograms using Euclidean (d_{x2}) and Kolmogorov–Smirnov (d_{KS}) distances, and (b) median intensity level within anatomical regions of interest.

Results: Leave-one-out validation indicated that both matching schemes yield comparably high accuracies across the entire validation dataset. The local scheme outperforms the global scheme with marginally lower dissimilarities at both histogram level and intensity level. High visual similarity was observed when comparing the matched images to their aligned counterparts.

Conclusion: Both local and global schemes are robust and produce accurate intensity matching. While local scheme performs marginally better than the global scheme, it requires a predefined histogram dataset and is more time consuming. Thus, for offline standardization of the images, the local scheme should be preferred for being more accurate. For online standardization or when another system is involved, the global scheme can be used as a simple and nearly-as-accurate alternative. © 2018 The Authors. Medical Physics published by Wiley Periodicals, Inc. on behalf of American Association of Physicists in Medicine. [https://doi.org/10.1002/mp.13103]

Key words: histogram specification, image intensity, intensity standardization, intravascular optical coherence tomography (IVOCT)

1. INTRODUCTION

Cardiovascular diseases (CVDs) are the leading cause of death worldwide. Introduction of intravascular optical coherence tomography (IVOCT) has largely advanced understanding and treatment of one of the most common CVDs, the coronary artery disease. Design of IVOCT enables visualization of superficial tissue structures of the arteries with resolution as high as 5–10 μm. The wavelength of its light

source is around 1300 nm, which permits a relatively deep penetration into the vessel wall. The *intravascular* term indicates that the images are acquired from the inside of the blood vessel. For current commercial systems, this is achieved by inserting a catheter into the coronary artery, pushing away the blood by injecting a flush media and pulling it back through the lesion location. The catheter has been designed to emit near-infrared light towards the artery wall and to receive the back-propagated light. The received light is recorded as a

one-dimensional signal (A-line) containing the back-propagated intensities ordered by ascending distances to the catheter. By rotating the catheter tip with a constant speed, two-dimensional (2D) images can be generated. Centering at the catheter, each cross-sectional image contains about 500 A-lines from different directions. These A-lines are recorded as a raw image in polar coordinates, as shown in Figs. 1(b) and 1(e), or transformed into the Cartesian coordinates, as shown in Figs. 1(c) and 1(f). As the catheter is pulled back at a constant speed using a motorized pullback device, a stack of images, referred to as a *pullback* is acquired.

The use of IVOCT in clinical studies increases exponentially.² Because of its high resolution, IVOCT contributed to confirmation of pathological findings on progression of (neo) atherosclerosis by visualizing morphologies like intimal erosion, fibrous plaque, calcified nodule, lipid pool, macrophages distribution, intraluminal thrombus, etc.^{6–12} Attracted by the conspicuous clinical prospects, many efforts were paid to detection and characterization of IVOCT morphologies, such as fibrous, lipid-rich and calcified plaques,¹³ macrophages distributions,⁷ thrombus,¹⁴ side branches,^{15,16} struts^{17–19}, and struts embedding²⁰ with image intensities, and/or optical parameters.^{21,22} However, diversity of IVOCT data can limit comparison of the results, especially when intensity values are used. In particular, there is no consented

standard for the imaging range, unlike, for example, in computed tomography (CT), meaning that IVOCT images generated with different commercially available systems are typically characterized by different intensity ranges. The most commonly used commercial systems are the Illumien Optis from St. Jude Medical (St. Paul, MN, USA), which saves the raw data in a 16-bit format, and the Lunawave from Terumo (Tokyo, Japan), which saves the raw data in a 8-bit format.

As a concrete example, the Cartesian images from these two systems are shown in Figs. 1(c) and 1(f), and their polar counterparts are shown in Figs. 1(b) and 1(e). Images were acquired shortly after each other during the same intervention at the same location inside an artery of one patient. The histograms of the corresponding regions on the IVOCT images acquired by the two systems indicate different intensity ranges within the same tissue type. Furthermore, the different shape of these histograms suggests that relationship of intensities between these two systems is not simply linear. In fact, an exponential relationship has been observed in our previous work.²³

Most OCT studies used the same type of IVOCT system to guarantee high reproducibility. On the other hand, doing so limits the scope of developed applications, when the same method is applied to data from another vendor that has different intensity distribution, repeated validation is required. To improve efficiency of development, images need to be

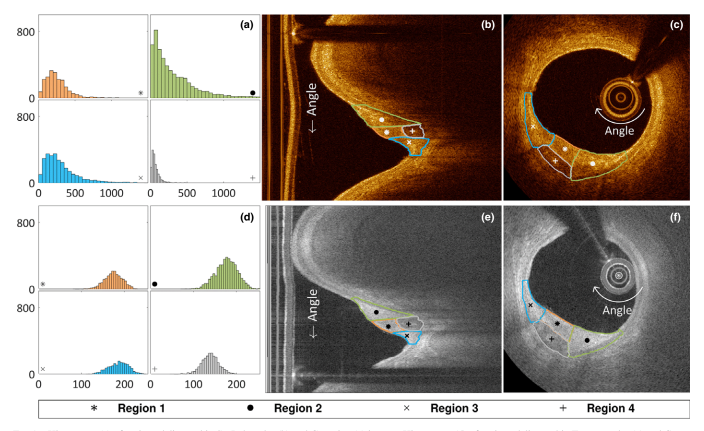


Fig. 1. Histograms (a) of regions delineated in St. Jude polar (b) and Cartesian (c) images. Histograms (d) of regions delineated in Terumo polar (e) and Cartesian (f) images. Side branches and lesions were used as landmarks to find the corresponding locations in St. Jude and Terumo images. Regions were separated according to their morphological appearance and visible borders and are correspondingly delineated in the polar images. [Color figure can be viewed at wileyon linelibrary.com]

standardized across devices. Only few papers on this topic have been published in the IVOCT field, whereas several papers on increasing the comparability of ophthalmological OCT images were published. In particular, a normalization approach was proposed for comparing images from two vendors. This approach involves three steps: density scaling and sampling, noise reduction, and amplitude normalization. It was later improved by integrating virtual averaging. This A-line normalization approach was shown to reduce the measurement difference and the appearance disparity.

In this work, we explore the possibility of converting the data between different OCT systems and propose a matching scheme with good generalization and minimal loss of detail. Our pretrained algorithm can also be used for intensity matching when the target data are not given. By doing this, when a method developed for data acquired with system-A needs to be evaluated with data acquired with system-B, we can modify the data from system-B to follow the intensity distribution of system-A, such that the method can be tested on the data from system-B with minimum modification. Such data conversion is referred to as the histogram modification. 28-31 The basic histogram matching theory has been proposed in the work of Hummel et al.²⁸ Since then, this study has been widely used as the fundamental theory in image modification studies at the histogram level. Later, the exact histogram specification (EHS) was proposed as a successful discrete solution to the model in practice.^{29–31} This approach was used to produce comparable measurements in ophthalmological OCT images generated with low signal strength to that generated with high signal strength³² and to compensate light attenuation in confocal microscopy.³³

Our main goal in this paper is to propose a framework for matching intensities in OCT images from different vendors using EHS. A straightforward approach would be to match intensities per pullback. We compare this *global* scheme to a *local* scheme that takes the local intensity variations into the consideration. All the analyses are conducted with raw polar images, whereas the Cartesian images are only used for the visualization of results. The *in vivo* patient data used in this study are unique in the sense that both St. Jude and Terumo pullbacks were specially acquired for this study. A more elaborate explanation of this is provided further in the manuscript.

The paper is structured as follows. For better understanding of the underlying principles and terminology, we explain both the model and the EHS in Section 2. In Section 3, the data and its processing are described, the distance measures are introduced, and the matching schemes are proposed and validated. Results are reported and discussed in Sections 4 to 6. Finally, in Section 7 we draw the conclusions.

2. THEORY AND TERMINOLOGY

Each 2D image can be represented as a matrix I(x,y), which is the discrete subsample of a bounded surface function f(x,y), where $\{(x,y)|0 \le x \le N, 0 \le y \le M\}$. The intensity function f(x,y) follows a distribution function $P_f(t)$ that

indicates the chance of f(x,y) being less or equal than t. Given two images $f_s \sim P_{f_s}(z_s)$ and $f_t \sim P_{f_t}(z_t)$, the goal of histogram transformation is to search for a mapping T, such that the composition is $T \circ f_s \sim P_{f_t}(z_t)$. In this work, f_s and f_t are referred to as the source and the target image, respectively. This search has been formulated by Hummel et al.²⁸ as an optimization problem:

$$\hat{T} = \arg \min_{T} \int_{a_{1}}^{a_{2}} \left[p_{T \circ f_{s}}(z) - p_{f_{t}}(z) \right]^{2} dz, \tag{1}$$

where $[a_1, a_2]$ defines the range of image intensities. Here, $p_X(z)$ is the probability density function (pdf) of image X, which is the derivative of the distribution function $P_X(z)$, and z is intensity level.

A unique monotonic solution of this model was given in Ref. [28]:

$$\tilde{T}(z_s) = P_f^{-1}(P_{f_s}(z_s)), p_{f_s}(z_s) > 0, \text{ for } \forall z_s.$$
 (2)

In practice, p_X in both source and target spaces is usually estimated as the normalized histogram vector: $p_X = \{p_i | \sum p_i = 1, i \in \{0, \dots, 2^{L_0} - 1\}\}$, where L_0 denotes the maximum gray level of the images, p_i denotes the frequency of the image intensity corresponding to the interval $[z_i, z_{i+1})$. Conventionally, the interval is referred to as a bin; $\{z_i\}$ are the bin edges; the average of every two adjacent bin edges $c_i = (z_i + z_{i+1})/2$ is the bin center, and p_i is the bin value. We define the bin edges for the i_{th} bin as $[c_i - 0.5, c_i + 0.5)$ in this work.

To ensure that the distribution function P_X is monotonically increasing, the following approximation is used:

$$P_{j} = \frac{c_{j} - z_{j}}{z_{j+1} - z_{j}} p_{j} + \sum_{i=0}^{j-1} p_{i}.$$
 (3)

This approximation is equivalent to interpolating $P_j(z)$ for $z \in [z_j, z_{j+1})$ with a piecewise-linear function, the slope of which is p_j and the intercept is $\sum_{i=0}^{j-1} p_i$. Using this monotonic approximation, \tilde{T} can be estimated with Eq. (2). However, this estimation only shifts bins centers, and splitting of bins is not possible. This becomes especially problematic when the source and the target images are within different intensity range, such as transforming 8-bit images to 16-bit images or the other way around. Since the bins cannot be split, information can only be retained based on the image that is represented by less bins. Attempts have been made to include local information²⁸ (local mean, entropy, etc.) into the objective function as a "context-aware" term, but doing so introduces more parameters, and the transformed images tend to be blurred.

The aforementioned issues can be overcome by a technique called exact histogram specification (EHS). $^{29-31}$ In the following sections, we will use the previously introduced notations. For an image of size $M \times N \times P$ with gray values in $\{0,1,\ldots,2^{L_0}-1\}$, the histogram divides $M \times N \times P$ pixels into 2^{L_0} bins. The fundamental idea behind EHS is to strictly order all the pixels, such that the ordered pixels can be

divided according to any given target histogram. This usually cannot be done by using intensities only, thus auxiliary information needs to be introduced.

Local means and wavelet coefficients have been reported to be successful source of auxiliary information for achieving strict ordering. In a more recent study, a moderately improved performance in nature images has been reported with a proposed auxiliary term involving three hyperparameters. Compared to the other two aforementioned approaches, the local-means approach better copes with noise and involves no hyperparameters, thus it is chosen in this study as the noise level of OCT images is known to be high.

Considering the intensity at each pixel as the first scale of the local mean, for each pixel i we calculate a vector of multiscale local means with increasing window size $\mu = [\mu_1(i), \mu_2(i), ..., \mu_K(i)]$, K being the number of scales. Consequently, all the pixels in the image can be ordered lexicographically with a relational operator defined as

$$\{i \prec j | \mu_{\ell}(i) = \mu_{\ell}(j)_{\forall \ell < \ell_0}, \mu_{\ell_0}(i) < \mu_{\ell_0}(j)\}.$$
 (4)

As result of using this approach, pixels in one image are expected to be ordered strictly by using just a few scales. It was observed that K = 6 scales are usually enough to arrange all the pixels in a strictly ascending order.³⁰ Once the strict ordering is achieved, the pixels can be easily grouped again following the histogram defined in the target image.

In practice, the EHS is often used to estimate the target image(s) given the source image(s) and the target histogram. In our case, however, the target histogram is unknown and should first be defined using a group of matched source and target images. Thus, our task is twofold: (a) To estimate the target histogram(s) from the matched images, and (b) To apply the estimated histogram(s) to a new source image. The following section describes our method in full detail.

3. MATERIALS AND METHODOLOGY

As it was previously elaborated, the key step for using EHS is to define the target histogram.

3.A. Global and local matching schemes

To determine the target histogram, the most straightforward approach is to use the overall histogram generated with all the images. However, using the global histogram as a reference might result in information loss as some of the less represented tissue structures might get overshadowed in the global histogram. Using local histograms might help resolving this issue.

Therefore, we introduce a *Global* scheme and a *Local* scheme for determination of the target histogram. For applying the Local scheme, each pullback is split into smaller sections (our pullback alignment and splitting algorithm is described in the following section). For providing a formal definition, we denote the database of matched histogram pairs as $\mathcal{H} = \{(H_1^k, H_2^k) | k = \overline{1, N_H}\}$, where H_1^k denotes the

histogram of kth section in the original space (e.g., St. Jude), H_2^k denotes its counterpart in the target space (e.g., Terumo), and N_H is the size of the database.

For matching a given image with histogram H_1^* , the Global scheme uses one overall histogram as the target histogram, that is,

$$H_2^G = \sum_{k=1}^{N_H} H_2^k. (5)$$

The Local scheme determines the target histogram for each section as the second component of the database entry whose first component is the most similar to H_1^* :

$$H_2^L = H_2^{\operatorname{argmin}_{k = \overline{1N_H}} d(H_1^*, H_1^k)}.$$
 (6)

Here, $d(H_i, H_j)$ denotes dissimilarity between two histograms H_i and H_j . The remainder of this section describes creation of the histogram database and introduces the dissimilarity measures.

3.B. Data description and alignment

In our case, the target histogram is estimated using a dataset of matched images. As the final goal of this study is to transform OCT images both from intensity space of St. Jude system (St. Jude space) to that of Terumo system (Terumo space) and the other way around, the target histograms in both St. Jude and Terumo spaces need to be determined. To achieve this, eight in vivo pullbacks were acquired from left anterior descending artery (LAD) of two different patients, marked as A and B. The data acquisition strictly followed the clinical guideline of Sakurabashi Watanabe Hospital (Osaka, Japan), and the analysts from Leiden University Medical Center (Leiden, The Netherlands) were blinded from all patients' information. For each patient, IVOCT pullbacks from the same vessel segment were acquired shortly after each other with St. Jude and Terumo systems, before and after the stent implanting procedure. As a result, four pairs of corresponding St. Jude-Terumo pullbacks were made available for the study. The St. Jude pullbacks were acquired at a pullback speed of 36 mm in 180 frames per second with a frame interval of 0.2 mm, and the Terumo pullbacks were acquired at a pullback speed of 40 mm in 158 frames per second with a frame interval of 0.25 mm. Raw polar images as provided by the vendors are used in this study. St. Jude images are 16-bit "linear" with a transversal pixel size of 0.0050 mm, and Terumo images are 8-bit "log-like" compressed with a transversal pixel size of 0.0049 mm.

Due to different distal and proximal locations in two corresponding pullbacks, images within overlapping part of the vessel should be matched. We determine the start and the end frames of overlapping part by searching for identical side branches that are closest to the proximal and the distal parts of the pullbacks. Since we primarily aim at analyzing the tissue region, images with stent struts points are excluded (more elaborate discussion about this is provided in Section 6).

Compared to the histogram of the entire overlapping part of pullbacks, histograms of smaller sections may preserve more local structural information. Therefore, we further split the overlapping part into small sections using key frames defined by two main types of anatomical landmarks (as shown in Fig. 2): the side branches and the tissue types with identical patterns, for exmaple, culprit lesion visible in Fig. 1(c) and 1(f). For the poststenting pullbacks, the proximal and distal edges of the stent struts are also considered to be crucial landmarks. Key frames were identified independently by two experienced IVOCT readers, using QCU-CMS (Quantitative Coronary Ultrasound—Clinical Measurement Systems; Leiden University Medical Center, Leiden, The Netherlands), which is the research version of QIVUS (Quantitative IntraVascular UltraSound; Medis, Leiden, The Netherlands). Only landmarks with consensus were selected.

Even though the catheter is pulled back with a uniform speed, the number of frames within a certain section can be affected by the heartbeat cycle, slight bending of the artery, interaction between the catheter and the artery wall, and so on. Due to this effect, finer splitting or frame-to-frame matching is not possible without introducing an interpolation error.

Table I gives an overview of the aligned data. Patient-A had no stent planted beforehand, while Patient-B did have one. Hence, the number of frames without stent struts in Patient-B is smaller than in Patient-A. Overall, the pullbacks were aligned with 23 key frames. Within these frames, regions of clinical interest (ROI) were delineated based on their appearance and visible borders according to the consensus; see Fig. 1. In total, we have 18 corresponding pairs of sections and 38 ROIs. After correspondence between the sections had been established, the database of matched histogram pairs was generated using foreground regions bounded by the lumen border and 1 mm behind. This corresponds to the depth of 200 pixels in St. Jude polar image and

204 pixels in Terumo polar image. The lumen border is detected by the QCU-CMS software in an automated manner.

3.C. Dissimilarity between histograms

The Euclidean and the Kolmogorov–Smirnov distances are used for comparing two histograms. They are first used in the analysis of the local variations and, after that, for evaluation of matching schemes.

For calculating the distances, the histograms are normalized to obtain the probability density functions (pdf's) and the corresponding cumulative density functions (cdf's). The Euclidean distance for measuring dissimilarity between two pdf's p and q is defined as

$$d_{x2}(\mathbf{p}, \mathbf{q}) = (\mathbf{p} - \mathbf{q}) \cdot (\mathbf{p} - \mathbf{q})', \tag{7}$$

the discrete approximation of which is equivalent to the objective function in Eq. (1). The Kolmogorov–Smirnov distance measures dissimilarity between the distribution functions (also known as cdf's) P and Q:

$$d_{KS}(\boldsymbol{P}, \boldsymbol{Q}) = \max_{i} |P_i - Q_i|. \tag{8}$$

3.D. Validation

Leave-one-out validation is used to compare the Global and Local schemes. One of the four pullback pairs is left out in turn, and the target histogram is determined with the other three. Consequently, the intensities of the left-out pullback are matched. As part of each leave-one-out experiment, median intensities within each ROI are compared in both target and matched images. The target and matched median values are shown in the scatter plot and compared in the Bland–Altman plot. The scatter plot can show whether two groups of

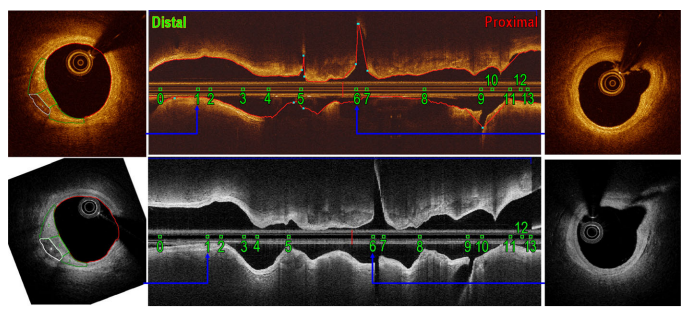


Fig. 2. The landmarks for aligning the pullbacks. [Color figure can be viewed at wileyonlinelibrary.com]

TABLE I. Data description.

Patient	Treatment	St. Jude					Terus	mo	I d	C4:	DOI	
		PL ^a (mm)	All ^b	Max ^b	Min ^b	PL ^a (mm)	All ^b	Max ^b	Min ^b	Landmark	Section	ROI
A	Prestenting	108	507	80	10	168	389	88	9	14	13	32
	Poststenting	108	69	52	17	147	53	41	12	4	2	4
В	Prestenting	75	40	21	19	152	33	19	14	3	2	1
	Poststenting	75	12	12	12	111	10	10	10	2	1	1
Total			628	80	10		485	88	9	23	18	38
ROI	Adventitia		Calcification		Fibrous		IML ^c	Lipid		Neointima		Total
No.	4		11		10		4	3		6		38

^aPL: total length of the entire pullback.

data are linearly related and how much the trend line deviates from the diagonal line, which is the ideal case indicating that two groups of data match perfectly. The Bland–Altman plot shows the average vs the difference of each two compared values, which reveals the difference of two groups of data more explicitly. The mean and the standard deviation values of the difference are also called the systematic difference and the random error. With a properly low random error, the closer the systematic difference to zero level is, the more likely that two groups of data are originating from the same distribution. For reference, also the key frames are matched using the target histograms extracted directly from their corresponding aligned frames.

4. RESULTS

We calculated the distances between all the histograms in two spaces. Figure 3 gives a general overview of all the dissimilarities. In the scatter plot of distances in Fig. 4, we observed a linear trend and further performed the regression analysis. The statistical results show that the dissimilarities in two spaces are correlated significantly with P < 0.005. This validates the assumption and supports the Local matching scheme.

Results of the leave-one-out validation for both St. Judeto-Terumo $(S \rightarrow T)$ and Te-ru-mo-to-St. Jude $(T \rightarrow S)$ images are reported in Figs. 5 and 6 and summarized in Table II. For a section containing 70 images, generating the histograms takes 10 s. For the Local scheme, the time for searching in our database of 34 histograms using our Matlab (MathWorks, R2016a with Statistics and Machine Learning Toolbox) implementation is 1.25 s. Figure 5 shows the estimated target probability profile (normalized histogram) together with that determined by the aligned data (in red color; will be further referred to as the aligned histogram). The corresponding cumulative probability plots are shown as well. Both d_{x2} and d_{KS} are shown as bar charts in Fig. 6, and the numbers are reported in Table II. The distances between matched and reference distributions for both Global and Local schemes are low and robust; see Fig. 6 and Table II. The average distance

of all four validation experiments (*Aver* in Table II) indicates that, in general, the Local scheme outperforms the Global scheme.

Scatter plots and Bland–Altman plots of the median intensities within ROIs are shown in Fig. 7. When comparing intensities in St. Jude space, an increasing trend is observed in the absolute systematic error; see Figs. 7(b), 7(e), 7(h), 7(k). Following the conventional statistical procedure for Bland–Altman analysis,³⁴ the St. Jude intensities were compared in the logarithm scale; see Figs. 7(c), 7(f), 7(i), 7(l). Without an obvious trend in Terumo space, the intensities are compared in the linear space; see Fig. 7(a), 7(d), 7(g), 7(j).

For the reference matching between the corresponding key frames, the systematic differences are -4.50 for S \rightarrow T and 0.15 for $T \rightarrow S$; see Figs. 7(a) and 7(c). This difference is the most likely to be caused by the selection of data and is independent of the scheme used. Using this intrinsic systematic difference as a reference, the absolute systematic differences yielded by the Local scheme are relatively small: $9.67 (S \rightarrow T)$ and 0.07 (T \rightarrow S) for d_{x2} , 7.92 (S \rightarrow T) and 0.06 (T \rightarrow S) for d_{KS} , compared to the Global scheme: 10.26 (S \rightarrow T), 0.12 $(T \rightarrow S)$. Taking into consideration the described intrinsic difference, the Local scheme yields absolute values closer to zero for both $S \rightarrow T$ and $T \rightarrow S$ matching. All comparisons in Bland-Altman analysis suggest that the Local scheme outperforms the Global scheme for EHS-based intensity transformation in IVOCT images between St. Jude and Terumo systems. The images matched with the target histogram determined by the Global scheme are shown in Fig. 8. The matched images, see Figs. 8(e)-8(h), 8(m)-8(p), and the corresponding aligned images, see Figs. 8(a)-8(d), 8(i)-8(l), show comparable intensity levels in both St. Jude and Terumo spaces.

5. COMPARING ATTENUATION COEFFICIENT VALUES USING BOTH ST. JUDE AND TERUMO IMAGES

One of the postprocessing steps in IVOCT data analysis is the estimation of the attenuation coefficients, which are defined as the distinction rate of light passing through a

^bAll, Max, and Min: the total, maximal, and the minimal numbers of frames in the aligned sections.

cIML: intima-media layer.

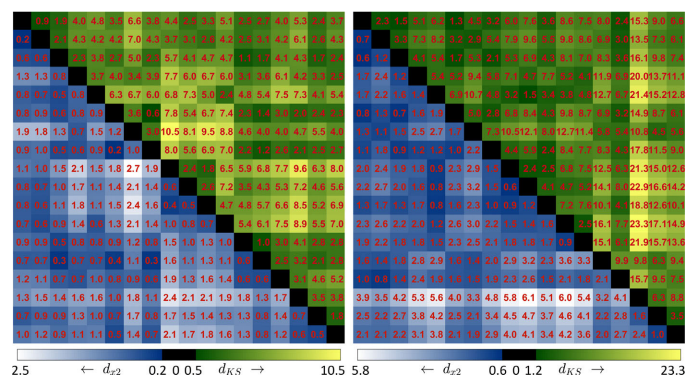


Fig. 3. The distances between all the histograms in the database for St. Jude (left) and Terumo (right) spaces. All the distances were multiplied by a factor of 100 for presentation purposes. In each distance map, d_{x_2} and d_{KS} are shown in the lower and upper triangles, respectively. [Color figure can be viewed at wileyonline library.com]

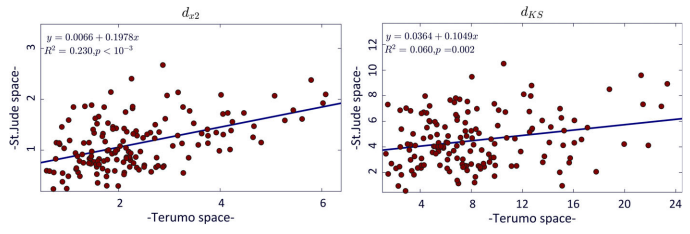


Fig. 4. Linear regression of distances of all the histograms in our database for St. Jude and Terumo. [Color figure can be viewed at wileyonlinelibrary.com]

volume of tissue with a unit of μm^{-1} . It is considered to be a key feature for identification of different tissue types in the arterial wall. In our previous work,³⁵ we reported the depthresolved (DR) estimation using St. Jude images. However, applying this estimation directly to Terumo images results in a different range of values. Therefore, we applied this estimation approach to the matched Terumo images (in St. Jude range) to validate the assumption that proposed matching scheme facilitates generalization of the attenuation estimation algorithm developed for St. Jude data to Terumo images. The results described above show comparable performance for both schemes. We further illustrate performance of the

Global scheme for the estimation of attenuation coefficient using the DR method.

The attenuation was estimated both in the matched and original Terumo images. The median values within ROIs were compared to those estimated using St. Jude images. The paired *t*-test at 5% significance level was used with a null hypothesis that the mean difference between two sets is zero.

Figure 9 shows the Bland–Altman plots of the comparison of median attenuation coefficient in ROIs in (matched/original) Terumo images and in corresponding ROIs in St. Jude images. In the plot presented in Fig. 9(a), original Terumo images were directly used for

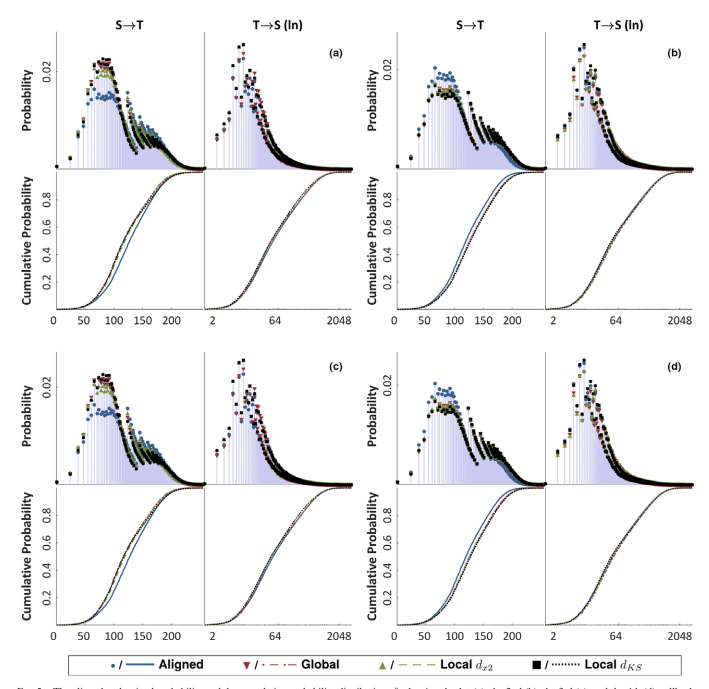


Fig. 5. The aligned and trained probability and the cumulative probability distributions for leaving the 1st (a), the 2nd (b), the 3rd (c), and the 4th (d) pullback out. Target profiles for both St. Jude $(S \rightarrow T)$ and Terumo $(T \rightarrow S)$ images are shown. For the latter case, the *x*-axis is shown in the logarithm scale (ln) for the sake of better visualization. [Color figure can be viewed at wileyonlinelibrary.com]

the estimation. Comparing to those estimated using the corresponding St. Jude images, the points are not evenly distributed around the zero-line, and both systematic difference and the range of random error are high. Paired t-test indicates that the mean difference is significantly different from zero with P < 0.001, and the 95% confidence interval (CI) is [0.74,1.42]. In Fig. 9(b), the matched Terumo images were used for the estimation. In this case, the difference with the values estimated using the St. Jude images is much smaller, which is indicated by lower systematic difference and random error range,

and the points are more randomly distributed around the zero-line. For this case, paired *t*-test gives P = 0.320 with a 95% CI of [-0.10,0.30].

6. DISCUSSION

Clinical significance of IVOCT structures has been reported extensively in clinical research. The use of IVOCT for the analysis of CADs grows exponentially. For more efficient analysis with minimum manual intervention, automated methods for tissue quantification, characterization, and

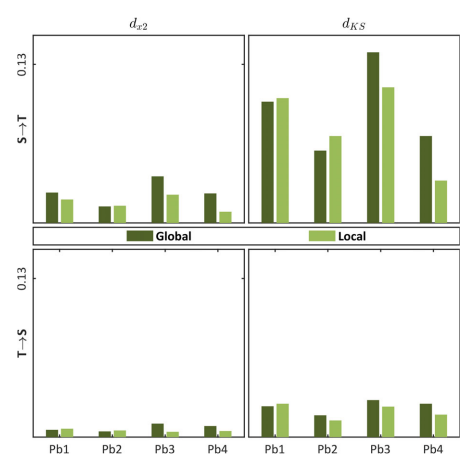


Fig. 6. The bar chart of the distances for leave-one-out experiment for matching St. Jude to Terumo $(S \rightarrow T)$ space and Terumo to St. Jude $(T \rightarrow S)$ space. "Pb" = "pullback". [Color figure can be viewed at wileyonlinelibrary.com]

TABLE II. Results of leave-one-out cross validation.

			$S \rightarrow T$					T→S					
		Pb1	Pb2	Pb3	Pb4	Aver ^a	Pb1	Pb2	Pb3	Pb4	Aver ^a		
$\overline{d_{x2}}$	Global	0.0248	0.0135	0.0381	0.0241	0.0251	0.0058	0.0046	0.0110	0.0090	0.0076		
	Local	0.0192	0.0141	0.0231	0.0091	0.0164	0.0069	0.0053	0.0042	0.0049	0.0053		
d_{KS}	Global	0.0994	0.0593	0.1398	0.0711	0.0924	0.0252	0.0179	0.0303	0.0273	0.0252		
	Local	0.1022	0.0724	0.1110	0.0347	0.0801	0.0276	0.0137	0.0246	0.0183	0.0210		

^aThe average performance of all the four leave-one-out experiments. Bold indicates the best performance.

classification are needed. However, a lack of standardized image intensities can increase the difficulty of designing algorithms or restrict the possibility of generalization of algorithms developed for one specific imaging system. Therefore, standardizing image intensity is a crucial processing step to speed up the development and validation of methods for intravascular tissue analysis.

This study aims at exploring a proper scheme to match IVOCT image intensities with the local-mean EHS technique. In our case, the most essential step is to determine the target histogram. Preliminary statistical analysis suggests that distances in both spaces are significantly correlated. Based on this analysis, we propose a Local matching scheme and

compare it with the Global scheme. Target histograms determined with both schemes turned out to be successful in matching IVOCT intensities at relatively low cost. In this, Local scheme marginally outperforms the Global scheme, which is in line with the results of our preliminary statistical analysis. Moreover, the attenuation estimation experiment presented in Section 5 illustrates benefits of using the Global scheme in practice by showing that it, in particular, largely improves the compatibility of the estimated attenuation coefficient across vendors.

Significant variation in the histograms of different sections is caused by many reasons, the difference in tissue composition being the major factor. As it has been reported in Ref.

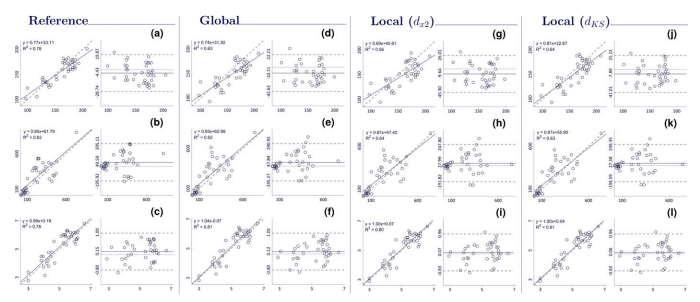


Fig. 7. Scatter and Bland–Altman plots for comparing the median intensities within ROIs in target and matched frames (n = 38). Reference intensities, matched according to the aligned key frames (a,c), compared to that matched with the Global scheme (d,f), the Local scheme with d_{x_2} (g,i) and the Local scheme with d_{KS} (j,l). In the scatter plots, the regression line is shown as the solid line and the dashed line indicates the y = x line. In the Bland–Altman plots, the x-axis and the y-axis are the average and the difference of the target and matched intensities, respectively. The solid line indicates the mean error, and the dashed lines mark the 95% confidence interval. The matched intensities are compared in Terumo (a,d,g,j), St. Jude (b,e,h,k), and logarithmic St. Jude (c,f,i,l) spaces, respectively. [Color figure can be viewed at wileyonlinelibrary.com]

[5], tissue types are mainly visually assessed by recognizing bright speckle, presence of following shadow, sharpness of border, etc. Quantitative results confirm that these image structures yield variations in the histogram. Our previous study³⁶ demonstrated that image intensities can also be affected by position of the catheter. This effect can cause a large variation in the distances and can, thus, explain the low \mathbb{R}^2 in the reported statistical analysis.

The Bland–Altman analysis shows a systematic difference in median intensities within selected regions of interest, even when the exact histogram is specified using the matched key frames. This intrinsic variation may be caused by many factors, such as the data alignment bias, intrinsic difference in systems for data acquisition, etc. Although this difference is relatively low, it should be accounted for in future applications of this method.

For clinical application, both schemes can also be easily embedded as an independent function for automated intensity standardization. For the global scheme, the target histogram needs to be saved within the program. During calculation, this target histogram is loaded and used in EHS. For the local scheme, the database of histogram pairs needs to be saved. During calculation, the target histogram needs to be searched in the database and then it is used in EHS. Furthermore, the proposed experimental setup is not limited to the two considered IVOCT systems and can be used to standardize image intensities between other OCT systems or even for other modalities, for example, MRI. Matching images to St. Jude or Terumo images can also further speed up the validation of newly developed IVOCT systems. As long as the order of intensities of different structures in both systems is consistent, strict ordering can be applied to insure that the exact histogram can be specified.

In this work, we developed and presented a framework for minimizing intensity variation between two different IVOCT systems. At the same time, there might be variations caused by differences between systems from the same vendor, differences between catheters for the same imaging machine, and even differences in the pullbacks acquired using the same hardware and catheter. Since the images were calibrated by design during the acquisition, we expect these variations to be small

Bare metal stents (BMS) were implanted in two out of the four pullbacks used in this particular study. Due to their high light reflectance, the BMS struts appear on the images as saturated bright spots with dark shadows behind them. They disturb the intensity distribution. Since this work mainly focused on matching images of the arterial tissue, images with stent struts were deliberately excluded from the analysis.

6.A. Limitation

The data used in this study are specially generated for construction of this intensity matching framework. Performing (virtually) simultaneous acquisition with two IVOCT systems is not done in clinical practice. Therefore, the amount of data used in this study is limited, which is a major limitation of this work. However, these data are unique and more representative for the analysis on intensity matching than *ex vivo* and animal data. Furthermore, results of leave-one-out validation show reasonable robustness through pullbacks and patients. At the same time, we acknowledge that extending the histogram database might potentially improve the histogram

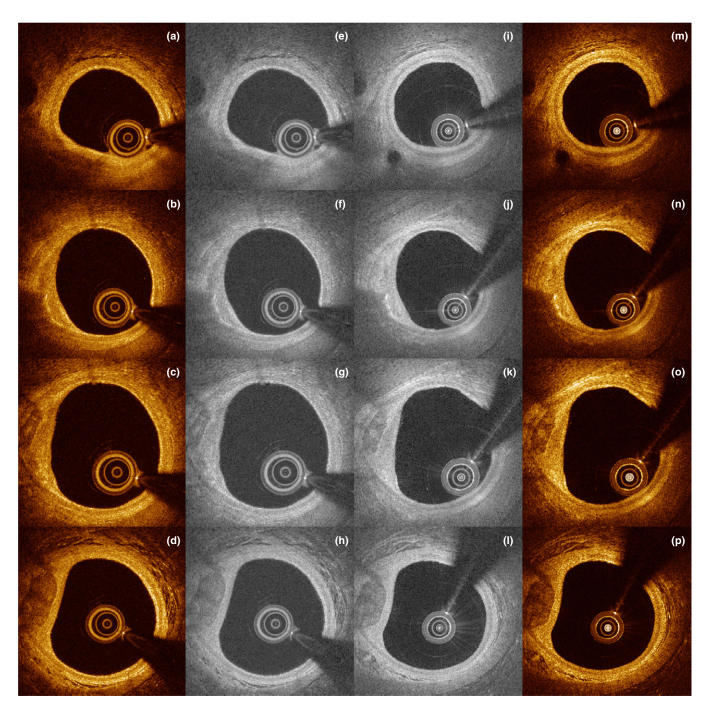


Fig. 8. Images matched using the Global scheme. Raw St. Jude images (a-d) were mapped to the Terumo intensity space (e-h). Raw Terumo images (i-l) were mapped to the St. Jude intensity space (m-p). [Color figure can be viewed at wileyonlinelibrary.com]

matching as using a larger database will most likely lead to more reliable estimation of the target histograms for both Global and Local schemes.

6.B. Future work

This histogram matching scheme will be used for standardization of IVOCT image intensities as a crucial first step for further quantification. Our future work will focus on comparing the outcome of existing quantification methods, such as attenuation estimation, quantification of the degradation of bioresolvable struts and differentiation of neointima, to data acquired by different IVOCT systems.

Once the matching framework is extensively validated, this approach can be routinely used as a preprocessing step for data standardization. The standardized images can be used for the development of universal algorithms for segmentation (of, for example, fibrous cap of TCFAs) or tissue analysis (e.g., for estimation of attenuation coefficients).

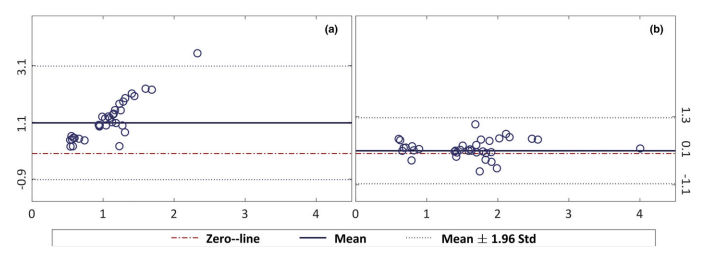


Fig. 9. Bland—Altman plot of the comparison between estimated attenuation coefficients in ROI's in (matched/original) Terumo images and in corresponding ROI's in St. Jude images. (a) Attenuation values were estimated directly using original Terumo images. (b) The values were estimated using the Terumo images matched to St. Jude intensity space. Here "Std" standard deviation". [Color figure can be viewed at wileyonlinelibrary.com]

In this study, we did not include the stented segments of the pullbacks due to the high reflectance of the metallic struts. However, for future work the database could be extended with the stented segments of the pullbacks by excluding individual strut points and their shadows from the images (rather than excluding the entire frame).

7. CONCLUSION

In this work, we presented our contribution to the construction of an intensity standardization framework for IVOCT images. We further contribute to the validation of two proposed schemes in the framework with data acquired by two of the most commonly used IVOCT systems in clinical research. Both local and global schemes are robust and produce accurate intensity matching. While local scheme performs marginally better than the global scheme, it requires a predefined histogram dataset and is more time consuming. Thus, for offline standardization of the images, the local scheme should be preferred for being more accurate. For online standardization or when another system is involved, the global scheme can be used as a simple and nearly-as-accurate alternative.

ACKNOWLEDGMENTS

This research was supported by the China Scholarship Council (No. 201206130062).

CONFLICTS OF INTEREST

The authors have no conflicts to disclose.

REFERENCES

- Mozaffarian D, Benjamin EJ, Go AS, et al. Heart disease and stroke statistics 2016 update. Circulation 2016;133:e38–360.
- Bouma BE, Villiger M, Otsuka K, Oh W-Y. Intravascular optical coherence tomography [invited]. Biomed Opt Express. 2017;8:2660–2686.
- Prati F, Guagliumi G, Mintz GS, et al. Expert review document part 2: methodology, terminology and clinical applications of optical coherence tomography for the assessment of interventional procedures. *Eur Heart* J. 2012;33:2513–2520.
- Prati F, Regar E, Mintz GS, et al. Expert review document on methodology, terminology, and clinical applications of optical coherence tomography: physical principles, methodology of image acquisition, and clinical application for assessment of coronary arteries and atherosclerosis. *Eur Heart J.* 2010;31:401–415.
- Tearney GJ, Regar E, Akasaka T, et al. Consensus standards for acquisition, measurement, and reporting of intravascular optical coherence tomography studies: a report from the international working group for intravascular optical coherence tomography standardization and validation. *J Am Coll Cardiol*. 2012;59:1058–1072.
- Ino Y, Kubo T, Tanaka A, et al. Difference of culprit lesion morphologies between ST-segment elevation myocardial infarction and non-ST-segment elevation acute coronary syndrome. *JACC Cardiovasc Interv.* 2011;4:76–82.
- MacNeill BD, Jang IK, Bouma BE, et al. Focal and multi-focal plaque macrophage distributions in patients with acute and stable presentations of coronary artery disease. J Am Coll Cardiol. 2004;44:972–979.
- Tanaka A, Imanishi T, Kitabata H, et al. Morphology of exertiontriggered plaque rupture in patients with acute coronary syndrome: an optical coherence tomography study. *Circulation*. 2008;118:2368– 2373.
- Tanimoto T, Imanishi T, Tanaka A, et al. Various types of plaque disruption in culprit coronary artery visualized by optical coherence tomography in a patient with unstable angina. Circ J. 2009;73:187–189.
- Taruya A, Tanaka A, Nishiguchi T, et al. Vasa vasorum restructuring in human atherosclerotic plaque vulnerability: a clinical optical coherence tomography study. *J Am Coll Cardiol*. 2015;65:2469–2477.
- Tian J, Hou J, Xing L, et al. Significance of intraplaque neovascularization for vulnerability: Optical coherence tomography study. *J Am Coll Cardiol*. 2012;59:E1439.
- Uemura S, Ishigami KI, Soeda T, et al. Thin-cap fibroatheroma and microchannel findings in optical coherence tomography correlate with subsequent progression of coronary atheromatous plaques. *Eur Heart J.* 2012;33:78–85.
- Yabushita H, Bouma BE, Houser SL, et al. Characterization of human atherosclerosis by optical coherence tomography. *Circulation*. 2002;106:1640–1645.

^{a)}Author to whom correspondence should be addressed. Electronic mail: J.Dijkstra@lumc.nl.

 Kume T, Akasaka T, Kawamoto T, et al. Assessment of coronary arterial thrombus by optical coherence tomography. Am J Cardiol. 2006;97:1713-7.

4170

- Karanasos A, Tu S, van Ditzhuijzen NS, et al. A novel method to assess coronary artery bifurcations by oct: cut-plane analysis for side-branch ostial assessment from main-vessel pullback. Eur Heart J Cardiovasc Imaging 2015;16:177–189.
- Cao Y, Jin Q, Chen Y, et al. Automatic identification of side branch and main vascular measurements in intravascular optical coherence tomography images. In: 2017 IEEE 14th International Symposium on Biomedical Imaging (ISBI 2017). Melbourne: VIC; 2017:608–611.
- Adriaenssens T, Ughi GJ, Dubois C, et al. Automated detection and quantification of clusters of malapposed and uncovered intracoronary stent struts assessed with optical coherence tomography. *Int J Cardio*vasc Imaging. 2014;30:839–848.
- Wang A, Eggermont J, Dekker N, et al. Automatic stent strut detection in intravascular optical coherence tomographic pullback runs. *Int J Car*diovasc Imaging. 2013;29:29–38.
- Wang Z, Jenkins MW, Linderman GC, et al. 3-d stent detection in intravascular OCT using a bayesian network and graph search. *IEEE Trans Med Imaging*. 2015;34:1549–1561.
- Sotomi Y, Tateishi H, Suwannasom P, et al. Quantitative assessment of the stent/scaffold strut embedment analysis by optical coherence tomography. *Int J Cardiovasc Imaging*. 2016;32:871–883.
- Ughi GJ, Adriaenssens T, Sinnaeve P, Desmet W, D'hooge J. Automated tissue characterization of in vivo atherosclerotic plaques by intravascular optical coherence tomography images. *Biomed Opt Expr.* 2013;4:1014– 1030.
- van Soest G, Koljenović S, Bouma BE, et al. Atherosclerotic tissue characterization in vivo by optical coherence tomography attenuation imaging. *J Biomed Opt.* 2010;15:11105.
- Liu S, Eggermont J, Nakatani S, Lelieveldt BPF, Dijkstra J. Light intensity matching between different intravascular optical coherence tomography systems. *Proc SPIE*. 2016a;9689;96893D–96897D.
- Chen C-L, Ishikawa H, Wollstein G, et al. Individual a-scan signal normalization between two spectral domain optical coherence tomography devices. *Invest Ophthalmol Vis Sci.* 2013b;54:3463–3471.

- Chen C-L, Ishikawa H, Wollstein G, Bilonick RA, Kagemann L, Schuman JS. Virtual averaging making nonframe-averaged optical coherence tomography images comparable to frame-averaged images. *Transl Vis Sci Technol.* 2016;5:1.
- Chen C-L, Ishikawa H, Ling Y, et al. Signal normalization reduces systematic measurement differences between spectral-domain optical coherence tomography devices. *Invest Ophthalmol Vis Sci.* 2013a;54:7317

 7322.
- Chen C-L, Ishikawa H, Wollstein G, Bilonick RA, Kagemann L, Schuman JS. Signal normalization reduces image appearance disparity among multiple optical coherence tomography devices. *Transl Vis Sci Technol*. 2017:6:13
- Hummel RA. Histogram modification techniques. Comput Graph Image Process. 1975;4:209–224.
- Nikolova M, Wen Y-W, Chan R. Exact histogram specification for digital images using a variational approach. J Math Imaging Vis. 2013;46:309–325.
- Coltuc D, Bolon P, Chassery JM. Exact histogram specification. *IEEE Trans Image Process*. 2006;15:1143–1152.
- Wan Y, Shi D. Joint exact histogram specification and image enhancement through the wavelet transform. *IEEE Trans Image Process*. 2007;16:2245–2250.
- Chen C-L, Ishikawa H, Wollstein G, et al. Histogram matching extends acceptable signal strength range on optical coherence tomography images. *Invest Ophthalmol Vis Sci.* 2015;56:3810–3819.
- Stanciu SG, Stanciu GA, Coltuc D. Automated compensation of light attenuation in confocal microscopy by exact histogram specification. *Microsc Res Tech.* 2010;73:165–175.
- Euser AM, Dekker FW, le Cessie S. A practical approach to Bland-Altman plots and variation coefficients for log transformed variables. *J Clin Epidemiol.* 2008;61:978–982.
- Liu S, Sotomi Y, Eggermont J, et al. Tissue characterization with depthresolved attenuation coefficient and backscatter term in intravascular optical coherence tomography images. J Biomed Opt. 2017;22:1–16.
- Liu S, Eggermont J, Wolterbeek R, et al. Analysis and compensation for the effect of the catheter position on image intensities in intravascular optical coherence tomography. *J Biomed Opt.* 2016b;21:126005.