Digital simulation of staining in histopathology multispectral images: enhancement and linear transformation of spectral transmittance

Pinky A. Bautista
Yukako Yagi
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Pinky A. Bautista and Yukako Yagi
Harvard Medical School, Department of Pathology, Massachusetts General Hospital, Boston, Massachusetts 02114

Abstract. Hematoxylin and eosin (H&E) stain is currently the most popular for routine histopathology staining. Special and/or immuno-histochemical (IHC) staining is often requested to further corroborate the initial diagnosis on H&E stained tissue sections. Digital simulation of staining (or digital staining) can be a very valuable tool to produce the desired stained images from the H&E stained tissue sections instantaneously. We present an approach to digital staining of histopathology multispectral images by combining the effects of spectral enhancement and spectral transformation. Spectral enhancement is accomplished by shifting the $N$-band original spectrum of the multispectral pixel with the weighted difference between the pixel’s original and estimated spectrum; the spectrum is estimated using $M < N$ principal component (PC) vectors. The pixel’s enhanced spectrum is transformed to the spectral configuration associated to its reaction to a specific stain by utilizing an $N \times N$ transformation matrix, which is derived through application of least mean squares method to the enhanced and target spectral transmittance samples of the different tissue components found in the image. Results of our experiments on the digital conversion of an H&E stained multispectral image to its Masson’s trichrome stained equivalent show the viability of the method. Keywords: hyperspectral imaging; image analysis; image processing; image understanding; multispectral enhancement; digital staining. Paper 11757 received Dec. 14, 2011; revised manuscript received Mar. 27, 2012; accepted for publication Apr. 2, 2012; published online May 4, 2012.

1 Introduction

The potential utility of multispectral imaging for various medical imaging applications has been investigated\(^\text{1}\) in pathology, particular interest is focused on the possible application of this imaging modality to improve tissue detection, classification and visualization\(^\text{2}\) in contrast to the conventional RGB-color image acquisitions systems, which use wideband filters, multispectral imaging uses $N$ narrowband filters to capture images across wide spectral range. The utilization of narrowband filters enables a multispectral image acquisition device to capture spectral features that are not normally captured with RGB-color image acquisition devices.

Chemical staining of tissue sections is an important part in clinical pathology diagnosis. Staining produces clearer visualization of the different tissue structures. From a stained tissue section, the tissue’s architecture and the morphological characteristics of its cells structures could easily be observed. The digital simulation of the effect of chemical staining, also known as digital staining, on histopathology images using multispectral features was explored.\(^\text{3,4}\) Digital staining aims at transforming the original image, which could be unstained or stained, into an image with the desired staining patterns by employing digital processing techniques. It produces images which do not only possess improved colorimetric differences between faintly discriminated tissue structures, but also the visualization inherent to the effect of chemical staining to the tissue. Hematoxylin and eosin (H&E) stain is currently the most popular for routine staining. To corroborate the pathologist’s initial findings on the H&E stained tissue sections, special stains and/or immunohistochemical stainings (IHC) are requested. Digital staining can be utilized to visualize the staining reactions of the H&E stained tissue structures to other types of dye without necessarily resorting to chemical staining. From the digitally stained images additional diagnostic information can be collected. The successful implementation of digital staining could therefore enhance the efficiency of the clinical diagnosis workflow while reducing the cost of chemical staining.

Digital staining methods for histopathology multispectral images have been introduced.\(^\text{5,6}\) The methods involve two basic processes: 1. spectral classification; and 2. spectral transformation. The digital staining method in Ref.\(^\text{7}\) showed promise in transforming H&E stained images to their Masson’s trichrome stained images equivalent. However, it was not as effective in differentiating tissue structures which have very subtle colorimetric difference. This was thought to be caused either by the limitation of the spectral classification itself or by the process by which the digital colorization was implemented.

Mitsui et al.\(^\text{8}\) proposed a spectral enhancement scheme to improve the visualization of obscured skin patterns. A similar approach was applied to histology images to enhance tissue structures which acquire similar H&E staining patterns.\(^\text{9}\) Initial experimental results showed the capability of the enhancement method to capture salient spectral differences. To circumvent the limitations of the previous digital staining method, the application of spectral enhancement combined with spectral transmittance...
transformation was introduced. Application of the proposed method to H&E stained tissue images resulted in the vivid discrimination between tissue structures that share similar colorimetric attributes, particularly the discrimination between smooth muscle and collagen fiber. While spectral enhancement increased the spectral color difference between the object of interest and background, application of spectral transformation converted the H&E stained spectral color of the tissue structures to the colors specific to their reaction to the specified stain, i.e., Masson’s trichrome. In this paper, we extended the discussion of the digital staining method presented in Ref.4. In the enhancement process, the original spectrum of the pixel at band $n$, $W_n$, $n = 1, 2, \ldots , N$, is shifted by a real-valued factor, which is the spectral residual-error of the multispectral pixel weighted by a constant. The spectral enhancement can be expressed as follows:

$$t_e = t_o + We.$$  

(2)

The variables $t_o$, $t_e$, and $e$ are all $N \times 1$ column vectors which denote the pixel’s enhanced spectral transmittance, original transmittance, and its spectral residual-error, respectively. The entries of the $N \times N$ matrix $W$ serve as weighting factors to the spectral residual-error at different bands. The parameters $W$ and $e$ will be discussed in the next sections.

2.3.1 Principal component analysis and spectral error

Principal component analysis (PCA) is a popular choice for multispectral/hyperspectral image compression and for spectral feature extraction. In the analysis, the original spectrum is estimated by considering only the first $M \ll N$ dominant principal component (PC) vectors, which were calculated from a chosen spectral dataset.

The objects in an image can generally be categorized into: 1. background (or objects of non-interest); and 2. object of interest. Let $T_e$ be a $P \times N$ matrix containing the original spectral transmittance samples of the background and object of interest, where $P$ is the total number of spectral samples and $N$ is the spectral dimension.

$$T_e = \begin{bmatrix} t_e(1,1) & t_e(1,2) & t_e(1,3) & \ldots & t_e(1,N) \\ t_e(2,1) & t_e(2,2) & t_e(2,3) & \ldots & t_e(2,N) \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ t_e(p,1) & t_e(p,2) & t_e(p,3) & \ldots & t_e(p,N) \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ t_e(P,1) & t_e(P,2) & t_e(P,3) & \ldots & t_e(P,N) \end{bmatrix} \quad (3)$$

The element $t_e(p,n)$ of the matrix $T_e$ above represents the transmittance value of the $p$th samples at the $n$th spectral band where $p = 1, 2, \ldots , P$ and $n = 1, 2, \ldots , N$. In multispectral image compression applications, the $M \ll N$ principal component (PC) vectors are normally derived from the covariance of the samples in $T_e$. The $M$ PC vectors are determined based on how much of the variance in the spectral samples in $T_e$ are captured. The accumulated variance captured by the $M$ PC vectors can be estimated using their respective eigenvalues:

$$\psi = \frac{\sum_{n=1}^{N} y_n}{\sum_{n=1}^{N} \bar{y}_n}.$$  

(4)

where $\psi$ denotes the accumulated variance, $y_n$ is the eigenvalue of of the $n$th PC vector, $\bar{y}_n = 1 \ldots N$. The original spectrum of a multispectral pixel can be reconstructed with minimum error when the first $M$ PC vectors captured most of the original spectral data variance:

$$\hat{t}_o = \sum_{m=1}^{M} a_m v_m + \hat{t}_o,$$  

(5)

$$a_m = v_m^T (t_o - \hat{t}_o).$$  

(6)

where $a_m$ is the coefficient of the $m$th PC vector, $v_m$; $\hat{t}_o$ denotes the estimate of the original transmittance $t_o$; $\hat{t}_o$ is the average of the spectral samples in $T_e$; and $e$ is the $N$-dimensional spectral residual-error, which is defined as follows:

$$e = t_o - \hat{t}_o.$$  

(7)
Let $T_g \subseteq T_r$ be a $P_g \times N$, where $P_g$ represents the total number of spectral samples of the background objects. For the current spectral enhancement method, the PC vectors are instead derived from the spectral samples in matrix $T_g$ rather than from $T_r$. That is, the spectral samples of the object of interest are not considered from the spectral set from which the PC vectors are drawn.

### 2.3.2 Weighting matrix $W$

In Ref. [5], the $N \times N$ weighting matrix $W$ in Eq. (2) was designed as follows:

$$[W]_{ij} = \begin{cases} k & i = j \\ 0 & \text{otherwise} \end{cases},$$

where $k$ is a real value. Equation (3) implies that all the spectral bands are equivalently weighted. That is, there are no specified bands for enhancement. In Ref. [5], the band for enhancement was identified from the spectral residual-error plot of the object of interest. The band at which the spectral residual-errors of the object of non-interest or background objects are close to zero and that of the object of interest was large was chosen as the band for enhancement. This condition was found to be necessary to isolate the effect of the enhancement to the objects of interest and minimize the spectral color shifts among the background objects. Hence, when spectral samples of the image objects are available, a more robust approach would be to specify the bands for enhancement at which the spectral residual-errors of the object of interest are identified to be much larger than the errors of the background objects. This does not only ensure that only the spectra of the objects of interest will be modified at the specified bands by the spectral enhancement process, but this also minimizes the effect of spectral variations in the original spectral data to the digital staining results in general. If we let $b \subseteq \lambda = \{\lambda_1, \lambda_2, \ldots, \lambda_M\}$ a vector containing the different bands at which the spectral residual-error of the object of interest is large while those of the background objects are small, we can rewrite Eq. (2) in the following form:

$$[W]_{ij} = \begin{cases} k & i \in b, j \in b \\ 0 & \text{otherwise} \end{cases}. \quad (9)$$

We referred to the elements of the vector $b$ in Eq. (9) as the bands for enhancement.

### 2.4 Spectral Transformation

In the transformation of spectral transmittance we assumed that the target transmittance value of the $p$th spectral sample at the $n$th spectral band, $t_i(p, n)$, is equal to the weighted summation of its reference transmittance, which is the original or enhanced transmittance of the object, in the whole spectral band.

$$t_i(p, n) = \sum_{j=1}^{N} t_r(p, j) q(j, n), \quad (10)$$

where $n = 1 \ldots N$; $t_r(p, j)$ is the transmittance value of $p$th reference spectral sample at $j$th band; and $q(j, n)$ is the element of the $N \times N$ transformation $Q$ at the $j$th row and $n$th column. With the appropriate values of the elements of the matrix $Q$, the reference spectrum can be converted to its target spectral configuration smoothly using Eq. (11).

The solution for the transformation matrix $Q$ can be determined using the least mean squares method. Let $T_r$ be a $P \times N$ matrix that contains the target spectral transmittance samples of the reference spectral transmittance samples in $T_r$. The matrix form of Eq. (11) can be written as:

$$T_r = T_r Q, \quad (11)$$

and the linear solution for $Q$ can be given by:

$$\hat{Q} = T_r^+ T_r, \quad (12)$$

where $T_r^+$ is the pseudoinverse of $T_r$. Using the result of Eq. (12), the reference spectrum (or original spectrum) can be converted to its target spectral configuration by the following:

$$\hat{t}_r = \hat{Q}^T t_r \quad (13)$$

where $t_r$ and $\hat{t}_r$ are $N \times 1$ column vectors representing the reference and its transformed spectrum, respectively; and $\hat{Q}^T$ is the transpose of $\hat{Q}$ in Eq. (12).

### 2.5 Digital Staining Scheme

Combining Eqs. (6), (9) and (11), we can rewrite Eq. (6) in the following form:

$$t_e = [W(I - VV^T) + I] t_r + W[I - VV^T] t_o, \quad (14)$$

where $I$ is an $N \times N$ identity matrix, and $V$ an $N \times N$ matrix whose $j$th column corresponds to the $j$th PC vector:

$$[V]_{ij} = \begin{cases} v_j & j \leq M \\ 0 & \text{otherwise} \end{cases} \quad (15)$$

The enhanced versions of the original transmittance samples resulting from the application of Eq. (14) are used instead of the original spectral samples in finding for the solution of the transformation matrix $Q$. With this, Eq. (11) becomes:

$$T_r = T_r Q, \quad (16)$$

where $T_r$ is the $P \times N$ matrix that contains the enhanced spectral samples. The linear solution for $Q$ as expressed in Eq. (12), still holds. Replacing $t_r$ in Eq. (11) with $\hat{t}_r$, we will have:

$$\hat{t}_r = \hat{Q}^T \hat{t}_r \quad (17)$$

Then, the digitally stained spectrum of a pixel, $\hat{t}_r$, can be determined by combining Eqs. (16) and (17).

$$\hat{t}_r = \hat{Q}^T [W(I - VV^T) + I] t_r + \hat{Q}^T W[I - VV^T] t_o. \quad (18)$$

Equation (18) illustrates that the current digital staining approach can be implemented by matrix manipulation. Since spectral classification is no longer necessary this makes the implementation of the present approach more straightforward compared to the previously proposed methods in Refs. [6] and [7]. Using the equation above, the original stained transmittance of a tissue structure, $t_r$, can be converted to its target spectral transmittance configuration, $\hat{t}_r$, by scaling and shifting. The original transmittance, $t_r$, is scaled by the product between the
$N \times N$ transformation matrix $Q^T$ and the principal component vectors, $V$ weighted by the elements of the $N \times N$ matrix $W$. The spectral color of the scaled transmittance is shifted by the average spectrum of the background objects, denoted by $I$, which is scaled by the weighted product between the matrices $W$, $Q^T$, and $V$. The matrices $V$ and $Q^T$ are determined offline and require the first and second stained spectral samples of the different tissue components, e.g., H&E and Masson’s trichrome. The entries for $W$ are set to zero by default. The decision as to which elements should be set to non-zero values depends on the identified bands for enhancement.

### 2.6 RGB-Color Visualization

To view the transformed spectrum of Eq. (18) on ordinary display, it is converted to its RGB-color. An $N \times N$ spectral transmittance can be converted to its RGB-color using the CIE XYZ color matching functions. Let $F$ be the $3 \times N$ matrix of the color matching function, $E$ as the $N \times N$ diagonal matrix of the illumination spectrum. The $XYZ$ tristimulus values of the digitally stained spectrum, $t_i$, are calculated as follows:

$$ z = FE t_i, \quad (19) $$

where $z$ is the $3 \times 1$ vector containing the $XYZ$ values, $z = [X, Y, Z]^T$. The RGB color values can be determined using the $3 \times 3 XYZ$ to RGB conversion matrix, $C$$

$$ \alpha' = CZ, \quad (20) $$

$$ \alpha = \text{ROUND}(\alpha'^{1/2.2}), \quad (21) $$

where $\alpha$ is the $3 \times 1$ RGB vector whose values are clipped into [0,255]. This procedure results in RGB data that meets the sRGB standard.

The digital staining method introduced in this work is summarized by the block diagram in Fig. 1. The spectral transmittance of a stained multispectral pixel is first calculated using Eq. (18). Then, spectral enhancement is applied by utilizing Eq. (18) using the $M$-PC vectors, which were determined using the spectral samples of the background objects, i.e., objects of non-interest, together with other pre-defined enhancement parameters. The enhanced spectrum is then transformed using the $N \times N$ transformation matrix $Q$, Eq. (17), which was determined using the reference and target spectral datasets, e.g., H&E and Masson’s trichrome stained spectra. RGB visualization of the transformed spectrum then follows, Eqs. (18)–(21). The process is repeated for all the image pixels.

### 3 Experiments and Results

#### 3.1 Multispectral Image Acquisition

In this work we used two sets of slides. Each set is composed of a pair of H&E and Masson’s trichrome stained slides that belong to the serial sections of a liver tissue. From these sets we captured 10 sets of multispectral images at 20x. Since we captured similar areas from both slides, the images in a set share similar structural composition. Of the 10 multispectral-image sets, we selected 5 sets from which we identified six different tissue components namely the nucleus, cytoplasm, red blood cells (RBC), collagen fiber, smooth muscle, and duct; the labeling of these tissue components were referred to a pathologist for correctness. We then collected the H&E and Masson’s trichrome spectral samples for these tissue components, and also for the white areas; the white areas are areas in the images which are devoid of any tissue structures.

#### 3.2 PC Vectors and Spectral Residual-Errors

The $P \times N$ matrix $T_\alpha$ contains the H&E spectral samples of all the identified tissue components while the $P \times N$ matrix $T_\beta$ from which we derived the $M$ PC vectors for spectral enhancement contains the H&E the spectral samples of the non-collagen fiber tissue components only (or the background objects), which includes the nuclei, cytoplasm, red blood cells (RBC), smooth muscle, duct, and the spectral samples of the white areas. The effective number of PC vectors, $M$, was determined using Eq. (17). The plot of one minus the accumulated variance with respect to the number of PC vectors is shown in Fig. 2. The vertical axis corresponds to one minus the log of the accumulated variance for a given number of PC vectors indicated on the horizontal axis. From this plot, we can see that the variance starts to taper at $M = 5$ PC vectors. We can also observe that there is not much difference in the variance between $M = 5$ and $M = 6$ PC vectors. This implies that most of the variance in the original spectral data in $T_\beta$ are captured by the first five PCs, and the spectra of the non-collagen fiber tissue structures can be estimated with reduced error. The plots in Fig. 3 are the average spectral residual-errors of the different tissue-components in $T_\alpha$. We can see that the spectral residual-error of the collagen fiber in Fig. 3(b) is markedly different from the rest. This demonstrates that the spectral residual-error may represent the spectral samples of the collagen fiber that were not considered in the composition of $T_\alpha$, which we used to calculate the PC vector, and allows for the enhancement of these residual-error spectra at chosen wavelengths.

#### 3.3 Digital Staining Processes

To demonstrate the effect of the spectral enhancement and the spectral transformation processes in the proposed digital
staining method, we considered the H&E spectral samples of the different tissue components in $T_r$. The average H&E transmittance spectra of these tissue components are shown in Fig. 4(a). The spectral samples from which these average spectra were calculated are translated into their RGB-color and they are illustrated by the color patches at the top of the spectral plots. The color of each patch characterizes the H&E staining reaction of a tissue structure. Eosin stained structures are stained pink to red, while the hematoxylin stained ones, as such as the nuclei, are stained blue to purple. Figure 4(a) shows that from an H&E stained section we can easily discriminate between eosin and hematoxylin stained structures.

### 3.3.1 Spectral enhancement

We can observe in Fig. 4(a) that differentiation between collagen fiber and smooth muscle, which are both stained with eosin dye, is not as straightforward as the differentiation between eosin stained and hematoxylin stained tissue components, such as the differentiation between nuclei and collagen fiber. In order to improve the H&E spectral color difference between the collagen fiber and smooth muscle, we utilized the spectral enhancement of Eq. (9). As mentioned previously, the collagen fiber is the current object of interest and thus the subject for enhancement. Hence, we need to identify the bands for enhancement in Eq. (9) which would emphasize this structure in the enhanced image. Referring to the spectral residual-error configuration of the collagen fiber, as shown in Fig. 4(b), we considered the bands at which the non-collagen fiber spectral errors are small while that of the collagen fiber are distinctly large. We then selected $b = [550 \text{ nm}, 555 \text{ nm}]$. Trial and error experiments were done to determine the values of the weighting factors for these bands to produce the RGB images which exhibit the staining patterns of a Masson’s trichrome stained image. Based on the initial results of the experiments, we set the weighting factors for these bands to $k = 10$. The effect of the spectral enhancement using these bands is illustrated by the spectral plots at the top panel in Fig. 4(b). Each plot represents the average spectrum of a particular tissue component. It is evident that when the bands for enhancement are chosen appropriately, only the spectra of the objects interest, i.e., collagen fiber, are modified by the enhancement. The RGB-colors of the enhanced spectral samples are indicated by the $5 \times 5$ pixels color patches above the plot. We can observe that while the patches for the smooth muscle remains pinkish the hue of the collagen fiber shifted from pinkish to light pinkish, which in effect increases the overall color difference between these structures.

### 3.3.2 Spectral transformation

The purpose of spectral transformation is to convert the original or enhanced transmittance of a tissue component to the spectral configuration associated to its reaction to a particular stain. In our work, the spectral transformation was achieved by designing an $N \times N$ transformation matrix $Q$ using the enhanced H&E and Masson’s trichrome stained spectral samples of the different tissue components by least mean squares method, Eq. (10). The spectral samples represent seven different classes of tissue components. To minimize the effect of intra-class spectral
variations, the spectral average of each class, e.g., nuclei, cytoplasm, etc., was instead considered to find the solution for $Q$. Taking this into consideration, Eq. (16) reduces to the following form:

$$
\begin{bmatrix}
t_t(1, 1) & t_t(1, 2) & \ldots & t_t(1, N) \\
t_t(2, 1) & t_t(2, 2) & \ldots & t_t(2, N) \\
\vdots & \vdots & \ddots & \vdots \\
t_t(c, 1) & t_t(c, 2) & \ldots & t_t(c, N)
\end{bmatrix}
= \begin{bmatrix}
t_e(1, 1) & t_e(1, 2) & \ldots & t_e(1, N) \\
t_e(2, 1) & t_e(2, 2) & \ldots & t_e(2, N) \\
\vdots & \vdots & \ddots & \vdots \\
t_e(c, 1) & t_e(c, 2) & \ldots & t_e(c, N)
\end{bmatrix} \left[ Q_{N \times N} \right], \tag{22}
$$

where $t_t(c, 1)$ and $t_t(c, 1)$ are the average transmittance of the $c$'th class at band $i = 1, 2, \ldots, N$ of the Masson’s trichrome and the H&E enhanced spectral transmittance samples. The $N \times N$ matrix $Q$ was then derived using the Pseudoinverse function as implemented by Matlab.

Figure 4 demonstrates the effect of the spectral transformation. While it is not very direct to distinguish the collagen fiber from the smooth muscle on the basis of their H&E stained spectra, we can observe a notable difference between their transformed spectra. In contrast to the spectral enhancement wherein only the spectral samples of the collagen fiber were modified at the selected bands for enhancement, in spectral transformation all the spectral samples, regardless of their classes, underwent changes in their spectral configurations. The RGB color visualization of the transformed spectral samples is shown at the panel above the spectral plots. Here, we can readily observe the color difference between collagen fiber and smooth muscle. The average plots of the physically-stained spectral samples and their corresponding RGB color patches are also provided in Fig. 4. If we refer the RGB color patches in Fig. 4(c) to 4(d) we can see that some of the spectral samples were mislabeled as a result of the spectral transformation—the RGB colors acquired by these samples were different from what they were supposed to exhibit. However despite the occurrence of spectral mislabeling we can still observe the delineation between smooth muscle and collagen fiber better compared to the original spectral data, Fig. 4(a), or to the result of the spectral enhancement, Fig. 4(b).

3.4 Digitally-Stained Histopathology Images

The original H&E stained images along with the results of the digital staining and the physically stained Masson’s trichrome...
images are shown in Fig. 5. The original H&E stained images, results of digital staining, and the physically stained Masson’s trichrome stained images are shown at the left, middle and right panels, respectively. The H&E stained image at the topmost row represents one of the five training images we used in the experiments, and the images at the succeeding rows represent the test images. The results of the digital staining at the middle panels show similarity to the actual Masson’s trichrome stained images on their right, particularly with respect to the differentiation between the eosin stained tissue areas, e.g., collagen fiber and cytoplasm areas.

Masson’s trichrome stain is used to obtain clear visualization of fibrous tissue in the diagnosis of liver diseases. We cropped similar areas that contain both the collagen fiber and smooth muscles from the original H&E and Masson’s trichrome stained images, and from the digitally stained Masson’s trichrome image. The cropped images are shown in Fig. 6. The original H&E and Masson’s trichrome stained images are shown at the leftmost and rightmost panels, and the result of the digital staining is shown at the middle panel. While the distinction between collagen fiber and smooth muscle in the original H&E stained image is not as obvious as it is in the Masson’s trichrome stained image, this has been improved after the application of digital staining. The digitally-stained image at the center panel share similar characteristics to the physically stained Masson’s trichrome image on its right in two aspects: 1. the differentiation between the collagen fiber and smooth muscle is better observed; and 2. the acquired colors of these tissue structures.

4 Discussion
The digital staining method we presented in this paper consisted of two general processes: (1) spectral enhancement, and (2) spectral transformation. Linear transformation of transmittance spectra works best when the spectral data in both the reference and target spectral transmittance are linearly separated. In the previous works spectral classification was considered as the solution to address the spectral overlaps in some transmittance
However, even with the application of spectral classification, tissue structures with similar colorimetric attributes were not effectively discriminated. Figure 7 shows the original and enhanced transmittance spectra of the different tissue components, which share similar H&E stained spectral attributes, i.e., cytoplasm, smooth muscle and collagen fiber, projected onto the PC axes. The original H&E spectral clusters in Fig. 7(a) and 7(b) show that there is no clear-cut separation between collagen fiber and smooth muscle. However the cluster plots of the enhanced spectra in Fig. 7(c) and 7(d) show improved separation between these two structures, especially the projection of the enhanced spectra onto PC2 and PC3 axes, Fig. 7(d). The plot illustrates that the spectral residual-errors of the tissue structures contain information which could improve the visualization of H&E stained tissue structures not normally observe with the naked eyes, particularly the differentiation between smooth muscle and collagen fiber.

Fig. 6 Areas containing both collagen fiber and smooth muscle. These areas were cropped from the original 1434 × 1050 images. (a) Original H&E stained image; (b) result of digital staining; and (c) physically stained Masson’s trichrome stained image. The collagen fiber could hardly be differentiated from the original H&E stained image, however application of digital staining results in their improved differentiation in which the smooth muscles were digitally stained with red color and the collagen fibers blue similar to their color when stained with Masson’s trichrome.

Fig. 7 Original and enhanced transmittance spectra of the eosin stained tissue components, i.e., cytoplasm, collagen fiber, and smooth muscle, projected onto the principal component axes. (a) to (b) original spectra and (c) to (d) enhanced spectra. The original spectra of collagen fiber and smooth muscle overlap to certain degree. Application of spectral enhancement minimizes the overlaps.
The use of principal component (PC) vectors to reconstruct the spectrum of a pixel is an established method. In this approach, the optimum number of PC vectors for used in the reconstruction depends on the application requirement. In the present experiment, the total number of PC vectors was determined such that the spectral reconstruction errors of the H&E stained spectral samples belonging to the background objects are minimized. With the current image samples, the background objects consisted of five classes of tissue components including the nucleus, cytoplasm, RBC, smooth muscle and duct; these are basically the non-collagen fiber tissue components. Included also in the category of background objects are the white areas which do not contain any tissue structures. We identified the first five PC vectors to reconstruct the spectra of these tissue components based on total amount of variance these vectors captured, Fig. 2. The different PC vectors are displayed in Fig. 3. To illustrate the possible correlation between the PC vectors and the staining characteristics of the spectral samples, the plots of the H and E dyes absorption spectra are also shown in Fig. 4. The first PC vector describes the luminance of the spectral samples. The second PC vector reveals spectral samples with variations in the eosin and hematoxylin dye amounts. The third PC vector demonstrates spectral samples which vary in the short wavelength and within the eosin absorption band. It also describes samples which moderately vary within the absorption band of hematoxylin dye. The fourth PC vector indicates spectral samples which prominently vary within the hematoxylin band, and the fifth PC vector represents spectral variations mainly within the eosin band. Here we see that each of the PC vectors captures particular staining characteristics of the spectral samples. Our experimental results also show that with five PC vectors we could isolate the spectral differences between collagen fibers and other eosin stained tissue structures. While the first two PC vectors captured around 98% of the total variance, they do not necessarily encompass the staining characteristics of the spectral samples. The diverse spectra found in a stained tissue section can be attributed to the absorbing elements in the tissue itself or to the change in the absorption coefficient of some tissue components. Since the resulting spectral characteristics of some H&E stained tissue components are attributed by the tissue and the stain itself, superposition of the H and E stain spectra to reconstruct the spectra of these stained tissue components may not completely capture the staining characteristics of these tissue components.

Figure 5 shows the normalized plots of the H and E dye absorption spectra and the spectral residual-error of the collagen fiber. The shape of the spectral residual error closely resembles the eosin dye absorption spectrum with the absorption peak shifted to a longer wavelength. The shift can be linked to the dye-protein interaction wherein the magnitude of the shift is proportional to the amount of protein concentration. Thus, it can be thought that the difference between the eosin-stained collagen fiber and smooth muscle lies in the amount of their protein concentrations.

4.1 Limitations and Future Work

The digital staining process involves some parameters that have to be appropriately specified or computed in order to obtain the ideal results. Spectral enhancement involves the determination of the number of PC vectors as well as the bands for enhancement, while spectral transformation requires the derivation of the transformation matrix. The number of PC vectors and the bands for enhancement could be affected by the characteristics of the spectral samples used to derive them. Hence, the number of PC vectors and the bands for enhancement we used in the present experiments may not hold true for all cases. Possible ways to produce consistent enhancement result are: 1. to standardize the staining conditions of the input images; and 2. to replace the use of the N-dimensional spectral residual-error in the spectral enhancement. The staining correction method proposed in Ref. 12 can be employed in the pre-processing of the images to obtain standardize staining conditions. And the N-dimensional spectral residual-error can be replaced with a one-dimensional variable such that the bands for enhancement do not depend on the spectral residual-error configurations, i.e., where the error peaks. Application of the spectral transformation process using the transformation matrix results in digitally stained images which have similar staining patterns to the physically stained Masson’s trichrome images. However, the colors of the digitally stained tissue components are not yet visually indistinguishable from the real Masson’s trichrome stained ones. An investigation on the more effective...
ways to implement spectral transformation will be part of our next work.

Aside from Masson’s trichrome stain, there are other special stains which are employed to highlight other tissue structures. That is, although it has been shown that the spectral features of H&E stained tissue structures contain information relevant to improving the visualization of collagen fiber and smooth muscle, which are effectively visualize using the Masson’s tri-chrome special stain, further investigations are still needed to firmly establish whether the information contained in the original H&E spectral feature is complete to warrant the digital conversion of an H&E stained image into an image which displays the staining patterns of any particular special stain. Furthermore, the digital conversion of an unstained histopathology image to its stained image counterpart may require other spectral features and the extension of the multispectral filter sensitivity beyond the visible spectrum.

5 Summary and Conclusions

In this paper we introduce a digital staining method for histopathology images using multispectral imaging. In the method we utilize the spectral features of a multispectral image and combine the effects of spectral enhancement and spectral transformation to convert an H&E stained image to its Masson’s tri-chrome stained equivalent. The method can also be extended to convert H&E stained images to images possessing the staining patterns of other special stains.

Digital staining goes beyond tissue classification. In addition to discriminating the structures in an image, it offers the kind of visualization inherent to chemically stained tissue sections. The success of this technology will alleviate the cost of staining while providing pathologists immediate access to the special stained images of the physically stained H&E stained tissue section. As whole slide imaging technology continue to evolve, it would be possible in the future to integrate digital staining to whole slide image scanning.

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References