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Abstract. We report a second derivative multispectral algorithm for quantitative assessment of cutaneous tissue oxygen saturation (StO₂). The algorithm is based on a forward model of light transport in multilayered skin tissue and an inverse algorithm for StO₂ reconstruction. Based on the forward simulation results, a parameter of a second derivative ratio (SDR) is derived as a function of cutaneous tissue StO₂. The SDR function is optimized at a wavelength set of 544, 552, 568, 576, 592, and 600 nm so that cutaneous tissue StO₂ can be derived with minimal artifacts by blood concentration, tissue scattering, and melanin concentration. The proposed multispectral StO₂ imaging algorithm is verified in both benchtop and *in vivo* experiments. The experimental results show that the proposed multispectral imaging algorithm is able to map cutaneous tissue StO₂ in high temporal resolution with reduced measurement artifacts induced by different skin conditions in comparison with other three commercial tissue oxygen measurement systems. These results indicate that the multispectral StO₂ imaging technique has the potential for noninvasive and quantitative assessment of skin tissue oxygenation with a high temporal resolution. (*© 2015 Society of Photo-Optical Instrumentation Engineers (SPIE)* [DOI: 10.1117/1.JBO.20.3.036001]

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1 Introduction

Tissue oxygen saturation (StO_2) is a relative measure of the amount of oxygen carried by hemoglobin in biological tissue. Measuring StO₂ will provide important information about tissue viability and help clinicians to make critical decisions in many clinical procedures, such as wound healing management, plastic surgery, and organ transplantation. Various methods, such as blood oxygen level dependent magnetic resonance imaging, pulse oximetry, phosphorescence, electroparamagnetic resonance, and microelectrode, have been used for detecting tissue oxygen content and hemodynamics.¹⁻³ Among these oximetry techniques, optical oximetry is able to monitor tissue StO₂ continuously and noninvasively by detecting characteristic spectra of oxy-hemoglobin and deoxy-hemoglobin.⁴ Commonly used optical oximetry devices are based on contact and singlepoint measurements of spectral reflectance or transmission in the visible and near-infrared wavelength ranges.^{2,5–8} This operation mode may not be suitable for some clinical applications where a sterile environment is required. For example, chronic wound management requires the accurate assessment of wound margin and appropriate detection of tissue viability without contamination to the disease area. Therefore, a noncontact method is preferred for cutaneous tissue StO₂ assessment. In addition, tissue StO₂ assessment in an imaging mode offers several clinical advantages over that of single-point detection. Considering that StO₂ varies spatially in not only suspicious lesions but also healthy tissues,⁹⁻¹² imaging tissue StO₂ distribution will help to characterize tissue heterogeneities, identify embedded lesions, and guide therapeutic procedures. Multiple multispectral, hyperspectral, and other advanced imaging techniques have been investigated in recent years for noncontact imaging of skin tissue StO_2 .^{13–17} These techniques collect skin tissue reflectance at multiple wavelengths, unmix the absorption components of oxy-hemoglobin and deoxy-hemoglobin, and derive the tissue StO₂ map by various reconstruction algorithms.^{15,18,19} However, tissue StO₂ maps acquired by various optical oximetry techniques are only relative indicators of tissue metabolism and viability. Accurate interpretation of tissue optical properties into absolute oxygenation levels is challenged by many experimental, physical, and biological limitations.²⁰⁻²² In this paper, we define various sources that cause deviations of the measured StO₂ values away from their actual values as artifacts. A variety of artifacts contribute to the lack of accuracy in tissue StO_2 measurements. The static artifacts introduce bias in StO_2 measurements. The dynamic artifacts (including motion artifacts) introduce temporal variations and fluctuations in StO₂ measurements. The static artifacts can be further divided into at least three categories. First, the physical principle limitations for an optical oximetry technique, such as the scattering of light in biologic tissue and the wavelength-dependent depth of light penetration in tissue, hinder the accurate interpretation of physiologic parameters from optical measurements. Second, variations in imaging hardware and test conditions, such as nonuniform illumination and specular reflectance conditions, introduce further artifacts in StO₂ measurement. Finally, intrapatient and interpatient variations in tissue physiologic conditions and chromophore concentrations also significantly affect the accuracy of StO₂ imaging. For example, skin pigments and chromophores other than oxy-hemoglobin and deoxy-hemoglobin contribute to heterogeneous background absorption at different

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skin sites and interpatient variation in skin color.²¹ Previous studies also show that oxy-hemoglobin, deoxy-hemoglobin, and scattering vary significantly for patients in different ethnic groups or for the same patient at different skin sites.^{23,24} In this paper, we focus on a multispectral imaging algorithm that may effectively reduce the StO₂ mapping artifacts introduced by other skin chromophores. At the current status of research, we have not studied the systemic methods for effective elimination of motion artifacts and other types of artifacts yet.

In an effort to minimize the artifacts for cutaneous tissue StO₂ measurement, various algorithms have been explored. Seo et al. used a linear model of oxy-hemoglobin and deoxyhemoglobin with a modulation of scattering and melanin to approximate the absorption spectrum of skin tissue in the wavelength range from 520 to 585 nm.²⁵ Stamatas et al. calculated the melanin concentration, corrected the absorption spectrum, and then it fitted into the absorption spectra of oxy-hemoglobin and deoxy-hemoglobin.^{26,27} Mansfield et al. analyzed the skin reflectance spectrum by fuzzy C-means clustering and then applied a four-term linear regression fitting with oxy-hemoglobin, deoxy-hemoglobin, offset, and slope terms.^{28–30} Sowa et al. reduced the bias from melanin by applying an orthogonal subspace projection method for hyperspectral image classification.^{31,32} Nishidate et al. determined the optical path lengths of different absorbers by inverse Monte Carlo simulation and applied the results to a multiple regression algorithm to find the fitting coefficients for oxy-hemoglobin, deoxy-hemoglobin, and melanin.³³ Jakovels and Spigulis used multiple Gaussian and exponential functions to fit the absorption spectra for oxy-hemoglobin, deoxy-hemoglobin, and melanin, and then approximated the skin tissue optical density spectrum from 500 to 700 nm by nonlinear least square approximation.¹⁵ Despite these advances, no systemic study has been carried out for quantitative study and evaluation of reliable algorithms for cutaneous tissue StO2 measurement at different skin conditions. Consequently, many existing hyperspectral and multispectral imaging devices can only provide relative information about tissue StO₂ vulnerable to skin conditions.

This paper reports a multispectral algorithm for noncontact, quantitative and high temporal resolution imaging of cutaneous tissue StO₂ with reduced measurement artifacts in comparison with conventional imaging methods. The multispectral algorithm is studied numerically in a simplified skin tissue model that only considers lipid and primary chromophores, such as oxy-hemoglobin, deoxy-hemoglobin, and melanin. Other chromophores, such as glucose, keratin, and bilirubin, are neglected at the current stage of research. The algorithm is also validated experimentally in a phantom system that consists of whole blood, lipid, and India ink. India ink is used to simulate different skin types since it has been adopted by many researchers to mimic the absorption properties of melanin.^{34,35} In Sec. 2.1, reflectance spectra of skin tissue at different StO₂, scattering, blood, and melanin conditions are simulated by an empirical light transport model with a semi-infinite boundary condition. A parameter of the second derivative ratio (SDR) is derived as the function of tissue StO₂ based on the simulated reflectance spectra and the method of wide-gap second derivative spectroscopy. An optimal set of wavelengths is defined so that the SDR value is monotonically determined by skin tissue StO₂ regardless of the tissue condition variations. In Sec. 2.2, the proposed multispectral imaging method is verified in tissue-simulating phantoms, where bovine blood, intralipid, and ink are mixed at different concentrations. Clinical feasibility of imaging skin tissue StO₂ is demonstrated on a healthy human subject following a postocclusive reactive hyperemia (PORH) protocol. During the PORH procedure, multispectral images are acquired at the specific set of wavelengths and the tissue SDRs are derived at each pixel for continuous mapping of tissue StO₂. To demonstrate that our StO₂ algorithm is less vulnerable to skin condition variations, 10 human subjects with different skin types were recruited and their skin StO₂ was measured by our multispectral imaging system. Statistical analysis shows that skin type does not significantly affect the StO₂ measurement by our system. To test the effect of additional background absorption on cutaneous StO₂ measurement, a portion of a human subject's forearm is painted with India ink and multispectral images are acquired within and outside the painted skin area at six designated wavelengths. The reconstructed StO₂ map is compared with that of a commercial OxyVu hyperspectral imaging system (HyperMed Imaging, Inc.).³⁰ Reconstructed StO₂ values within and outside the painted area are compared with those by the OxyVu system. The experimental results are discussed in Sec. 3, followed by conclusions drawn from these results.

2 Materials and Methods

2.1 Imaging Algorithms

Multispectral imaging of skin tissue StO_2 is achieved by successive algorithms of forward simulation and inverse reconstruction. The forward model simulates light transport in semi-infinite biologic tissue at different blood, scattering, melanin, and StO_2 conditions, with tissue parameters calibrated by Monte Carlo simulation. The inverse model reconstructs tissue parameters based on the analytical expression of SDRs at the designated wavelengths.

2.1.1 Forward simulation

Skin has a physiologic structure that can be divided into seven layers. Table 1 lists major tissue parameters of each skin layer, including thickness (d), volume fraction of hemoglobin in blood (r), and concentrations of blood (C_{blood}), water (C_{water}), fat (C_{fat}) , and melanin (C_{melanin}) .^{36–38} Traditionally, Monte Carlo modeling is used to simulate light transport in multilayered biologic tissue.³⁹ However, this method is time-consuming and is not appropriate for inverse reconstruction. Therefore, a simplified diffuse reflectance model with a semi-infinite boundary condition is introduced.⁴⁰ The absorption and scattering properties in this simplified model are averaged across different skin layers in order to fit in the semi-infinite boundary condition, while the multilayer tissue condition was used in Monte Carlo modeling. First of all, the absorption of any skin layer, μ_{ai} (i = 1to 7), is denoted as the sum of absorptions from different absorbers, including oxygenated-hemoglobin (μ_a^{HbO}), deoxygenatedhemoglobin (μ_a^{Hb}) , melanin (μ_a^{mel}) , water (μ_a^{water}) , fat (μ_a^{fat}) , and a baseline without any absorbers (μ_a^{base}):

$$\mu_{ai} = \mu_{ai}^{\text{HbO}} + \mu_{ai}^{\text{Hb}} + \mu_{ai}^{\text{mel}} + \mu_{ai}^{\text{water}} + \mu_{ai}^{\text{fat}} + \mu_{ai}^{\text{base}},$$
(1)

where

$$\mu_{ai}^{\text{HbO}} = 2.303 \varepsilon^{\text{HbO}} \cdot \text{StO}_2\left(\frac{C_{\text{Hb}}}{\text{M.W.}_{\text{Hb}}}C_i^{\text{blood}}\right),\tag{2}$$

Table 1 Properties of seven skin layers obtained and calculated from previous publications.^{36–38} *d* is the thickness of the layer; *r* is the volume fraction of hemoglobin in blood; C_{blood} , C_{water} , C_{fat} , and $C_{melanin}$ are the volume fractions of blood, water, fat, and melanin in each layer; *g*, *n*, and μ_s are anisotropy factor, refractive index, and scattering coefficient of each layer measured at 633 nm.

Skin layers	$d(\mu m)$	C_{blood}	r	$C_{\rm water}$	C_{fat}	$\mathcal{C}_{melanin}$	g	n	$\mu_s~(\mathrm{mm^{-1}})$
Stratum corneum	20	0	0	0.05	0	0	0.86	1.5	100
Living epidermis	80	0	0	0.2	0	0.02 (Caucasians); 0.13 (Asians); 0.3 (Africans)	0.8	1.34	45
Papillary dermis	160	0.04	0.1114	0.5	0	0	0.9	1.4	30
Upper blood net dermis	100	0.3	0.1114	0.6	0	0	0.95	1.34	35
Reticular dermis	1400	0.04	0.1114	0.7	0	0	0.8	1.4	25
Deep blood net dermis	100	0.1	0.1114	0.7	0	0	0.95	1.38	30
Hypodermis	28,140	0.05	0.1114	0.7	0.6	0	0.75	1.44	

$$\mu_{ai}^{\text{Hb}} = 2.303 \varepsilon^{\text{Hb}} \cdot (1 - \text{StO}_2) \left(\frac{C_{\text{Hb}}}{\text{M.W.}_{\text{Hb}}} C_i^{\text{blood}} \right), \tag{3}$$

$$\mu_{ai}^{\text{mel}} = \varepsilon^{\text{mel}} \cdot C_i^{\text{mel}},\tag{4}$$

$$\mu_{ai}^{\text{water}} = \epsilon^{\text{water}} \cdot C_i^{\text{water}} \cdot (1 - r_i C_i^{\text{blood}}), \tag{5}$$

$$\mu_{ai}^{\text{fat}} = \varepsilon^{\text{fat}} \cdot C_i^{\text{fat}} \cdot (1 - r_i C_i^{\text{blood}}), \tag{6}$$

$$\mu_{ai}^{\text{base}} = \varepsilon^{\text{base}} (1 - C_i^{\text{blood}} - C_i^{\text{mel}} - C_i^{\text{fat}}) (1 - r_i C_i^{\text{blood}}).$$
(7)

In the above equations, M.W._{Hb} and C_{Hb} are the molecular weight and total concentration of hemoglobin, respectively. ϵ^{HbO} , ϵ^{Hb} , ϵ^{water} , and ϵ^{fat} are the extinction coefficients of the corresponding absorbers obtained from previous studies.^{41,42} ϵ^{mel} is the absorption coefficient of melanin:³⁸ $\epsilon^{\text{mel}} = 1.7 \times 10^{12} \lambda^{-3.48} \text{ (cm}^{-1})$. ϵ^{base} is the absorption coefficient of skin free of any absorbers:⁴³ $\epsilon^{\text{base}} = 7.84 \times 10^8 \lambda^{-3.255} \text{ (cm}^{-1})$.

The absorption of the first layer stratum corneum is different from the other layers:^{36,37}

$$\mu_a^{\text{stratum}} = (1-3 \times 10^{-4} \lambda + 0.125 \varepsilon^{\text{base}}) \times 0.95 + \varepsilon^{\text{water}}$$
$$\times 0.05 \text{ (cm}^{-1}). \tag{8}$$

The overall absorption of the whole skin is calculated by taking the weighted average of the absorption of all layers.

$$\mu_a = \sum_{i=1}^7 \mu_{ai} \cdot f_i, \tag{9}$$

where f_i is defined as the weight factor for the absorption of a specific layer contributing to the overall absorption $(\sum_{i=1}^{7} f_i = 1)$. In general, the weight factor of the top layer is greater than that of the bottom layer because light attenuates exponentially with the tissue depth. These weight factors are determined by matching the simulated skin reflectance by the diffuse transport model with that by Monte Carlo modeling, as described in Appendix A. Different weight factors are derived for different skin conditions. The reduced scattering coefficient of the whole skin μ'_s is approximated by the following equation based on *in vivo* measurements:^{23,44}

$$\mu'_{s} = 1.1 \times 10^{12} \lambda^{-4} + 73.7 \lambda^{-0.22} \text{ (cm}^{-1}\text{)}.$$
(10)

The diffuse reflectance R is calculated by the following approximation algorithm:⁴⁰

$$R = R_{s} + \frac{(1 - R_{s})(1 - s)(1 - b_{1}s)}{1 + b_{2}s},$$

in which $s = \sqrt{\frac{\mu_{a}}{\mu_{a} + \mu_{s}'}}.$ (11)

In the above equations, $R_s = (n-1)^2/(n+1)^2$ is the specular reflectance; *n* is refractive index (n = 1.4 for typical biologic tissue); $b_1 = -0.34 - 1.1g + 0.68g^2$ and $b_2 = 4.11 - 2.77g + 2.27g^2$ are two constants; *g* is an anisotropy factor.

Finally, since our experiment used an optical filter with a bandwidth of 7 nm (Sec. 2.2.1), the resultant reflectance spectrum was corrected accordingly. The transmission of the filter with a bandwidth FWHM is assumed to be a Gaussian shape:

$$G(\lambda_0 - \lambda) = \frac{2\sqrt{\ln 2/\pi}}{\text{FWHM}} \cdot e^{-4\ln 2 \cdot \left(\frac{\lambda - \lambda_0}{\text{FWHM}}\right)^2}.$$
 (12)

The reflectance spectrum $R(\lambda_0)$ measured using this filter was calculated as the convolution of $G(\lambda_0 - \lambda)$ and the original reflectance spectrum $R(\lambda)$.

$$R(\lambda_0) = R(\lambda) \otimes G(\lambda_0 - \lambda) = \int_{\lambda_0 - 3\sigma}^{\lambda_0 + 3\sigma} R(\lambda) \cdot G(\lambda_0 - \lambda) d\lambda.$$
(13)

The above forward algorithm is used to simulate diffuse reflectance measurements at 450 skin conditions, including the combinations of five blood concentration levels ($C_{blood} = 0.8, 0.9, 1, 1.1, 1.2$ of normal C_{blood}), three melanin concentrations ($C_{melanin} = 0.02, 0.13$, and 0.3 in living epidermis, corresponding to Caucasians, Asians, and Africans), five scattering

levels ($\mu'_s = 0.8, 0.9, 1, 1.1, 1.2$ of typical μ'_s), and six oxygen saturation levels (StO₂ = 0, 20%, 40%, 60%, 80%, 100%). The diffuse reflectance R as a function of λ , C_{blood} , C_{melanin} , μ'_s , and StO₂ is simulated in the wavelength range from 500 to 650 nm. Figures 1(a), 1(b), and 1(c) show the skin reflectance spectra at three melanin concentrations and different levels of StO2, scattering, and blood concentration, respectively. According to Fig. 1, the dc offset of the skin reflectance of all wavelengths is affected by melanin concentration, scattering, and blood concentration. Among these three factors, the effect of melanin is over 10 times more significant than those of the other two factors. On the other hand, the shape of the spectrum is mainly determined by StO₂, regardless of variations in scattering, blood concentration, and melanin concentration. This observation indicates the possibility of using multispectral imaging to analyze the shape of reflectance spectra and mapping tissue StO₂ with a minimal measurement bias from other tissue components.

2.1.2 Reconstruction of tissue StO₂

A multispectral second derivative algorithm is developed to reconstruct skin tissue StO₂ based on the reflectance measurements at six designated wavelengths. The algorithm follows consecutive steps of calculating the absorption-scattering ratio, deriving wide-gap second derivative, obtaining an analytical expression of SDR, and curve fitting for StO₂ calculation. First, the ratio between absorption and reduced scattering coefficients ($r = \mu_a/\mu'_s$) is calculated by transforming Eq. (11):

$$r = \frac{\mu_a}{\mu'_s} = \frac{s^2}{1 - s^2},\tag{14}$$

where $b_1 \cdot s^2 - (b_1 + R_1 \cdot b_2 + 1) \cdot s + (1 - R_1) = 0$ and $R_1 = (R - R_s)/(1 - R_s)$.

After that, a wide-gap second derivative spectroscopic method is used to derive the second derivative of absorption-scattering ratio $rr(\lambda)''$ as a function of the wavelength λ and the wavelength gap $\delta \lambda$:⁴⁵

$$rr(\lambda)^{\prime\prime} = [r(\lambda + \delta\lambda) + r(\lambda - \delta\lambda) - 2r(\lambda)]/(2\delta\lambda).$$
(15)

The SDR is then expressed as the ratio between $rr(\lambda_1)''$ and $rr(\lambda_2)''$:

$$SDR(\lambda_1, \lambda_2) = \frac{rr(\lambda_1)''}{rr(\lambda_2)''} = \frac{r(\lambda_1 + \delta\lambda) + r(\lambda_1 - \delta\lambda) - 2r(\lambda_1)}{r(\lambda_2 + \delta\lambda) + r(\lambda_2 - \delta\lambda) - 2r(\lambda_2)}.$$
 (16)

After substituting the absorption-scattering ratio *r* with μ_a/μ'_s and neglecting higher-order terms as described in Appendix B, one can derive the approximated analytical expression of SDR(λ_1, λ_2):

$$\operatorname{SDR}(\lambda_{1},\lambda_{2}) = \frac{\mu_{s}'(\lambda_{2})}{\mu_{s}'(\lambda_{1})} \cdot \frac{\left[\left(\varepsilon^{\operatorname{HbO}}\right)_{\lambda_{1}}^{\prime\prime} - \left(\varepsilon^{\operatorname{Hb}}\right)_{\lambda_{1}}^{\prime\prime}\right]\operatorname{StO}_{2} + \left(\varepsilon^{\operatorname{Hb}}\right)_{\lambda_{1}}^{\prime\prime}}{\left[\left(\varepsilon^{\operatorname{HbO}}\right)_{\lambda_{2}}^{\prime\prime} - \left(\varepsilon^{\operatorname{Hb}}\right)_{\lambda_{2}}^{\prime\prime}\right]\operatorname{StO}_{2} + \left(\varepsilon^{\operatorname{Hb}}\right)_{\lambda_{2}}^{\prime\prime}}.$$
(17)

According to Eq. (17), SDR(λ_1, λ_2) varies with skin tissue StO₂ and the scattering coefficients at wavelengths λ_1 and λ_2 . In the case when these two wavelengths differ from each other by <10 nm, $\mu'_s(\lambda_1) \approx \mu'_s(\lambda_2)$ and SDR (λ_1, λ_2) is primarily determined by tissue StO₂. The wavelength pair of (λ_1, λ_2) can be optimized for one-to-one mapping between SDR and StO₂ with a minimal bias induced by different skin conditions. To identify the optimal wavelength pair, all the possible SDRs are calculated at different skin conditions for the wavelength pairs ranging from 500 to 650 nm and with the wavelength difference of 8 and 10 nm, respectively. The calculated SDRs are screened and the optimal wavelength set is found to be $\lambda_1 = 568$ nm, $\lambda_2 = 576$ nm, and $\delta \lambda = 24$ nm. According to Eq. (16), in total, six wavelengths are involved in the calculation of SDR $(568,576)_{(\delta\lambda=24)}$: 544, 552, 568, 576, 592, and 600 nm. Table 2 lists the SDR(568, 576) values at 12 representative skin tissue conditions selected from all the 450 conditions as described at the end of Sec. 2.1.1. The SDR level at 20% StO₂ is $0.872(\pm 0.004)$ averaged from 450 tissue conditions of different levels of blood concentration, melanin concentration, and scattering. Similarly, the SDR level at 80% StO₂ is $0.396(\pm 0.003)$. The overall relationship between SDR and StO_2 can be approximated by the following monotonic function:

$$StO_2 = 100(-1.4SDR^3 + 4.82SDR^2 - 5.66SDR + 2.38).$$
 (18)

Figure 2 plots the above monotonic function where skin tis-

sue StO₂ can be derived by taking the reflectance measurements (c) 0.5 Scattering levels Blood concentration levels



Fig. 1 Spectral reflectance of skin tissue at different conditions simulated by a forward algorithm. (a) Reflectance spectra at six StO_2 levels, three melanin concentrations, and fixed blood and scattering conditions. (b) Reflectance spectra at five scattering levels, three melanin concentrations, and fixed StO_2 and blood concentration conditions. (c) Reflectance spectra at five blood concentration levels, three melanin concentration levels, three melanin concentrations, and fixed StO_2 and scattering conditions.

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Table 2 Second derivative ratio $[SDR(568,576)_{(\delta\lambda=24)}]$ of 12 skin tissue conditions. The definitions of skin conditions were described in Sec. 2.1.1.

Different skin tissue conditions								
Blood level	Melanin level	Scattering level	StO ₂ (%)	SDR				
0.8	0.13	1	20	0.870				
1.2	0.13	1	20	0.872				
1	0.02	1	20	0.877				
1	0.3	1	20	0.868				
1	0.13	0.8	20	0.872				
1	0.13	1.2	20	0.871				
0.8	0.13	1	80	0.395				
1.2	0.13	1	80	0.397				
1	0.02	1	80	0.400				
1	0.3	1	80	0.392				
1	0.13	0.8	80	0.396				
1	0.13	1.2	80	0.396				

at the designated wavelengths (i.e., 544, 552, 568, 576, 592, and 600 nm) and calculating the SDR (568, 576) using Eqs. (15) and (16).

2.2 Experiments

2.2.1 Multispectral imaging system

A multispectral imaging system with a hyperspectral imaging capability is constructed for quantitative imaging of skin tissue StO₂. A Hamamatsu ORCA ER deep cooling CCD camera (Hamamatsu, Bridgewater, New Jersey) with a camera lens (Edmund optics, Barrington, New Jersey) is used for image acquisition. A Varispec liquid crystal tunable filter (Cambridge Research Inc., Cambridge, Massachusetts) with a working wavelength range of 400 to 720 nm and a bandwidth of 7 nm is mounted in front of the camera for multispectral imaging. An OSL1 fiber light source (Thorlabs, Newton, New Jersey) provides broadband optical illumination to the test object. A user interface is programmed by Labview Professional Development System (National Instruments, Austin, Texas) for equipment control, synchronization, and image acquisition tasks.

For quantitative assessment of skin tissue StO₂, multispectral images are acquired at 544, 552, 568, 576, 592, and 600 nm on skin tissue and on a National Institute of Standards and Technology traceable white diffuser. The reflectance map at each wavelength $R(\lambda)$ is calculated by taking the ratio of the skin and the diffuser measurements pixel by pixel. After that, the SDR value is calculated using Eqs. (15) and (16). Finally, the StO₂ map of skin tissue is obtained by Eq. (18).

Since the measurement of reflectance $R(\lambda)$ required multispectral images of both skin and the white diffuser, the intensity



Fig. 2 Oxygen saturation of skin StO₂ versus SDR(568,576)_($\delta\lambda$ =24). The error bar in the plot indicates the variation due to different blood concentration, melanin concentration, and scattering of skin.

of the light source has to be stable during the acquisition of these two groups of images. Thus, the light source was turned on for 10 min before any measurement was taken. An "auto exposure" function was developed in the Labview user interface and the camera exposure time was calibrated on the white diffuser to ensure that the gray level of the 8-bit image was always 235 at each wavelength.

2.2.2 Algorithm verification on tissue-simulating phantoms

The StO₂ algorithm is verified using a liquid blood phantom. The phantom is prepared by mixing fresh bovine blood, India ink (Sanford, Bellwood, Illinois), and 10% Intralipid (Henry Schein Inc., Melville, New York) in phosphate buffered saline (PBS, Fisher Scientific, Newton, New Jersey). First, a base phantom is prepared by mixing 2.5 mL of blood, 4 mL of Intralipid, and 30 µL of India ink in PBS solution to a total volume of 200 mL. Different skin tissue conditions are simulated by adding additional recipe components to the base phantom. For example, different levels of blood concentration, scattering, and skin darkness are simulated by adding 20% more blood, 20% more Intralipid, and 100% more India ink to the base phantom, respectively. The pH level of the phantom is adjusted to 7.4 by adding NaOH and HCl. The oxygenation level of the phantom is adjusted by the dropwise addition of sodium hydrosulfite at a concentration of 0.1 g/mL.

The phantom StO₂ is measured by the multispectral imaging method as described in the previous section and compared with a hyperspectral imaging method as described in Ref. 25. To eliminate the measurement artifact induced by ambient oxygen, the blood phantom is placed in a cardboard container and ventilated with argon gas. During the test, eight oxygenation plateaus ranging from ~100% to ~0% are achieved by the dropwise addition of sodium hydrosulfite. At each oxygenation plateau, the blood phantom is mixed to homogeneity by a stir bar. Hyperspectral images are acquired from 500 to 600 nm at a spectral resolution of 2 nm. Among them, images at 544, 552, 568, 576, 592, and 600 nm are used to reconstruct the StO₂ map following our multispectral imaging algorithm. In comparison, a hyperspectral imaging algorithm uses the whole reflectance spectra to reconstruct tissue StO_2 .

2.2.3 Algorithm verification on a human subject

The multispectral imaging algorithm is verified by human subject testing following a clinical protocol approved by the Ohio State University IRB (Protocol No: 2010H0017). The test measures the forearm tissue StO2 of a healthy male volunteer in response to a postocclusive reactive hyperemia (PORH) procedure.⁴⁶ The PORH protocol consists of a preocclusive baseline period of 2 min without compression, a supersystolic occlusion period of 2 min with a compression load of 50 mmHg above the systolic pressure applied to the arm by a pressure cuff, and a reactive hyperemia period of 2 min without compression. During the PORH test, multispectral images are acquired continuously. In the meantime, deep tissue oxygen saturation and skin oxygen tension on the same arm are measured by an OxiplexTS tissue spectrophotometer (ISS Inc., Urbana Champaign, Illinois) and a TCM transcutaneous oxygen monitor (Radiometer, Denmark), respectively. The skin tissue StO₂ levels obtained by multispectral imaging are compared with the OxiplexTS and the TCM measurements.

2.2.4 Measuring StO₂ of in vivo skin of different skin types

In order to evaluate the effect of skin type on oxygenation measurement, a total of 10 subjects with different skin types were recruited for continuous imaging of skin StO₂ following the PORH protocol. Among these 10 subjects, five are Caucasians and light tanned Hispanos, who have the lowest melanin levels; three are Asians with medium melanin levels; and two are darkly pigmented Asians with the highest melanin levels. The test procedure of each subject was the same as that described in Sec. 2.2.3. The influence of skin types on multispectral imaging of skin oxygenation was evaluated statistically and compared with the OxiplexTS tissue spectrophotometer and the TCM transcutaneous oxygen monitor measurements.

2.2.5 Measuring StO₂ of in vivo skin with additional background absorption

We have also tested *in vivo* whether the proposed multispectral imaging algorithm is able to reduce the StO_2 measurement bias owing to background absorption variations. A portion of the subject's forearm within the field of view is painted with 1% India ink dissolved in ethanol to simulate the darker skin color. Multispectral images are acquired with the field of view covering both the ink-painted area and the surrounding normal tissue. In comparison, the same experiment is repeated at the same skin location on the same subject using an OxyVu hyperspectral imaging system. The reconstructed StO_2 maps by these two imaging systems are compared side by side.

3 Results and Discussion

The multispectral imaging algorithm is verified by benchtop testing on a tissue-simulating phantom with different oxygenation levels. At each oxygenation level, multispectral images are acquired at the wavelengths of 544, 552, 568, 576, 592, and 600 nm. The diffuse reflectance measurements are averaged within three regions of interest (ROIs) for multispectral reconstruction of StO_2 . Figure 3 plots the reconstructed StO_2 values versus those derived from the previously reported



Fig. 3 Oxygen saturation of blood phantom measured by multispectral algorithm proposed in this paper versus that measured by hyperspectral imaging method of Ref. 25.

hyperspectral method.²⁵ According to the figure, our multispectral algorithm yields the StO_2 values linearly correlated with the hyperspectral algorithm. Moreover, StO_2 measurements at four simulated skin conditions (i.e., standard recipe, +20% lipid, +20% blood, and +100% ink) fall into a diagonal line, indicating that our StO_2 reconstruction algorithm is not vulnerable to variations in skin tissue parameters, such as blood concentration, scattering, and background absorption.

The multispectral imaging algorithm is also verified by human subject testing. Figure 4 shows the gray-scale image of the tested arm and the reconstructed StO₂ maps during the PORH procedure. According to the figure, the skin StO₂ level is in the range of 40 to 50% at the baseline condition before vascular occlusion [Fig. 4(b)], which is coincident with the previous study.⁴⁷ As the blood vessel is occluded, the skin StO₂ level drops to <10% [Fig. 4(c)]. After the occlusion is released, the skin StO₂ level increases to >60% due to hyperemia [Fig. 4(d)]. At the end of the test, the StO₂ level resumes its baseline value [Fig. 4(e)]. The StO₂ dynamics detected during a PORH procedure is consistent with the previous report.¹⁷

The StO₂ dynamics of cutaneous tissue during a PORH procedure is compared with simultaneous measurements of deep tissue oxygenation and transcutaneous tissue oxygen tension. Figure 5(a) illustrates the test protocol where a baseline period of 2 min is followed by a suprasystolic occlusion period of 2 min, and then a reactive hyperemia period of 2 min. Figure 5(b) plots the corresponding dynamic changes of skin tissue StO₂ calculated by averaging 10 selected ROIs in the StO₂ oxygenation map as shown in Fig. 4. Figures 5(c) and 5(d) are simultaneous measurements of deep tissue oxygenation and transcutaneous tissue oxygen tension. According to Fig. 5, the dynamic changes of cutaneous tissue StO₂ coincide well with those of deep tissue oxygenation and transcutaneous oxygen tension.

Skin tissue StO_2 images were acquired on 10 human subjects with different skin types to evaluate the robustness of our multispectral imaging technique at different skin conditions. The tests followed the same PORH protocol as described before. Figure 6 shows the averaged skin StO_2 levels for 10 subjects measured by our multispectral imaging system under the normal condition without vascular occlusion pressure. For each human subject, 10 measurements were carried out to



Fig. 4 Picture of the arm (a). Skin oxygenation map before (b), during (c), after vascular occlusion (d), and at the end of the test (e).



Fig. 5 Postocclusive reactive hyperemia (PORH) test protocol (a), and the history of cutaneous tissue oxygenation (b), deep tissue oxygenation (c), and transcutaneous tissue oxygen tension (d) measured by multispectral imaging, tissue oximeter, and TCM device, respectively.

generate 10 oxygen saturation maps. In each oxygen saturation map (500 × 500 pixels), 10 ROIs of 25 × 25 pixels were randomly drawn in the skin area without hair. In order to avoid measurement artifacts due to arm curvature, these ROIs were all selected away from the edge of the arm. Then the StO₂ for a subject was calculated by averaging StO₂ values from a total of $10 \times 10 = 100$ ROIs. In order to statistically evaluate the influence of skin darkness on StO₂ measurement, an analysis of variance (ANOVA) test was performed. The resultant *p* value is 0.29, indicating that skin type does not significantly affect the StO₂ measurement for our multispectral imaging system.



Fig. 6 Averaged skin StO_2 of 10 subjects under normal conditions measured by multispectral imaging with our algorithm. The subjects were arranged in the order of skin melanin levels with 1 being the lowest to 10 being the highest. Fitzpatrick scale of each subject is also listed.

Figure 7 shows the averaged skin tissue StO2 (%) acquired by our multispectral imaging system, the deep tissue StO_2 (%) acquired by the Oxiplex tissue oximeter, and the transcutaneous oxygen tension pO2 (mmHg) acquired by the TCM device during the PORH procedure on 10 subjects. The error bars indicate the variations among different subjects. During the baseline period (first 2 min), the skin StO₂ (%) measured by multispectral imaging system is $42.5(\pm 6.49\%)$; the deep tissue StO₂ (%) measured by the tissue oximeter is $55.0(\pm 13.6\%)$; and the transcutaneous oxygen tension pO2 (mmHg) measured by the TCM oxygen monitor is $62.0(\pm 19.7\%)$. These data show that the % standard deviation of StO_2 (%) measured by our system is much less than that of the other two measurements. Furthermore, the same ANOVA test was performed on the other two measurements to statistically evaluate the influence of skin type. The p values are both <0.01 for deep tissue StO_2 (%) measured by the tissue oximeter and transcutaneous oxygen tension pO2 (mmHg) measured by the TCM oxygen monitor. The above result shows that the measurement by our multispectral imaging system is less biased by skin type than those of the other two oximetry devices.



Fig. 7 Measurements of 10 subjects of different skin types during PORH: (a) average skin StO₂ (%) measured by our system, (b) deep tissue StO₂ (%) measured by Oxiplex tissue oximeter, and (c) transcutaneous oxygen tension pO_2 (mmHg) measured by TCM oxygen monitor.

The effect of additional background absorption on cutaneous StO_2 measurement is also tested on a human subject. A portion of the subject's forearm is painted with ink to simulate a darker skin. Cutaneous StO_2 maps are acquired by both our multispectral imaging system and an OxyVu hyperspectal imaging system. Figures 8(a) and 8(b) show the photographic image of the forearm tissue with the central area painted with ink and the corresponding StO_2 map reconstructed by our multispectral algorithm. Two square-shaped ROIs are selected within and outside the ink-painted area, with the cutaneous StO_2 levels averaged as 41.6 and 38.1%, respectively, resulting in a measurement difference of 3.5%. In comparison, the same test

performed by an OxyVu system on the same arm [Figs. 8(c) and 8(d)] shows an averaged cutaneous StO_2 level of 62.7% within the ink-painted skin area and 43.0% outside the area, resulting in a measurement difference of 19.7%. This result presumably indicates that the proposed multispectral algorithm effectively reduces the artifact of additional background absorption on cutaneous StO_2 measurement.

The multispectral imaging technique reported in this paper was developed primarily for the noncontact measurement of cutaneous tissue StO_2 since the proposed wide-gap second derivative spectral imaging algorithm is sensitive to tissue oxygenation changes. Considering that tissue oxygenation is



Fig. 8 Oxygenation of skin partially painted with ink. (a) and (c) Picture of the arm with a portion of skin painted with ink. (b) Oxygenation map generated by our multispectral imaging system. (d) Oxygenation map generated by a commercial hyperspectral imaging system.

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an important indicator of skin tissue viability, the proposed StO₂ imaging technique may be used in many clinical applications, such as plastic surgery, organ transplantation, and wound healing. The limitation of this imaging technique is that it may eliminate the lower-order heterogeneities of skin tissue, leading to the diminished structural features. By projecting different patterns of illumination, defining different imaging algorithms, and choosing different wavelength sets, it is also possible to use the multispectral imaging system for noncontact imaging of skin tissue hemoglobin concentration, scattering, and melanin concentration. Characterization of these additional functional parameters may help to identify angiogenesis, bleeding, size change of cell nucleus, and other tissue anomalies. A Monte Carlo model was used to simulate light propagation in skin tissue. Our Monte Carlo simulation result showed that the averaged interrogation depth is 1 to 2 mm and is as deep as reticular dermis, depending on the blood concentration, melanin, and scattering level in the skin. The simulation results were then fit into a semi-infinite diffusion model in order to derive an analytical expression of SDR for effective reconstruction of tissue StO₂. The above approximations may introduce a systemic error in StO₂ reconstruction. First of all, the Monte Carlo model assumed homogenous tissue properties in each layer while typical skin tissue has a lateral heterogeneity. Second, the diffusion approximation assumes a turbid medium where the reduced scattering coefficient is much greater than the absorption coefficient. This is not always true in multilayer skin tissue. Finally, both the epidermis and the dermis layers of skin are very thin, leading to the breakup of diffuse approximation for light with a short penetration depth. In order to minimize the effect of these artifacts on the accuracy of StO₂ measurement, our multispectral oxygen imaging algorithm defined the correlation between the Monte Carlo model and the analytical solutions in advance based on the simulated data.

Two major measurement artifacts involved in our in vivo experiments of skin tissue StO2 imaging are motion artifact and the artifact caused by the arm curvature. In our study, the motion artifact was minimized by keeping the subject's arm steady during the entire test period since a complete cycle of multispectral imaging at six wavelengths took around 2 s. An image processing algorithm was used to examine whether the arm location was maintained stable among the acquired six images. The measurement was considered invalid if there is a shift of the arm location. In terms of the artifact caused by the arm curvature, it mainly presents at the edge of the arm, as shown in Fig. 4. Currently, we do not have an effective calibration method to correct this artifact. To further reduce this artifact, we may consider profilometry correction using a tissue-simulating phantom⁴⁸ or by wrapping a paper mask on the curve arm for calibration.⁴⁹

Although wide-gap second derivative spectroscopy has been proposed for tissue oxygenation measurement for a while,⁴⁵ we are the first to implement it for multispectral imaging of cutaneous tissue StO₂. Compared with probe technologies that provide single-point measurements of oxygenation, our multispectral imaging technique has the advantage of noncontact measurement as well as less intrapatient variability since it provides oxygenation mapping of an area of skin. Compared with other hyperspectral imaging systems, our technique has the advantage of higher temporal resolution since the StO₂ reconstruction algorithm requires multispectral images at only a small number of wavelengths. We have previously explored

the SDR algorithm based on four wavelengths.⁵⁰ In comparison with our previous work, six wavelengths are selected in our current imaging algorithm so that more spectral characteristics of oxy- and deoxy-hemoglobin can be captured in SDR calculation. These six wavelengths are optimized for not only improved sensitivity to tissue oxygenation changes but also clinical utility and engineering feasibility. Currently, our multispectral imaging system is able to achieve an overall temporal resolution of 2 s per measurement and a typical postprocessing speed of 30 s per measurement, depending on the spatial resolution of the StO₂ map. These figures can be significantly improved by optimizing and customizing the engineering design of the imaging system.

In this study, skin StO_2 of 10 subjects with different skin types was measured to evaluate the effect of skin type on our multispectral measurement of oxygenation. Since the reflectance spectra of all StO_2 levels are almost overlapped with each other at the high melanin level, it is possible that the measurement bias is hard to reduce for patients with very dark skin. Extensive human tests with different skin colors are necessary in order to evaluate the robustness and clinical utility of the proposed imaging algorithm.

4 Conclusion

This paper reports a multispectral imaging algorithm for cutaneous tissue StO₂ mapping with minimal bias induced by skin conditions. A mathematical model is first established to simulate skin tissue reflectance spectra at different conditions, including skin oxygen saturation, blood concentration, tissue scattering, and melanin concentration. Based on the simulation results, the SDR parameter is derived and the optimal wavelength set of 544, 552, 568, 576, 592, and 600 nm is obtained for multispectral reconstruction of tissue StO₂ regardless of tissue variations. The proposed multispectral imaging algorithm is verified by benchtop and in vivo experiments. The benchtop experiment shows that the reconstructed StO₂ map is not vulnerable to variations in blood concentration, scattering, and background absorption levels. The in vivo experiment shows that the proposed multispectral imaging algorithm is able to monitor cutaneous tissue StO_2 at a sampling rate of 0.5 Hz. The test of 10 human subjects with different skin types shows that skin type does not have significant effect on our StO₂ measurement. We also show that the reconstructed StO₂ map by the multispectral imaging algorithm is less affected by skin background absorption compared with the OxyVu hyperspectral imaging system. Our experimental results indicate that the proposed multispectral imaging technique has the potential for quantitative imaging of skin tissue StO2 dynamics with minimal artifacts at different skin conditions. Further in vivo validation tests are necessary before implementation of the proposed multispectral imaging algorithm in a clinical setting.

Appendix A: Absorption Weight Factors Determination by Matching Diffuse Transport Model and Monte Carlo Model

As described in Eq. (9) in Sec. 2.1.1, a weight factor f was used to calculate the averaged overall absorption of the whole skin. This weight factor indicates how the absorption of a specific layer contributes to the overall absorption of skin. The weight factor was found by matching the skin spectral reflectance simulated by the diffuse transport model with the reflectance simulated by the Monte Carlo model. The diffuse transport model has been described in Sec. 2.1.1, and the Monte Carlo model is described here.

A Monte Carlo simulation of photon migration in a sevenlayer tissue model was used to simulate spectral reflectance of skin of different conditions. A Monte Carlo Simulation Package, developed in Ref. 39, was used with the following input parameters for each tissue layer: thickness d, refractive index n, anisotropy factor g, absorption $\mu_a(\lambda)$, and scattering $\mu_s(\lambda)$. The values of d, n, and g of each layer are listed in Table 1. The values of n and g here are measured in 633 nm and they are considered as constants for approximation in this simulation. The absorption coefficient $\mu_a(\lambda)$ of each layer was calculated using the same algorithm as in the diffuse model [Eqs. (1) to (8)]. The scattering $\mu_s(\lambda)$ of each layer, which was calculated differently compared to the diffuse model, is explicitly described here.

The scattering coefficient μ_s of each layer measured at 633 nm is listed in Table 1. But there is no published data of μ_s versus λ for each skin layer. Thus, the following method was used to approximate the wavelength-dependent μ_s for each skin layer. Tseng et al.²³ reported that the reduced scattering coefficient of *in vivo* skin can be expressed as

$$\mu'_{S}(\lambda < 600 \text{ nm}) = a_{1}\lambda^{-b_{1}}(\text{cm}^{-1}) \quad (b_{1} = 1.35 \text{ to } 1.6), \quad (19)$$

$$\mu'_{s}(\lambda > 600 \text{ nm}) = a_{2}\lambda^{-b_{2}}(\text{cm}^{-1}) \quad (b_{2} = 1.00 \text{ to } 1.15).$$
(20)

These two equations were solved using the μ_s (633 nm) values listed in Table 1 to get parameters a_1 and a_2 (fixed $b_1 = 1.5$ and $b_2 = 1.07$) for each skin layer.

The scattering of the hypodermis, which is very different from other layers, is given by 44

$$\mu_s' = 1050.6 \cdot \lambda^{-0.68}. \tag{21}$$

Finally, the *d*, *n*, *g*, $\mu_a(\lambda)$, and $\mu'_s(\lambda)$ of each layer were fed into the Monte Carlo simulation package. 200 K photons were used for a single simulation. The skin spectral reflectance of different skin conditions are simulated using the above Monte Carlo method.

Then the weight factors for seven skin layers in the diffuse transport model will be determined by matching the skin reflectance simulated by the diffuse model with the reflectance simulated by the Monte Carlo method. This problem can be represented by the nonlinear least square optimization problem below. The optimal weight factors are found when the square error between the reflectance of two models is minimized.

Decision variable: weight factor
$$f_i$$
 $(i = 1 \text{ to } 7)$
Objective function: Minimize $\sum_{\lambda} {R[\mu_a(f_i)] - R_{MC}}^2$
Constraints: $\sum_i f_i = 1$ and $0 \le f_i \le 1$ $(i = 1 \text{ to } 7)$

 $R[\mu_a(f_i)]$ is the spectral reflectance calculated using the diffuse model, and $R_{\rm MC}$ is the spectral reflectance simulated using the Monte Carlo method.

This optimization problem was solved by a trust-regionreflective algorithm using MATLAB[®]. Different weight factors are derived for different skin conditions.

Appendix B: Analytical Expression of $SDR \sim StO_2$

Taking the second derivative of $r(=\mu_a/\mu'_s)$ as a function of wavelength λ , one will get

$$\frac{\mathrm{d}^2 r}{\mathrm{d}\lambda^2} = A \cdot \frac{\mathrm{d}^2 \mu_a}{\mathrm{d}\lambda^2} + B \cdot \frac{\mathrm{d}\mu_a}{\mathrm{d}\lambda} + C \cdot \mu_a,\tag{22}$$

where

$$A = (1/\mu'_s); \quad B = -(2/\mu'^2_s) \left(\frac{\mathrm{d}\mu'_s}{\mathrm{d}\lambda}\right);$$
$$C = \frac{1}{\mu'^2_s} \left[\frac{2}{\mu'_s} \left(\frac{\mathrm{d}\mu'_s}{\mathrm{d}\lambda}\right)^2 - \frac{\mathrm{d}^2\mu'_s}{\mathrm{d}\lambda^2}\right].$$

Since $\mu'_s = a\lambda^{-4} + b\lambda^{-0.22}$ ($a = 1.1 \times 10^{12}$; b = 73.7) for *in vivo* skin, one can find that the second and third terms in Eq. (22) are much smaller than the first term. Thus,

$$\frac{\mathrm{d}^2 r}{\mathrm{d}\lambda^2} \approx \frac{1}{\mu_s'} \cdot \frac{\mathrm{d}^2 \mu_a}{\mathrm{d}\lambda^2}$$

Since $\mu_a = \sum_{i=1}^7 \mu_{ai} \cdot f_i$, where $\mu_{ai} = \mu_{ai}^{\text{HbO}} + \mu_{ai}^{\text{Hb}} + \mu_{ai}^{\text{mel}} + \mu_{ai}^{\text{hat}} + \mu_{ai}^{\text{base}}$, substituting each absorption term using Eqs. (2) to (7), one will get

$$\begin{split} \mu_a \ &= F^{\text{Hbt}} \cdot \varepsilon^{\text{HbO}} \cdot \text{StO}_2 + F^{\text{Hbt}} \cdot \varepsilon^{\text{Hb}} \cdot (1 - \text{StO}_2) \\ &+ F^{\text{mel}} \cdot \varepsilon^{\text{mel}} + F^{\text{water}} \cdot \varepsilon^{\text{water}} + F^{\text{fat}} \cdot \varepsilon^{\text{fat}} + F^{\text{base}} \cdot \varepsilon^{\text{base}}, \end{split}$$

where

$$\begin{split} F^{\text{Hbt}} &= \left(2.303 \frac{C_{\text{Hb}}}{\text{M.W.}_{\text{Hb}}} \sum_{i=1}^{7} C_{i}^{\text{blood}} \cdot f_{i}\right), \\ F^{\text{mel}} &= \sum_{i=1}^{7} C_{i}^{\text{mel}} \cdot f_{i} \\ F^{\text{water}} &= \sum_{i=1}^{7} C_{i}^{\text{water}} \cdot (1 - r_{i}C_{i}^{\text{blood}}) \cdot f_{i}, \\ F^{\text{fat}} &= \sum_{i=1}^{7} C_{i}^{\text{fat}} \cdot (1 - r_{i}C_{i}^{\text{blood}}) \cdot f_{i} \\ F^{\text{base}} &= \sum_{i=1}^{7} (1 - C_{i}^{\text{blood}} - C_{i}^{\text{mel}} - C_{i}^{\text{fat}})(1 - r_{i}C_{i}^{\text{blood}}) \cdot f_{i}. \end{split}$$

The absorption of fat and water are small terms. Thus,

$$\mu_a \approx F^{\text{Hbt}} \cdot \varepsilon^{\text{HbO}} \cdot \text{StO}_2 + F^{\text{Hbt}} \cdot \varepsilon^{\text{Hb}} \cdot (1 - \text{StO}_2) + F^{\text{mel}} \cdot \varepsilon^{\text{mel}} + F^{\text{base}} \cdot \varepsilon^{\text{base}}.$$

Thus,

$$\frac{\mathrm{d}^2 r}{\mathrm{d}\lambda^2} \approx \frac{1}{\mu_s'} \left[F^{\mathrm{Hbt}} \frac{\mathrm{d}^2 \varepsilon^{\mathrm{HbO}}}{\mathrm{d}\lambda^2} \cdot \mathrm{StO}_2 + F^{\mathrm{Hbt}} \frac{\mathrm{d}^2 \varepsilon^{\mathrm{Hb}}}{\mathrm{d}\lambda^2} \cdot (1 - \mathrm{StO}_2) \right. \\ \left. + F^{\mathrm{mel}} \frac{\mathrm{d}^2 \varepsilon^{\mathrm{mel}}}{\mathrm{d}\lambda^2} + F^{\mathrm{base}} \frac{\mathrm{d}^2 \varepsilon^{\mathrm{base}}}{\mathrm{d}\lambda^2} \right].$$

It is found that $(d^2 \varepsilon^{\text{mel}})/(d\lambda^2)$, $(d^2 \varepsilon^{\text{base}})/(d\lambda^2) \ll (d^2 \varepsilon^{\text{HbO}})/(d\lambda^2)$, $(d^2 \varepsilon^{\text{HbO}})/(d\lambda^2)$ [since $\varepsilon^{\text{mel}}(\lambda)$ and $\varepsilon^{\text{base}}(\lambda)$ have an exponential decay shape]. Thus,

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$$\frac{\mathrm{d}^2 r}{\mathrm{d}\lambda^2} \approx \frac{F^{\mathrm{Hbt}}}{\mu'_s} \left[\frac{\mathrm{d}^2 \varepsilon^{\mathrm{HbO}}}{\mathrm{d}\lambda^2} \cdot \mathrm{StO}_2 + \frac{\mathrm{d}^2 \varepsilon^{\mathrm{Hb}}}{\mathrm{d}\lambda^2} \cdot (1 - \mathrm{StO}_2) \right].$$
(23)

Thus, the second derivative ratio (SDR) is

$$SDR = \frac{rr(\lambda_{1})''}{rr(\lambda_{2})''} = \frac{\mu'_{s}(\lambda_{2})}{\mu'_{s}(\lambda_{1})} \cdot \frac{[(\epsilon^{\text{HbO}})'_{\lambda_{1}} - (\epsilon^{\text{Hb}})'_{\lambda_{1}}]StO_{2} + (\epsilon^{\text{Hb}})'_{\lambda_{1}}}{[(\epsilon^{\text{HbO}})'_{\lambda_{2}} - (\epsilon^{\text{Hb}})'_{\lambda_{2}}]StO_{2} + (\epsilon^{\text{Hb}})'_{\lambda_{2}}'}.$$
 (24)

This is the approximated analytical expression of SDR for any λ_1 and λ_2 .

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