In vivo time-resolved autofluorescence measurements to test for glycation of human skin

Jennifer Blackwell
Kamal M. Katika
Laurent Pilon
University of California, Los Angeles
Mechanical and Aerospace Engineering Department
Los Angeles, California 90095

Abstract. We present an evaluation of time-resolved fluorescence measurements on human skin for screening type 2 diabetes. In vivo human skin is excited with a pulse diode at 375 nm and pulse width of 700 ps. Fluorescence decays are recorded at four different emission wavelengths: 442, 460, 478, and 496 nm. Experiments are performed at various locations, including the palms, arms, legs, and cheeks of a healthy Caucasian subject to test single-subject variability. The fluorescence decays obtained are modeled using a three-exponential decay. The variations in the lifetimes and amplitudes from one location to another are minimal, except on the cheek. We compare the fluorescent decays of 38 diabetic subjects and 37 nondiabetic subjects with different skin complexions and of ages ranging from 6 to 85 yr. The average lifetimes for nondiabetic subjects were 0.5, 2.6, and 9.2 ns with fractional amplitudes of 0.78, 0.18, and 0.03, respectively. The effects of average hemoglobin A1c (HbA1c) from the previous 4 yr and diabetes duration are evaluated. While no significant differences between the fluorescence lifetimes of nondiabetic and diabetic subjects are observed, two of the fractional amplitudes are statistically different. Additionally, none of the six fluorescence parameters correlated with diabetes duration or HbA1c. One of the lifetimes as well as two of the fractional amplitudes differ between diabetic subjects with foot ulcers and nondiabetic subjects.

Keywords: skin autofluorescence; time-resolved fluorescence; advanced glycated end-products; diabetes; foot ulcers.

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1 Introduction

Fluorescence is the physical phenomenon in which light is emitted by a substance as a result of excited electrons returning to their ground states after absorption of excitation light. Biological tissues contain several endogenous fluorophores such as nicotinamide adenine dinucleotide (NADH), aromatic amino acids such as tryptophan, and structural proteins including collagen and elastin. The optical properties of these fluorophores respond to the environment and metabolic status of the tissue, thus making fluorescence spectroscopy a valuable tool to study the state of biological tissues. Fluorescence spectroscopy consists of measuring the fluorescent intensity emitted by fluorophores after being excited by light at a suitable wavelength and can be performed in either a steady-state or a time-resolved manner.

Steady-state measurements typically involve measuring either the fluorescence intensity over a range of wavelengths for a fixed excitation wavelength (emission spectra) or the fluorescence intensity at a particular wavelength for a range of excitation wavelengths (excitation spectra). Time-resolved measurements consist of measuring the lifetime of the fluorophores and are much more complex than steady-state spectral measurements due, in part, to the instrumentation required. Steady-state measurements are relatively easy to perform. However, they present some limitations. First, they are intensity dependent and therefore are sensitive to excitation light intensity, optical losses in the experimental setup, and absorption and scattering by tissue. Second, the fluorescence intensity is integrated over time and, therefore, information associated with the dynamics of the fluorescence process is lost. Time-resolved measurements, on the other hand, capture the transient decay, which depends on the relative concentrations and lifetimes of the fluorophores contributing to the fluorescence signal. They are also extremely sensitive to the local biochemical environment such as the pH and oxygenation, which may differ in healthy and diseased tissue. In addition, the fluorophore lifetimes are independent of absolute intensity and so do not change with variations in excitation intensity or optical losses from hemoglobin absorption. Finally, fluores-
cence lifetime measurements enable the discrimination of fluorophores with overlapping emission spectra but different lifetimes.\(^3\) For example, various tryptophan residues in proteins have similar emission spectra but different lifetimes and thus can be distinguished by time-resolved fluorescence measurements.\(^4\)

Time-resolved fluorescence techniques can be performed in either the frequency domain or the time domain.\(^5\) Frequency-domain techniques involve exciting the sample with a sinusoidally modulated source, which could be a continuous wave laser or a flash lamp, and measuring the phase shift of the emitted light at multiple excitation frequencies to compute the fluorescence lifetime(s). Alternatively, time-domain measurements typically use ultrashort pulsed light to excite the sample and measure the emitted fluorescence as a function of time, thus enabling one to compute the lifetime of the fluorophore. Note that the pulse width must be of the same order of magnitude as or shorter than the fluorescence lifetime. Time-domain measurements are preferred in a clinical setting, because they can be obtained in short acquisition times compared to frequency-domain measurements\(^7\) and they are not affected by ambient light at the collection site.\(^8\)

The detection of the diabetic state has become an important issue recently as the number of people with diabetes is rising and approximately one-third of people affected by type 2 diabetes in the United States are undiagnosed.\(^9\) The current methods of diagnosis consist of detecting glucose levels in the blood or plasma.\(^9\) Even with screening, the sensitivity is much less than ideal.\(^7\) Once diabetes has been diagnosed, glucose levels and hemoglobin A1c (HbA1c) are currently utilized to follow the efficacy of treatment. The latter is a measure of percent glycated hemoglobin and reflects average glycemic control over the antecedent 3 to 4 months.\(^8\) As a test of short-term glycation, HbA1c does not adequately reflect the accumulation of glycated tissue over a period of years, which may result in diabetic complications. To directly measure tissue glycation, a punch biopsy must be taken,\(^9\) making it difficult to determine a patient’s risk for complications. Thus, a simple, rapid method of screening is being pursued to enable early diagnosis of the disease and to follow the efficacy of therapy and the effects of long-term hyperglycemia on tissue.

Spectroscopy-based methods are attractive for diabetes screening and monitoring because they can noninvasively detect some markers of disease progression. For example, the accumulation of advanced glycation end-products (AGEs) in tissue can potentially be observed via spectroscopy. AGEs are the final products of the complex chemical reaction, known as the Maillard reaction, between sugars such as glucose and proteins, lipids, and nucleic acids.\(^9\) While the accumulation of these in tissues is a natural process and serves as a marker of aging,\(^10\) elevated blood glucose levels in diabetic individuals result in a more rapid accumulation of AGEs. Some AGEs, such as pentosidine, are known to fluoresce and their fluorescence parameters due to age, skin color, and gender was also tested. Also, 18 diabetic subjects with foot ulcers were compared with 18 age-matched healthy subjects to test for differences due to the development of complications. Finally, the possible origin of these fluorescence lifetimes is discussed. This is the first study to report the application of time-resolved fluorescence to the detection of diabetes and/or monitoring for the development of diabetic complications. This work is of particular importance to studies on skin autofluorescence measurements for noninvasive glucose sensing\(^11\) and the design of implantable glucose sensors based on fluorescence.\(^12\) It would also serve as a baseline for using time-resolved autofluorescence of human skin to monitor other diseases and conditions such as skin cancer, wounds, and ulcers.

### 2 Current State of Knowledge

#### 2.1 Time-Resolved Fluorescence Measurements on Tissue

**In vivo** time-resolved fluorescence measurements have come into prominence only recently and a limited number of studies have been reported in the literature, which is summarized in Table 1. For example, Pradhan et al.\(^16\) used fluorescence lifetime measurements to study breast tissue. They made use of laser pulses of 100 fs pulse width at 310 nm and a streak camera to obtain a 2-D map of the fluorescence lifetimes. It was shown that time-resolved fluorescence measurements at an emission wavelength around 340 nm can be used to distinguish malignant tumors from nonmalignant breast tissues.

Fluorescence lifetime measurements have also been used to study tissue during endoscopy. Glanzmann et al.\(^17\) described a time-resolved fluorescence instrument based on pumped laser excitation sources and a streak camera coupled to a spectrograph as a detector. The instrument was used to measure the fluorescence decays of endogenous fluorophores and of δ-aminolevulinic acid (ALA)-induced protoporphyrin IX (PPIX) in an excised human bladder with carcinoma. The authors were able to correctly distinguish healthy tissue, which appeared abnormal in steady-state spectroscopy, from cancerous tissue. The instrument was then used to perform in situ characterization during endoscopy to obtain time-resolved spectra of tissue fluorescence of human bladder, bronchi, and esophagus.\(^18\) Further studies were conducted by the same group\(^19\) to test the viability of fluorescence lifetime as a new contrast parameter between normal and malignant tissue in...
Table 1: Experimental studies of time-resolved fluorescence spectroscopy on human tissues.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Excitation Wavelength (nm)</th>
<th>Emission Wavelength (nm)</th>
<th>Spectroscopy Method</th>
<th>Sensing Region</th>
<th>In Vivo/In Vitro</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>337.1</td>
<td>N.A.</td>
<td>time domain using an avalanche photodiode (APD) and fast oscilloscope</td>
<td>human skin</td>
<td>in vivo</td>
</tr>
<tr>
<td>16</td>
<td>310</td>
<td>340 and 440</td>
<td>time domain using a streak camera</td>
<td>breast tissue</td>
<td>in vitro</td>
</tr>
<tr>
<td>17</td>
<td>337.1, 425, 476, 500</td>
<td>500–680</td>
<td>time domain using a streak camera</td>
<td>human bladder</td>
<td>in vitro</td>
</tr>
<tr>
<td>18</td>
<td>337.1, 480</td>
<td>375–515, 525–615</td>
<td>time domain using a streak camera</td>
<td>bladder, bronchi, esophagus</td>
<td>in vivo</td>
</tr>
<tr>
<td>19</td>
<td>406</td>
<td>430–680</td>
<td>time domain using a streak camera</td>
<td>bronchii</td>
<td>in vivo, endoscopy</td>
</tr>
<tr>
<td>20</td>
<td>514.5</td>
<td>550–650</td>
<td>frequency domain</td>
<td>bronchii</td>
<td>in vivo, endoscopy</td>
</tr>
<tr>
<td>21</td>
<td>337, 400</td>
<td>550</td>
<td>time domain using a gated intensified diode array</td>
<td>esophagus</td>
<td>in vivo, endoscopy</td>
</tr>
<tr>
<td>22-24</td>
<td>446</td>
<td>510–700</td>
<td>time domain using TCSPC</td>
<td>human ocular fundus</td>
<td>in vivo</td>
</tr>
<tr>
<td>25,26</td>
<td>337.1</td>
<td>370–510</td>
<td>time domain using a transient digitizer</td>
<td>aorta, and its components</td>
<td>in vitro</td>
</tr>
<tr>
<td>27,28</td>
<td>405</td>
<td>N.A.</td>
<td>time domain using time gating</td>
<td>skin tumors</td>
<td>in vivo</td>
</tr>
<tr>
<td>29</td>
<td>415</td>
<td>N.A.</td>
<td>time domain using time gating</td>
<td>rabbit knee joint and human tooth</td>
<td>in vivo</td>
</tr>
<tr>
<td>30-32</td>
<td>750–850</td>
<td>N.A.</td>
<td>time domain using TCSPC</td>
<td>human skin</td>
<td>in vivo</td>
</tr>
</tbody>
</table>

the bronchi. However, they were unable to obtain a contrast in the fluorescence lifetime or the spectrum between normal and moderately dysplastic tissue in the spectral region between 510 and 650 nm. Mizeret et al. 20 also studied the feasibility of performing endoscopic fluorescence lifetime measurements using frequency modulation techniques in real time in a clinical setting. The instrument made use of an excitation source at 514.5 nm and a detection unit consisting of CCD units with image intensifier tubes. Fluorescence lifetime values for healthy bronchial tissue were reported. Pfefer et al. 21 also performed in vivo fluorescence lifetime and spectral measurements on patients undergoing routine endoscopic surveillance for Barrett’s esophagus. A time-domain instrument consisting of a laser pulsing at 337 nm with a pulse width of 3 ns and an APD coupled to an oscilloscope was used in the experiments. The excitation wavelengths used were 337 and 400 nm and the emission was collected around 550 nm. Results showed that low-risk tissue and high-grade dysplasia could not be differentiated based on the observed data.

Furthermore, Schweitzer et al. 22,23 performed fluorescence lifetime studies on the human ocular fundus. They made use of TCSPC techniques with a pulsed laser of 300-ps pulse width at a wavelength of 457.9 nm and various emission wavelength ranges. They obtained a lifetime of 1.5 ns in the parapapillary region of the eye and a lifetime of 5 ns in the optical disc, which they attributed to collagen. They also showed that breathing 100% oxygen affects the fluorescence lifetimes and speculated that the lifetime of fluorophores present in the retinal pigment epithelium depends on the oxygen supply.

Fluorescence lifetime measurements have also been performed on tissue in vitro. For example, Maarek et al. 24 used time-domain in vitro fluorescence lifetime measurements and spectroscopy for diagnosis of atherosclerotic lesions. The instrument consisted of an excitation laser at 337 nm and a pulse width of less than 1 ns and a gated imaging system. An exogeneous marker was used to induce selective accumulation of PPIX in proliferative tissue. Tumors were identified using lifetime imaging due to the longer fluorescence lifetime.
of PPIX (18 ns) compared to that of healthy tissue (≈10 ns). Siegel et al.29 also used FLIM techniques, utilizing a pulsed laser at 415 nm and a pulse width less than 100 fs and a gated optical intensifier to image the autofluorescence from various biological tissue in vitro, such as animal tissue, knee joints, and human teeth. Using this instrument they were able to distinguish between tendon, bone, and surrounding tissue, which was not otherwise possible with white light and steady-state fluorescence measurements. Similarly, they showed that various regions in an extracted human tooth can be clearly distinguished using FLIM unlike reflectance and steady-state fluorescence measurements.

Other studies have performed time-resolved fluorescence measurements on human skin. Konig et al.,30 Konid and Riemann,31 and Riemann et al.32 used 3-D autofluorescence lifetime imaging and investigated various tissues of patients with psoriasis, nevi, dermatitis, basalioma, and melanoma in vitro as well as the skin of healthy subjects in vivo. Their system was able to achieve a subcellular spatial resolution and a temporal resolution of 250 ps. The authors analyzed the fluorescence decays from images obtained at various depths of the skin and reported lifetimes of around 1.85 ns at the stratum corneum and a lifetime of about 2.4 ns at a depth of 50 μm within the stratum spinosum. The same group also reported using multiphoton imaging of skin at an excitation wavelength of 740 nm with a 150-ps resolution.33 A biexponential fit was used to recover a fast lifetime of 0.4 ns and a slow lifetime of 2.4 ns at the stratum spinosum in the epidermis. Additionally, Pitts and Mycek3 described a time-domain instrument with both spectral and temporal resolution to measure fluorescence lifetimes and emission spectrum in vivo and in vitro. The instrument was based on a pumped dye laser with a pulse width of 4 ns as an excitation source and an APD along with a 1-GHz oscilloscope as a detector. Fluorescence lifetime measurements were performed on human skin in vivo at an excitation wavelength of 337 nm and an emission wavelength of 460 nm. The time-resolved data were fitted to a double-exponential decay, yielding lifetimes of 0.938 and 5.3 ns. The longer lifetime was attributed to the emission from collagen. The instrument was also used to measure fluorescence lifetimes of biomolecules present in human cells.

### 2.2 Optical Noninvasive Screening of Type 2 Diabetes

A few groups have applied spectroscopic studies to the noninvasive screening of type 2 diabetes. For example, Meierwaldt et al.34 observed steady-state autofluorescence measurements of skin on the arm and lower leg of both diabetic and healthy subjects. Diabetic and healthy subjects were age- and sex-matched. Using excitation spectrum between 300 and 420 nm and emission spectrum between 420 and 600 nm, they found a positive correlation between skin autofluorescence intensity with age, diabetes duration, and mean HbA1c levels from the previous year. The fluorescence intensity also correlated well with the concentrations of AGEs in tissue biopsies, such as pentosidine and carboxymethyllysine. This demonstrated the capability of steady-state fluorescence measurements to distinguish diabetics from healthy subjects. The study’s primary focus was monitoring disease progression and evaluating the patient’s risk for diabetic complications. In a later study, the same group showed that autofluorescence also correlated with severity of diabetic neuropathy35 and vascular damage in type 2 diabetic subjects.36 However, the effect of skin color was not addressed in these studies as nearly all subjects recruited were Caucasian. In fact, the researchers reported that data from dark-skinned subjects had to be discarded, because of nearly complete absorption of excitation light by the skin.36 Otherwise, the steady-state measurements must be adjusted to account for skin complexion.

Brown et al.37 clinically evaluated the use of a near-IR (NIR) spectroscopic device for its ability to correctly identify diabetic and nondiabetic subjects. The NIR wavelength region under observation in the study was from 1250 to 2500 nm. The diffusely reflected radiation was collected and the tissue absorbance was retrieved. Data from both diabetic and healthy subjects were collected and the device was found to have a sensitivity and specificity of 77.7% and 70%, respectively. Thus, this system’s sensitivity and specificity are comparable to the fasting plasma glucose (FPG) test. Although a definite chemical or physiological explanation for the difference between diabetic and nondiabetic patients is yet to be elucidated, they speculate that structural changes in skin due to chronic hyperglycemia contribute to the differentiation between diabetic and nondiabetic subjects. However, the same group’s previous work38 suggested that NIR spectroscopy does not detect AGEs directly.

Recently, Maynard et al.39 reported using a spectroscopic device for type 2 diabetes screening utilizing steady-state fluorescence as well as diffuse reflectance measurements. The fiber optic probe illuminated skin from the forearm of subjects using LED lights centered at 375, 405, 420, 435, and 460 nm. A spectrometer and CCD array then collected the light reflected and/or emitted from the skin and detected it. The spectra were corrected for skin pigmentation, hemoglobin content, and light scattering as well as age to obtain an “intrinsic” dermal fluorescence spectra. These data were then used to determine a likelihood that a subject belonged to the abnormal class, or that they were either prediabetic or diabetic. This method of screening was compared to the FPG test and A1C testing. Assuming a clinically relevant specificity of 77.4%, their method showed a sensitivity advantage of 28.8 and 17.1% versus FPG and A1C testing, respectively. The group also reported that the sensitivity of their device was not significantly different between subjects of lighter skin and those with of darker skin. Thus, their study demonstrates the promise of steady-state fluorescence for type 2 diabetes screening if factors such as skin color and age can be adjusted for in measurements. Time-resolved fluorescence may hold promise in diabetes detection by distinguishing different fluorophores emitting at the same wavelength and potentially being able to detect AGEs directly.

### 3 Materials and Methods

#### 3.1 Experimental Apparatus

The study reported here made use of the TCSPC technique to measure fluorescence lifetimes of skin. A schematic of the experimental setup is shown in Fig. 1. Excitation and emission light were channeled by means of liquid light guides attached to a sensing head. Liquid light guides were used instead of commonly used fiber optics as their transmission
and coupling efficiency are higher compared with fiber optics. Both light guides had a diameter of 8 mm, and the excitation light guide was held 1.59 cm above the skin. The receiving end of the collection liquid light guide was held abut against the excitation liquid light guide and angled at 45 deg to the normal. The sensing head also served to shield the sample surface from ambient light. A total surface area of 6.41 cm² was exposed to excitation light.

The excitation source used was a light emitting diode or LED (PLS 370 diode, PicoQuant GmbH) with its center wavelength at 375 nm with a spectral width of around 10 nm. The full width half maximum of the diode pulses at a repetition rate of 2.5 MHz and average power of 4.8 μW per pulse was 664 ps. The LED was driven by a diode driver (PDL-800 B diode driver, PicoQuant GmbH, Berlin, Germany) at a repetition rate of 5 MHz for all measurements, producing an estimated energy fluence rate of 2.07 × 10⁻⁶ W/cm² at the skin surface for a pulse energy of 2.65 pJ and pulse width of 664 ps. A liquid light guide designed for the UV to visible part of the spectrum (77628, from Spectra-Physics, U.S.A) carried light from the LED to the sensing head. Another liquid light guide designed for the visible part of the spectrum (77631, from Spectra-Physics, U.S.A), was used to collect the fluorescence emission and carry it to a monochromator (SpectraPro-150, Acton Research Corporation, U.S.A.). A photomultiplier tube (PMT) assembly (PMA-M 165 from PicoQuant GmbH) was coupled to the monochromator on the other end. The monochromator slits on the entry and exit of the light path were adjusted manually to maintain a bandwidth of 3 nm.

Finally, for data acquisition, the “sync” output signal from the diode driver and the “start” signal from the PMT assembly were fed to the respective channels on the data acquisition board (TimeHarp 200 from PicoQuant GmbH), via standard 50-Ω coaxial cables. The “sync” output from the diode driver provided the signal to synchronize the timing electronics on the data acquisition board mounted on a personal computer (PC). The time resolution of our device was estimated to be 200 ps.

### 3.2 Experimental Procedure and Parameters

The measurement procedure and parameters were arrived at by conducting preliminary studies on the palms, arms, cheeks, and feet of a 31-yr-old male Caucasian subject. The fluorescence lifetimes were measured at four different emission wavelengths: 442, 460, 478, and 496 nm. The first wavelength was chosen based on studies reporting that collagen fluorescence exhibited a maximum at around 440 nm when excited by light at 370 nm, corresponding to collagenase-digestible collagen cross-links, including AGEs on collagen. The remaining wavelengths were arbitrarily chosen to cover the spectral range up to 500 nm.

To measure the fluorescence lifetimes, the monochromator was set to the emission wavelength of interest and the monochromator slits were adjusted to match that particular wavelength. The probe was then placed on the skin surface and the LED was turned on.

The PMT shutter was opened once the sensing head was firmly placed on the skin surface and then the histograms were collected using the TimeHarp software. The probe was held in contact with the palm until 1500 to 5000 counts were acquired in the peak channel. This number was arrived at by considering a trade-off between the duration of the experiments and the signal-to-noise ratio. The duration of a single measurement ranged from 5 to 20 min, based on the sensing location and emission wavelength. It was observed that the time taken for each measurement was shortest on the palm, possibly due to the low concentration of melanin. Moreover, higher emission wavelengths required longer durations to achieve these counts due to a lower intensity of the fluorescence. Figure 2 shows a typical sample of in vivo time-resolved fluorescence measurement on human skin along with the χ² and associated residuals indicating the goodness of the fit. For an acceptable fit, the χ² must be around 1.0 and the residuals should be randomly distributed around 0.

The instrument response function (IRF) was measured by placing the probe on a Teflon block for the same number of maximum photon counts of at least 1500 as in the in vivo measurements, with the PMT collecting at 403 nm. The deconvolution software FluoFit version 4.0 was used to recover the lifetimes from the measured fluorescence decays. This software utilized the Marquardt-Levenberg algorithm with a weighting factor W(i) defined as

\[
W(i) = 1/\sqrt{D(i)},
\]

where \(D(i)\) is the number of counts contained in channel \(i\). An attempt was made to describe the decays with a single-exponential, a biexponential, and a triexponential model. Only the biexponential and the triexponential models gave a value of \(\chi^2\) around 1.0, which indicated a good fit. However, the biexponential model was rejected in favor of the more complicated triexponential model due to the presence of oscillations in the plot of the residuals compared to more random residuals for the triexponential model, as illustrated in Fig. 2. Thus, the fluorescence decay could be modeled as...
\[ F(t) = A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2) + A_3 \exp(-t/\tau_3), \]

where \( A_i \) and \( \tau_i \) are the fractional amplitude and lifetime, respectively, for decay \( i \). Additionally, note that initial guess lifetimes were entered to enable faster convergence. A total of 75 subjects participated in this study. The population characteristics for all subjects are shown in Table 2. The palms were chosen as the sensing location for these subjects due to the shorter duration of the experiments as well as ease of access. The first measurement at 442 nm was performed on the left palm and subsequent measurements at higher wavelengths were performed alternately on the right and left palms.

In addition to time-resolved fluorescence measurements, mean HbA1c values from the previous 4 yr were recorded for diabetic subjects by obtaining medical records with Institutional Review Board (IRB) approved consent (UCLA IRB #04-12-012-02). Subjects were also recruited from the local VA hospital with VA hospital IRB approved consent (VA IRB #PCC 2005-030449). All consent forms were signed prior to testing and collection of medical record data. All available data for each subject from the previous year were averaged and two subsets of these data were compared, one with “high” average HbA1c values and the other with “low” average HbA1c values. Data on duration of diabetes were obtained directly from the subject being tested.

### 4 Results and Discussion

Table 3 presents a statistical summary of lifetimes \( \tau_1 \), \( \tau_2 \), and \( \tau_3 \) and the corresponding fractional amplitudes \( A_1 \), \( A_2 \), and \( A_3 \) for six measurements on the left palm of the same healthy subject at an emission wavelength of 442 nm. These experiments were performed over a period of 1 month at different times of the day. The coefficient of variation, which represents the variation in measurements, is less than 5% in all cases.

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**Table 2** Population characteristics of healthy and diabetic subjects recruited for this study.

<table>
<thead>
<tr>
<th></th>
<th>Male/Female</th>
<th>Age Range (Yr)</th>
<th>African-American</th>
<th>Asian</th>
<th>Caucasian</th>
<th>Hispanic</th>
<th>Persian</th>
<th>South Asian</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy</td>
<td>22/15</td>
<td>10–85</td>
<td>4</td>
<td>6</td>
<td>18</td>
<td>5</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Diabetic</td>
<td>33/5</td>
<td>6–80</td>
<td>10</td>
<td>1</td>
<td>17</td>
<td>10</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
except for the parameters $\tau_1$ and $A_3$, for which it reaches 7 and 15%, respectively. This shows that measurements performed with this device are highly reproducible for a single subject. Additionally, six consecutive measurements, each lasting 10 min, were performed on the same hand within an hour. No noticeable changes in lifetimes, fractional amplitudes, or acquisition times were measured, suggesting that photobleaching was not induced during the measurements.

A set of three experiments were also performed on the inner arm of the same subject over the same period of time and at an emission wavelength of 442 nm. Then, the Student’s $t$-test was used to test differences in the fluorescence lifetimes obtained on the arm and the palm for this subject. A two-sample $t$-test assuming unequal variances was performed. In all cases, a level of significance of $p < 0.05$ was used. It was found that there were no significant differences in the fluorescence lifetime $\tau_1$. However, the $t$-test seemed to indicate that there was a difference in the fluorescence lifetimes $\tau_2$ and $\tau_3$ obtained on the arm and the palm at an emission wavelength of 442 nm. Finally, a measurement was also taken on the cheeks and the feet of the same subject. Figure 3 shows the fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and their corresponding fractional amplitudes $A_1$, $A_2$, and $A_3$ obtained as a function of emission wavelength and sensing location, namely, the palms, the arms, the cheeks, and the feet. The error bars in the plots of the fluorescence lifetimes represent asymptotic standard errors in the fluorescence lifetimes obtained using the Fluofit software package. Figure 3 indicates that the variations in the lifetimes from one location to another are minimal except on the cheek. Moreover, it was found in the course of the measurements that the time taken for each measurement was the least on the palm, possibly due to the low concentration of melanin. Therefore, the palm was chosen as the sensing location for all the other subjects.

Figure 4 shows the raw data of the fluorescence lifetimes and their corresponding amplitudes at an emission wavelength of 442 nm for 37 healthy subjects and 38 diabetic subjects of various ages, genders, and skin complexion obtained at the VA and UCLA. The error bars represent asymptotic standard errors in the fluorescence lifetimes obtained using the Fluofit software package. To further analyze the fluorescence lifetime data, various factors varying from one individual to another were considered, namely, (1) skin complexion (2) gender, and (3) age. The statistical analysis used was the Student’s $t$-test assuming unequal variance. Based on this test, a two-tailed $p$ value, defined as the probability that the difference between two groups randomly occurred during the experiment, was assumed to be statistically significant.

### 4.1 Effect of Skin Complexion and Gender

Figure 5 shows the fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and their corresponding amplitudes $A_1$, $A_2$, and $A_3$ as a function skin complexion for healthy subjects at an emission wavelength of 442 nm. The figure indicates that the fluorescence lifetimes do not significantly vary for different skin complexities. Using the Fisher’s exact test, the fluorescence lifetimes and fractional amplitudes of healthy subjects with dark complexion and light complexion were compared. The light-complexioned group consisted of 10 Caucasian subjects, age 42 ± 12 yr, while the dark-complexioned group was composed of 10 Hispanic, Persian, South Asian, and African-American subjects, age 39 ± 20 yr. Based on these test results, no statistically significant difference in average fluorescence lifetimes and fractional amplitudes exists between these two groups ($p > 0.05$ in all cases). Note that measurements took longer for dark skin. However, skin complexion had no effect on the lifetimes and fractional amplitudes.

Moreover, Fig. 6 shows the fluorescence lifetimes and amplitudes as a function of gender for healthy subjects in the same age group at the same emission wavelength of 442 nm. Again, there are no noticeable trends in the fluorescence lifetimes with respect to gender. The comparison of the fluorescence lifetimes and fractional amplitudes of 15 healthy female subjects, average age 42 ± 12 yr, and 22 healthy male subjects, average age 48 ± 20 yr, did not produce statistically significant differences ($p > 0.05$ in all cases). Thus, Figs. 5 and 6 as well as the statistical analysis suggest that fluorescence lifetimes and fractional amplitudes do not vary with skin complexion and gender. The same conclusions were made for other emission wavelengths (data not shown).

### 4.2 Effect of Age

Finally, an attempt was made to model the fluorescence lifetimes of healthy subjects shown in Fig. 4 as a function of age using linear regression. The square of the correlation coefficient $R^2$ was less than 0.03 for all three lifetimes, thus indicating that variations in fluorescence lifetimes with age cannot be described by a linear model (data not shown). Moreover, the small regression coefficients in all the cases also indicate that the three lifetimes do not depend on the age of the subject. Student’s $t$-tests were performed on all six parameters (fluorescence lifetimes and fractional amplitudes) comparing young and old healthy subjects. In addition, 15 subjects in both the young and old groups were compared with average ages of 31 ± 7 and 63 ± 11 yr, respectively. No statistically significant difference was detected for any of the parameters ($p > 0.05$). Thus, the lifetimes and fractional amplitudes obtained for healthy subjects are statistically independent of age, gender, and skin complexion. Table 4 presents the mean values and the standard deviations of the fluorescence lifetimes and their corresponding fractional amplitudes for healthy subjects, providing reference values for healthy human skin. This
Fig. 3 Effect of sensing location on fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and fractional amplitudes $A_1$, $A_2$, and $A_3$ as a function of emission wavelength for a 31-yr-old Caucasian male subject at excitation wavelength of 375 nm.
Fig. 4 Effect of age on the fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and fractional amplitudes $A_1$, $A_2$, and $A_3$ for healthy and diabetic subjects at excitation and emission wavelengths of 375 nm and 442 nm, respectively.
Fig. 5 Effect of skin complexion on the fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and fractional amplitudes $A_1$, $A_2$, and $A_3$ as a function of age for healthy subjects at excitation and emission wavelengths of 375 nm and 442 nm, respectively.
Fig. 6 Effect of gender on the fluorescence lifetimes $\tau_1$, $\tau_2$, and $\tau_3$ and fractional amplitudes $A_1$, $A_2$, and $A_3$ as a function of age for healthy subjects at an excitation and emission wavelengths of 375 nm and 442 nm, respectively.
clearly demonstrates the advantage of time-resolved fluorescence over steady-state fluorescence in that it is insensitive to age and skin complexion.

### 4.3 Effect of Diabetes

Table 4 summarizes the experimental data collected for 18 healthy subjects and 27 age-matched diabetic subjects. Their fluorescence lifetimes and fractional amplitudes were compared by means of an unpaired Student’s t-test. Only the fractional amplitudes $A_1$ and $A_2$ produced statistically significant differences in the means of the diabetic and the healthy patients. This was based on the criteria that $p < 0.05$ for the difference between average values to be considered significant. Additionally, the relative contributions of each lifetime to the overall intensity of diabetic and healthy subjects were compared using the Student’s t-test. The fractional contribution of a single lifetime to the steady-state intensity can be defined as:

$$I_i = (\tau_i A_i) / \sum_{j=1}^{n} (\tau_j A_j).$$

However, no statistically significant differences were found for any of the three fluorescence lifetimes considered. The same conclusions were reached for the other emission wavelengths considered (data not shown).

Further testing was performed on only the diabetic subjects, comparing duration of diabetes as well as average HbA1c values over the past 4 yr. To test the effect of diabetes duration, the diabetic subjects were divided into two groups with an average of $11 \pm 5$ yr since diagnosis in the shorter duration group and $24 \pm 7$ yr in the longer duration group. However, no statistically significant difference was found between these two groups when performing the unpaired Student’s t-test. Similarly, any correlation with average HbA1c from the previous 4 yr was tested by dividing the diabetic subjects into two groups: one with a low average HbA1c of $6.9 \pm 0.7\%$ and higher average HbA1C of $9.8 \pm 0.9\%$. No significant difference was detected between these two groups ($p > 0.05$ for all cases).

Additionally, 18 diabetic subjects with foot ulcers, aged $62 \pm 8$ yr, were compared with 18 age-matched healthy subjects, average age $57 \pm 15$ yr, using the Student’s t-test. Diabetic subjects with foot ulcers were specifically chosen because many diabetic subjects who develop ulcers also have other comorbidities associated with diabetes.\(^{42,43}\) Thus, this group represents subjects whose AGE accumulation is expected to be much higher than other diabetic subjects. Not only were statistically significant differences noted between the two groups for fractional amplitudes $A_1$ and $A_2$, but also differences in lifetime $\tau_1$ between the two groups was also statistically significant, albeit with a $p$ value of 0.038. All the test results are presented in Table 5. Given the long half-life of collagen in skin\(^{34}\) ($\sim 15$ yr), one can expect changes in the time-resolved fluorescence properties of skin to correlate more with the occurrence of diabetic complications than with HbA1c values. This measurement would indicate the long-
term accumulation of AGEs and other oxidative damage. The rate at which tissue accumulates AGEs is also dependent on various genetic or metabolic factors.\(^{45}\) Therefore, not all diabetic subjects are as susceptible to the development of complications, in spite of poor blood glucose control and high HbA1c levels. The development of complications cannot be predicted by conventional treatment monitoring such as blood glucose measurements and HbA1c tests. Finally, nine diabetic subjects without foot ulcers were compared with 18 age-matched healthy subjects using the Student’s t-test. No statistically significant differences between the two groups were detected in any of the parameters except in amplitude $A_2$ ($p$ value of 0.045). This indicates that the differences in the amplitude $A_1$ and $\tau_1$ between 18 healthy and 27 diabetic subjects reported earlier might be mainly due to the diabetic subjects with foot ulcers.

While a previous study by Meerwaldt et al.\(^{34}\) showed that steady-state autofluorescence measurements correlated with age, diabetes duration, and mean HbA1c of the previous year, this study did not find any significant relationships between the fluorescence lifetimes and amplitudes and these variables. Meerwaldt et al.\(^{34}\) used a polychromatic excitation source emitting at wavelengths ranging from 300 to 420 nm and fluorescence was measured over the entire window of 420 to 600 nm. This is in contrast with this study, where the excitation source was a monochromatic diode emitting at 375 nm (spectral width of around 10 nm) and fluorescence was measured at 442, 460, 478, and 496 nm with spectral width of 3 nm. It is speculated that these differences in the excitation wavelength might be the source of the different conclusions obtained in the two studies. This study should be repeated for a different excitation wavelength between 300 and 420 nm and emission wavelength between 420 and 600 nm to corroborate results by Meerwaldt et al.\(^{34}\)

4.4 Identification of Fluorophores

In an earlier study on fluorescence lifetime measurements on human skin,\(^{33}\) two fluorescence lifetimes of 0.4 and 2.4 ns at the stratum spinosum were reported using multiphoton excitation at 740 nm, equivalent to single-photon excitation at 370 nm. In the presented study, similar values were found when the decays were fit to a triexponential model. This suggests that the fluorophores with the high fractional amplitudes $A_1$ and $A_2$ in our measurements originate, at least in part, from the stratum spinosum. Our measurements are detecting an- other fluorophore as well, perhaps deeper in the tissue, which would explain the lower contribution of this fluorophore to the overall fluorescence signal. The similarities between our results and the previously reported data help to validate our findings in spite of using different multieponential models. Moreover, the shorter two lifetimes $\tau_1$ around 0.4 ns and $\tau_2$ around 2.7 ns observed in our measurements were similar to the lifetimes of free and protein-bound NADH reported in literature.\(^{31,46}\) Indeed, Schneckenburger et al.\(^{46}\) studied the autofluorescence from cultured endothelial cells and determined the fluorescence lifetime of free and protein-bound NADH to be between 0.4 and 0.5 and between 2.0 to 2.5 ns, respectively. In addition, König and Riemann\(^{31}\) also reported a fluorescence lifetime of 2.7 ns at a depth of 50 μm within the skin, which could possibly be attributed to bound NADH.

Based on these studies, it is speculated that the two shorter fluorescence lifetimes $\tau_1$ and $\tau_2$ correspond to free and protein-bound NADH, respectively. This suggests that the differences in fractional amplitudes $A_1$ and $A_2$ observed between diabetic and healthy subjects are related to the metabolic states of their tissues rather than directly detecting the presence of glycation products. However, as the presence of glycation products is known to increase tissue oxidative stress,\(^{47,48}\) they may have an indirect effect on our measurements.

Furthermore, it is speculated that the third observed average lifetime $\tau_3$ equal to 9.2 ns for healthy subjects corresponds to collagen cross-links formed\(^{34,38}\) by AGEs. However, our data on diabetic subjects seem to contradict this speculation. The longest lifetime $\tau_3$ and the corresponding fractional amplitude $A_3$ are not significantly different between diabetic and healthy subjects. Given the known correlation between AGEs and diabetics and skin steady-state autofluorescence,\(^{34}\) one would expect the fractional amplitude of the third lifetime to increase with increasing AGE accumulation, if the lifetime could be attributed to AGEs. With elevated concentration of AGEs in the skin, their fluorescence signal would be stronger and, thus, contribute a higher fractional intensity. Further in vitro experimentation is required to precisely determine the origin of this lifetime.

5 Conclusions

We presented the design and implementation of an experimental apparatus based on TCSPC assembled to measure autofluorescence lifetimes of human skin. With an excitation wavelength of 375 nm and pulse width of 700 ps, fluorescence was detected at wavelengths of 442, 460, 478, and 496 nm. The device was tested on various locations of a 31-yr-old healthy male Caucasian subject as well as the palms of 37 healthy subjects and 38 diabetic subjects. The following conclusions were drawn:

1. In tests on all subjects, the fluorescence decay was fit to a multieponential model with three average fluorescence lifetimes of 0.5, 2.6, and 9.2 ns associated average fractional amplitudes of 0.78, 0.18, and 0.03, respectively, for nondiabetic subjects.

2. A statistical analysis of the fluorescence lifetimes obtained on the inner arm, palm, feet, and the cheeks of a 31-yr-old healthy male Caucasian subject showed differences between sensing locations for $\tau_1$ and $\tau_3$, while there were no differences in $\tau_1$.

3. It was found that the fluorescence lifetimes measured on the palms were independent of the skin complexion and the gender of the subjects.

4. The only statistically significant differences between diabetic and nondiabetic subjects were between fractional amplitudes $A_1$ and $A_2$ of the two groups.

5. No differences in any parameters were observed between subjects due to HbA1c values or diabetes duration.

6. A comparison between diabetic subjects with foot ulcers and healthy subjects showed statistically significant differences not only in fractional amplitudes $A_1$ and $A_2$, but also in lifetime $\tau_3$.

7. Finally, the only parameter that showed a statistically significant difference between the diabetic subjects without
foot ulcers and age-matched healthy subjects was $A_2$. The fluorescence lifetime data of healthy subjects is of particular interest as a baseline especially for using fluorescence lifetimes of NADH to noninvasively sense glucose concentrations as well as monitoring diseases and skin conditions such as skin cancer, wounds, and ulcers.

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