Optical coherence tomography to investigate optical properties of blood during coagulation

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Abstract. This study investigates the optical properties of human blood during the coagulation process under statics using optical coherence tomography (OCT). OCT signal slope (OCTSS) and 1/e light penetration depth ($d_{1/e}$) were obtained from the profiles of reflectance versus depth. Results showed that both OCTSS and $d_{1/e}$ were able to sensitively differentiate various stages of blood properties during coagulating. After 1 h clotting, OCTSS decreased by 47.0%, 15.0%, 13.7%, and 8.5% and $d_{1/e}$ increased by 34.7%, 29.4%, 24.3%, and 22.9% for the blood samples at HCT of 25%, 35%, 45%, and 55%, respectively. The slope of $d_{1/e}$ versus time ($S_r$, $\times 10^{-4}$ mm/s), associated with clot formation rate decreased from 6.0 ± 0.3, 3.7 ± 0.5 to 2.3 ± 0.4 with the increasing of HCT from 35%, 45%, to 55%. The clotting time ($t_c$) from the $d_{1/e}$ evolution curves was estimated to be 1969 ± 92 s, 375 ± 12 s, 455 ± 11 s, and 865 ± 47 s for the blood of 25%, 35%, 45%, and 55%. This study demonstrates that the parameters ($t_c$ and $S_r$) from the variations in $d_{1/e}$ had better sensitivity and smaller standard deviation. Furthermore, blood hematocrit affecting backscattering properties of blood during coagulation was capable of being discerned by OCT parameters. It is concluded that OCT is a potential technique to quantify and follow the liquid-gel transition of blood during clotting.

Keywords: optical coherence tomography; hematocrits; blood coagulation; optical coherence tomography signal slope; 1/e light penetration depth.

1 Introduction

Blood coagulation is a natural process that can be both beneficial and life-threatening for the human body. It allows the maintenance of hemostasis after vascular injury, but it can also cause deep vein thrombosis and heart stroke. Inappropriate blood coagulation represents the leading cause of morbidity and mortality in the United States, accounting for roughly 40% of all deaths. Most existing clinical tests of hemostasis such as the prothrombin time, partial thromboplastin, and activated clotting time tests, are performed on plasma containing all the coagulation factors. They are assays designed to identify specific defects in clot formation, but cannot identify hypercoagulability. These assays are limited by the variability in reagents, methodologies, and artificial conditions of the assay that do not precisely reproduce the physiologic event. Furthermore, they do not provide any information about the quality of the clot or the dynamics of its formation.

During the last decades, many disciplines of applied research have tried to understand the complex phenomenon of the blood clotting process, namely mechanical, optical, and ultrasound characterization methods for in vitro tests. The viscosity and elasticity detection methods produce variable results depending on the shape of the fibrin clumps, i.e., the quantity or viscosity of the fibrin and on the strength of the magnetic field of the element. Furthermore, it is impossible to detect coagulation unless the viscosity increases to above a specific level. The turbidity detection method involves mixing the plasma specimen and coagulation reagent, and then measuring changes in the transmitted light or the scattered light during the process of the plasma coagulating. Small changes in the transmitted or scattered light can be detected even with a low level of fibrin, and it is hence free from the shortcoming of the viscosity detection method. Nevertheless, few in vitro tests that are also used in clinical practice are performed on blood samples. Recent studies have revealed an important need to develop accurate and global standard in vitro coagulation tests using blood samples. These new tests should complement existing standard tests performed with plasma samples. Due to the nature of acoustic waves, ultrasound measurement techniques have the major advantages of allowing the use of blood samples. Measurements of ultrasound backscattering, sound velocity, and ultrasound attenuation have been used to analyze the process of blood coagulation and clot formation. In optics, blood coagulation was studied in a special beaker by analyzing the properties of the reflected waves. The blood samples were illuminated with a yttrium aluminum garnet laser and coagulation was characterized from the refractive index and absorption coefficient.

For more than a decade, another growing field of research that allows to assess the microstructures of soft biologic tissues has been in development, namely optical coherence tomography (OCT), which relies on short temporal coherence interferometry and measures the optical path and intensity of back-reflected near-infrared light. Recently, the blood properties have been widely studied using OCT and Doppler OCT (ODT) due to their high resolution, real time, and noninvasive capabilities. OCT backscattering signal and ODT variance imaging has been extensively utilized in characterizing some properties of blood, such as hematocrit (HCT), red blood cell (RBC) aggregation,
blood optical clearing, and blood photocoagulation in ours and other groups.17, 23–30

Because the OCT technique measures in-depth intensity distribution with high resolution, changes in the in-depth distribution of the blood-scattering coefficient or refractive index are reflected in changes in the OCT signal24, 27, 28. Thus, because blood coagulation may introduce local changes in their optical properties (scattering coefficient and local and mean refractive indices), one can monitor and quantify the coagulation process by analyzing the OCT signal slope (OCTSS)31 that reflects the least scattering of blood to some degree and 1/e light penetration depth (d1/e)32 that more accurately represents multiple scattering of blood. To better assess properties of blood during coagulation, the OCT signal slope and 1/e light penetration depth were attempted to be applied to follow the process of blood coagulation and clot formation from blood samples of various hematocrits ranging from 25% to 55%. Measurements were performed using a 1310 nm OCT. The effect of hematocrit associated with blood properties during coagulating was investigated and discussed. The main purpose of the present study was to investigate the feasibility of OCT to provide a description of the coagulation process of human blood.

2 Materials and Methods

2.1 Materials

The fresh whole blood was taken from healthy volunteers aged from 22 to 27 yrs. The blood sample was stabilized by 3.2% sodium citrate solution with the volume ratio of 9:1 for anticoagulation purpose. The whole blood was centrifuged to separate the erythrocytes from the plasma. The concentrated erythrocytes were washed twice with a phosphate buffer (pH = 7.4). The blood samples with various HCTs ranging from 25% to 55% were obtained by reconstituting the concentrated erythrocytes with the separated plasma. The coagulation process was initiated by adding calcium chloride (0.25 M with the volume of 1/9). All measurements were carried out within 12 h after the blood collection.

2.2 OCT measurements

A delay line OCT system with a 1310 nm central wavelength and 40 nm bandwidth light source (18 mW) was used. The light source yields an 18 μm axial resolution in free space, or approximately 13 μm within tissue if the mean refractive index of bulk tissue is assumed to be 1.38; this determines the axial imaging resolution of the system. The transverse resolution was measured at 15 μm. For the scanning speed of the reference arm at 100 A-scans, an OCT image of 400×100 pixel size requires about 2 s.32

Experiment 1 was performed to measure the OCTSS and d1/e of the reconstituting blood samples during coagulation. A rectangular glass cuvette of ~5 mm height (thickness) and 10 mm × 40 mm area was used as a blood sample holder. The sample holder was placed almost perpendicular to the probing beam, but tilted at a small angle of ~5 deg. relative to the cell surface to ensure that only light scattered or reflected by blood constituents was detected and the specular reflection that occurs at the glass-fluid interface was eliminated from the detection system. When the blood sample was added with CaCl2 solution as a coagulation agent, the OCT imaging of the blood sample was acquired immediately. The OCT images were taken every ~12 s, until 60 min after sampling. A cross-section imaging was obtained by taking 100 A-scans over the lateral length of 5 mm and the axial depth of 2.5 mm.

Experiment 2 was carried out to measure changes in refractive index of the plasma separated from the whole blood during coagulation. A glass tube with an inner diameter of 0.3 mm was used as a sample container. OCT images were taken every 5 min for a total of 30 min after sampling.

Data processing was illustrated in Fig. 1. Digital registration of the tilted blood surface because of the tilt cuvette in the OCT system. When the blood sample was added with CaCl2 solution as a coagulation agent, the OCT imaging of the blood sample was acquired immediately. The OCT images were taken every ~12 s, until 60 min after sampling. A cross-section imaging was obtained by taking 100 A-scans over the lateral length of 5 mm and the axial depth of 2.5 mm.

Data processing was illustrated in Fig. 1. Digital registration of the tilted blood surface because of the tilt cuvette in the OCT
Fig. 2 OCT images (a) and corresponding OCT in-depth reflectance profiles (b) recorded from the blood sample at HCT = 45% at 0, 20, 40, and 60 min during coagulation.

profiles based on an edge-detection algorithm was conducted to obtain a flat surface [Fig. 1(a)]. Quantitative data were obtained by averaging the signal intensity across the lateral imaging range as a function of depth. Every 100 A (depth) scans (covering the lateral length of 5 mm) were averaged to one depth profile in the OCT imaging. The OCTSS was obtained by linear fitting of the region from the blood surface to the depth of 0.27 mm of the averaged depth profile. The region in the blood, where the signal was linear and underwent minimal alterations, was selected as shown in Fig. 1(b). A best fit exponential curve in depth was applied to the averaged and normalized signal from which the corresponding 1/e light penetration depth was derived [Fig. 1(c)]. The group refractive index of plasma was obtained from the equation $n_g = z_{opt}/z_{phy}$. In this study, the physical length ($z_{phy}$) of all samples equals ~0.3 mm, i.e., the thickness of the sample container, which was determined by OCT. The optical path length ($z_{opt}$) of the plasma samples is measured by subtracting the axial position of the lower interface and that of the upper interface between the sample and the glass tube.

3 Results and Discussion

Figure 2 shows the representative OCT images [Fig. 2(a)] and the corresponding OCT in-depth reflectance profiles [Fig. 2(b)] recorded from the blood sample of a 26-year-old woman with HCT of 45% at 0, 20, 40, and 60 min after coagulation. There was slight movement of the boundary between blood plasma and RBCs layers with time [Fig. 2(b)]. The sedimentation rate of 0.06 ± 0.02 mm/h for the clotting blood after 60 min coagulation was much smaller than that for the control (2.16 ± 0.11). The latter fits well the normal sedimentation rates for women (2 to 15 mm/h). The fibrin formed from fibrinogen in the blood plasma during blood coagulation may impede the RBC
sedimentation. Normal blood sedimentation might cause a problem for accurate measurements of blood coagulation because sedimentation might change the optical properties as well, such as the increase of the 1/e light penetration depth.

Thanks to the slight sedimentation property of the blood during coagulation, we could accurately calculate the OCT signal slope and relative 1/e light penetration depth of the blood samples from the OCT in-depth reflectance profiles with the algorithm used in our previous works of Refs. 24, 27, 28, and 33. Figures 3(a) and 3(b) respectively, present the typical evolution of the OCTSS and d_1/e as a function of time acquired from the blood sample at HCT of 35% as an example. The mean OCTSS tended to decrease from 1.20 ± 0.03 to 1.04 ± 0.06 after 60 min coagulation. The entire process of blood coagulation relative to variations of the OCTSS, shown in Fig. 3(a), was divided into two stages. The results indicated that the OCTSS decreased faster during the first stage (0 to 1100 s) than during the second stage (1100 to 3600 s). We noted that the OCTSS measurement was not very accurate (large variation) by the time-domain OCT system. We expect better accuracy with a frequency domain OCT system that is able to reduce effectively the random noise.24, 25 Nevertheless, the decreased OCTSS with blood coagulation was found.

The variations in d_1/e were represented by convex curve characterized by two stages [Fig. 3(b)]. The first stage corresponded to a rapid increase in penetration depth, and the second stage corresponded to a very slow increase with a horizontal asymptote. The curves of the variations in the penetration depth can be linked to changes in state of the medium and coincide with the two stages of a liquid to a solid/gel state. Before CaCl_2 solution was added in the blood sample, the medium was in a liquid state. During stage 1 (0 to 395 s), the medium is turning into a gel state as the transformation of fibrinogen into fibrin through a coagulation cascade reaction. In this stage, d_1/e increased rapidly. To better characterize variations of blood properties during coagulation and clot formation, the clotting time (t_c) was defined as the period of time where 1/e light penetration depth curve started to be stabilized. For the blood of HCT of 35%, the clotting time was estimated to be about 395 s as indicated in an arrow in Fig. 3(b). In the following stage (396 to 3600 s), variations in d_1/e were small.

Blood is a scattering system that consists of scattering particles, for example RBCs and the surrounding media, i.e., plasma. The refractive index mismatching between RBC cytoplasm and blood plasma is the major source of light scattering in blood.21 Following the addition of CaCl_2 solution, a subsequent series of reactions including the conversion of plasma protein prothrombin into enzyme thrombin will be immediately initiated. Thrombin then catalyzes a reaction to allow several polypeptides to split from molecules of large rod-shaped plasma protein fibrinogen. The still-large fibrinogen remnants then bind to each other to form fibrin. During the blood coagulation process, the size, dimensions, and shape of the red blood cells almost do not change. The changes in the scattering resulted from the changes in its surrounding environment, i.e., plasma. During coagulation, the fibrinogen in the plasma transformed to coarse fibrin (the size from 1 nm to 1 μm). The refractive index of the plasma increased with the fibrin formation from 1.34 to 1.40 (Fig. 4). The refractive index of the red blood cells was measured to be 1.42. The reduced refractive index mismatching between the red blood cells and the plasma led to the decrease in the blood scattering, thus to the decrease of OCTSS and the enhancement of the light penetration depth. Eventually, as most erythrocytes were trapped in the fibrin meshwork, the clot was formed and the blood tended to be solidified. The transformation led to
Fig. 5 OCT signal slope as a function of time obtained from coagulating blood of (a) 25%, (b) 35%, (c) 45%, and (d) 55% hematocrit. The OCT signal slope decreased with time for all blood samples. Changes in OCTSS decreased with the increase of hematocrit. The coagulation process of the blood at 25% was abnormal.

Figures 5 and 6, respectively, demonstrate the values of OCTSS and $d_{1/e}$ acquired from blood samples at different HCTs of 25%, 35%, 45%, and 55% as a function of time. For comparison, the values for the blood sample of 35% were also included as Figs. 5(b) and 6(b). Similar to the blood of 35%, the OCTSS for the other three blood samples decreased with time and totally reduced by 47.0%, 15.0%, 13.7%, and 8.5% for the blood samples at 25%, 35%, 45%, and 55% after 60 min coagulation. The variations in the OCTSS against time could also be divided into two stages: the point was at approximately 900 s for the blood samples of 45% [Fig. 5(c)] and 800 s for 55% [Fig. 5(d)]. The results indicated that a decrease in the OCTSS during the first stage was faster than that during the second stage for 45% and 55%, as well as 35%.

Two distinct stages of the evolution of $d_{1/e}$ were also individualized for the samples at 45% [Fig. 6(c)] and 55% [Fig. 6(d)] on each curve: 1. a strong increase in $d_{1/e}$, corresponding to the liquid-gel transition of whole blood, and 2. a plateau with small variations. The clotting time ($t_c$) was approximately 375 ± 12 s, 455 ± 11 s, and 865 ± 47 s for the blood at HCTs of 35%, 45%, and 55%, respectively. Moreover, the parameter $S_r$, defined as the slope of $d_{1/e}$ within the period of time where $d_{1/e}$ increased dramatically following the induction of blood coagulation, were calculated to characterize dependence of scattering and attenuation properties of the coagulating blood on different hematocrits. In accordance with the increase of blood HCTs from 35%, 45%, to 55%, the $S_r$ ($\times 10^{-4}$ mm/s) tended to decrease from 6.0 ± 0.3, 3.7 ± 0.5 to 2.3 ± 0.4. These results also consistently indicate that the clotting time was longer for the blood of higher HCT than that of lower HCT.

Unlike the blood samples at HCTs of 35%, 45%, and 55%, the blood at 25% had different variation evolution in OCTSS [Fig. 5(a)] and $d_{1/e}$. There were three stages on each curve:
regarding OCTSS 1. a plateau (0 to 900 s), 2. a rapid decrease in OCTSS (900 to 1800 s), and 3. another plateau (1800 to 3600 s). The curve regarding $d_{1/e}$ [Fig. 6(a)] included the first stage (0 to 282 s) with strong variations, the second stage (283 to 1969 s) with moderate variations, and a plateau with no or very small variations (1969 to 3600 s). The clotting time was approximately 1969 s that was largely delayed compared with that for the blood at 35%, 45%, or 55%. These results indicated a slower coagulation for the blood at very low HCT. The abnormality in optical properties of the 25% blood, which is consistent with the previous findings resulted from the blood at an abnormal hematocrit. The hematocrit, defined as the ratio of the volume of packed red blood cells to the volume of whole blood, is normally 40% to 50% in men and 35% to 45% in women. The hematocrit changes in certain circumstances or disease, for patients with blood disorders, during dialysis, in an emergency room environment, and during surgery. For example, if the HCT is higher, he or she may have high red blood cell proliferation problem. On the contrary, it can be seen in people who suffer from anemia. Recent studies have revealed that blood coagulation is HCT dependent in which patients with an abnormally lower HCT might interfere with the capability of blood coagulation for bleeding, and that with an abnormally higher HCT of the blood is more likely to be associated with the increasing risk with the formation of thrombosis or embolism. Our results demonstrate that it is possible to recognize abnormality in the coagulation rate by OCT detection.

Fig. 6 $1/e$ light penetration depth as a function of time obtained from coagulating blood of (a) 25%, (b) 35%, (c) 45%, and (d) 55% hematocrit. The clotting time ($t_c$) was indicated as an arrow symbol. The clotting time was increased with the increase of hematocrit except for the blood at HCT = 25%.

4 Conclusions

In this paper, the OCT technique has demonstrated the efficiency to follow liquid to solid transition during blood clotting. Measurements of OCT signal slope and relative $1/e$ light penetration depth permitted description of the dynamics of the blood clotting process. OCT parameters such as $t_c$ and $S_r$ made it possible to describe the temporal sequence of two stages of the blood clotting process and the clot growth rate. Furthermore, the parameters were proposed to discriminate between different hematocrits. Certainly, more studies, such as the establishment of a unified relationship between clot characterization and quantitative OCT parameters, need to be carried out. Future investigations with standardized blood samples and with different blood coagulation protocols, particularly to explore the extrinsic pathway, will provide a more accurate evaluation of the potential of this technique.

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