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Experimental and Imaging Techniques for Examining Fibrin Clot Structures in Normal and Diseased States

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Abstract

Fibrin is an extracellular matrix protein that is responsible for maintaining the structural integrity of blood clots. Much research has been done on fibrin in the past years to include the investigation of synthesis, structure-function, and lysis of clots. However, there is still much unknown about the morphological and structural features of clots that ensue from patients with disease. In this research study, experimental techniques are presented that allow for the examination of morphological differences of abnormal clot structures due to diseased states such as diabetes and sickle cell anemia. Our study focuses on the preparation and evaluation of fibrin clots in order to assess morphological differences using various experimental assays and confocal microscopy. In addition, a method is also described that allows for continuous, real-time calculation of lysis rates in fibrin clots. The techniques described herein are important for researchers and clinicians seeking to elucidate comorbid thrombotic pathologies such as myocardial infarctions, ischemic heart disease, and strokes in patients with diabetes or sickle cell disease.

Introduction

Injury to a blood vessel's endothelial lining is repaired through the hemostatic response, or the formation of a blood clot. When blood permeates into the extracellular matrix, tissue factors activate platelets in the blood stream that facilitate initiation of the coagulation cascade. The key mechanical component of this healing process is the fibrin matrix, composed of fibrin fibers that are highly elastic, and can sustain large forces. Many researchers have studied the formation structure, and function of fibrin extensively in the past decades. Patients with diseases such as diabetes mellitus and sickle cell have an increased risk of developing thrombotic complications such as myocardial infarctions, ischemic heart disease, and strokes. Over 2 million people are newly diagnosed with diabetes mellitus each year in the United States. There are two types of diabetes: Type I, where the body fails to produce adequate amounts of insulin, and Type II, where the body becomes resistant to insulin. Among diabetic patients, cardiovascular disease (CVD) is the cause for 80% of the morbidity and mortality associated with the disease.

Sickle cell disease (SCD) is a genetic blood disorder that affects more than 100,000 people in the United States. SCD is a point-mutation disease that causes red blood cells to become crescent-shaped, making it difficult for the cells to pass through the blood vasculature. Both of these disease states increase the chances of developing atherothrombotic conditions in the body. One of the reasons for this is a result of altered fibrin structure and function in diseased states.

In both diabetes and sickle cell disease, there is hypercoagulation and hypofibrinolysis activity that induces atherothrombosis and cardiovascular disease (CVD) as compared to healthy patients. It is known that hypofibrinolysis promotes atherosclerosis progression and engenders recurrent ischemic events for patients with premature coronary artery disease. In the current manuscript, we investigated the role of fibrin physical properties in this particular setting. Fibrin clot structures in non-diseased patients are composed of thin fibers, larger pores, and generally less dense. The increased porosity and less dense fibrin clots in healthy patients have been found to facilitate fibrinolysis. In hyperthrombotic conditions such as diabetic and sickle cell disease, there is an increase in fibrinogen production, causing the fibrinogen concentration to increase from normal levels of 2.5 mg/ml in healthy patients to 5-13. Fibrin clots formed in diabetic patients have been found to be less porous, more rigid, have more branch points, and denser when compared to healthy, non-diabetic patients. The altered fibrin structure is a result of glycation mechanisms that occur in the proteins involved in clot formation. Nonenzymatic (irreversible) glycation occurs when glucose molecules bind to lysine residues on the fibrinogen molecule, which inhibits human factor XIIIa (FXIIIa) from properly cross-linking glutamine and lysine residues.

1. Collect 40-120 ml of blood from donors in 10 ml heparinized vacutainer tubes. Start PBMC (peripheral blood mononucleated cell) isolation within 4 hr of collection. During this time, keep blood at RT. 

2. Dilute whole blood 1:1 in ice-cold (4 °C) sterile PBS.

3. Pipette 10 ml of ice-cold hydrophilic polysaccharide into an empty, sterile 50 ml conical tube. Gently transfer diluted whole blood on top of the hydrophilic polysaccharide.

4. Use a 25 ml pipette in slow setting and pipette blood along the side of the tube very slowly. Ensure that the blood is not mixed with the hydrophilic polysaccharide.

5. Centrifuge blood samples for 30 min at 400 x g at 4 °C. If available, reduce the brake speed of the centrifuge to half of maximum for better separation after centrifugation.

6. Aspirate off the plasma/platelet fraction of the centrifuged blood sample. Carefully aspirate off the hydrophilic polysaccharide layer directly above the packed RBC layer.

7. Collect 8-10 ml of packed RBCs and dilute 1:1 with unsterile 0.9% NaCl saline.

2. Confocal Microscopy Evaluation of Glycated Clot Structures (Simulated Diabetic Clots)

1. Defrost human fibrinogen and fluorescently labeled human fibrinogen conjugate at 37 °C.

2. Pipette the following into a 0.5 ml graduated microcentrifuge tube.
   1. For the healthy, normal fibrinogen clots, mix human fibrinogen with 10% fluorescently labeled human fibrinogen conjugate in a 50 mM Tris/100 mM NaCl buffer at a concentration of 5 mg/ml.
   2. For artificially glycated fibrinogen (to simulate diabetic clots), incubate human fibrinogen and 10% fluorescently labeled fibrinogen conjugate in a 5 mM solution of glucose dissolved in 50 mM Tris/100 mM NaCl for a resulting concentration of 6.8 mg/ml.

3. Cover the micro-centrifuge tubes using an opaque material to avoid exposure to light. Incubate the tubes in a 37 °C water bath for 48 hr.

4. Make a chamber on a glass microscope slide by affixing a thin adhesive on 2 sides of the slide.

5. After the 48 hr incubation period, prepare reagents for fibrin formation by defrosting FXIIIa and human alpha thrombin at RT.

6. Dilute FXIIIa to 80 U/ml in 5 mM CaCl₂ buffer.

7. Dilute thrombin to 4 U/ml in 5 mM CaCl₂ buffer.
   1. For normal clots, ensure that the final concentrations of fibrinogen, FXIIIa, and thrombin are 2.5 mg/ml, 20 U/ml, and 1 U/ml, respectively at a volume of 50 μl.
   2. For the glycated clots, ensure that the final concentrations of fibrinogen, FXIIIa, and thrombin are 3.4 mg/ml, 20 U/ml, and 1 U/ml, respectively at a volume of 50 μl.

8. Immediately pipette 30 μl of the fibrin solution into the chamber formed by the adhesive.

9. Place a 22 mm x 22 mm glass cover slip with thickness of 0.15 mm on top of the 2 layers of adhesive. Seal the open sides with clear adhesive to prevent the fibrin clot from drying out. Ensure that the adhesive does not interact with the fibrin clot.

10. Allow the fibrin clots to polymerize at 21-23 °C for 2 hr before confocal imaging.

11. Take confocal microscopy images of the clots using a confocal microscope with a 40X/1.30 Oil M27 lens with a 488 nm Argon laser.

NOTE: The following protocol adheres to the guidelines set by the Institutional Review Board (IRB) at Georgia Tech.
3. Confocal Microscopy Evaluation of Fibrin Clots Containing RBCs from Sickle Cell Patients

1. Defrost human fibrinogen and fluorescently labeled human fibrinogen conjugate at 37 °C.
2. Pipette the following into a 0.5 ml graduated microcentrifuge tube.
   1. For the healthy, normal fibrin clot, mix human fibrinogen with 10% fluorescently labeled human fibrinogen conjugate in a 50 mM Tris/100 mM NaCl buffer at a concentration of 5 mg/ml.
   2. For the clots containing SCD RBCs, mix human fibrinogen and 10% fluorescently labeled human fibrinogen conjugate in a 50 mM Tris/100 mM NaCl such that the resulting concentration of the fibrinogen is 10 mg/ml.
3. Label the isolated RBCs (both normal and those from sickle cell patients obtained from the RBC isolation protocol) using a cell-labeling solution.
   1. Suspend cells at a density of 1 x 10^6 per ml in any chosen serum-free culture medium.
   2. Add 5 μl/ml of the cell suspension to the cell-labeling solution. Mix well by gentle pipetting gently.
   3. Incubate for 20 min at 37 °C.
   4. Centrifuge the labeled suspension tubes at 1,500 rpm for 5 min at 37 °C.
   5. Remove the supernatant and gently re-suspend the cells in 37 °C medium.
   6. Repeat the wash procedure (3.3.4 and 3.3.5) two more times.
4. Dilute FXIIIa to 80 U/ml in 5 mM CaCl_2 buffer.
5. Dilute thrombin to 4 U/ml in 5 mM CaCl_2 buffer.
   1. For normal clots, ensure that the final concentrations of fibrinogen, FXIIIa, and thrombin are 2.5 mg/ml, 20 U/ml, and 1 U/ml, respectively at a volume of 50 μl.
   2. For the clots containing SCD RBCs, ensure that the final concentrations of fibrinogen, FXIIIa, and thrombin are 5 mg/ml, 20 U/ml, and 1 U/ml, respectively at a volume of 50 μl.
6. Pipette 10 μl of the labeled RBCs into the fibrin solution. Pipette 30 μl of the fibrin with RBCs onto the microscope glass (refer to slide preparation for glycated clots).
7. Allow the fibrin to polymerize at 21-23 °C for 2 hr before confocal imaging.
8. Take confocal images of the clots using the confocal microscope, and utilize excitation lasers with wavelengths of 488 nm and 633 nm to excite the fluorescently labeled fibrin and RBCs.

4. Confocal Microscopy Evaluation of Fibrinolysis Rates in Glycated Clot Structures

1. Repeat the confocal microscopy protocol of glycated clots (Protocol Step 2) for the fibrinolysis rate evaluation in glycated clots.
2. Do not seal the ends of the chamber with clear adhesive. Allow the fibrin clots to polymerize for 1.5 hr at 21-23 °C in a sealed enclosure containing 250 ml of water to prevent dehydration.
3. After polymerization, obtain images of the clots using the confocal microscope.
4. Pipette plasmin into the open end of the chamber and disperse the plasmin through the clot via capillary action.
   1. For the normal and glycated fibrinolysis experiments with normal concentrations, pipette 25 μl at 200 μg/ml of plasmin into the opening of the chamber.
   2. For the glycated clots with reduced concentrations, pipette 25 μl at 50 μg/ml into the opening of the chamber.
5. Use the time series feature (see Figure 1) in the confocal microscope software to capture real-time video footage of the plasmin lysing the clot.
   1. Click acquisition mode and then select "Time Series." Select the number of cycles and the recording time interval.

5. Calculating Fibrinolysis Rates

1. Determine the lysis rate by setting an area (2,500 μm^2) and calculating the elapsed time required to dissolve a fixed area of the clot. Calculate the lysis rate using the following equation (Figure 2):
   \[
   \text{Plasmin lysis rate} = \frac{\text{clot area}}{\text{time to lyse the clot area}}
   \]

6. Statistical Analysis of Fibrinolysis Rates

1. Analyze the lysis data using statistical analysis software. Perform a one-way ANOVA and a post hoc Tukey-Kramer test\(^{44,45}\).

**Representative Results**

Confocal Microscopy Analysis of Glycated Fibrin Clot Structures

The confocal microscopy images of normal and glycated clots are presented in Figure 3. Confocal microscopy analysis of the normal and glycated clots reveals that glycated clots are denser with smaller pores than the normal clots both with and without the addition of FXIIIa during
clot polymerization. In Figure 3A and 3B, there is a lower fibrin concentration which created a less dense structure than the glycated clots with higher fibrin concentration (Figures 3C and 3D).

The greater density observed (number of fibers per unit area) in the glycated clots is a result of the increased fibrinogen concentration that occurs in people with diabetes. In the normal clots, the fibrin fibers were more linear and formed longer fibers upon polymerization with FXIIIa, which is likely a result of the FXIIIa forming cross-links between the fibers. In the case of the glycated clots, the fibers appeared to be shorter, even with addition of FXIIIa. This is likely the result of the polymerization process that was altered due to the presence of glycated regions on the fibrinogen molecule. In addition, from analyzing Figure 3, it is likely that the fiber thickness was also reduced as a result of glycation, which is likely why the fluorescence from the glycated fibers appear to have a lower intensity.

Confocal Microscopy Analysis of Fibrin Clots Containing RBCs from Sickle Cell Patients

The confocal microscopy images of the fibrin clot structure of clots with normal and RBCs from sickle cell patients are presented in Figure 4. As predicted, the confocal images of the fibrin clots containing RBCs from sickle cell patients had denser structures than those of healthy fibrin clots. Figure 4A shows a homogenous distribution of fibrin and normal RBCs. However, in Figure 4B there are clumps of fibrin and voids where no fibrin network formed. The RBCs from sickle cell patients were not homogenously dispersed, but rather were embedded in the fibrin clumps. The increased density (number of fibers per unit area) was a result of the increased fibrinogen concentration that occurs as a result of hypercoagulation in SCD patients. Furthermore, the natural tendency of the RBCs from sickle cell patients to form aggregates engendered clumps of RBCs that became interwoven into the fibrin network. This resulted in the formation of aberrant clot structures that are markedly different in morphology than those in normal patients. From Figure 4B, it is observed that RBCs from sickle cell patients caused greater heterogeneity in the fibrin sample and also caused a wider spatial distribution of RBCs in the fibrin sample, when compared to the healthy fibrin clots with normal RBCs. In contrast, the healthy fibrin clots with normal RBCs displayed a more homogenous dispersion of cells throughout the fibrin network.

It is unknown how much HbS (sickle hemoglobin) was polymerized in the erythrocytes from sickle cell patients and is contained in the samples. What is known, however, is that the presence of any amount of HbS causes greater aggregation of cells in number of HbS molecules per unit volume vs. normal HbA molecules. Pan et al. 46 have demonstrated this phenomenon in a study where they compared the aggregation/clustering effects of deoxy-HbS, and, for comparison, of oxy-HbS and oxy-normal adult hemoglobin, HbA. The clusters formed within a few seconds after solution preparation and their sizes and numbers remained relatively steady for up to 3 hr. One of the conclusions from the study was that both oxy-HbS and deoxy-HbS molecules clustered at a higher number density than oxy-HbA (normal) molecules as a function of temperature. Pertaining to the current study, this means that the RBCs from sickle cell patients were likely to cluster together, and trapped fibrin as it was being polymerized from the fibrinogen precursor.

Confocal Microscopy Lysis Rate Analysis of Normal and Glycated Clots

It was hypothesized that the fibrinolysis rates would be greater for normal fibrin clots compared to glycated fibrin clots with decreased plasmin concentration. To test this hypothesis, confocal microscopy was used to assess the lysis rates of fibrin clots with the addition of plasmin. The average (mean) lysis rates of normal fibrin with the addition of 200 µg/ml of plasmin was 43.2 ± 26.5 µm²/sec and the average (mean) lysis rates of glycated fibrin with the addition of 200 µg/ml of plasmin was 29.7 ± 11.2 µm²/sec. For the glycated fibrin clot lysed with 50 µg/ml of plasmin, the average lysis rate was 12.8 ± 3.1 µm²/sec. A total of 10 samples each (30 samples) were used to determine the average and standard deviation values. These values are presented in Figure 5. A p-value of 0.05 was used to determine significance between the independent experimental groups. Analysis of the fibrinolysis rates demonstrated that the value for the normal clots compared to the glycated (simulated diabetic) clots with decreased plasmin were significantly different.
Figure 1. Sample screenshot from confocal microscope software detailing method for obtaining real-time, continuous imaging of fibrin clot lysis. Please click here to view a larger version of this figure.

Figure 2. Example confocal image with fixed area used to calculate lysis rate of fibrin clot sample. Please click here to view a larger version of this figure.
Figure 3. Confocal images of healthy and glycated (diabetic) fibrin clots. Green fluorescence represents the fibrin network. The unglycated fibrin represents clots formed in healthy patients, whereas glycated fibrin represents clots formed in diabetic patients. (A) Unglycated fibrin without the addition of FXIIIa. (B) Unglycated fibrin with FXIIIa. (C) Glycated fibrin without FXIIIa and (D) Glycated fibrin with FXIIIa. The red scale bar indicates 50 μm. Please click here to view a larger version of this figure.

Figure 4. Confocal images of healthy fibrin clots with normal RBCs and fibrin clots with RBCs from sickle cell patients. Green fluorescence represents the fibrin network and magenta represents labeled healthy and RBCs from sickle cell patients. (A) Healthy RBCs in normal fibrin concentration with FXIIIa and (B) Increased fibrin with FXIIIa and RBCs from a sickle cell patient. The white scale bar indicates 50 μm. The white circles in image (B) indicate recognizable sickled RBCs. Please click here to view a larger version of this figure.
Figure 5. Fibrinolysis rates of normal fibrin clots compared to glycated (diabetic) fibrin clots and glycated clots with reduced plasmin concentration. Statistical significance was determined based on a one-way ANOVA and a post hoc Tukey-Kramer multiple comparison test. The ANOVA analysis showed that there was a significant difference between lysis rate means. The Tukey-Kramer test showed that there was a significant difference in means between the normal clots and clots with decreased plasmin with a p<0.05. The * indicates a significant difference from normal clots.

Discussion

To obtain meaningful data about the structure of clotting mechanisms in disease states, it is important to isolate the factors involved in clotting to determine the effects of the proteins and cells in these conditions. This protocol was developed for the purposes of investigating the structure of the fibrin clot in diabetic and SCD states in vitro.

It is necessary to understand the mechanisms involved in fibrin formation and fibrinolysis in disease states since altered conditions cause hypercoagulation, atherothrombosis, and CVD. From the confocal images obtained in this experiment, it is evident that hypercoagulation activity occurs in diabetic and SCD states, which causes an increased fibrinogen concentration. An increased fibrinogen concentration results in a denser fibrin network with more branch points and smaller pores throughout the network. In fibrin networks of SCD patients, not only is there more fibrin present, but the aggregating nature of the erythrocytes from sickle cell patients also changes the structure of the network as it entraps fibrin into clusters. The conglomeration of RBCs from sickle cell patients also reduces porosity of the fibrin clusters in the network, but appears to increase void concentration in the overall network. The results of the fibrinolysis analysis of the glycated clots supported the hypothesis of the lysis rate in glycated clots being lower with reduced plasmin than that of healthy, normal clots. However, the results did not provide conclusive evidence about lysis rates of glycated clots with normal concentrations of plasmin. Thus it can be concluded that for diabetic patients with normal plasmin concentration, there may be sufficient fibrinolysis activity to degrade fibrin clots in an efficient manner.

For the experiments performed in this research study, isolated proteins and RBCs were used, instead of whole blood and plasma. By using separated proteins to synthesize the fibrin clots, the causes for altered structures between the normal vs. diseased clots could be determined. This greatly reduced the possibility of having other plasma proteins interfere with the structure and morphology of the fibrin clots. However, by using isolated proteins, it is crucial to store and handle the cells and proteins properly, as they are more vulnerable to cell death and protein degradation.

The confocal microscopy imaging technique chosen for this study was due to the ability to obtain images of the fibrin clot structures while retaining the natural thrombotic properties of the cells and proteins. Confocal microscopy has also been shown to give improved contrast and optical resolution of the specimen with reduced noise compared to traditional light microscopy. The laser light is directed into a pinhole that blocks unfocused light from entering. Utilization of confocal microscopy obviated the need for the addition of fixative chemicals; thus the cells and proteins were able to maintain their natural protein activity of forming fibrin clots. As a result the cells, proteins, and enzymes functioned in their native state, allowing the use of confocal microscopy for capturing images and video of real-time activity. With confocal microscopy, 3D images of the fibrin clots were captured and videos of plasmin digestion of fibrin clots were obtained in real-time. When adding plasmin to the clot to capture a time series video of the clot lysis process, it was important to use capillary action via plasmin to homogenously disperse the enzyme through the fibrin matrix. This was done to ensure that the added plasmin concentration was accurate and allowed the enzyme to lyse the entire clot, in contrast to the local area that the plasmin was added.

The method and the results obtained in this study were similar to a study done by Dunn et al, who found that clot lysis rates were significantly lower in diabetic subjects than control subjects. In both studies, there were significant differences in lysis rates of diabetic (glycated) clots; however, the current protocol may accurately represent diabetic states because of the reduced plasmin levels. Additionally, in this research study fibrinolysis was captured continuously in real-time using the confocal microscope, in contrast to analyzing discrete sections at different time intervals. Wooton et al. have investigated fibrin clot lysis and employed a lysis front velocity (cm/s) to investigate the rate of lysis for clots. They used confocal image analysis in their study to determine the front velocity. In the current study, instead of a lysis front velocity, a lysis area
velocity was used (because of the way the image was analyzed on the confocal). To further validate the method developed in our study, the authors in 48 have described a mathematical model of fibrin clot lysis by plasmin, in which they prescribe the lysis front velocity as a function of the effective rate constant of fibrinolysis. The equation is given as:

\[
\frac{dx}{dt} = \frac{1}{\rho_{Fn}} \left( \frac{k_{eff} K A C_{pm}}{1 + K A C_{pm}} \right).
\]

Here, \( \frac{dx}{dt} \) represents the effective rate constant of fibrinolysis, \( K_A \) is the adsorption constant, \( C_{pm} \) is the plasmin concentration, and \( \rho_{Fn} \) is the density of fibrin. If the measured quantity \( \frac{dA}{dt} \) (measured from the confocal analysis in the current study) is utilized and the chain rule is employed, this leads to:

\[
\frac{dA}{dt} = x \frac{dy}{dt} + y \frac{dx}{dt}.
\]

Assuming only one dimension is changing per unit time leads to:

\[
\frac{dA}{dt} = \frac{dx}{dt}.
\]

Thus the effective rate constant of fibrinolysis for the experiment leads to:

\[
k_{eff} = \rho_{Fn} \frac{1}{y} \frac{dA}{dt} \left( \frac{1 + K A C_{pm}}{K A C_{pm}} \right).
\]

This relationship infers that the effective rate constant of fibrinolysis is a function of the plasmin concentration \( C_{pm} \) and the area lysis rate \( \frac{dA}{dt} \).

This technique is also important for the study of thrombosis in SCD patients, since there is little known about fibrin and clotting mechanisms with respect to this disease. There have been very few studies conducted on the effects of SCD on the structure of fibrin clots; thus the confocal images collected from this study provides rare visualization of the clot structure with incorporated sickled RBCs.

To compare the differences in means between the groups, use a one-way ANOVA and a post hoc Tukey-Kramer test. A one-way ANOVA is a statistical test used to determine a statistically significant difference in the means between independent groups. The Tukey-Kramer post-test is used to compare the means from each independent group and determine which groups’ means are statistically different from each other. Before performing a one-way ANOVA, there are six assumptions that must be met to ensure that the results are valid. The six assumptions are: a continuous dependent variable (lysis rate), an independent variable consisting of two or more independent categorical groups (normal, glycated, and glycated with reduced plasmin), and independently observed data in each group such that one event occurring does not affect another event (This assures that there is no overlap in data.), the dataset not containing any significant outliers, normally distributed data verified by the Shapiro-Wilk statistical test, and homogeneity of variances in the data verified by the Levene statistical test. Once these assumptions were met, a one-way ANOVA and a post hoc Tukey-Kramer test were performed with a p-value of 0.05 as the threshold for determining statistical significance. For a rigorous detail of the one-way ANOVA and post-hoc Tukey-Kramer test, refer to 44,45.

Overall, this method allows for imaging of normal and abnormal (due to diseased states) fibrin clots in their native states. The method also provides details for real-time imaging of clot lysis as well as statistical analysis of lysis rates. The ease with which the thrombus can be produced and imaged in vitro is useful for a wide range of applications in cellular and tissue engineering.

Disclosures

The authors have nothing to disclose.

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