Material Science and Engineering with Advanced Research

Comparative Study of Hemostatic Agents Using Thrombelastography

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Abstract

Hemorrhage is one of the leading causes of death in civilian and military trauma. Thrombelastography (TEG) quantitatively measures the viscoelastic changes of whole blood during clotting, from the beginning of coagulation to the end with fibrinolysis. In this study, we successfully developed and used a TEG method to compare a well-known Chinese herb medicine (i.e., Yunnan Baiyao) for hemorrhage control with other commercial hemostatic products (e.g., Celox™ and QuikClot™) and chitosan in the form of dry powder or dispersion. When tested in the powder form, the herbal material outperformed Celox™ in the form of dry powder or dispersion. In addition, random and block chitosan in the form of dry powder or dispersion. The ultimate objective is to develop a more effective, inexpensive, easy to store and use hemostatic agent.

Keywords: Hemostatic agent, Thrombelastography, Hemorrhage control.

Introduction

Hemorrhage control is vital for clinical outcome after surgical treatment and prehospital trauma injuries. Both systemic and local strategies have been developed for management of surgical and traumatic bleeding. The former typically involves hemostatic resuscitation of blood products and coagulation factor concentrate [1], while the latter uses numerous hemostatic biomaterials [2]. Advances in hemostatic materials have been made in the past few years given the significant interests in hemorrhage control on the battlefield [3]. Chitosan-based Celox™ and HemCon™, and zeolite-based QuikClot™ are among the few most effective hemostatic products [4]. However, each product has their limitation due to either the side effects, e.g., thermal damage to the tissues by QuikClot™ [5] or the lack of efficacy for hemorrhage control in an injury with limited vessel access [4].

Yunnan Baiyao is a Chinese remedy clinically used for stopping bleeding from surgery [6], diseases such as advanced cancer and ulcer [7,8] and combat trauma [9]. It has been administered both topically and orally. Given its other medical effects, e.g., anti-inflammation, it has become a multipurpose remedy.

Thrombelastography (TEG) has been widely used in clinical settings for monitoring coagulopathy [10,11] and guide hemostatic resuscitation in trauma [12]. In contrast, its use as a research tool for evaluating the effectiveness of hemostatic agents, especially those insoluble in blood, is limited [13]. In this study, we successfully applied a TEG method to test different hemostatic agents in the form of dry powder or dispersion. The ultimate objective is to develop a more effective, inexpensive, easy to store and use hemostatic agent.

Materials and Methods

Hemostatic Agents

Yunnan Baiyao was made by Yunnan Baiyao Group Co. Ltd (Yunnan, China). QuikClot™ was provided by Z-Medica Corporation (CT, USA). Celox™ was kindly provided by SAM Medical Products (TX, USA). Three types of medical grade chitosan derived from different natural resources were purchased. Chitosan derived from giant squid cartilage with a molecular weight larger than 1,000,000 and deacetylation degree higher than 90% in a powder form was purchased from Arabio Co., Ltd. (Guui-Dong, Gwangin-Gu, Korea) for initial comparison with Yunnan Baiyao, QuikClot™ and Celox™. In addition, random and block chitosan was purchased from Biosyntech Canada Inc. (Laval, QC, Canada). Both were derived from crustacean shells, the former had a molecular weight of 223,000 and deacetylation degree of 95.1% and the latter had a molecular weight of 3,410,000 and deacetylation degree of 94.5%. The hemostatic properties of the three types of chitosan were compared with the commercial products.

Thrombelastography

Figure 1(a) shows a TEG™ 5000 hemostasis analyzer...
The machine measures the viscoelastic properties of blood as it clots under a low shear. It has two channels; for each channel, a pin suspended by a torsion wire is immersed in whole blood or plasma in a plastic cup made of acrylic polymer with a smooth interior surface. The cup oscillates back and forth constantly at a set speed through an arc of 4°45° every 5 seconds. The torque of the cup is transmitted to the pin, via the fibrin strands in the blood clots as coagulation proceeds, and to the torsion wire for conversion by a mechanical-electrical transducer to an electrical signal, which can be monitored by a computer [14]. The measurement is graphically represented as a characteristic shape profile over time (Figure 1c), from which the following parameters can be derived to provide main information about the coagulation and fibrinolysis: 1) reaction time ‘R’ or time to first clot formation, which is related to plasma clotting factors and circulating inhibitory activity; 2) coagulation time ‘K’ or time to a specific level of clot strength, which is associated with the activity of the intrinsic clotting factors, fibrinogen and platelets; 3) rate of clot formation ‘α angle’ or rapidity of fibrin cross-linking, which is a main function of platelets, fibrinogen and plasma components residing on the platelet surface; 4) maximum amplitude ‘MA’ or maximum clot strength, which is a direct function of the maximum dynamic properties of fibrin and platelet number and function; 5) time to reach MA ‘TMA’; and 6) fibrinolysis at 30 min ‘LY30’ or the rate of amplitude reduction 30 min after MA, which relates to fibrinolysis. In our study, three key TEG parameters were used for studying the effects of pro-coagulant agents [15]: 1) time to fibrin initiation (R); 2) rapidity of fibrin cross-linking (α); 3) maximum clot strength (MA).

To perform a TEG measurement, the analyzer was first calibrated through electronics testing and quality control according to manufacturer's protocol. A powder material at a various amount (0.5-2.0 mg) was mixed with 340-µL citrated human blood from healthy volunteers in a TEG sampling cup pre-warmed to 37°C. Blood coagulation was initiated by adding 20 µL of 0.2 M CaCl2 to the cup and the measurement was run immediately until all interested parameter values were obtained. The percentage difference for each parameter was calculated as follow: \((P_{ha} - P_0)/P_0\), where \(P_{ha}\) and \(P_0\) are the parameters measured with or without addition of the hemostatic agent, respectively. +/- was used to denote either positive or negative effects on blood coagulation. For example, negative changes in R and positive changes in α and MA indicate an enhancement of clot formation and strength. The magnitude suggests the extent of the effects.

For the dispersion formulation, the material was suspended in a pH 7.4 Tris buffer and 20-µL dispersion was mixed with 320-µL citrated human blood from healthy volunteers in a TEG cup pre-warmed to 37°C, followed by adding 20 µL of 0.2 M CaCl2. The percentage difference for each parameter was calculated using the above equation.

**Statistical Analysis**

Data were presented as mean ± standard deviation and compared using a two-tailed t test with 95% confidence to identify significantly different groups.

**Results**

We confirmed that the TEG method is applicable in demonstrating the clotting effect of a known hemostatic agent, QuikClot™[16], which we used as a positive control in this study (Figures 2-4). Figures 2-4 also showed the hemostatic properties of a first-aid herbal powder, YunanBaiyao. Specifically, the herbal material,
when added as a powder in the amount of 0.5 to 2 mg, reduced R by 50-70% (Figure 2), increased α by 20-60% (Figure 3), but had a minimal effect on MA (Figure 4). In contrast, QuikClot™ reduced R by 40-50%, increased both α and MA by 30-100% and 7-20%, respectively. Celox™ and chitosan led to less decrease in R (-6 to 5% and -10 to -40%), less increase in α (-4 to 36% and 0.6 to 78%), and MA (0.7 to 13% and -5 to 21%).

Figure 5 shows dose-response effects of the herbal medicine on blood coagulation when added in dispersion. It appears that the hemostatic effects increased with the dose as indicated by R. In addition to the similar effects on R and α as its powder form, the herbal material in dispersion also increased MA by 15%, being more procoagulant than its powder form which had a minimal effect on MA.
Figure 4: Effects of the amount of hemostatic agents added to blood on clot strength MA. Data are expressed as mean± SD (n=3–6). *, + Different from Chinese remedy and QuikClot™, respectively (p<0.05).

Figure 5: Effects of the amount of Chinese remedy added in dispersion to blood on coagulation. Data are expressed as mean±SD (n=3). * Different from the lowest amount (p<0.05).

Figure 6 compares the effects of the hemostatic agents on the blood coagulation when added as dispersion in a buffer. The herbal remedy showed the most procoagulant effects as measured by reduced R time by 51%, increased rate of clot formation by 78% and increased maximum clot strength MA by 14%, significantly larger than the corresponding values for QuikClot™ (33%, 14% and -10%), Celox™ (0%, -7% and 0%) and chitosan (33%, -19% and -13%). Compared to their powder forms, all hemostatic agents except the herbal material became less procoagulant in the dispersion form.

In addition, Figure 7 compares the commercial hemostatic products (QuikClot™ and Celox™) with various chitosan biomaterials in powder forms. Specifically, chitosan powders were as effective as Celox™ as indicated by the TEG parameters, but less effective than QuikClot™, which is in agreement with their hemostatic mechanism mainly by tissue adhesion. As far as the onset of clot formation (R) is concerned, the block chitosan outperformed Celox™, but not in terms of clot formation rate (α) and strength (MA). No significant differences were observed among the chitosan agents while block chitosan appears more effective than...
squid chitosan, likely due to their differences in chitosan source and physical structure. Furthermore, as implied by reduced R time (approximately 20% decrease), all chitosan materials exhibited hemostatic activities, perhaps due to their induced formation of the coagulum of red blood cells and platelet aggregation as reported by many investigators [17].

Discussion
The hemostatic agents in this study are very different in their compositions and physiochemical properties and expected to have different hemostatic effects. This has been confirmed using TEG, a method that has been used to characterize various aluminosilicate materials for hemostasis [18]. It should be noted that the observed hemostatic effects of all agents, could not be attributed to an increase in blood viscosity, because in a control study where no calcium was added, no coagulation effects (i.e., no changes in blood viscosity) were observed in terms of R, α and MA values.

Different herb medicines have been used for hemorrhage control such as notoginseng [19]. The observed effects of Yunnan Baiyao...
on TEG variables are consistent with shortened bleeding and clotting time as measured by various techniques in vitro and vivo [20]. Compared to other commercial hemostatic materials in powder forms, the herbal material outperformed Celox™ and chitosan as indicated by all three TEG parameters. As far as the clotting time (R) is concerned, the herbal material appears more effective than QuikClot™, but less effective in terms of clot formation rate (α) and strength (MA).

It is interesting that the formulation of the herbal remedy (dry powder vs. dispersion) affects its hemostatic properties. The initial material appeared as a homogenous powder, however, it contained granular particles of various sizes and clusters at a microscopic level. When dispersed in water the opaque conglomerated particles became singular and transparent [9].

Several mechanisms for hemostasis induced by the herbal material have been reported. Activation of blood platelets was observed to cause the release of platelet constituents and subsequent hemostatic effects [22]. Microchemical analysis has confirmed the herb medicine contains the preponderance of polysaccharide structure and significant amount of calcium [9]. The main active components in Yunnan Baiyao are saponins [23] although a total of 34 compounds were recently identified using liquid chromatography hybrid ion trap time-of-flight mass spectrometry [24]. Further identification of active ingredients in this herbal remedy and investigation of their hemostatic mechanisms are warranted to develop a superior hemostatic agent.

Chitosan has well-documented bioactivities, such as antibacterial [25] and hemostatic properties [26,27,17]. The hemostatic potential of chitosan has long been recognized [27,28], however, its hemostatic potency reported in the literature varies significantly which could be due to the differences in physiochemical properties [17] as confirmed in this study.

Various mechanisms may contribute to its hemostatic properties and be postulated to be via vasoconstriction and the rapid mobilization of red blood cells, clotting factors, and platelet to the site of the injury, but the exact hemostatic mechanism of chitosan is still under study [26]. In one study where chitosan (Mw=80,000, deacetylation degree=80%) particle with 2.8 and 6.2 mm suspended in phosphate-buffered solution (PBS, pH 7.2) at a concentration of 30 mg/ml reduced blood coagulation time and enhanced platelet aggregation [26]. Another study showed that chitosan (Mw=50,000, deacetylation degree >90%) significantly promoted platelet adhesion and aggregation through enhancing expression of platelet glycoprotein IIb/IIIa complex on platelet membrane surfaces and increasing intracellular calcium level in platelets [29]. Scanning and transmission electron microscopy revealed that red blood cells bound to chitosan particles and became cup-shaped [30]. Similar to QuikClot™, chitosan can also promote blood coagulation by its hydration leading to concentrated coagulation factors. A recent study showed that the hemostatic efficacy of chitosan was associated to its water binding capacity [31].

Conclusion
Different hemostatic agents were successfully evaluated and compared using TEG. When tested in a dispersion form, which may be easier to use (as a spray) than in a powder form, the herbal material provided better hemostasis than the other materials, where the coagulation actions of the latter were reduced in the dispersion form. The sources and methods for chitosan preparation could affect its hemostatic properties. The observed procoagulant effects of the hemostatic agents could be attributed to different physiochemical and biological mechanisms. Further investigations of the herbal material and chitosan may be useful to pursue a potentially better substance for hemorrhage control.

References