Calibrated Automated Thrombin Generation in Frozen-Thawed Platelet-Rich Plasma to Detect Hypercoagulability

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Abstract
To enhance the practical applicability of the calibrated automated thrombogram (CAT) we investigated whether frozen-thawed platelet-rich plasma (ft-PRP) can be used to assess the function of the protein C inhibitory pathway, while preserving the natural phospholipid composition. Recalcified ft-PRP triggered with 0.5 pM recombinant human tissue factor shows a median thrombin potential of 1,779 nM·min, against 1,576 nM·min for fresh PRP. To obtain ~70% inhibition, 6.7 nM activated protein C (APC) has to be added, instead of 25 nM in fresh PRP; so the relative APC resistance of PRP appears to depend upon the presence of intact platelets. Factor VIII, added to normal ft-PRP to obtain a concentration of 3.3 U/ml, increases the thrombin potential in the presence of APC 1.5-fold, from 524 to 808 nM·min, in keeping with previously published increases in thrombotic risk in patients with high factor VIII levels. We conclude that thrombography in ft-PRP, with and without added APC, can be used to assess known risk factors for thrombosis, which allows the design of large clinical studies aimed at proving the relationship between thrombin potential and clinical outcome.

Introduction
Assessing the individual thrombotic risk by analysis of genetic polymorphisms and coagulation factors still leads to considerable discrepancies between the general risk associated with these abnormalities and the risk experienced by the individual patient. Measuring the effect of the combined genetic, acquired and environmental factors on the function of the hemostatic system ('phenotyping') is a more promising approach [1]. Thrombin activity measured in clotting plasma is a promising tool for such phenotyping [2]. We introduced the calibrated automated thrombogram (CAT) in our clinical laboratory in order to investigate the suitability of this technique for clinical diagnosis and treatment. As a first result, we demonstrated in platelet-rich plasma (PRP), stimulated with a low concentration (0.5 pM) of recombinant human tissue factor (rh-TF), acquired resistance to activated protein C (APC) in patients with lupus anticoagulants despite a prolongation of the lag time of the thrombin burst [3].

Given the essential role of platelets in blood coagulation [4, 5], it is of paramount importance to include platelets in the system tested. To avoid artifacts due to accidental platelet activation, this requires the use of carefully prepared PRP, tested within 90 min after venipuncture. This, however, is not easily accomplished in large multicenter studies that are required to ascertain correlations between clinical and laboratory data. We therefore inves-
tigated in how far frozen-thawed PRP (ft-PRP) would be a convenient alternative to fresh PRP. To track down APC resistance, our tests included the effects of adding APC and thrombomodulin (TM) and we also investigated the effect of addition of factor VIII, a well established and common risk factor for thrombosis, per se and via decreased sensitivity to APC [6].

**Materials and Methods**

**Proteins and Reagents**

Bovine serum albumin (BSA) and all chemicals were from Sigma (St. Louis, Mo., USA), rh-TF, relipidated and not containing polybrene or calcium, was a kind gift from Dade Behring (Marburg, Germany). The fluorogenic substrate Z-Gly-Gly-Arg-AMC was from Bachem (Bubendorf, Switzerland). Human factor VIII was provided by LFB (Lille, France). The calibrator with a constant, known, thrombin-like activity was from Synapse BV (Maastricht, The Netherlands). Soluble rh-TM (rhs-TM) composed of all the extracellular domains of TM without chondroitin sulfate chains [7] was a kind gift of Asahi (Japan). Human APC was prepared in-house as described previously [8]. For the clot-based assay [9], we used the STA coagulometer from Diagnostica Stago (Asnières, France) with the automated APTT reagent from Organon Teknika (Akzo-Nobel, The Netherlands) as a trigger.

**Blood and Plasma**

Blood from healthy volunteers was obtained by venipuncture with minimum trauma via a 21-gauge needle and collected into 0.129 mol/l sodium citrate (9:1 vol/vol). PRP was prepared by centrifugation at 194 g for 10 min at room temperature. Platelet counts were measured using a Micros 60 ABX model (Montpellier, France) and adjusted to 150 x 10⁶ platelets/l in PRP by addition of autologous platelet-poor plasma (PPP), which was obtained by double centrifugation at 2,500 g for 10 min. Exclusion criteria were factor V 506Q polymorphism or medical treatment. All individuals exhibited normal values for routine clotting tests, clotting factor levels and platelet counts. Factor VIII levels were measured with a one-stage clotting assay using a commercially available plasma deficient in factor VIII. PRP was used within 90 min after venipuncture, and an aliquot was frozen at -80°C. Frozen PRP was thawed at 37°C for 15 min and tested within 30 min thereafter.

**Fluorogenic Measurement of Thrombin Activity**

The thrombograms were recorded at 37°C according to Hemker et al. [10, 11] in a microtiter plate fluorometer (Fluoroskan Ascent, ThermoLabSystems, Helsinki, Finland) equipped with a 390/460 filter set for excitation and emission and using the dedicated software program (Thrombinoscope, Synapse BV). Round-bottom 96-well Greiner blue plates (reference No. 650204) were used. Coagulation was triggered by recalcification in the presence of 0.5 pM rh-TF in 300 µl Hepes, 140 mM NaCl, pH 7.35, containing 5 g/l BSA (HBS buffer) and 15 µl of HBS buffer or APC or rhs-TM in HBS buffer were added to 80 µl of PRP. Experiments were carried out in triplicate. A calibrator well in which 20 µl of a mixture of 2.5 mM fluorogenic substrate in Hepes 20 mM, pH 7.35, containing 60 g/l BSA and 100 mM CaCl₂ was dispersed in each well by the instrument, and the fluorescent signal was recorded with a measurement interval of 15 s. The molar amount of thrombin present in clotting plasma was calculated from the signal of wells in which thrombin is generated and the calibration signal as previously described [2]. The total amount of thrombin activity (i.e. the thrombin potential) was assessed as the area under the curve.

**Statistical Analysis**

The comparison between thrombin potentials obtained with 'fresh' PRP and ft-PRP was assessed by means of the two-sided Wilcoxon test for pairs. The correlation between the factor VIII level and the thrombin potential was analyzed by the Spearman rank correlation coefficient.
Results

Thrombin Potential in Fresh PRP and ft-PRP
As expected, the burst of thrombin generation started earlier, and maximal thrombin activity (the thrombin peak) was higher with ft-PRP than with fresh PRP. There was a slight but significant increase (~10%) in the endogenous thrombin potential with ft-PRP as compared with fresh PRP (fig. 1). In our population of 20 normal controls, the median values of thrombin potential were 1,576 and 1,779 nM·min for PRP and ft-PRP, respectively (p < 0.005).

Inhibition of Thrombin Potential by APC
In PRP preparations from healthy volunteers, the addition of APC led to a concentration-dependent decrease in thrombin potential. Similar inhibition patterns were observed with added, exogenous APC and endogenous APC, formed during thrombin generation via rhs-TM (fig. 2a) added to PRP. A ~70% decrease in thrombin potential was observed with 25 nM APC or with 10 nM rhs-TM. In ft-PRP, this inhibition is obtained with ~6.7 nM exogenous APC or ~12 nM rhs-TM (fig. 2). A batch-to-batch comparison of three in-house APC preparations showed superimposable thrombograms both with
Fresh PRP and ft-PRP (fig. 3). Similarly, low intra-individual variations were observed with fresh PRP and ft-PRP (fig. 4).

**Effect of Addition of Purified Factor VIII to Normal Plasma**

Our data confirmed an association between factor VIII level and sensitivity to APC, as assessed with the original, clot-based assay and our in-house APC preparation, in patients referred to the specialized clinical unit for the detection of thrombophilia (fig. 5). We then measured the effect on thrombin generation in normal PRP of added purified human factor VIII at levels encountered in thrombosis-prone patients (95 percentile). As factor VIII increased, the thrombin potential increased, and the sensitivity to APC decreased (fig. 6), both in fresh PRP and in ft-PRP ($p = 0.045$ for each of the 4 experimental conditions). Factor VIII, added to normal ft-PRP to obtain a concentration of 3.3 U/ml, increases the thrombin potential in the presence of APC 1.5-fold, from 524 to 808 nM.min.
Fig. 6. Dependence of the thrombin potential on FVIII concentration in the absence or presence of an APC system. 

- Experiments were performed with fresh PRP (○) and ft-PRP (■) in the absence or presence of 25 and 6.7 nM APC for PRP (○) and ft-PRP (■), respectively. Results are the mean of three tests with iterative blood samples of the same normal subject.
- Typical thrombograms with ft-PRP in the presence of 6.7 nM APC. Factor VIII concentrations from bottom to top: 0.8, 1.3, 1.8, 2.3, 2.8, and 3.3 U/ml.

Discussion

In a previous study, we have shown that for measuring the effect of lupus anticoagulants the sensitivity of the test is increased if the platelet phospholipids of the patient himself are used rather than exogenously added phospholipids [3]. Like any physiological function test, thrombin generation in fresh PRP requires the presence of the patient. For practical purposes it is more convenient to work on stored material; therefore, in the present study, we investigated whether ft-PRP can be used to assess the effect of the protein C system on thrombin generation.

The time course of thrombin differs between fresh PRP and ft-PRP, in that the thrombin burst is earlier as well as higher in ft-PRP, although the thrombin potential is not much higher (fig. 1). This is probably due to the fact that after freezing and thawing, with its concomitant membrane damage, the exposure of procoagulant phospholipids is no longer dependent upon the physiological membrane scrambling of the intact platelet [13]. Since TF pathway inhibitor (TFPI) is the key regulator of the initiation phase of thrombin generation [14], the observation that the thrombin potential is somewhat higher with ft-PRP than with fresh PRP could be explained by a lesser effectiveness of TFPI when the bulk of activated factor X is rapidly produced.

As in PPP with added PL and in fresh PRP [11], ft-PRP also showed low intra-individual (day-to-day) variations. The distribution of thrombin potential values confirms a relatively large interindividual variability.

For measuring APC sensitivity, either exogenous APC or endogenous PC, which is activated in the presence of added TM, can be used. In the latter case not only APC sensitivity of factors V and VIII is tested but also the APC forming system. APC might be preferred since it is a natural soluble protein, which is easily obtained in a reproducible manner (fig. 3). TM, in contrast, is a transmembrane protein that is currently used as a recombinant soluble form lacking the transmembrane domain and its physiological surroundings. Whatever the source of APC, this inhibitor needs a phospholipid surface for its activity, the optimal composition of the surface being different from the optimal composition for procoagulant reactions, so that using the patients’ own platelet phospholipids gives additional information on the equilibrium of the two processes in vivo.

With respect to the concentration of APC used, a peak circulating APC level of 23 nM was reported in baboons.
infused with thrombin [15]. Since the concentrations of protein C and prothrombin in the baboon are in the normal human range [16], we can speculate that a similar circulating level of APC is reached in human plasma at the site where thrombin is formed; we therefore chose 25 nM, which gave a convenient 70% inhibition of thrombin generation in normal controls. In ft-PRP we then sought the concentration that gave a similar inhibition.

Our data show an increased sensitivity to APC in ft-PRP as compared with PRP. Taube et al. [17] reported progressive APC resistance with increasing platelet activation, including a freeze-thawing cycle. This is only an apparent discrepancy because Taube et al. [17] based their conclusion on the use of clotting times. Also, in our study, we found that the response to APC, expressed as a ratio of the lag times before the burst of thrombin (roughly equivalent to the clotting time) in the presence or absence of APC, was lower with ft-PRP than with PRP (data not shown). This again emphasizes the importance of measuring the whole thrombin generation curve rather than clotting times only.

Perhaps more intriguing is the similar, or even slightly decreased sensitivity of ft-PRP to endogenous APC formed during thrombin generation in the presence of added TM. It has been demonstrated that platelets release an inhibitor to APC during platelet activation [18] and a time-dependent formation of a complex between this inhibitor and APC has been reported [19]. Also, phosphatidyethanolamine has been shown to be essential for optimal APC activity [20]. It is highly probable that frozen-thawed platelets, contrary to intact platelets, provide phosphatidyethanolamine right at the beginning of the reaction, thus enabling APC to engage in anticoagulant activity immediately. On the other hand, an inhibitor that is normally secreted in the neighborhood of the platelet surface is probably much diluted in the bulk of ft-PRP, and consequently the time-dependent neutralization of APC becomes less efficient.

Regarding the effect of factor VIII, it is well established that elevated levels of factor VIII are associated with a thrombotic risk [21, 22] and decreased APC sensitivity [6]. Here we report that the thrombin potential in the presence of APC increases about 50% with increasing factor VIII levels. Of note, the clot-based assay seems to underestimate the impact of factor VIII, since the decrease in the APC sensitivity ratio is only 10% (from approximately 2.5 to 2.3, for plasma levels of factor VIII between 1 and 3 U/ml). Furthermore, using thrombography, the effect on the lag time (~clotting time) was of the same order of magnitude as that seen with the original clot-based assay (data not shown), and much less conspicuous than the effect on thrombin potential.

We conclude that with the CAT methodology thrombography can be conveniently implemented in clinical research. It appears that ft-PRP retains several features of fresh PRP, so that it may be used for the detection of APC resistance if it is impossible to work in the vicinity of the patient. We also noted that known risk factors for thrombosis, e.g. factor VIII (this study) or lupus anticoagulants [3], definitely affect this test, and thus it is apparently a promising and attractive candidate to assay the ensemble of genetic and other effects on the hemostatic system determining the individual thrombotic risk.

Acknowledgments

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