Calibrated Automated Thrombin Generation Measurement in Clotting Plasma

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Abstract

Calibrated automated thrombography displays the concentration of thrombin in clotting plasma with or without platelets (platelet-rich plasma/platelet-poor plasma, PRP/PPP) in up to 48 samples by monitoring the splitting of a fluorogenic substrate and comparing it to a constant known thrombin activity in a parallel, non-clotting sample. Thus, the non-linearity of the reaction rate with thrombin concentration is compensated for, and adding an excess of substrate can be avoided. Standard conditions were established at which acceptable experimental variation accompanies sensitivity to pathological changes. The coefficients of variation of the surface under the curve (endogenous thrombin potential) are: within experiment 3%; intra-individual: <5% in PPP, <8% in PRP; interindividual 15% in PPP and 19% in PRP. In PPP, calibrated automated thrombography shows all clotting factor deficiencies (except factor XIII) and the effect of all anticoagulants [AVK, heparin(-likes), direct inhibitors]. In PRP, it is diminished in von Willebrand’s disease, but it also shows the effect of platelet inhibitors (e.g. aspirin and abciximab). Addition of activated protein C (APC) or thrombomodulin inhibits thrombin generation and reflects disorders of the APC system (congenital and acquired resistance, deficiencies and lupus antibodies) independent of concomitant inhibition of the procoagulant pathway as for example by anticoagulants.

Introduction

The thrombin generation curve (thrombogram) is an old and established tool in blood coagulation research [1–4]. It reflects much, if not all, of the overall function of the blood clotting system [5]. In platelet-poor plasma (PPP), it shows all clotting factor deficiencies with the exception of factor XIII [6–8], and it is sensitive to the action of oral anticoagulation, heparins of all types [9], direct thrombin inhibitors [10] and in fact all anticoagulant drugs tested. Hyperprothrombinemia [11] and lack of antithrombin [12], protein S or protein C as well as activated protein C (APC) resistance increase thrombin generation. Thrombin generation and a derivative test, the end level of α2-macroglobulin-thrombin (α2-M-T) in the ‘serum’ of defi-
brinated plasma, have been shown to be sensitive indicators of congenital [13] and acquired [14–16] defects in the APC pathway.

In platelet-rich plasma (PRP), the thrombogram also reveals the part played by platelets and hence the effect of von Willebrand factor [17], hypofibrinogenemia [17], thrombopenia [18] and Glanzmann’s [17] and Bernard-Soulier’s thrombopathies [Béguin, submitted] as well as the effect of anti-platelet drugs [19, 20], GPIIb/IIIa antagonists included [21]. Agents that increase platelet reactivity (e.g. arachidonic acid, epinephrine and collagen [22]) accelerate thrombin generation in PRP.

Despite its potential value for clinical diagnosis, drug monitoring and epidemiology, thrombin generation has not found universal application, primarily because of technical reasons: with the established subsampling techniques it takes about 1 h for a skilled laboratory worker to acquire one to four curves. The use of a slow-reacting fluorogenic substrate enables continuous measurement of thrombin generation in fibrinogen-containing media which makes it possible to use PRP and to test the influence of platelets on thrombin generation [23, 24]. However, it raises the question of how to convert the velocity of a fluorescence change (dF/dt) into a thrombin concentration [25]. For various reasons, during the course of the experiment, the relationship between dF/dt and thrombin concentration varies as fluorescence increases.

We calculate thrombin activity as a function of time by comparing the fluorescent signal from the thrombin-generating sample to that from a known stable concentration of thrombin in the coagulating sample in real time during the experiment. Using a 96-well plate fluorescent reader, up to 48 samples can thus be run.

### Materials and Methods

**Chemicals**

Recombinant relipidated tissue factor (rTF) not containing polybren or Ca²⁺ was a kind gift from Dade Behring (Marburg, Germany). Phospholipid vesicles consisted of 20 mol% phosphatidylethanolamine and 60 mol% phosphatidylcholine. Heparin-buffered saline comprised 20 M NaCl, 0.02% NaN₃. The reconstituted tissue factor was added to a heparinized plasma sample to prevent coagulation of the plasma sample upon use. The material is lyophilized and used as a calibrator after reconstitution with sterile water.

**Preparation of the Calibrator**

In order to obtain a stable thrombin-like activity in plasma, we used α₂ macroglobulin-thrombin complex (α₂M-T). A crude α₂M-containing fraction was isolated from citrated bovine plasma by precipitation with 12% (w/v) PEG-20,000; the pellet was dissolved in 1/10 vol 100 M NaCl, 20 M Hepe (pH 7.9) (modified from Barnett [27]). To this solution we added 12 μM bovine prothrombin, 6 M CaCl₂, 50 μM phospholipid vesicles (20% brain phosphatidylserine, 80% egg yolk phosphatidylcholine), 5 nM bovine factor Xa and 0.78 nM bovine factor Va. The mixture was stirred for 30 min at room temperature and then kept overnight at 4°C. The formed fibrin was removed and the preparation was applied to a Sepharclcolom equilibrated with 100 M sodium citrate, 20 M Hepe (pH 7.4) and 0.02% NaN₃. The α₂M-T that had formed eluted as a sharp peak before contaminating proteins.

The concentration of α₂M-T is measured by its activity towards S2238 and adjusted to the activity of 600 nM human thrombin; 100 nM bovine antithrombin and 2 U/ml heparin are added in order to prevent coagulation of the plasma sample upon use. The material is lyophilized and used as a calibrator after reconstitution with sterile water.

**Automated Fluorogenic Measurement of Thrombin Generation under Standard Conditions**

The thrombograms are measured in a 96-well plate fluorometer (Ascent reader, Thermolabsystems OY, Helsinki Finland) equipped with a 390/460 filter set (excitation/emission) and a dispenser. Immulon 2HB, round-bottom 96-well plates (Dynex) are used.

Each experiment needs two sets of readings, one from a well in which thrombin generation takes place (TG well) and a second one from a well to which the calibrator has been added (CL well). The color of the plasma can influence the fluorescence intensity, therefore
Fig. 1. The effect of constant thrombin activity on the fluorescent signal production in plasma. a The fluorescent traces of a range of α2M-T concentrations with the tangents at t = 0 (initial velocity). b Relation between initial velocity and α2M-T concentration. c Difference between tangents and curves of the top frame plotted as a function of the measured fluorescence.

Table 1. Kinetic constants of thrombin and α2M-T

<table>
<thead>
<tr>
<th>Kinetic constants</th>
<th>α2M-T in buffer</th>
<th>α2M-T in plasma</th>
<th>Thrombin (buffer)</th>
<th>Thrombin (plasma)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Km, μM</td>
<td>292 ± 44</td>
<td>310 ± 34</td>
<td>305 ± 46</td>
<td>280 ± 57</td>
</tr>
<tr>
<td>kcat·M⁻¹·s⁻¹</td>
<td>0.95 ± 0.1</td>
<td>0.89 ± 0.1</td>
<td>1.86 ± 0.26</td>
<td>ND</td>
</tr>
</tbody>
</table>

n = 12. The Km of thrombin in plasma was measured indirectly as the Ki of the inhibition of thrombin decay.

Experimental Results

From Fluorescent Intensity to Thrombin Activity

Free thrombin activity rapidly decays in plasma, so, in order to investigate the effect of constant thrombin activity on substrate consumption in plasma we used the α2M-T complex that splits the signal substrate like thrombin does but that is not affected by plasma inhibitors. Under test conditions, its amidolytic activity in plasma, as tested by subsampling on S2238, remained constant for over 3 h (results not shown). The kinetic constants of α2M-T were compared to those of thrombin (table 1). The Km of thrombin and α2M-T is identical within the limits of experimental error. Therefore, the variation in the reaction velocity with decreasing substrate concentration will...
be practically identical. The turnover velocity of $\alpha_2$M-T ($k_{cat}$) is half that of thrombin. To compensate for this difference, we express henceforward the concentration of $\alpha_2$M-T in ‘thrombin equivalents’, i.e. as the concentration of free thrombin that gives the same reaction velocity, i.e. at half the actual concentration.

Figure 1a demonstrates that constant $\alpha_2$M-T activities do not cause a constant rate of fluorescence increase. The initial rates, measured as the tangents to the curves at $t = 0$, are proportional to the enzymatic activity added (fig. 1b). The curvature is due to substrate consumption and to the non-linearity of fluorescence intensity with increasing concentration of fluorescent molecules (the ‘inner filter effect’) [28]. Both effects increase with the amount of product formed and hence with the level of fluorescence. We calculated the difference between the ideal fluorescence intensity ($F_i$), i.e. the value that would have been obtained experimentally if the initial velocity would have been maintained and the experimentally measured fluorescence ($F_e$), i.e. the vertical distance between the bold and the thin lines in figure 1a. The plot of $F_i - F_e$ against $F_e$ superimposes for the lines obtained at the six different thrombin activities (fig. 1c), demonstrating that the deviation from the initial velocity is indeed only dependent on the level of fluorescence. Therefore, every fluorescence level requires a different calibration factor to convert the velocity of the fluorescence increase ($dF/dt$) into thrombin activity (in nM). This calibration factor can be read from the graph obtained with a known calibrator concentration.

The calibration factor is not only a function of the fluorescence level attained, it also depends upon the instrument used, the age of lamp and filters, and the color of the plasma; therefore, each group of experiments that is carried out with the same plasma requires its own calibration experiment.

**Influence of the Fluorescent Substrate on the Course of Thrombin Generation**

The more substrate is present, the more thrombin molecules are occupied by the substrate and the less are they available for interaction with the natural antithrombins [29]. This effect is not corrected for by calibration, because the calibrator does not react with antithrombins. Therefore, the measured thrombogram increases with increasing concentrations of the fluorescent substrate (fig. 2a). When the amount of free thrombin is calculated (= total thrombin \cdot [Km/(Km + [substrate])], the influence of the substrate concentration becomes negligible (fig. 2b), showing that the fluorogenic substrate has little or no influence on the mechanism of prothrombin conversion.

**The Effect of Fibrinogen**

Previous techniques for automatic determination of the thrombogram used chromogenic substrates and required defibrination with coagulating snake venom enzymes in order to prevent turbidity changes [29]. We observed that the removal of fibrinogen has a profound influence on the thrombogram: In the absence of fibrinogen, the fluorescent technique shows a lower thrombin peak and a higher $\alpha_2$M-T end-level than in its presence (fig. 3a). By adding increasing concentrations of fibrinogen to afibrinoginemic plasma, fibrinogen dose-dependently decreases the amount of thrombin found in a subsampling experiment and increases the amount of thrombin in a fluorometer experiment (fig. 3b).
Fig. 3. The influence of fibrinogen on the thrombogram. a Course of amidolytic activity (expressed as thrombin equivalents) in normal pooled plasma before (A) and after (B) defibrination. Means ± 2 SD, n = 24. b ETP of the plasma from an afibrinoginemic patient with increasing amounts of fibrinogen added. ● = Fluorogenic method; ○ = subsampling method.

**Standard Conditions**

In PPP we chose a final concentration of 5 pM of rTF together with 4 μM of procoagulant phospholipids. At this concentration of rTF, thrombin generation is dependent upon factors VIII and IX (fig. 4) but not on factor XI, the influence of which is only seen at TF concentrations below 1 pM [8, and results not shown]. In the 3 pM range and above, thrombin generation in PPP is not critically dependent on the rTF concentration, whereas around 1 pM it is. rTF does not normally carry enough procoagulant phospholipids to ensure optimal prothrombin conversion. Thrombin generation is phospholipid dependent in the 0 to 2 μM range and reaches a plateau at ~3 μM. At ~10 μM, the contact activation properties of negatively charged PPL start to play a role [8], we therefore used 4 μM PPL.

In the absence of traces of rTF, the lag phase of thrombin generation in PRP is long and variable, probably influenced by endogenous TF and contact activation. Adding 0.5 pM (final concentration) of rTF largely abolishes this variability. It is essential that no PL is added; thus the test is critically dependent upon the PL coming from the activated platelets. Under these conditions, thrombin generation is dependent upon the platelet number in the range of 0–100 × 10⁹/l; above 200 × 10⁹/l dependence is minimal. In our experience, at 150 × 10⁹/l a useful equilibrium establishes between the independence of small fluctuations in the platelet count and the sensitivity to platelet inhibitors.

In PRP, under these conditions, thrombin generation is dependent upon platelet function (and hence on the level of von Willebrand factor [7]) as well as upon the concentration of factors VIII, IX and XI; the effect of activated factor VII is readily seen (results not shown).

The thrombogram reflects the action of all anticoagulants tested, alone or in combination. In the case of heparin, release of platelet factor 4 causes the inhibitory effect to be significantly smaller in the presence of platelets than in PPP (fig. 5). In the case of oral anticoagulation, a concomitant inhibition of the protein C system is readily seen.

**Normal Values and Variability**

The experimental variability in the main parameters of the thrombogram is shown in table 2. (The lag time is arbitrarily defined as the moment that 10 nM thrombin is formed.) The variation measured in identical parallel experiments in one 96-well plate is low (3–5%) and even lower when an occasional apparently aberrant curve, usually due to the presence of a floating bubble or accidental miss-pipetting, is left out.

The intra-individual CV was measured by making thrombograms of 4 healthy subjects once in 9 consecutive weeks. The CV of the endogenous thrombin potential (ETP) for each individual was around 8% (7.4, extremes 6–11). The interindividual variability determined based on the means of the 9 individual experiments was 17.5%. The combination of relatively stable day-to-day values and large interindividual variation is further illustrated in
fig. 4. Influence of TF concentration on thrombin generation in normal and hemophilia A and B plasma. From left to right: Normal plasma; hemophilia A plasma; hemophilia B plasma. rTF concentrations in decreasing peak height: 15, 10 and 5 pM rTF.

fig. 5. Influence of heparin administration on thrombin generation in PPP and PRP. At t = 0, 7,000 aXa units of enoxaparin were injected subcutaneously. Thin line = PPP, t = 0; bold line = PPP, t = 4 h; ++: PRP, t = 0; ∞∞ = PRP, t = 4 h. Time axis truncated at t = 30.

fig. 6. Intra- and interindividual variability in the PRP thromboagram. From the experiments used to establish the intra- and interindividual variability, the curves are given of three authors, determined four times at 2-week intervals. The line type is varied per individual. ○ = Mild von Willebrand factor deficiency.

Influence of Added APC and TM

Figure 7 demonstrates that thrombin generation is considerably decreased by the addition of APC or TM. PRP is significantly less sensitive to APC than PPP is. The sensitivity to TM does not show this large difference between PPP and PRP. The degree of inhibition observed is dependent upon both the TF concentration and the concentration of APC and TM. Under our standard conditions the inhibitions shown in table 3 are observed. Because the effect of APC or TM is compared to the same plasma to which no activator of the PC system is added, its effects can be observed independent of concomitant inhibition of thrombin generation as for example under antithrombotic treatment and/or in the presence of lupus anticoagulants [16].

Discussion

Measuring the hemostatic function of the blood is important both in bleeding and thrombotic disorders. Nevertheless, there is no universal test of this function available to the clinician. Clotting times of different types do not indicate a thrombotic tendency of unknown origin or mild hemostatic disorders. Different clotting assays are needed for different types of anticoagulant (oral anticoagulation/heparin), and for some types of antithrombotic therapy no adequate function test is available (e.g. heparin likes).
The thrombin generation test is a time-honored tool in coagulation research [1–4], and its wide scope is easily understood from the focal place of thrombin in the hemostatic-thrombotic mechanism [5]. On the one hand, scores of factors influence its formation, and on the other hand, it has numerous actions on blood and vessel components. The ensemble of pro- and anticoagulant factors determines thrombin generation in PPP, where a standard amount of procoagulant phospholipids is normally added. In PRP, thrombin generation is also dependent upon platelet function because platelets provide the necessary phospholipids, clotting factor V and, presumptively, clotting factor receptors.

Thrombin generation in PRP thus reflects the major part of the physiological clotting system. Ideally, the contribution of white and red cells and the vessel wall should be included, and flow conditions created. To date, this is not yet technically possible although the vessel wall can be partly represented by adding TF and TM.

In the older literature thrombin generation is used both as a diagnostic test and as a tool in basic research [1–4, 18]. Its use in modern, large-scale, clinical, epidemiological or pharmacological studies is hampered by its extreme labor-intensiveness. With the conventional subsampling techniques, even when optimally streamlined [30], it takes about 1 h of a skilled technician to obtain one to four thrombograms. Therefore, we developed the test presented here, which allows a throughput of up to 48 single thrombin-time graphs per hour in an automated procedure.

In this test, the thrombin concentration in the sample is calculated from the conversion velocity of a slow-reacting fluorogenic thrombin substrate. The principle of this approach has been developed with chromogenic substrates [29]. Unlike optical density measurements, the fluorescent signal is not influenced by the turbidity of the emerging clot, so fibrinogen-containing plasma and PRP can be used.

With fluorogenic substrates, however, constant thrombin activity does not cause a constant increase in the output signal, and thus the first derivative of the signal produced does not directly reflect thrombin concentrations. As seen in figure 1, the velocity of the fluorescence increase diminishes as the level of fluorescence increases. It is also seen that (i) the initial velocities are proportional to the amount of enzyme added (fig. 1b) and (ii) that the decrease in signal production is a function of the level of fluorescence but not of time or of the thrombin activity per se (fig. 1c). The two phenomena that cause this non-linearity are (i) non-linearity of fluorescence with the concentration of fluorescent molecules [28] and (ii) substrate consumption.

In practice, the effect of substrate consumption – with any substrate – cannot be overcome by adding the substrate in excess. The more substrate is added, the more thrombin molecules are bound to it and thus protected from the action of antithrombin. This not only prolongs the experiment beyond reasonable limits but also increases substrate consumption and consequently the inner filter effect.

**Fig. 7.** Effect of the protein C system on normal plasma. **a** PPP. **b** PRP. Drawn line = Control; ++ = TM added (10 nM); ○○ = APC added (10 nM in PPP, 6 nM in PRP). Inhibition of the ETP: PRP-TM: 64%; PRP-APC: 29%; PPP-TM: 76%; PPP-APC: 87%.
Moreover, the substrate is a competitive inhibitor of physiological feedback reactions and thus may interfere with the development of prothrombinase: figure 2 shows that at the concentration of substrate used in our standard procedure, no significant inhibition of thrombin formation is observed.

The effect of non-linearity cannot be solved by comparing thrombin generation in an experimental sample to that in standard plasma. In an inhibited sample less product is formed than in the standard so substrate consumption and the inner filter effect are low and more signal will be obtained per unit of thrombin activity than in the control. Thus inhibition is underestimated. Inversely, in samples where the activity is higher than normal, the mean signal production per unit thrombin activity will be higher than in the control so that activation is also underestimated. From the data in figure 1 we calculated the magnitude of such errors, both as a percentage of the normal control and as a percentage of the actual value and found that they transgress acceptable limits as soon as the deviation from the standard becomes significant (fig. 9).

**Fibrinogen**

In subsampling experiments, where the thrombin concentration in the fluid phase is determined, 30–50% more thrombin is found in defibrinated plasma than in plasma from which the fibrin has not been removed [31]. In fluorescence experiments, significantly more thrombin activity is found in the presence of fibrin than in its absence (fig. 3). Therefore, thrombin bound to fibrin must be co-estimated in the fluorescent experiment. This is not without importance because it has been shown that thrombin bound to fibrin can activate factors V, VIII and XI as well as platelets [31]. Also, antithrombin (and hence heparins) are reported not to inhibit fibrin-bound thrombin, whereas heparin cofactor II (and thus dermatan sulfate) and direct thrombin inhibitors do [32–34]. Our previously reported study showing that in PRP the presence of fibrinogen is essential for normal thrombin generation [17] has

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been confirmed with the fluorogenic technique presented here (results not shown).

Another important effect of fibrinogen is that it diminishes the amount of α2M-T formed during the experiment (fig. 3). The high end-level attained in defibrinated plasma (or in the presence of polymerization inhibitors, results not shown) suggests that polymerizing fibrin prevents the interaction between thrombin and α2-M. Therefore, correction of the α2M-T end-level is minimal, and the traces of amidolytic activity obtained during the experiment practically represent the course of free thrombin. This implies that the use of the α2M-T end level as a means to estimate ETP [29, 15] is not only dependent on the level of α2-M in the plasma but also on the level of fibrinogen. For unknown reasons, the experimental error in defibrinated plasma is higher than in non-defibrinated plasma (fig. 3b).

**Standard Conditions**

The form and magnitude of the thrombogram is dependent upon the trigger used. An excess of tissue factor (>1 nM, as in a thromboplastin time estimation) will cause the mechanism of prothrombins formation to take the extrinsic pathway, i.e. involve factors VII, V, X and TF pathway inhibitor. Decreasing the TF concentration changes the pathway of thrombin formation to include first factors VIII and IX and then factor XI [for a review, see ref. 36]. In the absence of any TF but with sufficient procoagulant phospholipids added, contact activation (‘intrinsic pathway’) will cause a burst of thrombin formation after ~10 min. Even though plastics are used throughout, contact activation cannot be entirely avoided unless blood is taken on citrate-containing corn trypsin inhibitor (results not shown). In samples with a high content of circulating TF [37], thrombin generation can be seen even in the presence of corn trypsin inhibitor (results to be published).

If in PPP the test is run without added phospholipids, thrombin generation becomes dependent upon non-sedimentable PL present in the plasma, i.e. upon the presence of circulating procoagulant microparticles, which in the first place help to check pre-analytical conditions. Variable amounts of PL can be introduced by the use of vacuum tubes, by insufficient centrifugation, by storing PRP, especially in the cold. Consequently, the conditions of centrifugation as given in the methods section should be strictly observed.

Thrombin generation is dependent upon the concentrations of PL and TF added. In the lower ranges, the relation is practically linear, to approach an upper limit at higher values. The concentrations chosen in our standard procedure are such that the upper ranges of thrombin generation are attained but not exceeded. Thus, the method is insensitive to small experimental variations in trigger, whereas an excess of trigger will not overrule modest inhibitions. The number of platelets was chosen on the basis of the observation that between 0 and 75 × 10^9 platelets/l there is an almost linear increase in thrombin generation with increasing platelet count, which levels off subsequently to reach a plateau near 150 × 10^9 platelets/l. Higher platelet numbers hardly increase thrombin generation [18, 38], but thrombin generation is less sensitive to the influence of platelet-based mechanisms and their inhibitors.

**Normal Values, Normal Effects of Addition of APC or TM**

The normal values under our standard conditions are shown in table 2. Of note, the thrombin peak is much lower in PRP than in PPP, whereas the ETP is not significantly different. This is most likely explained by the fact that platelet procoagulant activity gradually develops in the course of the activation process, whereas in PPP the added procoagulant phospholipids are present at recalcification, i.e. at zero time.

The inhibitory effect of APC and TM on the thrombin generation curve (table 3) is significantly diminished in dysfunctions of the APC system, such as the use of oral contraceptives or factor V Leiden [to be published, 39]. A comparison of thrombin generation in the presence or absence of TM or APC enables to assess the function of the APC system even if the coagulation process is impaired. In figure 8, for example, thrombin generation is diminished by oral anticoagulation, which concomitantly affects the APC mechanism, resulting in a diminished response to the addition of TM. It also has been shown that in the presence of the lupus anticoagulant severe resistance to APC and TM is accompanied by a considerably prolonged lag time (clotting time) [16].

The experimental variation is within acceptable limits (table 2) and similar between the three instruments at two locations used in the experiments reported here. In our experience, the fluorescence intensity of an identical product solution (AMC) can differ severalfold between fluorometers, even if they are of the same type. This variation is compensated for by the use of the calibrator.

The variation in the population is much higher than the day-to-day variation per individual. Thus, some individuals generate more thrombin than others. A correlation between the clotting factor level and the occurrence
of venous and/or arterial thrombotic disease has been reported for various clotting factors [40, 41], which strongly suggests a relationship between the thrombin generation capacity and thrombotic disease. Measurement of overall thrombin generation enables the assessment of simultaneous effects of numerous small changes, which may either reinforce each other (e.g. in pregnancy [42]) or compensate each other (e.g. a bleeding tendency and factor VLeiden [43–45]).

**Hemorrhagic Diatheses**

It has been shown before [for a review, see ref. 5] that thrombin generation is diminished in hemophilia and rare clotting disorders [6–8]. Also, in PRP, it is diminished if von Willebrand factor is low [17] and increases upon DDAVP treatment [7]. As can be seen from figure 4 and 6, calibrated automated thrombography shows these effects as well as subsampling techniques do. In an accompanying article by Turecek et al. [46] it is shown that in hemophilia, the effect of inhibitor bypassing therapy can be seen in uncalibrated fluorescence experiments, thus enabling the quantification by calibrated automated thrombography.

**Anticoagulant Treatment**

We hypothesize that in anticoagulant therapy diminished thrombin generation is essential. All anticoagulant drugs have this effect in common although they may achieve it in completely different ways: decreased prothrombin and prothrombin conversion (AVK), increased thrombin inactivation (heparin) or direct inhibition (hirudin, argatroban and melagatran). There is no single clotting test that reacts similarly to these different types of anticoagulants. Clotting tests measure the length of the initiation stage of thrombin formation, i.e. the time during which no significant amount of thrombin (<5 nM) is formed. Clotting occurs at the end of this initiation phase and is complete when >95% of all thrombin is still to appear. That the amount of thrombin forming in the clot is essential for thrombosis and hemostasis can be derived from comparisons of laboratory and clinical observations. Heparin, for example, has a minimal effect on the TF-induced clotting time ("quick time") but has an undeniable antithrombotic effect. Defects in the protein C system have a recognized prothrombotic effect, nevertheless this system only comes into action after APC is formed by the TM-thrombin complex, i.e. in the later stages of thrombin formation, long after the clot has formed. In figures 7 and 8, the effect of added TM is seen in the second half of the thrombin generation curve. In fact the use of clotting times is restricted to those cases where the length of the initiation phase indicates the amount of thrombin to come, which is far from being a general rule.

Some observations that are liable to be of clinical importance will readily escape our attention unless thrombin generation curves are made. It is logical, for example, that AVK treatment impairs the function of the initiation stage of thrombin formation.
Alternative Methods Depending on Clot Formation

Several methods exist that measure the moment of clot formation and ensuing changes in the mechanical or optical properties of the clot and/or retraction of that clot through activated platelets, such as thrombelastography [47], resonance thrombography: [48], roTEG coagulation analysis [49], the sonoclot analyzer [50], measurement of platelet-mediated force development during clot formation [51] and free oscillation rheometry [52]. The results of such methods sometimes show a superficial similarity to thrombin generation curves [53]. These methods can often be applied to whole blood, which is an important practical advantage.

The disadvantage of clot-based methods is that they depend upon thrombin formation as well as on properties such as fibrinogen content and quality, fibrin polymerization, crosslinking and fibrinolysis [49]. These two aspects may counteract each other, e.g. in a febrile patient (high fibrinogen) under anticoagulant or other antithrombotic treatment (e.g. aspirin). Clot formation is only one of the many functions of thrombin and not necessarily the most important one, as can be judged from the mild bleeding syndrome in hypofibrinemia. Through selection of the right substrate and comparing the fluorescent signal to that of a calibrator with fixed activity we can obtain a correct estimate of the course of thrombin concentration in clotting plasma. Fibrinogen as a substrate is quickly exhausted and the signal from its product – i.e. one or the other mechanical or optical property – cannot be unequivocally related to the amount of thrombin formed.

References


