Inhibition of Activated Protein C by Platelets

S. M. Jane, C. A. Mitchell, L. Hau, and H. H. Salem
Department of Medicine, Monash Medical School, Prahran, Victoria, Australia

Abstract

Activated protein C (APC), an anticoagulant that acts by inactivating Factors Va and VIIIa, is dependent on a suitable surface for its action. In this study we examined the ability of human platelets to provide this surface and support APC-mediated anticoagulant effects. The activity of APC was examined in three systems: the Factor Xa calcification time of Al(OH)₃ adsorbed plasma, studies of thrombin generation in recalcified plasma, and assessment of the rate of inactivation of purified Factor Va. In comparison with phospholipid, intact platelets required significantly greater concentrations of APC to achieve a similar degree of anticoagulation. When washed platelet membranes were substituted for intact platelets, adequate support of APC was observed and the anticoagulant effect was similar to that obtained with phospholipid. Platelet releasate obtained by stimulation of platelets with thrombin and epinephrine contained an inhibitor that interfered with the ability of phospholipid and washed platelet membranes to catalyze the anticoagulant effects of APC. A noncompetitive inhibition was suggested by Dixon plot analysis of the interaction between platelet releasate and APC. The activity of the platelet APC inhibitor was immediate and was not enhanced by heparin, distinguishing it from the circulating protein C inhibitor. The presence of this inhibitor in the platelet and its release with platelet stimulation emphasizes the procoagulant role of this cell.

Introduction

Protein C is a vitamin K-dependent serine protease that is activated by thrombin in the presence of thrombomodulin or, to a lesser extent, Factor Va (1-4). Activated protein C (APC)¹ is a potent anticoagulant that inactivates the coagulation cofactors Va and VIIIa in a calcium- and phospholipid-dependent reaction (5). The effects of APC are also enhanced by a second protein, protein S, which is thought to increase the affinity of APC to the phospholipid surface (6). Studies using unstimulated bovine platelets have demonstrated that APC had little or no influence on the rate of Factor Va inactivation in the absence of protein S (7).

The in vivo regulation of APC activity is poorly understood. Unlike other serine proteases in the coagulation system it is not inactivated by antithrombin III (8). A specific APC inhibitor was first suggested by Malar and Griffin (9) and subsequently purified and characterized by Suzuki et al. (10). This inhibitor covalently complexes with APC in a reaction accelerated 10-30-fold by large concentrations of heparin. The prolonged incubation time required for the neutralization of the amidolytic and anticoagulant activity of APC by this inhibitor suggests that it is probably not the sole physiological regulator of APC.

In this study we report the presence of an APC inhibitor in human platelets that, when released, interferes with the ability of platelet-mediated support of APC.

Methods

All chemicals were purchased from Sigma Chemical Co. (St. Louis, MO) except the following: NaCl, CaCl₂, Na₂HPO₄, NaH₂PO₄, 2H₂O, and EDTA (Ajax Chemical Co., Sydney, Australia); K₂HPO₄ and D-glucose (Mallinkrodt, Inc., St. Louis, MO); SDS (Pierce Chemical Co., Rockford, IL); acrylamide (Bio-Rad Laboratories, Richmond, CA); and H-D-Phe-pipocetyl-Arg-p-nitroanilide (S2238; Kabi AB, Stockholm, Sweden).

The assay buffer referred to throughout this manuscript was 20 mM Tris, pH 7.4, containing 0.15 M NaCl and 0.1% gelatin. The platelet washing buffer, pH 6.5, contained 0.113 M NaCl, 4.3 mM K₂HPO₄, 4.3 mM Na₂HPO₄, 24.3 mM NaH₂PO₄·2H₂O, and 5.5 mM glucose. Coagulation factors were of human origin. Protein C (11), thrombin (12), antithrombin III (13), Factor V (14), prothrombin, Factor X (15), protein S (16), and thrombomodulin (17) were purified and activated as previously described. Protein C (1 mg/ml) was activated using 20 nM thrombin in the presence of 15 mM human thrombomodulin. The reaction buffer consisted of 20 mM Tris, pH 7.4, containing 0.15 M NaCl and 5 mM CaCl₂. The mixture was incubated at 37°C and aliquots removed at various time points and assayed for protein C activation using the chromogenic substrate S2238 as previously described (11). Protein C activation was usually completed after a 60-min incubation, at which time antithrombin III (150 μg/ml) and heparin (1 U/ml) were added. The reaction was incubated for a further 10 min at 37°C, allowing neutralization of all thrombin. The antithrombin III-heparin concentrations used in the highest concentration of APC were shown to have no effect on the results of coagulation studies.

Platelets were obtained from healthy volunteers. To minimize platelet activation the blood was collected in 5 mM EDTA, 3.8% tri-Na-citrate, 5 mM theophylline, and 0.2 μg/ml PGE₁, and the platelets prepared and washed as previously described (18). The final platelet preparation was resuspended in assay buffer and counted in duplicate using a hemocytometer. The platelets were used within 4 h of blood collection and unless otherwise specified were not subjected to any further activation before use in coagulation studies. However, since some degree of activation is virtually inevitable during preparation, nonactivated platelets cannot be assumed to be completely inactive. For the separation of the membranes from the cytosol, 2 ml of washed platelets (5 × 10⁷/ml) were disrupted by three cycles of freeze-thawing and the suspension subjected to maximum intensity sonication (Labsonic 15110, ultrasonic cell disruptor; B Braun, Melsungen A. G., West Germany) for 45 s at 4°C. The disrupted platelets were centrifuged at 100,000 g for 60 min at 4°C and the supernatant used as the cytosolic component. The membranes were washed twice in the platelet-washing buffer and resuspended in 2 ml of assay buffer.

1. Abbreviations used in this paper: APC, activated protein C.

Address correspondence to Dr. S. M. Jane, Department of Medicine, Alfred Hospital, Commercial Rd., Prahran, Victoria, Australia.

Received for publication 30 November 1987 and in revised form 2 August 1988.

0021-9738/89/01/222/05 $2.00
Volume 83, January 1989, 222-226
Platelet aggregation was performed using 400 μl of 5 × 10^7/ml platelets that were treated with 8 × 10^-6 M epinephrine and 20 nM thrombin. The platelets were stirred at 37°C in a Payton’s dual channel aggregometer module at 250 rpm for 10 min.

Three methods were used to assess the anticoagulant activity of APC: the Factor Xa recalcification time of Al(OH)₃ adsorbed plasma (19), thrombin generation in recalcified plasma, and studies of the rate of inactivation of purified Factor Va. The phospholipid used in these studies was of two types: a commercial rabbit brain phospholipid reagent (platelet, General Diagnostics, Morris Plains, NJ), or homogeneous single-lamellar phospholipid vesicles consisting of 75% phosphatidylycholine and 25% phosphatidylserine (PCPS) prepared as described (20).

Thrombin generation was assessed by incubating 100 μl plasma, 20 μM phospholipid, and/or 5 × 10^7/ml platelets, and varying concentrations of APC. The volume of the reaction was adjusted to 600 μl using the assay buffer, and thrombin formation was initiated by the addition of 8 mM Ca²⁺. At various time points 40-μl aliquots were removed and the amount of thrombin formed assayed by measuring the rate of hydrolysis of the chromogenic substrate S2238. The amidolytic reaction (600 μl) contained 0.2 mM S2238 in assay buffer. The concentration of thrombin formed was determined by reference to a standard curve constructed using known amounts of thrombin.

Factor V was assayed using a purified component assay as described (21).

Results

In preliminary experiments we investigated if platelets could substitute for phospholipid as a surface for the mediation of the anticoagulant activity of APC. These studies were performed using washed platelets (as detailed in Methods) that were not subjected to any specific activation before use in coagulation assays. Initial experiments were performed using a Factor Xa recalcification time of Al(OH)₃ adsorbed plasma and the concentration of platelets was adjusted to give a clotting time, that in the absence of APC, was comparable to that obtained using 20 μM phospholipid. In this assay, APC (20 nM) resulted in a 100-s prolongation of the clotting time in the presence of either the commercial phospholipid reagent or the homogeneous PCPS vesicles. When platelets were substituted for phospholipid, the same concentration of APC produced an insignificant prolongation of the clotting time (2 s). Furthermore, platelets, when added to the phospholipid containing reaction, negated the effects of APC, and the prolongation of the clotting time was similar to that seen using platelets alone (Fig. 1). These results suggested that platelets failed to provide an adequate surface for APC anticoagulant activity and instead inhibited the effects of APC observed in the presence of phospholipids.

To validate the results obtained using the Factor Xa recalcification time, studies of the effects of APC on thrombin formation in recalcified plasma were performed. The advantage of this system is that it quantitates the rate of thrombin generation in contrast to the Factor Xa recalcification time which is an end point assay. The concentration of platelets used in these studies was adjusted to yield rates of thrombin formation comparable to those obtained using phospholipid. As in previous assays the platelets were not specifically activated before the experiments. The results are shown in Fig. 2, A-C. As seen, 2 nM APC caused a marked reduction in thrombin generation in the presence of the lipid reagent (Fig. 2 A). On the other hand, platelets nullified the activity of APC irrespective of the presence or absence of phospholipid (Fig. 2, B-C).

The effect of increasing concentrations of APC on the inhibitory potential of intact platelets is shown in Fig. 3. Using 5 × 10^7/ml platelets, a 15-fold greater concentration of APC was necessary to produce inhibition in thrombin generation comparable to that achieved by APC and phospholipid.

The observation that the platelet offered poor support for the effects of APC was confirmed by studies of the rate of inactivation of purified Factor Va (Fig. 4). In these experiments APC (0.125 nM) and phospholipid rapidly inactivated Factor Va with a rate of 1.7 U/min. In contrast, when 5 × 10^7/ml platelets were substituted for the lipid, the proteo-

![Figure 1. Comparison of platelet and lipid support of APC anticoagulant activity. APC anticoagulant activity was measured using a Factor Xa recalcification time of Al(OH)₃-ad sorbed plasma as described in Methods. Baseline clotting studies in the absence of APC were performed using 20 μM phospholipid (A), 5 × 10^7/ml platelets (C), or 20 μM phospholipid and 5 × 10^7/ml platelets (E). The effect of APC (20 nM) in each system is shown in B, D, and F, respectively.](image)

![Figure 2. (A) Effect of APC and phospholipid on thrombin generation in recalcified plasma. Thrombin generation in platelet-poor plasma was assessed using 20 μM phospholipid in the presence (●) or absence (○) of 2 nM APC. Thrombin formation was monitored using the chromogenic substrate S2238 as detailed in Methods. (B) Effect of APC and platelets on thrombin generation in recalcified plasma. Studies of thrombin generation were performed using 5 × 10^7/ml platelets in the absence (●) or presence (○) of 2 nM APC. Other details are as described for A. (C) Effect of APC on thrombin generation in the presence of phospholipid (20 μM) and 5 × 10^7/ml platelets. Thrombin generation was assessed as for A in the presence (●) or absence (○) of 2 nM APC.](image)
lytic effects of APC in concentrations up to 0.5 nM were negated and Factor Va remained stable. In the absence of platelets or phospholipid 0.5 nM APC slowly inactivated Factor Va at a rate of 0.28 U/min.

To determine whether APC was inhibited by a surface or internalized component of platelets, membrane and cytosol fractions were prepared as described in Methods. The time course of Factor Va inactivation by 0.125 nM APC and platelet membranes demonstrated rapid cleavage of Factor Va with a rate comparable to that achieved with phospholipid. The recombination of platelet cytosol with the membranes reproduced the APC inhibition observed in experiments where intact platelets were combined with phospholipid. These results suggest that the inhibitor is cytosolic and that washed platelet membranes can support APC-mediated Factor Va inactivation with rates comparable to those obtained with phospholipid.

We next examined the ability of the platelets to release this inhibitor. For this purpose 400 μl of washed platelets (5 x 10^7/ml) were aggregated as detailed in Methods. After irreversible aggregation the sample was subjected to centrifugation at 100,000 g for 25 min. The releasate was tested for its APC-inhibitory activity in a Factor Va inactivation system and the results compared with those obtained using supernatant from nonactivated platelets. The rate of inactivation of Factor Va by 0.125 nM APC and phospholipid was 1.7 U/min and was not significantly affected by the addition of supernatant from unactivated platelets (rate of inactivation = 1.65 U/min). However, when the releasate of activated platelets was used a significant inhibition of the rate of Factor Va inactivation occurred (rate = 0.23 U/min). These results, shown in Fig. 5, confirm that the APC inhibitor is contained within the secretory granules of the platelets and is released upon platelet activation.

The effect of increasing concentrations of APC on the inhibitory potential of platelet releasate was assessed (Fig. 6). In the presence of phospholipid the rate of Factor Va inactivation increased linearly with increasing APC concentrations. Addition of platelet releasate to this system resulted in an eightfold increase in the amount of APC necessary to achieve comparable rates of Factor Va inactivation. The effects of the platelet releasate were also assessed using two concentrations of Factor Va in the presence of APC. A Dixon plot of these results (Fig. 7) suggests that platelet releasate is a reversible noncompetitive inhibitor of APC.

The inhibition of APC by platelet releasate was not enhanced by heparin in concentrations up to 5 U/ml and was not overcome by the addition of human protein S in concentrations up to 60 nM.

The inhibitor appears to be a protein since the activity was destroyed by heating at 100°C for 10 min. It is not dialyzable using membranes with a cutoff of 10,000 D, and appears to be a high molecular weight molecule since it is excluded from a Sepharose 4B-CL gel filtration column, appearing in the void volume. The inhibitory activity is stable between pH 7 and 9.5. In buffers of pH 5 or less the activity was lost. In other studies we attempted to directly demonstrate that the inhibitor binds to or modifies the structure of APC. Different approaches were
Discussion

An integral part of the hemostatic mechanism is the assembly of activation complexes on negatively charged surfaces. Traditionally, phospholipid has been used as the in vitro model of these surfaces. In vivo, platelets and endothelial cells catalyze several coagulation reactions, playing an essential role in accelerating thrombin formation (12, 22). These cells do not merely serve as a source of phospholipid, but provide high affinity binding sites essential for the interaction of coagulation proteins (12, 23). This is aptly demonstrated in studies of the prothrombinase complex, where platelets catalyze thrombin formation 20-fold faster than saturating concentrations of phospholipids (12). Similarly, endothelial cells possess specific binding sites for Factor IXa IXa which participates in Factor IXa-VIII activation of Factor X. The nature of this cellular receptor is unclear, although recent studies suggest that a membrane protein may be involved (23).

Studies on activated protein C have highlighted the importance of a negatively charged phospholipid surface for maximal anticoagulant activity (5). The ability of platelets or endothelial cells to provide this support has so far not been fully delineated in man.

Our results confirm that the activity of APC is strongly dependent on the presence of phospholipids. We have demonstrated this using a Factor Xa recalcification time of Al(OH)₃ adsorbed plasma, assays of thrombin generation in recalcified plasma, and studies of the rate of inactivation of purified Factor Va. Surprisingly, intact platelets failed to effectively support the activity of APC in all three systems, and when mixed with phospholipid negated the effect of this surface.

Assays of thrombin formation in recalcified plasma indicate that in the presence of 5 × 10⁷/ml platelets a 15-fold greater concentration of APC was necessary to achieve an anticoagulant effect equivalent to that obtained using phospholipid. This same number of platelets was capable of generating thrombin in the absence of APC at rates comparable to those obtained using phospholipids.

In all these experiments the platelets were not specifically activated. However, it is likely that the washing procedure coupled with the early generation of thrombin during the coagulation reaction would have resulted in rapid platelet activation and the concomitant release of the APC inhibitor. This is supported by the observation that stimulation of the platelets with physiological agonists resulted in the release of APC-inhibitory activity, suggesting that it resides in the secretory granules. Dixon plot analysis of the interaction between the platelet APC-inhibitor and APC at two different concentrations of Factor Va suggested a noncompetitive inhibition. Meaningful calculation of the inhibitory constant for the reaction would be best performed when the inhibitor is purified.

Washed platelet membranes supported the anticoagulant activity of APC, with rates of Factor Va inactivation comparable to those obtained with phospholipid. These observations suggest that once released the APC inhibitor binds weakly to the platelet surface and becomes dislodged during the preparation of the membranes.

The fact that intact platelets inadequately support the effects of APC is not peculiar to man. In the bovine system Harris and Esmon (7) have reported that nonphysiological concentrations of APC (5 µg/ml) are required to saturate platelet binding sites. Furthermore, these high concentrations of APC resulted in rates of Factor Va inactivation significantly less than those obtained in the presence of phospholipids. These workers did not, however, explore the reasons for the difference between the activity of platelets and phospholipid.

The platelet APC inhibitor reported in our study differs from the circulating protein C inhibitor characterized by Suzuki et al. (10). Inhibitory activity did not require preincubation with APC and was not enhanced by heparin. Initial characterization of the platelet APC inhibitor suggests that it is a protein of high molecular weight. However, until the purification of this inhibitor is complete further comparison with the plasma protein C inhibitor must remain speculative.

In conclusion, the results of this study demonstrate that platelets contain an inhibitor to APC that is released during platelet activation. Although platelets can offer an efficient surface for thrombin formation they are unable to effectively catalyze the activity of APC, and hence should be regarded as a predominantly procoagulant surface.

Acknowledgments

Dr. Jane is a National Health and Medical Research Council Ph.D. student, and this manuscript represents part of his thesis.

This study was supported by a program grant from the National Health and Medical Research Council of Australia.

References


