Formation of Factor Va by Atherosclerotic Rabbit Aorta Mediates Factor Xa-catalyzed Prothrombin Activation

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Abstract

Vascular cell procoagulant activity may be important in the pathogenesis of atherosclerosis. In previous studies, we described the ability of the atherogenic metabolite homocysteine to activate endothelial cell Factor V, a key coagulation cofactor for thrombin generation. The present study was designed to investigate Factor V activity and Factor Xa-catalyzed prothrombin activation by control and atherosclerotic aorta from normal and hypercholesterolemic rabbits. Factor Xa generated ninefold more thrombin on atherosclerotic aortic segments than on control segments. Atherosclerotic segments activated [125I]prothrombin with Factor Xa in the presence of the thrombin inhibitor dansyl arginine-4-ethylpyridine amide and cleaved [125I]-Factor V. This suggests that increases in vessel-wall Factor V activity and Factor Xa-catalyzed prothrombin activation result from activation of vessel-wall Factor V. [125I]-Factor V peptides generated by atherosclerotic aorta were very similar in molecular weight to those generated by homocysteine-treated cells. When vascular endothelium was mechanically removed by brushing, atherosclerotic vessels still generated four- to fivefold more thrombin than control vessels. These data and results from immunochemical studies suggest that Factor V in atherosclerotic vessels is associated with both endothelium and other cells of the lesion. In contrast, Factor V in control vessels is associated primarily with endothelium. The increases in Factor V activity and thrombin formation in the blood vessel wall of hypercholesterolemic rabbits may contribute to the development of atherosclerosis and its complications.

Introduction

Endothelial cell injury is postulated to initiate atherosclerosis (1). Two conditions associated with atherosclerotic vascular disease are elevated serum levels of cholesterol (2) or homocysteine (3). In a previous study, we reported that homocysteine-treated endothelial cells exhibited increased procoagulant activity (4) as a result of activation of endogenous Factor V. Factor V associated with cellular surfaces serves as the receptor for Factor Xa; formation of the Factor Vα-Factor Xa complex results in the activation of prothrombin to the coagulation protease thrombin (5). The ability of perturbed endothelial cells to activate this important coagulation cofactor resulted in enhanced Factor Xa-catalyzed prothrombin activation and increased generation of thrombin (4). Thrombin formation is important, not only in the generation of fibrin, but also in the amplification of the coagulation mechanism (6).

These data led us to consider whether hypercholesterolemia would induce similar changes in vascular procoagulant activity, since exposure of endothelium to high concentrations of atherogenic lipoproteins (modified LDL and β-VLDL) results in abnormal endothelial cell morphology (7, 8). In fact, specific receptors for modified LDL (9) and β-VLDL (10) have been identified on cultured endothelium. Perturbation of endothelial cells by other stimuli, including endotoxin (11) and IL-1 (12), has been demonstrated to induce expression of procoagulant (tissue factor) activity. Expression of vascular procoagulant activity may, therefore, occur in hypercholesterolemia, leading to thrombin generation. Thrombin has been demonstrated to cause monocyte chemotaxis (13) and mitogenesis (14), endothelial cell retraction (15), and release of endothelial cell mitogens (16), events that could be involved in atherogenesis. Thus, thrombin may contribute to the development of the atherosclerotic lesion.

One approach to the study of atherogenic stimuli and endothelial cell procoagulant properties is to use an animal model of atherosclerosis. Although there are limitations in extrapolating from data obtained in these in vivo models to humans atherosclerosis, certain animal models of hyperlipidemia have provided important information relevant to the human form of this disorder (1, 17–20). In this study, we used a rabbit model of diet-induced atherosclerosis to investigate vessel-wall procoagulant properties. In particular, Factor V activity and Factor Xa-catalyzed prothrombin activation in thoracic aorta from control and hypercholesterolemic animals were examined. These coagulant properties were previously noted to be enhanced after in vitro exposure of endothelial cells to the atherogenic stimulus homocysteine (4). The data demonstrate that vessel segments from hypercholesterolemic animals exhibit increased Factor V activity and enhanced Factor Xa-catalyzed prothrombin activation. This procoagulant activity results primarily from induction of a vessel-wall activator of Factor V.

Methods

Materials. Trizma base, BSA (RIA grade), Russell's Viper venom, Protein A-Sepharose, and plasmas deficient in Factors VII-X and Factor V were obtained from Sigma Chemical Co. (St. Louis, MO). Sodium [125I]iodide was obtained from Amersham Corp. (Clearbrook,
IL). Iodogen was supplied by Pierce Chemical Co. (Rockford, IL). DME and other reagents for cell culture were obtained from the University of California, San Francisco Cell Culture Facility and had < 10 pg of endotoxin/ml. Calf serum was supplied by HyClone Laboratories, Inc. (Logan, UT). The thrombin inhibitor, dapsyl arginine-4-ethylpyperidine amide (DAPA),1 was provided by Dr. Philip Majerus (Washington University, St. Louis, MO). A human anti–Factor V IgG was prepared from the plasma of a patient with an acquired antibody. The IgG was isolated from plasma using a Protein A-Sepharose affinity column. This IgG (100 μg) could neutralize 95% of the coagulant activity of 1 μg of purified human plasma Factor V after a 2-h incubation at 37°C. Immunochemical studies to detect Factor V in atherosclerotic lesions were performed using this anti–human Factor V IgG and a Vectastain kit (Vector Laboratories, Inc., Burlingame, CA). Adult bovine aortic endothelial (ABAE) cells were cultured as previously described (4).

Factor X and prothrombin were purified from human plasma as described (21). Human Factor V was purified according to the methods of Kane and Majerus (22). Bradford’s technique was followed to assay protein concentrations (23). Prothrombin and Factor V were iodinated using the Iodogen technique (22). Standard clotting assays were used to measure Factor X activity (24), and Factor X was activated using Russell’s Viper venom coupled to Sepharose (25). Normal rabbit plasma was obtained by venipuncture using an anticoagulant solution containing 3.8% sodium citrate, 10 mM benzamidine, and 0.1 mM PMSF.

Hypercholesterolemic rabbit model. Male New Zealand White rabbits (3-3.5 kg) were fed either a standard diet (control: rabbit chow, Zeigler Brothers, Inc., Gardners, PA) or an atherogenic diet containing 0.5% cholesterol (NIH Zeigler 700-79) for 12-16 wk (26). 10 control rabbits and 10 rabbits on the high-cholesterol diet for this period were studied. Aortas from control and hypercholesterolemic rabbits were obtained after anesthesia and exsanguination. Atherosclerotic aorta generally exhibited diffuse fatty streak lesions, and the involved thoracic segments were taken for assay. Control segments were taken from similar locations of normal aorta. The vessels were placed in warmed, sterile DME containing 1% BSA.

Vessel-wall coagulation assays. Coagulant properties of the vessels were assessed using a Lucite template device (exposed vessel surface area ~ 0.75 cm²), similar to that previously used by our laboratory to study prothrombin activation by bovine vessels and Factor Xa (27). In this prothrombin activation assay, the following additions were made to the template well: 0.4 ml of warmed 20 mM Tris-HCl, 150 mM NaCl (Tris-buffered saline [TBS]), pH 7.4; 35 μg of purified prothrombin; and 0.05 ml of 100 mM CaCl₂. The reaction was initiated by adding 10 ng of Factor Xa. Total reaction volume was 0.5 ml. At intervals, aliquots were removed for thrombin assay (28). Replicate values obtained by this assay varied by ±10% (SEM).

To determine whether vascular tissue not exposed to template manipulation had similar coagulant properties, aortic arches were obtained from control and hypercholesterolemic rabbits. These segments were gently washed with medium to remove blood. The major blood vessels originating from the aortic arch and the root of the aorta were ligated to form a small pouch ~ 1-1.5 cm long. Prothrombin activation by these vessels and Factor Xa was assayed in the vessel segment by using 20% of the amounts of the components listed above; total volume was 0.1 ml. Thus, similar concentrations of reactants were used in studies with the template device and with aortic pouches.

Immunochemical techniques. Cross-reactivity between the anti–human Factor V IgG and rabbit Factor V was demonstrated by immunoblotting (29). Purified human Factor V and normal rabbit plasma were subjected to 5% nonreduced SDS-PAGE. Separated proteins were transferred to nitrocellulose filter paper. The total protein was then visualized with Ponceau S (30). In a duplicate blot, Factor V was detected using the anti–human Factor V IgG and an 125I-labeled second antibody. Visualization was by autoradiography.

The presence of Factor V in aorta from control and hypercholesterolemic rabbits was examined by peroxidase immunochemistry as described previously (31). 2-mm² segments of aorta were fixed for 4 h at 4°C with freshly prepared 4% formaldehyde in 0.15 M phosphate buffer, pH 7.4, and then infiltrated for 12 h with 18% sucrose in PBS. The tissue was then frozen in isobutane cooled with liquid nitrogen. 6-8-μm frozen sections were processed for immunocytochemical examination on gelatin-coated slides (31). 0.15% Triton X-100 was included in the initial blocking solution to make the cells more permeable. The primary antibody was human anti–human Factor V IgG. This antibody cross-reacted with rabbit Factor V. In control experiments in which the solution that contained the primary antibody was incubated with authentic human Factor V, reactivity with the rabbit aortic segments was completely blocked.

In certain experiments, endothelial cells were removed from aortic segments to determine the contribution of these cells to overall vessel-wall Factor V activity. Endothelium was removed by brushing the intimal surface with a moist camel-hair brush and rinsing the segment with PBS. Removal of ~ 95% of the endothelial cells was confirmed by light-level microscopy of the plastic-embedded sections (32).

Assays of Factor V (Vₐ) activity. Factor V (Vₐ) activity of control and atherosclerotic aortic segments was measured by freeze-thawing ~ 1-cm² punch biopsy samples of the vessel wall in 0.5 ml of TBS, followed by assay of Factor V activity using a clotting technique (33). Factor V activity was expressed as mU/μg of protein; 1 U of Factor V activity is equivalent to that present in 1 ml of pooled human plasma.

Two methods were used to determine whether atherosclerotic vessel segments contained Factor Vₐ activity. In the first method, cleavage of 125I-prothrombin by Factor Xa and either control or hypercholesterolemic aortic segments was determined in the presence or absence of the thrombin inhibitor DAPA (100 μM). The rationale for these experiments is that Factor Xₐ cannot cleave 125I-prothrombin in the presence of a thrombin inhibitor unless Factor V has been activated. ~ 1-cm² vessel segments were incubated in 0.5 ml of culture medium that contained 30 μg of 125I-prothrombin. After addition of 10 ng of Factor Xₐ, aliquots of the incubation were removed at intervals, subjected to nonreduced SDS-PAGE, dried, and processed for autoradiography (25) to determine the extent of 125I-prothrombin cleavage.

The second method was used to determine whether atherosclerotic vessel segments were able to activate exogenous Factor V. 300 ng of freshly prepared 125I-Factor V was added to ~ 1-cm² control or hypercholesterolemic aortic segments in 0.5 ml of DME-BSA. At intervals, the medium was removed and 0.1 ml of reduced SDS–gel buffer was added to the segments. Solubilized material associated with the vessel segments was analyzed by reduced SDS-PAGE and autoradiography.

Lipoprotein studies. Human LDL was isolated from normal plasma as described (34), and the lysine residues were modified by acetylation with dityke (35). Hyperlipoproteinemic plasma was obtained from cholesterol-fed rabbits for isolation of β-VLDL (35). To determine whether purified lipoproteins could affect Factor V activity of cultured endothelial cells, purified native and modified lipoproteins were incubated with confluent ABAE cells for 3–48 h in serum-free medium. The cells were processed as described for determination of Factor V activity (4).

Results

Factor Xₐ-catalyzed prothrombin activation by atherosclerotic aorta. Rabbits that had been fed a diet containing 0.5% cholesterol for 12–16 wk were routinely studied. As a result of this diet, serum cholesterol levels were > 1,000 mg/dl at this time, and typical fatty streak atherosclerotic lesions of the aorta were grossly visible. In contrast, serum cholesterol levels in control

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1. Abbreviations used in this paper: ABAE, adult bovine aortic endothelial cells; DAPA, dapsyl arginine-4-ethylpyperidine amide; PDGF, platelet-derived growth factor; TBS, Tris-buffered saline.
rabbits were < 100 mg/dl and no evidence of atherosclerotic disease was present, as determined by histologic examination.

Two methods were used to examine Factor Xa-catalyzed thrombin activation by atherosclerotic aortic segments. In one study, vessel segments were placed in a template device to conduct incubations; in the other method, an intact aortic arch was obtained and the reactions were conducted within the aortic lumen. Fig. 1 depicts a typical time course of thrombin generation by control and atherosclerotic vessels in the template device. Incubation mixtures included purified prothrombin, Factor Xa, and calcium. Typically, prothrombin activation by atherosclerotic vessels was enhanced eight- to ninefold over that produced by control vessels. Omission of Factor Xa or substitution of the zymogen, Factor X, resulted in a failure of the vessel segments to activate prothrombin. Thus, prothrombin activation by atherosclerotic vessels occurred via the Factor V-Va-Factor Xa pathway, and not by an activator of Factor X or prothrombin.

Experiments with aortic arch segments were conducted to ensure that the enhanced prothrombin activation was not an artifact that resulted from placing the vessel segments flat in the template device. These aortic arch segments obtained from hypercholesterolemic animals also activated nine times more prothrombin than did control aortic arches (data not shown), which demonstrated that the enhanced prothrombin activation was not an artifact.

The duration of hypercholesterolemia required for enhanced aortic Factor Xa-catalyzed prothrombin activation was also investigated. In these studies, we measured Factor Xa-catalyzed prothrombin activation by aortic segments from rabbits on the diet for 5 d, 12 d, 4 wk, 5 wk, and 6 wk. As in other experiments, vessel segments were selected from similar areas of the thoracic aorta. Aortic segments from some animals on the diet 4 wk had visible fatty streaks, but they did not generate more thrombin than control segments. In contrast, after 5 wk of hypercholesterolemia, atherosclerotic vessels routinely exhibited increased prothrombinase activity.

In two rabbits that had been on the high-cholesterol diet for 14 wk, focal atherosclerotic lesions of the thoracic aorta were present instead of the diffuse lesions commonly observed. Assays of Factor Xa-catalyzed prothrombin activation by these aortic segments revealed increased activity only in involved areas; assays using what grossly appeared to be uninvolved aortic segments generated a level of thrombin very similar to that seen in control aortic segments. The inability to detect thrombin generation by vessel segments either from hypercholesterolemic animals on the diet for < 4 wk or from focally uninvolved vessel segments may reflect insensitivity of our thrombin time assay to low concentrations of thrombin (the lower limit of detection was ~ 0.5 U/ml).

Experiments were also performed with segments of superior vena cava from control and hypercholesterolemic rabbits; atherosclerotic plaques were sometimes observed in this venous tissue of hypercholesterolemic rabbits. Factor Xa-catalyzed prothrombin activation by vena-caval segments was enhanced threefold compared with that seen with control venous segments.

**Factor V (V<sub>a</sub>) activity of atherosclerotic aorta.** Although our human anti-Factor V IgG could immunologically detect rabbit Factor V, it did not neutralize rabbit Factor V coagulant activity. Therefore, to correlate increased aortic Factor Xa-catalyzed prothrombin activation with vessel-wall Factor V activity, punch biopsy samples from control and atherosclerotic aortic segments were obtained from five rabbits of each group, and Factor V (V<sub>a</sub>) activity was measured by clotting assay (Table I). Atherosclerotic segments were diffusely involved with lesion. Compared with control aortic segments, atherosclerotic aortic biopsy samples possessed approximately three- to fourfold more Factor V (V<sub>a</sub>) activity; this increase was statistically significant (P < 0.05).

Previous studies with another atherogenic stimulus, homocysteine, indicated that activation of endogenous Factor V, and not increased cellular concentration of Factor V, was associated with enhanced Factor Xa-catalyzed prothrombin activation (4). To determine whether vessel-wall Factor V was activated in the rabbit model of hypercholesterolemia, we studied the ability of Factor Xa and control or hypercholesterolemic vessel segments to cleave [125I]-prothrombin in an assay containing the thrombin inhibitor DAPA. The rationale for these experiments, based on earlier studies, is that prothrombin activation by endothelial cells and Factor Xa requires thrombin to activate Factor V (4, 25). Fig. 2 is an autoradiogram demonstrating that with control aortic segments, Factor Xa cleaved [125I]-prothrombin in the absence of DAPA (left lane), but not in its presence (right lane). These data are similar to those obtained with normal cultured aortic endothelial cells (25). With atherosclerotic vascular tissue, Factor Xa cleaved more [125I]-prothrombin than did control vascular tissue in the

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**Table I. Vessel-Wall Factor V (V<sub>a</sub>) Activity in Control and Atherosclerotic Segments**

<table>
<thead>
<tr>
<th>Vessel segment</th>
<th>Factor V (V&lt;sub&gt;a&lt;/sub&gt;) activity mU/µg protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.08±0.02</td>
</tr>
<tr>
<td>Atherosclerotic</td>
<td>0.28±0.05</td>
</tr>
</tbody>
</table>

Control and atherosclerotic aortic punch biopsies (~ 1 cm²) were obtained, placed in 0.5 ml of TBS, and freeze-thawed before assay of Factor V (V<sub>a</sub>) activity. Atherosclerotic samples were obtained from rabbits maintained on the high-cholesterol diet for 12–16 wk. Each value represents the mean±SEM of five experiments done on aortic segments of five rabbits.

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**Figure 1.** Time course of thrombin generation by Factor Xa and either control or atherosclerotic aortic segments in the template device. Vessel-wall surface area was ~ 0.75 cm². Atherosclerotic segments were obtained from rabbits on the high-cholesterol diet for 14 wk. Prothrombin activation was measured by incubating 70 µg of prothrombin/ml and 20 ng of Factor Xa/ml in TBS containing 10 mM CaCl₂. At intervals, aliquots were removed for assay of thrombin concentration. Each value represents the mean±SEM of three experiments.

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Figure 2. Cleavage of ¹²⁵I-prothrombin by Factor X₄ and control or atherosclerotic aortic segments. ¹²⁵I-Prothrombin activation was performed as described previously in the presence or absence of the thrombin inhibitor DAPA (25). Aliquots were obtained after 20 min and processed for nonreduced SDS-PAGE, which was followed by autoradiography. The autoradiogram of the dried gel is shown. The band just above thrombin represents an intermediate product of prothrombin activation (25).

To demonstrate directly the presence of a Factor V activator in atherosclerotic aorta, ¹²⁵I-Factor V was incubated with vessel segments obtained from control or hypercholesterolemic rabbits. As shown in Fig. 3 A, significant cleavage of the single-chain protein resulted after incubation for 10 and 30 min with atherosclerotic but not with control vascular tissue. Prominent ¹²⁵I-Factor V₈ peptides with apparent molecular weights of 240,000, 180,000, 108,000, and 70,000–80,000 were observed. A comparison of the ¹²⁵I-Factor V₈ peptides produced by homocysteine-treated cultured endothelial cells (4) is shown in Fig. 3 B. The Factor V₈ peptides produced by both atherogenic stimuli had very similar molecular weights.

To determine the contribution of vascular endothelial cells to the increased vessel-wall Factor V activity, segments of control and atherosclerotic aorta were assayed for prothrombin activation before and after removal of the endothelium with a moist camel-hair brush. Removal of vascular endothelium from atherosclerotic aorta was associated with a 50% reduction in thrombin formation, while brushing of control vessel segments abolished thrombin formation (Fig. 4). Similar findings were observed in aortic segments obtained from three additional rabbits. Histologic examination of plastic-embedded sections made after the assays confirmed the removal of ~95% of the endothelium by brushing and the retention of ~80–90% of the endothelial cells in the nonbrushed segments (Fig. 5).

Immunochemical localization of Factor V antigen in control and atherosclerotic aorta. The ability of Factor X₄ to activate prothrombin after the removal of most endothelial cells from atherosclerotic vessel segments suggested that Factor V (V₈)-like activity might be associated with components of reduced SDS-PAGE and by autoradiography. The autoradiogram of the dried gel is shown. (B) Comparison of the ¹²⁵I-Factor V₈ peptides produced by atherosclerotic aorta or homocysteine-treated cultured endothelial cells. A Factor V standard and thrombin-generated Factor V are also depicted.

Figure 3. (A) Cleavage of ¹²⁵I-Factor V by control and atherosclerotic aortic segments. 300 ng of ¹²⁵I-Factor V/ml was incubated in DME that contained 1% BSA with ~1 cm² aortic segments obtained from control or hypercholesterolemic rabbits. At intervals, medium was removed and reduced SDS-gel buffer was added to the segments. Solubilized material associated with the vessel segments was analyzed by
Intact Atherosclerotic

Brushed

Inbtact _{-}{}ushed

Control

5 10 15 20
25 30
Time (min)

Figure 4. Reduction in vessel-wall Factor Xₐ-catalyzed prothrombin activation after removal of endothelium. Segments of control and atherosclerotic aortas were assayed for prothrombin activity before and after removal of endothelium with a moist camel-hair brush. Removal of endothelium from atherosclerotic aorta was associated with a 50% reduction in thrombin formation. Each value represents the mean of duplicate determinations.

Figure 5. Photomicrographs of 2-μm, toluidine-stained, plastic-embedded atherosclerotic aortic sections after prothrombin activation assays. After completion of the experiments described in Fig. 4, vessel segments were processed for histologic analysis as described under Methods. (A) Atherosclerotic aorta before brushing. (B) Atherosclerotic aorta with endothelium removed (“brushed”). EC, endothelial cell; FC, foam cell. × 150.

the blood vessel wall other than endothelial cells. To identify Factor V antigen in rabbit aortic tissue, it was first necessary to confirm the cross-reactivity of the anti–human Factor V IgG with rabbit Factor V. This was done with an immunoblot technique (Fig. 6), which clearly demonstrated that the anti–human Factor V IgG cross-reacted with rabbit Factor V. To assess the distribution of Factor V in atherosclerotic aorta, immunocytochemistry was performed using IgG from a patient with an autoantibody to Factor V. Factor V antigen (indicated as black peroxidase reaction product) was observed in both control (Fig. 7 B) and atherosclerotic aortic segments (Fig. 7 D). In control aorta, peroxidase reaction product was confined to the endothelium, whereas in atherosclerotic aorta, reaction product was also observed deeper within the intimal plaque. Preincubation of the anti–human Factor V IgG with purified human Factor V prevented the peroxidase reaction in both control and atherosclerotic segments (Fig. 7, A and C, respectively). (The hematoxylin counterstain results in promi-
Cells treated with homocysteine; lysates of treated cells exhibited a fourfold increase in Factor V activity, while intact cells enhanced Factor \( \text{Xa} \)-catalyzed prothrombin activation 9- to 10-fold (4).

The Factor V in control vessels was associated with endothelial cells exclusively, as demonstrated by immunocytochemistry. In contrast, Factor V in atherosclerotic vessels was associated with both endothelial cells and cells deeper within the atherosclerotic lesion (presumably smooth muscle cells, macrophages, or both). There are reports that the hypothesis that both macrophages and vascular smooth muscle cells may be responsible for Factor V staining in atherosclerotic lesions. Rothberger and McGee (36) have reported that rabbit alveolar macrophages secrete Factor V, and one of the authors (G. M. Rodgers) has recently demonstrated that cultured vascular smooth muscle cells synthesize and secrete Factor V (37). Factor V antigen was not detected in smooth muscle cells of control vessels (Fig. 7). Therefore, the synthesis of Factor V in smooth muscle cells in atherosclerotic lesions must be induced, perhaps in response to lipid accumulation, cell proliferation, or both. Alternatively, since Factor V is a plasma protein, its accumulation in the atherosclerotic lesion may result from increased permeability of the vessel wall (38). Previous ultrastructural localization studies of IgG in aortic tissue from atherosclerotic rabbits (39) have suggested that increased permeability or filtration may account for a portion of the increased Factor V antigen seen in our experiments. Hansson et al. studied the deposition of IgG in vascular tissue of rabbits with diet-induced atherosclerosis by using immunohistochemistry at the electron microscope level (39). Immunoreactive material was found only in the 5–8 \( \mu \)m of lesion adjacent to the endothelium. In our studies (Fig. 7), Factor V was observed throughout the lesions to a depth of \( \sim 1,400 \mu \)m. Therefore, some of the peroxidase staining in our experiments may represent increased vascular permeability. However, it is unlikely that the Factor V antigen found so much deeper within the lesion resulted solely from increased vascular permeability. Another possibility would be that cells within the atherosclerotic lesion (vascular smooth muscle cells and macrophages) also contributed to this Factor V antigen.

We have previously reported that incubation of cultured vascular endothelial cells with the atherogenic metabolite homocysteine-activated endogenous Factor V (4). The components of hypercholesterolemic plasma responsible for mediating the increased vessel-wall Factor V activity observed in the present study are unknown. Incubation of the atherogenic lipoproteins (LDL, acetoacetylated LDL, and \( \beta \)-VLDL) with cultured aortic endothelial cells failed to increase endogenous Factor V activity, even though cultured endothelial cells have receptors for these lipoproteins (9, 10) and even though these lipoproteins can be cytotoxic to cultured cells (40, 41). However, we cannot exclude a role for these lipoproteins in the increased expression of Factor V activity by atherosclerotic vessels.

The enhanced vascular Factor V activity and Factor \( \text{Xa} \)-catalyzed prothrombin activation seen in atherosclerotic vessel segments could result from increased amounts or activation of Factor V, or both. Both mechanisms have been found to account for increased Factor V activity of perturbed vascular endothelial cells. Annamalai et al. (42) reported that mechanical injury to human umbilical vein endothelial cells resulted in the expression or exposure of additional Factor V antigen, and
we described activation of endothelial cell Factor V by a homocysteine-induced vascular activator (4). The present data suggest that increased vessel-wall Factor V activity in hypercholesterolemic rabbits results from activation of endogenous Factor V. Although increased amounts of Factor V antigen were associated with cells within the atherosclerotic plaque, the importance of this source of Factor V to overall vessel-wall Factor V activity is uncertain. In our studies and in reports by others (43), vascular endothelium remains essentially intact in this animal model. The possibility cannot be excluded that subendothelial expression of Factor V contributes to vessel-wall Factor Xa-catalyzed prothrombin activation in this model. However, the ability of atherosclerotic vessel segments and Factor Xa to cleave $^{125}$I-prothrombin in the presence of a thrombin inhibitor, and the cleavage of $^{125}$I-Factor V by ath-

erosclerotic, but not by control vessel segments are more consistent with the hypothesis of Factor V activation. Nonetheless, both mechanisms may be operative in this model.

The $^{125}$I-Factor V peptides produced by atherosclerotic aortic segments have apparent molecular weights very similar to those generated by homocysteine-treated aortic endothelial cells (Fig. 3 B). These data suggest that both atherogenic stimuli, homocysteine and hypercholesterolemia, induce activation of Factor V by a similar mechanism, perhaps by a similar tissue protease. If this is the mechanism, previous studies with thrombin inhibitors in cultured endothelial cells (4), as well as the present data with DAPA (Fig. 2), indicate that this endothelial cell protease is not thrombin-like. The recent description of neutrophil-derived elastase that could activate Factor V (44), as well as the report of sulfhydryl protease activation of

**Figure 7.** Photomicrographs of hematoxylin-stained control and atherosclerotic aorta showing the immunocytochemical localization of Factor V antigen. Aortic segments from control animals are shown in A and B. Atherosclerotic aortic segments are shown in C and D. The sections shown in B and D were incubated with an anti–Factor V IgG and processed to demonstrate the presence of Factor V (see Methods). The sections shown in A and C were incubated with anti–Factor V IgG that had been neutralized with purified Factor V. The only dark staining seen in A and C is due to hematoxylin staining of nuclei; in B and D, both nuclei and Factor V immunoreaction product are visible. × 50.
Factor V (45), suggests that there may be a variety of cellular proteases that are important in regulation of coagulation by perturbed cells.

Thrombin cleavage of Factor V generates three peptides: an amino-terminal heavy chain, a carboxyl-terminal light chain (both function as Factor V₄ in the presence of calcium ions), and an activation peptide (22, 46). We speculate that the cellular protease(s) found in vascular tissue exposed to atherogenic stimuli cleaves Factor V much as thrombin does, producing a Factor V₄ species consisting of a heavy chain and a light chain that differ in molecular weight from those generated by thrombin. The observation that a similar number of Factor V₄ peptides are generated by thrombin and by the cellular protease supports this hypothesis. Future experiments, which will attempt to determine which of the Factor V₄ peptides generated by the cellular protease(s) bind to Factor X₄, may provide additional information on Factor V proteolysis in our model.

In the hypercholesterolemic animals, increased Factor V activity may lead to increased production of thrombin. Increased amounts of thrombin may contribute to atherogenesis, since thrombin has been shown to be both chemotactic and mitogenic for monocytes (13, 14). Thrombin has also been shown to cause endothelial cell retraction (15) and generation of platelet-derived growth factor (PDGF)–like mitogenic activity (16). The optimal concentration of thrombin required for these effects is 1–10 nM (0.1–1 U/ml) (13, 14, 16), an amount that the vessel wall or cultured vascular cells could generate (27).

Other investigators have observed evidence of thrombin-like activity in human atherosclerotic lesions. Shainoff and Page described the deposition of a modified fibrinogen species within the aortic intima of atherosclerotic vascular tissue (47). This modified fibrinogen had the solubility characteristics of a stabilized clot, but a fibrinopeptide content consistent with fibrinogen. The exact nature of this protein is uncertain. More recently, immunocytochemistry studies using MAbs specific for fibrin determinants have identified fibrin that is presumably derived from thrombin cleavage of fibrinogen within atherosclerotic human vascular tissue (48).

In the initial response-to-injury hypothesis (49), Ross and Harker proposed that the atherosclerotic lesion resulted from smooth muscle cell proliferation; this hypothesis required PDGF secretion from platelets adherent to denuded endothelium. However, such denuding endothelial cell injuries have not been demonstrated to occur before the development of the atherosclerotic lesion. A recent modification of this hypothesis (1) proposed that subtiler endothelial cell perturbation is sufficient to initiate atherogenesis. In this model, perturbed endothelial cells would secrete PDGF, initiating smooth muscle cell proliferation (1). Our data would be compatible with this mechanism, in that thrombin generation resulting from perturbed endothelium may be sufficient to initiate mitogen secretion and other critical events in atherogenesis.

Although there are differences in the pathogenesis of atherosclerosis in animal models and humans (17, 18, 50, 51), the hypercholesterolemic rabbit model shares several features with the human vascular disorder. As recently reviewed by Ross (1), examination of atherosclerotic lesions from several animal models, including hypercholesterolemic rabbits, revealed a cellular composition and histologic pattern similar to that observed in vessels from affected patients (52). These similarities included monocyte–endothelial cell interactions, monocyte migration into the vessel wall, the appearance of macrophage-derived foam cells, and formation of fibrous plaque by proliferating smooth muscle cells. We hypothesize that endothelial cell injury induced by hypercholesterolemia leads to expression of vessel-wall procoagulant (Factor V) activity that may mediate many of these events.

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