A Myeloma Paraprotein with Specificity for Platelet Glycoprotein IIIa in a Patient with a Fatal Bleeding Disorder

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Abstract

Impaired platelet aggregation, normal shape change, and agglutination and normal ATP secretion and thromboxane synthesis in response to high concentrations of thrombin or arachidonic acid were found in a patient with multiple myeloma and hemorrhagic tendency. The purified IgG, kappa or its F(ab')2 fragments induced similar changes when added in vitro to plateletrich plasma from normal subjects. In addition, the paraprotein inhibited adhesion to glass microbeads, fibrin clot retraction, and binding of radiolabeled fibrinogen or von Willebrand factor to platelets exposed to thrombin or arachidonic acid without affecting intraplatelet levels of cAMP. The radiolabeled paraprotein bound to an average of 35,000 sites on normal platelets but it bound to <2,000 sites on the platelets from a patient with Glanzmann’s thrombasthemia. Immunoprecipitation studies showed that the platelet antigen identified by the paraprotein was the glycoprotein IIIa. Furthermore, binding of radiolabeled prostaglandin E1 (PGE1) to resting platelets as well as binding of von Willebrand factor to platelets stimulated with ristocetin were entirely normal in the presence of patient’s inhibitor. These studies indicate that bleeding occurring in dysproteinemia may be the result of a specific interaction of monoclonal paraproteins with platelets. In addition, our data support the concept that the interaction of fibrinogen and/or von Willebrand factor with the platelet glycoprotein IIIb-IIIa complex is essential for effective hemostasis.

Introduction

Patients with dysproteinemia and hemorrhagic tendency often have hemostatic abnormalities and circulating monoclonal paraproteins (1). Most of these paraproteins have been identified as immunoglobulin inhibitors of phospholipids (2) or major components of the coagulation cascade (3–7); however, inhibitors of platelet function have also been reported (8–11). Platelet abnormalities in dysproteinemia include impaired platelet aggre-

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Methods

Preswollen microgranular DEAE cellulose was from Whatman, Inc., Clifton, NJ. Firefly luciferase and luciferin (Chronolume 395) were purchased from Chrono-Log Corp., Haverton, PA; disodium EDTA and 2-dithiothreitol (DTT) were obtained from Fisher Scientific Co., Pittsburgh, PA. EDTA was prepared as a 100-mM stock solution in distilled water (pH adjusted to 7.4 and 8.7 with NaOH). Pepsin (2,500 U/mg) was from Worthington Biochemical, Freehold, NJ; Nonidex P40 was obtained from Particle Data Inc., Elmhurst, IL and Triton X-100 was obtained from Amershamp Corp., Arlington Heights, IL. Acrylamide, N,N-methylene-bis-acrylamide, Coomassie Brilliant Blue R 250, and molecular weight standards were from Bio-Rad Laboratories, Richmond, CA; bovine serum albumin (Pentex fraction V) was from Miles Laboratories, Inc., Elkhart, IN. Hirudin, phenylmethylsulphonyl fluoride (PMSF), ADP, ATP, sodium dodecyl sulfate (SDS), Heps, lactoperoxidase (70–100 U/mg protein) agar, and prostaglandin E1 (PGE1) were from Sigma Chemical Co., St. Louis, MO. Agar and SDS were dissolved in distilled water immediately before each series of experiments; ADP, ATP, and hirudin were dissolved in distilled water and stored in small aliquots at −20°C; PGE1 was dissolved in ethyl alcohol and stored at −20°C. Thrombin (Parke Davis Co., Detroit, MI) was dissolved in distilled water and stored in small aliquots at −70°C. Protein A-Sepharose CL 4B and cyanogen bromide-activated Sepharose were from Pharmacia Fine Chemicals, Piscataway, NJ. Arachidonic acid (AA) and calcium ionophore A23187 were kindly provided by Dr. Renato Saggioratto (Menarini Diagnostici Firenze, Italy). Protosol, [14C]Jnulin, carrier-free 125I-sodium iodide, and [3H]PGE1 were from New England Nuclear, Boston, MA. Istagel scintillation fluid and Tri-Carb 2660 betacounter were from Packard Instrument Co., Inc., Downers Grove, IL. Antisera against human IgG/L chains of kappa and lambda type were from Behringwerke AG, Marburg, West Germany; murine monoclonal IgG to human IgG1, IgG2, and IgG4 were from Thermo-Genetics, S. Mauro Torinese, Italy, and alkaline phosphatase conjugated goat F(ab')2 fragments anti-mouse IgG were from Cappel Scientific Division, Malvern, PA. Highly specific antibodies to thromboxane B2 (TxB2) were from Seragen Inc., Boston, MA. Silicone oils (methyl-silicone 1.0 DC 200 and Hi-phenyl AA, arachidonic acid; cAMP, cyclic AMP; DTT, 2-dithiothreitol; GPlIIa, glycoprotein IIIa; GPlIb-IIIa, glycoprotein IIb-IIIa complex; Mv, relative molecular weight; PFP, platelet-free plasma; PGE1, prostaglandin E1; PMSF, phenylmethylsulfonyl fluoride; PRP, platelet-rich plasma; TxB2, thromboxane B2; vWF, von Willebrand factor.
silicone (125 DC 550) were from W. F. Nye, Specialty Lubricants, New Bedford, MA. Mixtures of DC 200 and DC 550 were prepared as previously described (12). Kits for the radioimmunoassay of cyclic AMP (cAMP) were from Beckton-Dickinson & Co., Oxnard, CA. Von Willebrand factor (vWF) purified, characterized, and labeled according to Ruggeri et al. (13) was kindly provided by Dr. A. Federici, Milano, Italy. Formaldehyde-fixed platelets; fibrinogen purification, characterization and labeling; TxB2 measurements; intraplatelet cAMP; platelet secretion of nucleotides; platelet fibrin clot retraction and binding to platelets of radiolabelled fibrinogen, PGE1, and vWF were handled as recently reported (12). For preparation of platelet-rich plasma (PRP), platelet-free plasma (PFP), and suspensions of washed platelets, nine volumes of blood were collected from the antecubital vein via a 19-gauge scalp vein needle into a plastic tube containing 1 vol of 3.8% trisodium citrate. PRP was obtained by centrifugation of the blood at 200 g for 15 min at room temperature and PFP was obtained by centrifuging fresh PRP in an Eppendorf centrifuge (Beckman Analytical, Milano, Italy) at 12,000 g for 5 min. For preparation of platelet suspensions, aliquots of PRP were cooled on ice for 30 min and EDTA was added to a final concentration of 5.6 mM. After centrifugation at 2,000 g for 6 min at 4°C in a centrifuge (J-6B; Beckman Instruments, Inc.), the platelets were resuspended in a volume of buffer (0.14 M NaCl, 20 mM Tris, 5 mM glucose, and 1 mM EDTA, pH 7.4) equal to that of the discarded plasma, centrifuged at 900 g for 6 min at room temperature, and resuspended at counts in the range of 1–5 × 10^10/ml in a Tyrode's buffer (0.14 M NaCl, 2.7 mM KCl, 1 mM CaCl2, 12 mM NaHCO3, and 0.4 mM Na2HPO4) containing 5 mM glucose; 3.5 mg/ml bovine serum albumin, and 10 mM Hepes, pH 7.4. Platelet counts were determined by phase-contrast microscopy. When trace amounts of [125I]fibrinogen or [125I]vWF were added to the PRP, >99% of these proteins were removed by the washing procedure. Further properties of platelets washed and resuspended in this manner have been previously reported (12). Suspending of washed platelets showed full aggregation only in the presence of 600 nM (final) of added fibrinogen. Therefore, this concentration of fibrinogen was used in all the aggregation studies in which washed platelets were employed. For both PRP and platelets in suspension, aggregation was measured as percent light transmittance 3 min after the addition of the stimulus in an ELVI 840 aggregometer (Elvi-Logos, Milano, Italy). Platelet aggregation on glass microbeads was determined using a pump system (Adaplet "S"; Mascia-Brunelli, Milano, Italy). For this study, 5–7 ml of blood were drawn from the antecubital vein through a 19-gauge butterfly infusion set into a series of 20 ml Terumo syringes containing heparin (50 μl of a 1-U/ml solution) or heparin plus patient's paraprotein (50 μg/ml). After gentle mixing, the syringes were placed in the infusion pump and adhesion to glass microbeads was determined and quantified according to the manufacturer's recommendations.

Purification of patient's paraprotein. Electrophoretic and immunoelectrophoretic studies showed the presence of a monoclonal IgG kappa peak in the serum of the patient. For further studies, patient's paraprotein was first isolated by nitrocellulose paper chromatography, then eluted with phosphate buffer, 0.015 M, pH 7.4, and dialyzed overnight at 4°C and finally applied to a 2 × 20-cm DEAE column pre-equilibrated with the same buffer. Elution was performed with 300–400 ml of phosphate buffer, pH 7.4, in a gradient from 0.015 to 0.30 M, as recommended by Fahey and Terry (14). Patient's paraprotein eluted from the column at low phosphate molarity (0.015–0.060). At this low molarity only immunoglobulins of the G type are eluted (14). Confirmation of the purity of the preparation was obtained when the eluted paraprotein was incubated overnight at 4°C in polystyrene micro-wells and the coated solid phase was allowed to react with monoclonal mouse immunoglobulins against human IgG1, IgG2, and IgG4. After 3 h of incubation at room temperature, purified goat anti-mouse F(ab')2 fragments coupled with alkaline phosphatase were added and, after an additional 3 h incubation at room temperature, p-nitrophenyl-phosphate was added. The reaction was stopped 30 min later by adding 3 N NaOH. These studies showed a strong colored reaction only in the wells in which anti IgG2 antibodies had been added. IgG4 paraproteins from the sera of two other patients with multiple myeloma and no bleeding tendency were purified and characterized in a similar fashion and used as a control throughout. Immunoglobulins G from the sera of normal subjects were purified according to Fahey and Terry (14) and IgG subclasses were separated from IgG3 in a protein A-Sepharose column according to Duhamel et al. (15). No quantitative or qualitative differences in the results were found when the paraproteins were purified as reported above or isolated from the serum on a protein A-Sepharose column (15).

F(ab')2 preparation. Purified paraprotein was dialyzed overnight at 4°C against a 0.2-M sodium acetate solution pH 4.1 after which freshly prepared pepsin (1 mg/ml) was added in an amount equal to 2% of the antibody's weight and the solution was incubated at 37°C. 18 h later, digestion was stopped by centrifuging the solution at 4,000 rpm for 10 min at 4°C, neutralizing the supernatant with NaOH 1 N, and dialyzing against citrate-phosphate buffer, pH 7. Residual Fc fragments and intact immunoglobulins were removed on a protein A-Sepharose column. Both patient's paraprotein and its F(ab')2 fragments showed a single band of 160,000 mol wt and 105,000 mol wt, respectively, when electrophoresed in a 7.5% SDS-polyacrylamide gel (16).

Paraprotein iodination and binding to human platelets. Intact paraprotein as well as its F(ab')2 fragments were labeled by the iodine monochloride method as previously described for murine monoclonal antibodies (17). Specific activities of these preparations were 4.3 × 10^6 cpm/μg for the intact paraprotein and 3.3 × 10^6 cpm/μg for F(ab')2 fragments. For time-dependent binding studies, aliquots (3 ml) of unlabeled platelets (1–5 × 10^10 platelets/ml) in PRP or in suspensions of washed cells were incubated at room temperature (22°C) with AA (10 μM) or microliter amounts of buffer. After a 3-min incubation, labeled paraprotein was added at a final concentration of 10 μg/ml and, at intervals between 1 and 15 min, 0.4 ml of the suspension was removed and layered onto 50 μl of silicone oil in a 0.5 ml of micro-Eppendorf tube. Free and platelet-bound ligand was separated by centrifugation for 2 min at 12,000 g in an Eppendorf centrifuge and counted in a Beckman DP 5,500 gamma-counter (DP 5,500; Beckman Instruments, Inc.). For dose-response studies, 500-μl aliquots of unlabeled platelets were incubated with 10 μM AA or equal volumes of buffer for 3 min, after which labeled paraprotein was added in final concentrations between 1 and 128 μg/ml. Binding was determined 1 min later. Specific binding was calculated by subtracting the binding measured in the presence of a 20-fold excess of unlabeled paraprotein from the total. This latter nonspecific binding represented 1.73±0.07% of the total radioactivity bound. Binding was not corrected for the radioactivity trapped as determined by adding trace amounts of 125I-inulin. Trapping never exceeded 0.5% of the total bound.

Effect of patient's paraprotein on aggregation, secretion, and binding of vWF to platelets. Increasing amounts (from 1 to 128 μg/ml, final) of patient's paraprotein or 500 μg/ml of control antibodies were incubated with 0.5 ml PRP (or washed platelets in suspension) that had been stirring at 1,000 rpm for 1 min at 37°C. After a further minute of stirring the agents were added and 1 min later, buffer (or fibrinogen, 600 nM final) was added. In parallel, unlabeled patient's paraprotein, in concentrations ranging from 1 to 128 μg/ml, were incubated with platelet suspensions in the absence of stirring at 22°C. After 1 min, the aggregating agent was added and after a further minute, [125I]fibrinogen (600 nM, final) or [125I]vWF (20 μg/ml, final) were added. Binding of the ligands was measured after 5 min for fibrinogen and after 45 min for vWF (12). When thrombin was the stimulus employed to expose binding sites for fibrinogen, hirudin (0.1 U/ml) was used to prevent clotting and was added simultaneously with fibrinogen. In other studies, platelet suspensions (1–5 × 10^9/ml) were incubated at room temperature for 1 min with 50 μg/ml of patient's paraprotein or buffer after which 10 μM thrombin, 10 μM AA, or 1.5 mg/ml ristocetin were added. 3 min later, increasing concentrations of [125I]fibrinogen (from 37.5 to 1,200 nM final) or [125I]vWF (from 1.25 to 40 μg/ml) were added, and at appropriate time-intervals (5 min for fibrinogen and 45 min for vWF), free and platelet-bound ligands were separated in silicone oil and counted. When ristocetin was the stimulus, formaldehyde-fixed platelets were used.

Studies on specificity of patient's paraprotein. Patient's paraprotein specificity was determined as described in detail elsewhere for a murine monoclonal antibody (17). Briefly, 6 × 10^9 washed platelets in 1.5 ml
Tyrode's buffer pH 7.4 were radiolabeled with 2 mCi of \(^{125}\text{I}\) by reacting with 2 mg of lactoperoxidase (18), washed, twice, resuspended in Tyrode's buffer containing Ca\(^{2+}\) (1 mM) or EDTA 10 mM (pH 7.4 or 8.7), and lysed at 4°C with 0.5% of Nonidet P40. 15 min after the addition of Nonidet P40, the lysis was centrifuged at 12,000 g for 15 min and aliquots of the supernatant (containing \(\sim 10^9\) cpm) were incubated at 4°C with 25 \(\mu\)l of patient's paraprotein. 18 h later the complexed antigen was bound to 25 \(\mu\)l of protein A-Sepahrose (0.2 g/ml) and after 6 h at 4°C the pellet was washed extensively with Tyrode's buffer and the bound protein was released by adding 3% SDS with or without 0.2 M DTT and heating to 100°C for 3 min. After centrifuging the sample at 12,000 g for 5 min, amounts of the supernatant (containing \(\sim 30 \mu\)g of protein) were electrophoresed on a 7.5% SDS polyacrylamide gel according to Laemmli (16). Molecular weight standards were run on each gel. Gels were run at 40 mA constant current until the front (visible as refractive index difference) reached the bottom of the gel. Running time was 4–5 h. Gels were then stained with Coomassie Blue, dried, autoradiographed, and identified after 1 wk of development at \(-70^\circ\text{C}\). The blue-stained bands of the molecular weight standards were used to determine the approximate relative molecular weight \((M_r)\) of the bands recorded on the autoradiograms. In another series of studies 5 \(\times\) 10^6 washed platelets were incubated at 4°C in a buffer containing 0.15 M NaCl, 0.02 M Tris, 2.5 mM EDTA, pH 7.4, 1 mM PMSF, 0.2 M DTT, and 1% Triton, pH 7.5. 15 min later, the solubilized platelet suspension was centrifuged at 40,000 \(\times\) g for 60 min at 4°C and the supernatant applied to a 1 \(\times\) 4-cm column containing 5 mg of patient’s paraprotein coupled to 4 \(\mu\)l of cyanogen-bromide-activated Sepharose. This column had been pre-equilibrated with the same buffer in which solubilized platelets were resuspended. After extensive washing of the column, the affinity-bound proteins were eluted with 5 ml of a 3% SDS solution, heated to 100°C for 3 min, and subjected to a 7.5% SDS polyacrylamide gel electrophoresis (16).

Clinical summary. The patient was a 62-yr-old white male in good health until December 1982, when he began to have bone pain and bleeding tendency (easy bruising, nose and gum bleeding) and noted an 8-kg weight loss. He had no past history of spontaneous bleeding or excessive blood loss following surgery. On physical examination he had petechiae of the arms and legs, diffuse bone pain, splenomegaly, and hepatomegaly without lymphatic involvement. His blood counts showed: hemoglobin, 7.5 g/dl; platelets, 110,000/\(\mu\)l; leukocytes, 10,500/\(\mu\)l, with 53% neutrophils, 2% bands, 35% lymphocytes, 2% atypical lymphocytes, and 8% monocytes. His SMA-12 (Sequential Multiple Analyzer, Technicon) as well as activated and nonactivated thromboplastin time, prothrombin time, thrombin time, fibrinogen, and fibrinogen degradation products were all within normal ranges. Skeletal roentgenograms showed diffuse osteolytic lesions. Bone biopsy showed a hypoplastic marrow infiltrated with plasma cells (>35%). There was an IgG kappa monoclonal paraprotein present in his serum (but not in urine) with a concentration ranging between 2.7 and 3.8 g/dl during the course of the illness. Stools were always 3+ guaiac positive. Endoscopy showed the presence of a gastric ulcer and the patient was given 400 mg cimetidine twice daily. This treatment reduced the bleeding but only partially. The patient died 8 wk after the diagnosis from massive gastrointestinal bleeding. At autopsy he had multiple myeloma, gastric ulcer, and evidence of external and internal (mainly gastrointestinal) hemorrhages.

Results

Patient’s platelet test. Results of the screening tests in PRP from the patient are shown in Table I. Platelet aggregation in response to ADP or epinephrine was almost suppressed and that in response to collagen was markedly reduced. In one experiment in which higher concentrations of collagen and ADP were employed, it was found that the response to collagen (5 \(\mu\)g/ml) was 46% of normal and the response to ADP (50 \(\mu\)M) was only 8% of normal. In contrast, agglutination in response to ristocetin and ATP secretion and TxB2 synthesis in response to thrombin or AA were within normal ranges. As with ADP, shape change in response to collagen was entirely normal (not shown).

Effect of the addition of patient’s paraprotein to normal PRP. Addition of the patient’s purified IgG kappa to normal PRP (adjusted with PFP to platelet counts of 90,000–120,000/\(\mu\)l) caused all the platelet abnormalities seen in the patient’s PRP. Furthermore, when added to the sample for at least 1 min before each test, the paraprotein also inhibited platelet adhesion to glass beads and fibrin clot retraction. Maximal impairment of platelet adhesion, aggregation, and clot retraction was achieved at paraprotein concentrations of 50 \(\mu\)g/ml. Results of seven sets of experiments in which these concentrations of paraprotein were employed are summarized in Table II. Up to 128 \(\mu\)g/ml the inhibitory effect of patient’s paraprotein was not accompanied by changes in platelet shape change (not shown) or intraplatelet levels of cAMP (73.2 + 21.7 pmol/10^6 cells in buffer-treated platelets vs 91.9 + 32.2 in platelets pretreated with 50 \(\mu\)g/ml of patient’s paraprotein, \(P > 0.05\)). Inhibition of ATP secretion and TxB2 formation in PRP depended on the dose of the agonist used, with marked effect at low doses and little effect when higher doses were employed. Similar results were found in suspensions of washed platelets (1 \(\times\) 10^9/ml) when 10–40 \(\mu\)M AA or 10–40 uM thrombin were employed. In addition, normal secretion of ATP (i.e., 3.2 ± 0.5 \(\mu\)M) was observed when unstimred PRP, pretreated with 50 \(\mu\)g/ml of patient’s paraprotein, was incubated

<table>
<thead>
<tr>
<th>Test</th>
<th>Patient</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aggregation by ADP (10 (\mu)M)</td>
<td>3.6±1.3</td>
<td>67.5±7.3</td>
</tr>
<tr>
<td>Aggregation by epinephrine (20 (\mu)M)</td>
<td>2.2±4.1</td>
<td>62.5±3.8</td>
</tr>
<tr>
<td>Aggregation by collagen (1 (\mu)g/ml)</td>
<td>27.9±5.4</td>
<td>65.7±6.7</td>
</tr>
<tr>
<td>Agglutination by ristocetin (1.5 mg/ml)</td>
<td>49.3±10.3</td>
<td>68.4±10.5</td>
</tr>
<tr>
<td>TxB2 synthesis by thrombin (1 U/ml)</td>
<td>416±100.7</td>
<td>378±97.2</td>
</tr>
<tr>
<td>TxB2 synthesis by AA (0.8 mM)</td>
<td>1700±398</td>
<td>1803±349</td>
</tr>
<tr>
<td>ATP secretion by thrombin (1.0 U/ml)</td>
<td>2.5±0.6</td>
<td>2.4±0.8</td>
</tr>
<tr>
<td>ATP secretion by AA (0.8 mM)</td>
<td>1.6±0.2</td>
<td>1.8±0.3</td>
</tr>
</tbody>
</table>

Platelet aggregation and agglutination were measured as percent maximal light transmission 3 min after the addition of the stimulus and adjusting the aggregometer so that PRP and platelet-poor plasma produced 10 and 90% light transmission, respectively; TxB2 synthesis was measured by radioimmunoassay in aliquots of stirred (1,000 rpm) samples 3 min after the addition of the stimulus and expressed as pmol TxB2/3 \(\times\) \(10^9\) platelets; secretion of ATP was determined in a lumig-agarrometer by using luciferin-luciferase reagent (50 \(\mu\)l of a 10 mg/ml solution) and expressed as \(\mu\)mol ATP secreted/\(10^9\) platelets. The values reported for the patient are means±SEM of three determinations; those for controls are composite data from seven PRP studies. A significant difference (\(P < 0.01\)) was always found when the data relative to the aggregation of platelets from the patient were compared with those from controls. In all the other determinations the \(P\) value was always >0.05.

Myeloma Paraprotein to Glycoprotein IIIa 159

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Table II. Effect of Addition of Patient’s Paraprotein to Normal PRP in Platelet Screening Tests

<table>
<thead>
<tr>
<th>Test</th>
<th>IgG&lt;sub&gt;i&lt;/sub&gt;</th>
<th>Buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelet adhesion</td>
<td>16.3±3.6</td>
<td>86.5±10.2</td>
</tr>
<tr>
<td>Clot retraction by thrombin (1 U/ml)</td>
<td>18.6±2.1</td>
<td>96.5±2.3</td>
</tr>
<tr>
<td>Aggregation by ADP (10 μM)</td>
<td>6.8±3.4</td>
<td>67.5±7.3</td>
</tr>
<tr>
<td>Aggregation by epinephrine (20 μM)</td>
<td>5.1±1.3</td>
<td>62.5±3.8</td>
</tr>
<tr>
<td>Aggregation by collagen (1 μg/ml)</td>
<td>12.4±5.8</td>
<td>65.4±6.7</td>
</tr>
<tr>
<td>Aggregation by AA (0.5 mM)</td>
<td>10.5±3.1</td>
<td>80.5±10.3</td>
</tr>
<tr>
<td>Agglutination by ristocetin (1.5 mg/ml)</td>
<td>58.3±6.5</td>
<td>68.4±10.5</td>
</tr>
<tr>
<td>TxB&lt;sub&gt;2&lt;/sub&gt; synthesis by thrombin (0.1 U/ml)</td>
<td>21.5±8.2</td>
<td>96.5±12.3</td>
</tr>
<tr>
<td>TxB&lt;sub&gt;2&lt;/sub&gt; synthesis by thrombin (1 U/ml)</td>
<td>400±78.3</td>
<td>378±97.2</td>
</tr>
<tr>
<td>TxA&lt;sub&gt;2&lt;/sub&gt; synthesis by AA (0.5 mM)</td>
<td>75.2±13.4</td>
<td>336.8±79.3</td>
</tr>
<tr>
<td>TxA&lt;sub&gt;2&lt;/sub&gt; synthesis by AA (0.8 mM)</td>
<td>1740±316</td>
<td>1803±349</td>
</tr>
<tr>
<td>ATP secretion by thrombin (0.1 U/ml)</td>
<td>0.7±0.3</td>
<td>1.7±0.4</td>
</tr>
<tr>
<td>ATP secretion by thrombin (1.0 U/ml)</td>
<td>2.7±0.1</td>
<td>2.4±0.8</td>
</tr>
<tr>
<td>ATP secretion by AA (0.5 mM)</td>
<td>0.5±0.3</td>
<td>1.7±0.2</td>
</tr>
<tr>
<td>ATP secretion by AA (0.8 mM)</td>
<td>2.0±0.4</td>
<td>1.8±0.3</td>
</tr>
</tbody>
</table>

Platelet adhesion to glass microbeads was determined in an Adeplat “S” system employing blood freshly drawn into syringes containing heparin or heparin plus patient’s paraprotein; fibrin clot retraction was calculated by relating the volume of serum extruded at 60 min to the total amount of the initial mixture (PRP + thrombin) and expressed as a percentage; methods employed for studying and measuring platelet aggregation, agglutination, TxB<sub>2</sub> synthesis, and ATP secretion are given in the legend of Table I. Each value is the mean±SEM of composite data from seven PRPs studied. Addition of the antibody had no effect (P always > 0.05) on the agglutination of platelets in response to ristocetin as well as on the secretion of TxB<sub>2</sub> in response to high concentrations of thrombin or AA. In all the other studies patient’s paraprotein significantly affected platelet function (P always < 0.01).

with 5 mM EDTA for 5 min before the addition of A23187 (50 μM). These data are similar to those obtained using platelets from thrombasthenic patients (12) and comparable to those obtained by incubating normal platelets with a murine monoclonal antibody to the glycoprotein IIb-IIIa complex (GPIIb-IIIa) (19) and were interpreted to suggest that the aggregation of platelets may act to enhance secretion of nucleotides and TxB<sub>2</sub> synthesis when low doses of agonists are used, whereas at higher doses secretion and TxB<sub>2</sub> formation are largely independent of aggregation. Similar results were found when equimolar concentrations of F(ab')<sub>2</sub> fragments of patient’s paraprotein were employed. In contrast, concentrations of the two control paraproteins or normal IgG<sub>i</sub> as high as 500 μg/ml did not affect platelet adhesion, aggregation, secretion, TxB<sub>2</sub> synthesis, and clot retraction.

Binding of 125I-patient’s paraprotein to platelets. Both in PRP and washed platelets in suspension binding of radiolabeled monoclonal paraprotein to unstimulated or stimulated platelets was complete in 1 min. Up to 15 min incubation, >97% of the labeled ligand could be displaced within 10 min by a 20-fold excess of unlabeled paraprotein. The extent of the binding was the same in PRP and platelets washed free of fibrinogen as well as in the presence or absence of EDTA (10 mM). Estimates of the affinity determined as recommended by Klots (20) and Coller et al. (19) showed that at saturating levels of added ligand (i.e., 10–15 μg/ml for the inflection point and 30–50 μg/ml for maximal binding) the paraprotein bound to an average of 35,523 sites/cell (1 SEM = 3,476) in PRP from normals (Fig. 1) (apparent dissociation constant, 16.4 ± 3.9 μg/ml) and to 1,989±284 sites/cell in PRP from a thrombasthenic patient. Laboratory studies on the platelets from this patient have been previously reported (12).

Effect of the addition of patient’s paraprotein of 125I-fibrinogen, 125I-vWF, and [3H]PGE<sub>1</sub> binding to platelets. Addition of purified IgG<sub>i</sub> kappa to washed platelets inhibited in a concentration-dependent fashion the binding of 125I-fibrinogen in response to thrombin or AA. Maximal inhibition was found at concentrations of added paraprotein of 50 μg/ml. Results of eight sets of experiments in which this concentration of inhibitor was employed are summarized in Table III and are similar to those found when higher concentrations of thrombin and AA (i.e., 40 μU/ml for thrombin and 40 μM for AA) were used. To further characterize inhibition of fibrinogen binding by patient’s paraprotein, platelets in suspension were incubated at room temperature with a single concentration of the inhibitor (50 μg/ml) or buffer. AA (10 μM) was added 1 min later and

![Figure 1](https://doi.org/10.1172/JCI112270)
fibrinogen binding was measured as a function of $^{125}$I-fibrinogen concentration employed. Analysis of the data by using double reciprocal plots was consistent with a noncompetitive inhibition of the binding by the paraprotein (Fig. 2). Selectivity of the inhibition of fibrinogen binding by the paraprotein was then determined by evaluating the effects of the inhibitor on the binding of PGE$_1$ and vWF to platelets. The former was studied because a murine monoclonal antibody (B59.2) to the platelet receptor for fibrinogen does not affect the binding of this prostaglandin to resting platelets (unpublished observations of the authors); the latter was chosen since the presumed receptor for fibrinogen on the platelet surface is also one of the binding sites for vWF on these cells (13, 21, 22). At concentrations between 50 and 128 μg/ml (final), purified monoclonal paraprotein did not affect the extent of the binding of radiolabeled PGE$_1$ to resting platelets in PRP. At saturating levels of added PGE$_1$ (i.e., at a final concentration of 300 nM), platelets preincubated for 1 min with 50 μg/ml of patient's paraprotein bound 0.28±0.07 pmol PGE$_1$/10$^6$ cells and platelets preincubated with equal microliter amounts of buffer bound 0.31±0.03 pmol PGE$_1$/10$^6$ cells ($P > 0.05$). When we studied the binding of vWF, we found that the paraprotein did not affect the interaction of this adhesive protein with platelets stimulated with ristocetin, while it impaired the binding in response to 10 mU/ml thrombin or 10 μM AA (Table III). Similar results were found when higher concentrations of thrombin or AA (i.e., 40 mU/ml for thrombin and 40 μM for AA) were used. Concentrations of control paraproteins or normal IgG$_1$ as high as 500 μg/ml did not affect binding of fibrinogen or vWF in response to AA or thrombin (not shown).

**Specificity of patient's paraprotein.** The immunoprecipitate formed by the reaction of patient's paraprotein with radiolabeled platelet membrane glycoproteins was solubilized in 3% SDS, reduced with 0.2 M DTT, and autoradiographed. As shown in Fig. 3 A, the inhibitor precipitated very close bands (apparent $M_r$, ~116,000) in the presence of Ca$^{2+}$ (1 mM) or EDTA (10 mM, pH 7.4). These bands were not present in immunoprecipitates in which platelets from a thrombocytopenic patient were used or in those in which platelets from normal donors were incubated with an IgG$_1$ paraprotein from a patient with multiple myeloma and no bleeding tendency. Specificity of patient's paraprotein was further confirmed in affinity chromatography studies. After reduction, two peptides of 120,000 apparent $M_r$ and 110,000 apparent $M_r$ were eluted in addition to a faint band of 65,000 $M_r$, comigrating with albumin in some samples to another band of ~40,000 $M_r$ comigrating with actin. These minor bands were not present in immunoprecipitates and did not interact with the paraprotein in immunoblots (24). However, it is worth mentioning that also the platelet antigens identified by the paraprotein in affinity chromatography as well as in immunoprecipitation studies were not recognized by the inhibitor in immunoblots, possibly because of the loss of the epitope during exposure to SDS and/or electrophoresis.

**Published values** (19, 23) for molecular weights of platelet

![Figure 2. Inhibition of fibrinogen binding to AA-stimulated platelets by patient's paraprotein. Suspensions of 1–5 × 10$^6$ platelets/ml were incubated at room temperature for 1 min with 50 μg/ml of patient's paraprotein (open circles) or buffer (closed circles) after which 10 μM AA was added. 3 min later increasing concentrations of $^{125}$I-fibrinogen (from 37.5 to 1,200 nM, final) were added and 5 min later free and bound ligand were separated in silicone oil and counted. Specific binding was determined by subtracting from the total bound the radioactivity bound in the presence of a 20-fold excess of cold fibrinogen. Non-specific binding never exceeded 15%. The data were then expressed in terms of double-reciprocal plots. Each point is the mean±SEM of four different platelet suspensions tested. Similar results were found when thrombin was the stimulus used.](image)

**Table III. Effect of Patient's Paraprotein on $^{125}$I-Fibrinogen and $^{125}$I-vWF Binding to Stimulated Platelets**

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>Buffer</th>
<th>IgG$_1$</th>
<th>Buffer</th>
<th>IgG$_1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thrombin (10 mU/ml)</td>
<td>10.1±1.6</td>
<td>3.1±0.6*</td>
<td>15.1±3.8</td>
<td>8.6±2.4*</td>
</tr>
<tr>
<td>AA (10 μM)</td>
<td>6.2±1.5‡</td>
<td>2.4±0.5*</td>
<td>3.9±0.2‡</td>
<td>1.21±0.2*</td>
</tr>
<tr>
<td>Ristocetin (1.5 mg/ml)</td>
<td>ND§</td>
<td>ND§</td>
<td>16.7±2.9</td>
<td>18.9±3.1</td>
</tr>
</tbody>
</table>

* P < 0.01 buffer vs. paraprotein.
‡ P < 0.01 thrombin vs. AA-induced binding (in the presence of buffer).
§ ND, not detectable.

Platelet suspensions (1–5 × 10$^6$/ml) were incubated for 1 min with 50 μg/ml of patient's paraprotein or buffer after which 10 mU/ml thrombin or 10 μM AA or 1.5 mg/ml ristocetin were added. 3 min later increasing concentrations of $^{125}$I-fibrinogen (from 37.5 to 1,200 nM final) or $^{125}$I-vWF (from 1.25 to 40 μg/ml) were added and at appropriate time intervals (5 min for fibrinogen and 45 min for vWF), free and platelet-bound ligands were separated in silicone oil and counted. When thrombin was employed to expose binding sites for fibrinogen, hirudin (0.1 U/ml) was used to prevent clotting. When ristocetin was the stimulus, formaldehyde-fixed platelets were used. In the presence of both buffer and patient's paraprotein fibrinogen binding reached saturation at ~600 nM, while vWF at ~20 μg/ml (final) of added protein. Each value is the mean±SEM of composite data from eight suspensions of platelets studied.
glycoproteins suggest that the antigen identified by the paraprotein was the GPIIb-IIIa complex. This complex has been shown to be maximally dissociated by EDTA at pH 8.7 or higher (24–26). Therefore, to determine whether the paraprotein was directed to a single glycoprotein or to the complex, platelets were labeled with $^{125}$I, washed twice in Tyrode’s buffer, and resuspended in the same buffer containing 10 mM EDTA pH 8.7. The immunoprecipitate formed by the reaction of the paraprotein with platelets processed in this manner is shown in Fig. 3 B. A single band was present with a molecular weight of $\sim 115,000$ under reducing conditions (DTT, 0.2 M) and $\sim 90,000$ mol wt under nonreducing conditions. The higher molecular weight upon reduction is characteristic of GPIIla (19, 23).

Discussion

Monoclonal paraproteins that induce hemostatic abnormalities have often been described in dysproteinemia. The immunoglobulin nature of these inhibitors has been established and their mechanism of action has been delineated (1–7). Platelet abnormalities in dysproteinemia have also been reported (8–11), but the mechanism of the effect of paraproteins in the impairment of platelet function is poorly understood. In a recent comprehensive review on this topic (1) it has been stated that “. . . abnormal monoclonal immunoglobulins occurring in dysproteinemias have no known physiologic specificity for the surface of the hemostatic components with which they react,” and that “. . . the adherence of the paraprotein nonspecifically to the platelet surface appears to interfere with normal platelet function.” Our findings suggest that impairment of platelet function in dysproteinemia may also be the result of a specific immunologic reaction of the paraprotein with platelets. Studies over the past years have suggested that monoclonal proteins are normal immunoglobulins produced in excessive quantities (27). However, antibody activity has been clearly demonstrated for only a few of them (28). The paraprotein that we have isolated from our patient has antibody activity biochemically and functionally comparable to that of an antibody isolated from a polytransfused thrombocytopenic patient (29, 30) or from patients with immune thrombocytopenia (31, 32).

Much evidence has been accumulated to indicate that GPIIb-IIIa is the receptor for fibrinogen and vWF on platelets exposed to naturally occurring aggregating agents and that the interaction of fibrinogen and vWF with this complex is essential for effective hemostasis as well as for a variety of in vitro platelet functions (21). Patients with Glanzmann’s thrombasthenia have a bleeding tendency (21) and platelets from these patients lack GPIIb-Illa (18, 21, 33) and exhibit negligible binding of fibrinogen after stimulation with ADP (30, 34–37) and deficient binding of vWF when exposed to thrombin (22). Estimates of the affinity indicate that normal platelets have $\sim 39,000–50,000$ binding sites for fibrinogen (12, 17, 19, 21, 34, 35), a figure comparable to the number of GPIIb-Illa complexes identified on the surface of normal platelets by specific monoclonal antibodies (17, 19, 38–41). GPIIb-Illa may form complexes with fibrinogen (42), and murine monoclonal antibodies to GPIIb and/or IIIa inhibit aggregation as well as binding of fibrinogen or vWF to platelets (13, 17, 19, 38–40). Screening tests (Table I) in platelets from this patient showed abnormalities comparable to those described in Glanzmann’s thrombasthenia (21). Most reports indicate that thrombasthenic platelets do not aggregate at all in response to collagen (21). However, studies by several investigators showed that small aggregates are, in fact, produced (43–46). Similar results are also obtained after in vitro addition to normal platelets of a monoclonal antibody to GPIIb-Illa complex (19). It is unclear whether these aggregates are the result of platelet-platelet interactions or rather the adhesion of several platelets to collagen fibrils, which can simulate the appearance of an aggregate. The in vitro addition of purified paraprotein to normal platelets
caused all the abnormalities observed in platelets from the patient and also impaired adhesion to glass beads, fibrin clot retraction, and binding of fibrinogen and vWF in response to thrombin or AA. Thus, it induced a thrombathenic-like state in normal platelets. In addition, the paraprotein bound to normal platelets (Fig. 1) but only negligibly to platelets from a thrombathenic patient; it reacted with GPIIb-IIIa on immunoprecipitates as well as on an antibody-coupled affinity column and it only inhibited the binding to platelets of agonists whose receptor is known to be GPIIb-IIIa.

The aggregation as well as the binding of fibrinogen and vWF to platelets requires exposure of the receptor, and thromboxane synthesis, nucleotide secretion, translocations of calcium ions across the membranes, and intracellular cAMP are thought to play a major role in this exposure (21, 47). Our data make it unlikely that patient’s paraprotein acts by affecting one or more of these basic events. The ability of platelets to bind fibrinogen involves platelet activation as well as the presence of intact GPIIb-IIIa complexes (21). Since the paraprotein only reacts with GPIIb (Fig. 3 B) and in detergent extracts GPIIb and GPIIIa from reversible heterodimer complexes (48, 49), one could postulate that patient’s paraprotein impairs fibrinogen and vWF binding to platelets by inhibiting the formation of the complex. Our data are not consistent with this possibility since at physiological pH more than one band can be detected on unstimulated intact platelets even in the presence of EDTA (Fig. 3 A). Therefore, it seems reasonable to conclude that patient’s paraprotein induces platelet abnormalities by sterically hindering the association of fibrinogen and vWF with the GPIIb-IIIa.

These data show that bleeding occurring in dysproteinemia may be the result of a specific interaction of monoclonal paraproteins with platelets and support the concept that the interaction of fibrinogen and/or vWF with the GPIIb-IIIa is essential for effective hemostasis.

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