Binding of Quinine- and Quinidine-dependent Drug Antibodies to Platelets Is Mediated by the Fab Domain of the Immunoglobulin G and Is Not Fc Dependent

Mark E. Smith, Diane M. Reid, Charlie E. Jones, James V. Jordan, Carol A. Kautz, and N. Raphael Shulman

Clinical Hematology Branch, National Institute of Diabetes, Digestive and Kidney Diseases,
National Institutes of Health, Bethesda, Maryland 20892

Abstract

The antibody domain controlling reactions between platelet membranes and drug-dependent (dd) antibodies from patients with thrombocytopenia induced by cinchona alkaloids was studied using F(ab')2, Fab, and Fc fragments made from purified dd-IgG. By direct binding radioimmunoassay (RIA) measurements, 20,000 to 50,000 antibody molecules bound per platelet equivalent of purified platelet membranes at apparent saturation with three different antibodies. F(ab')2 and Fab fragments bound to platelet membranes drug dependently but Fc fragments did not. The ability of dd-IgG fragments to compete with intact IgG was quantitatively measured by RIA and by complement fixation. F(ab')2 and Fab competed with intact IgG at an 8:1 and > 50:1 molar ratio, respectively, in RIA, and at a 1.6-3:1 and 44-75:1 ratio, respectively, by complement fixation assays. Fc did not compete with IgG in either assay. We conclude that the Fab domain supports attachment of dd antibody to the platelet surface.

Introduction

Drug-induced antibodies that cause thrombocytopenia attach to platelet membranes in a high affinity reaction only when the sensitizing drug or one of its analogues is present in solution (1-4). In vitro binding of the antibody to platelet membranes has been measured over drug concentrations of 10^-7 to > 10^-2 M (2-4), while in vivo binding, as evidenced by decreases in circulating platelets, has been observed at drug concentrations as low as 10^-9 M (5, 6). The highest concentrations of drug used in antibody binding experiments, 10^-3-10^-1.8 M, supported maximal attachment of antibody to platelets even when antibody concentrations were in the range of 10^-8-10^-9 M (3, 4) and cell membrane receptors were in the range of 10^-8 M. Thus, under conditions of a fixed ratio of antibody concentration to platelet concentration, antibody binding to platelets in vitro is saturable with increasing concentrations of free drug and remains so as drug is increased to at least 10^-2-fold greater than the concentration of either antibody or platelet binding sites.

Since haptenic inhibition of antibody attachment to platelets does not occur, the assumption that simple haptenic conjugation of the drug or a metabolite with the platelet surface creates a site for antibody attachment cannot be defended. Moreover, platelets exposed to drug and washed once in the absence of drug-dependent (dd) antibody are incapable of supporting antibody attachment (3, 4). To be consistent with these observations, it was proposed that antibody binds drug, resulting in an immune complex that can be adsorbed by platelet membrane binding sites (2, 3). Although antigen–antibody complexes generally are considered to be adsorbed by cell membrane Fc receptors through the activated Fc domain of the antibody, the platelet surface component that interacts with drug antibodies has been demonstrated to be the glycoprotein I complex, which has a reduced apparent molecular weight (Mf) of 150,000 (7). This receptor differs significantly from the platelet Fc receptors thus far described which, when reduced, have Mfs of 40,000 (8) and 30,000 (9). Moreover, if Fc receptors were involved, drug antibodies would be expected to bind to more than one cell type, but anti–platelet drug antibodies are exclusively specific for platelets (1, 2, 6, 10). Additionally, drug antibodies that react with granulocytes or erythrocytes show no binding to platelets (6, 10).

These observations indicate that the cell membrane binding site for drug antibodies contributes some of the specificity to the overall interaction. Since adsorption of a drug–antibody complex via the Fc domain does not appear to occur, one might expect dd antibodies to attach via the Fab domain. However, two groups of investigators have described unsuccessful attempts to demonstrate drug-mediated binding of F(ab')2 fragments prepared from dd-IgG (11, 12). Nevertheless, there is a possibility that the anti-IgG used in these experiments may not have reacted with F(ab')2; experiments to document the reactivity of the anti-IgG used were not mentioned. On the other hand, Staphylococcus aureus protein A has been used successfully to detect dd-IgG on platelet surfaces (10, 12-14); hence the Fc domain of the dd-IgG is accessible after binding. Thus, either the Fab domain is primarily involved in binding dd-IgG to the platelet surface or Fc-mediated binding does not interfere with protein A binding to the Fc region. That the latter may be possible is evident from observations on Fc receptor binding of IgG complexed to protein A (15).

We have generated data indicating that the Fab domain of the dd-IgG rather than the Fc domain supports attachment of the antibodies to cell membranes. The high affinity antibody–drug–platelet ternary complex appears to involve cooperative interaction among the three reactants, in that the affinity between any two of the reactants in the absence of the third is too low to be measured by techniques that have been applied to date.

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Address reprint requests to Dr. Shulman, CHB/NIDDK, NIH, Bldg. 10, Rm. 8C101, 9000 Rockville Pike, Bethesda, MD 20892. Dr. Smith's current address is Cancer Research Center, Columbia, MO 65203, and Dr. Jordan's current address is Christ Hospital, Chicago, IL 60453.

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1. Abbreviation used in this paper: dd, drug-dependent.
Methods

Patients. The antibodies studied were from three patients who developed severe thrombocytopenia while taking either quinine or quinidine therapeutically. dd antibody binding in each serum was initially quantitated by complement fixation assays as reported previously (2).

Production of purified IgG and fragments. 10 ml of whole serum containing dd antibody was dialyzed against 0.020 M potassium phosphate, pH 8.2, and applied to a 1.5 × 45-cm column of DEAE-Affigel blue (Bio-Rad Laboratories, Richmond, CA) equilibrated with the same buffer. The protein peak not retained by this column was composed of ~90% IgG and 10% transferrin based upon analysis of the pooled material by silver-stained sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) (16, 17). Fab fragments were prepared from this partially purified IgG by papain digestion (18) and gel filtration on Sephacryl S-300 (2.5 × 95 cm). Eluate fractions containing both Fab and Fc were applied sequentially to a protein A-Sepharose column then an anti-Fc–Sepharose column to remove the Fc fragments, yielding purified Fab fragment containing < 2% contaminants. The bound Fc fragments were eluted from these columns with 1.0 M acetic acid, pH 3.0. The fractions were immediately neutralized by the addition of sodium hydroxide and concentrated by ultrafiltration over an Amicon YM-10 ultrafiltration membrane. The Fab(1/2) fragment was prepared by pepsin digestion of the partially purified IgG (19). The crude digestion mixture was also applied to a Sephacryl S-300 gel filtration column as described above and subsequently residual IgG and degraded Fc fragments were removed by adsorption to a protein A–Sepharose column. The purity of all these fragments was determined by analytical SDS–PAGE (e.g., Fig. 7).

Platelet membrane preparation. Platelets were obtained from normal donors by platelet apheresis using citrate-anticoagulant. EDTA was added to the platelet-rich plasma to a final concentration of 5.0 mM and platelets were washed three times in phosphate-buffered saline (PBS), 5.0 mM EDTA. To prepare membranes, glycerol-loaded platelets were lysed by hypotonic shock and the membrane fragments were separated from intact platelets, cellular debris, and organelles by centrifugation through a discontinuous sucrose density gradient (20). The membrane preparation sedimented to the interface between the 0.25 M and 27% sucrose solutions. The material that sedimented through the 27% sucrose cushion was resuspended in 0.25 M sucrose and subjected to a second differential centrifugation. This procedure resulted in a greater recovery of the platelet drug–antibody receptor as has been previously described (7). Protein concentration in platelet membrane suspensions was determined using fluorescence measurements after derivitization with o-phthalaldehyde (21). Membranes were washed and resuspended in PBS and stored at -70°C. Each batch of platelet membranes was evaluated using radioimmunoassay (RIA) for PI(II) antigen (22). Specifically, glycoprotein IIb was purified by differential centrifugal extraction (23) in gel filtration on Sephacryl S-300 (24) and radiolabeled using a chloramine-T method (25). The immunounassay consisted of mixing this radiolabeled antigen with anti-PI(II) at molar equivalence and precipitating the resulting antigen–antibody complexes with formalin-fixed S. aureus to obtain the maximum value for percent binding in Fig. 1. Curves were then obtained using varying amounts of either intact platelets or platelet membranes to compete with the labeled glycoprotein for anti-PI(II), as shown in Fig. 1. By comparing the number of intact platelets with the amount of purified platelet membranes required to inhibit binding by 50%, a specific amount of platelet membrane protein could be converted to an equivalence of whole platelets based upon PI(II) antigen content.

dd binding to platelet membranes. All fragments of each antibody were assayed using a single preparation of platelet membranes to minimize the variation in receptor concentration that may arise from comparisons among individual platelet preparations. Membrane fragments were washed once and diluted in PBS, 0.5% bovine serum albumin (BSA), and 1.0 mM EDTA. The amount of platelet membrane protein equivalent to 1.5 × 10^7 platelets (~10 μg of protein) was incubated with purified IgG or the purified IgG fragments, with or without the addition of appropriate drug (1.0 mM final concentration) in a total volume of 0.15 ml. This mixture was incubated for 30 min at room temperature, centrifuged at 12,000 g for 5 min in a Beckman model 12 microfuge, and the pellet was washed three times in saline with or without the appropriate drug. An acid eluate was prepared from the final washed pellet by re-suspension in 266 μl of PBS, 0.5% BSA, plus 133 μl of 10 mM phosphoric acid, to give a final pH of 3.1. This mixture was incubated at room temperature for 10 min and again centrifuged at 12,000 g for 10 min, after which 52 μl of 2.5 M Tris, pH 10.0, was added to a 332-μl aliquot of the supernatant to bring the pH to ~ 7.0.

A "sandwich" RIA was used to measure eluted IgG (26). Various dilutions of the eluates in 40-μl aliquots were added to 100 μl of RIA buffer (10 mM Tris HCl, 0.5% BSA, pH 8.3) and then loaded onto wells of polystyrene microtiter plates that had been precoated with anti-human IgG (affinity-purified goat anti-human IgG, heavy chain–specific; Kirkegaard & Perry, Gaithersburg, MD). After incubation for 2–18 h at room temperature, the plates were washed with RIA buffer, and a radioiodinated second antibody (anti-human IgG from the same lot as the material with which the plate was originally coated) diluted in RIA buffer to contain 50,000–100,000 cpm/100 μl was added to each well. This labeled antibody was allowed to bind for 2 h at room temperature. The plates were then washed three times, and individual wells were cut and counted in a gamma counter. The quantitative results were determined from standard curves on control IgG, Fab(1/2), Fab, or Fc that were run with each determination. A typical set of standard curves is shown in Fig. 2. The raw data was analyzed by a modified logit transformation, where each curve was subjected to linear regression analysis. Standard curves spanning the range of 0.05–125 ng of IgG or its fragments were used to quantitate the total acid eluted immunoreactive material.

Quantitative complement fixation. Briefly, the dd-IgG was incubated with 8 × 10^7 platelets in the presence of 10 U of guinea pig complement with or without appropriate drug. After 30 min of incubation at 37°C the reaction was stopped, an aliquot of the suspension removed, and the complement activity remaining in the aliquot was quantitated using an indicator system of hemolsyn-sensitized sheep red blood cells. The assay procedure was identical to that described previously (27) for direct determination of IgG binding as well as for evaluation of competitive binding of non-complement-fixing IgG allo-antibodies in the presence of complement-fixing antibodies of the same specificity.

When competitive experiments were performed, the order of addition of reagents to the reaction mixture was as follows: veronal-buffered saline; platelets; IgG; IgG fragments [Fab(1/2), Fab, or Fc]; guinea pig complement; and drug.

Preparation of purified dd-IgG. A 50-ml sample of whole serum containing a cross-reacting (quinidine and quinine) dd-IgG was incubated with 96 mg of purified platelet membrane protein, which was equivalent to ~1.3 × 10^11 platelets, for 18 h at room temperature in 1.0 mM quinidine. The supernatant (adsorbed serum) of this incubation contained ~5% of the initial concentration of antibody. The membranes were

Figure 1. Standardization of purified platelet membranes. As described in Methods, purified platelet membranes and the intact platelets from which they were prepared were compared with respect to their ability to displace [125I]-labeled GPIIIa from anti-PI(II). The number of platelets or the amount of membrane protein capable of producing 50% competitive inhibition was taken as the equivalence point, allowing calibration of the platelet membrane preparation. 10 μg of purified platelet membrane equivalent to ~1.3 × 10^7 platelets.
commercial molecular weight standards (Bio-Rad Laboratories, Richmond, CA) were included on the gel for reference purposes.

Results

Binding of dd-IgG and dd fragments to platelet membranes. Fig. 3 shows the dose-dependent response of dd-IgG binding to washed platelet membranes using purified total serum IgG from three different patients, one having quinidine-dependent purpura, a second having quinidine-dependent purpura, and a third who showed sensitivity to both drugs. The amounts of purified IgG available limited evaluation to the values shown, which suggests, but may not assure, saturation in each case.

When IgG and its fragments were incubated with platelet membranes, binding occurred with IgG, Fab, and F(ab')2 only in the presence of the drug. Using equimolar amounts of IgG and its fragments, the amounts of each preparation that were eluted after binding to a standard sample of platelet membrane are shown in Fig. 4. IgG showed the maximum, F(ab')2 intermediate, and Fab the least binding. In contrast, binding of the Fc fragment was not dd, since the same amount bound whether or not the appropriate drug was present, as shown in Fig. 4 b.

Competitive inhibition of dd-125I-labeled IgG binding to platelet membranes. To determine whether F(ab')2, Fab, or Fc fragments prepared from dd-IgG bound to the same membrane site as dd-IgG, and to confirm the direct binding measurements, each fragment was tested as a competitor for radiolabeled purified dd-IgG using purified platelet membranes as a solid-phase re-

Figure 2. Standard curves for quantitation of IgG and IgG fragments in the solid-phase RIA. Known amounts of IgG (solid circles), F(ab')2 (triangles), Fab (squares), and Fc (open circles) fragments were added to immobilized affinity-purified polyclonal anti-human IgG coated on microtiter plates and then exposed to radiolabeled anti-human IgG. Values for the fragments are normalized and compared with the maximum binding of IgG under identical assay conditions.

washed three times in PBS containing 0.5% BSA and 1.0 mM quinine. The washed membranes were resuspended in 2.0 ml PBS, 0.5% BSA. The pH was then adjusted to 2.8 with dilute phosphoric acid, and the membranes, after incubation for 10 min in these acid conditions, were removed by centrifugation. The pH of the supernatant was immediately adjusted to 7.0 with dilute NaOH. This acid eluate contained 1.4 × 10^{-4} mol of IgG by RIA, of which 95% was dd. This percentage was determined by quantitating the maximum amount of IgG bound when an aliquot of eluate was exposed to an excess of platelet membrane in the presence of 1.0 mM quinidine.

For further purification, the eluate was then applied to a protein A-Sepharose column equilibrated in PBS, and the adsorbed dd-IgG was eluted in 1.0 M acetic acid, pH 2.8, pooled, and neutralized. Aliquots of 10 μg dd-IgG were radioiodinated with 125I using chloramine-T (25).

Acid elution and electrophoresis of platelet-bound radiolabeled F(ab')2. F(ab')2 was prepared by pepsin digestion of IgG separated from serum of patient III and labeled with 125I by the chloramine-T method (25). 1.5 × 10^7 platelets were mixed with sufficient 125I-F(ab')2 to saturate binding sites, assuming F(ab')2 attached to platelets in the same quantities as dd-IgG. Quinidine glucuronate was added to achieve a final drug concentration of 1 mM. Drug was omitted in the control mixture. After incubation at room temperature for 24 h, platelets were pelleted and supernatants removed. Pellets were washed twice in 0.85% saline with or without 1 mM drug, as in the original mixtures, and final pellets were resuspended in 0.375 cm^{-3} 0.85% saline. The pH was adjusted to 2.8 with 0.1 N HCl, and after 10 min platelets were pelleted and supernatants recovered and neutralized to pH 7 with 0.01 N NaOH. Dilutions of the radiolabeled F(ab')2 starting material and of the different elutes were electrophoresed, nonreduced, on a 7% SDS–polyacrylamide gel, which was subsequently stained with Coomassie Brilliant Blue, soaked in autoradiograph (National Diagnostics, Inc., Somerville, NJ), dried, and autoradiographed. Several nonradioactive IgG fragment preparations and

Figure 3. dd binding of purified IgG preparations to platelet membranes. Aliquots of purified IgG from serum of three patients having quinine or quinidine purpura were incubated with platelet membranes in the presence or absence of the appropriate drug. The IgG bound was determined by RIA of an acid elution obtained after a 1-h incubation. Values for each point are the average of triplicates from which background obtained with no drug, or the inappropriate drug was subtracted. Among triplicates the maximum deviation was 10% of the mean. In the absence of added IgG, the levels of platelet membrane-bound IgG were ~0.01 pmol per 10 μg of platelet membrane protein. This equals 0.1 fg per platelet equivalent, which is 10- to 20-fold lower than the amount of IgG associated with normal, washed, intact platelets assayed by the same technique. With the addition of IgG and no drug or the inappropriate drug, the value of adsorbed IgG was always <10% of that when the appropriate drug was present. IgG from patient I (open squares) was used with 1.0 mM quinine, from patient II (solid squares) with 1.0 mM quinidine, and from patient III (solid circles) with 1.0 mM quinine. (open circles) 1.0 mM quinine.
ceptron. As shown in Fig. 5, Fab and F(ab')2 fragments and IgG competitively inhibited binding of 125I-dd-IgG. The difference between Fab and the F(ab')2 with respect to their effectiveness in displacing 125I-IgG was consistent with affinity differences expected for monovalent and divalent binding (28). The Fc fragment caused no binding inhibition. Commercially obtained control Fab, F(ab')2, and intact IgG (Kirkegaard & Perry) also did not significantly inhibit binding of radiolabeled dd-IgG.

**Inhibition of dd-IgG-mediated complement fixation.** The ability of Fab and F(ab')2 fragments to interact with the dd-IgG binding sites was also observed using complement fixation as the measure of IgG binding. As shown in Fig. 6, both F(ab')2 and Fab significantly inhibited complement fixation by the parent IgG. A summary of the relative efficiency of inhibition of these IgGs by their respective fragments is shown in Table I. The molar excess of fragments required to produce 50% inhibition of maximum complement fixation by IgG, from which they were derived, is 1.6–3-fold for the F(ab')2 fragments and 44–75-fold for the Fab fragments.

**SDS–PAGE of eluted radiolabeled dd-F(ab')2.** Eluates of platelet membranes exposed to F(ab')2 in the presence or absence of drug were electrophoresed and autoradiographed as described in Methods.

As shown in Fig. 7, the eluate prepared with quinidine (Fig. 7, lanes 7 and 8) contained a high concentration of F(ab')2, while the eluate prepared without drug (Fig. 7, lane 9) had only barely detectable F(ab')2. The dd eluate also contained traces of material of ∼50,000 mol wt that contaminated the F(ab')2 preparation (Fig. 7, lanes 5 and 6). This contaminant is most likely Fab' fragment derived from the pepsin treatment of IgG that was not removed by the S. aureus protein A and anti–Fc columns.

**Discussion**

In an attempt to resolve the conflicting data concerning antibody domain involved in dd–antibody binding, our studies were done

**Table I. Inhibition of dd-IgG–mediated Complement Fixation by F(ab')2 and Fab Fragments**

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<th>dd-IgG</th>
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<tr>
<td></td>
<td>F(ab')2</td>
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<td>Molar ratio to IgG</td>
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<td>0.46</td>
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<tr>
<td>Patient II</td>
<td>0.42</td>
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<td>Patient III</td>
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Competitive complement fixation experiments were performed with dd-IgG and its F(ab')2 and Fab fragments from three different patients, as described in Methods, and as shown for patient II in Fig. 6. Molar quantities of reagents were calculated on the basis of the following molecular weights: IgG, 160,000; F(ab')2, 100,000; and Fab, 50,000.
using highly purified IgG, IgG fragments, and platelet membranes in four experimental systems. Direct quantitative binding of dd-F(ab')₂, dd-Fab, and dd-Fc fragments relative to intact dd-IgG was measured; two methods of determining inhibition of IgG binding by dd-IgG fragments were used; and membrane-bound ¹²⁵I-dd-F(ab')₂ was evaluated by qualitative SDS-PAGE. Results of these assays consistently demonstrated that only the Fab domain of the IgG is responsible for dd-IgG binding, and that the affinity of dd binding increases in the order of Fab, F(ab')₂, and IgG. Additionally, the observed differences in apparent affinity of the monomeric and dimeric fragments, as evaluated by differences in their ability to compete for dd-IgG binding strongly suggest that the Fab-mediated binding of dd-IgG to the platelet surface is divalent at the optimal drug concentrations used.

As shown in Fig. 5, an eightfold molar excess of F(ab')₂ over that of unlabeled dd-IgG was required to inhibit 50% of the binding of radiolabeled dd-IgG in direct binding experiments, and 50-fold M excess was required for 20% inhibition by Fab. In complement fixation experiments with the three antibodies evaluated in this study, 50% inhibition of dd binding by the parent IgG required a 1.6- to 3-fold molar excess of F(ab')₂ and a 44- to 75-fold excess of Fab. The molar ratio causing inhibition by complement fixation assays may have been lower than that observed by direct measurements because of the apparent requirement of nearest neighbor distribution of IgG to fix complement (2). This requirement may cause a disproportionate decrease in complement fixation relative to the number of molecules of IgG displaced. Fc fragments prepared from dd-IgG did not inhibit complement fixation by the parent dd-IgG even when present in a 100-fold molar excess.

Two possible explanations for the requirement of an excess molar concentration of dd-F(ab')₂ to compete with dd-IgG from which it was derived are the following: First, although the Fc portion of dd-IgG is not essential for high affinity binding, we cannot rule out the possibility that its presence contributes to the stability of the antibody (possibly the hinge region), and thereby may increase the stability of the antibody-drug-platelet complex. Also, minor structural changes in the F(ab')₂ and Fab fragments inherent in the enzymatic cleavage of parent IgG potentially could affect affinity of the overall reaction.

Although each binding curve in Fig. 3 seems to approach a different asymptotic value of saturation, this may be due to variations in apparent affinity between platelet receptor and the different dd-IgGs. Previous studies using whole serum and intact platelets from several different patients described similar results (4). Our results indicate that the number of dd-IgG molecules bound to the platelet at apparent saturation is 20,000–50,000, which is similar to the estimated 26,000–33,000 glycoprotein I₆b molecules on the platelet surface (29). A recent estimate of platelet receptor sites for dd-antibody binding was 36,000–161,000 per platelet (4). The range of values is higher than that obtained in the present study which may, in part, be related to our use of purified platelet membranes rather than whole platelets; i.e., when compared on the basis of protein content, purified membranes may contain only 77% the number of dd-antibody receptor sites as intact platelets. If the value for bound dd-IgG molecules exceeds GPIb complexes, this would imply that each complex may bind more than one dd antibody or that additional binding sites may exist.

It is not yet known which bimolecular reaction, i.e., that between antibody and drug, or cell and drug, occurs as an initial step or whether a quasi-trimolecular reaction takes place. Quantitation of the binding affinity and stoichiometry of the possible bimolecular reactions is difficult because of the above-cited indication that these reactions are of low relative affinity. Preliminary data on such binding in an erythrocyte dd-antibody system indicates the types of approaches that will be necessary to define the dd-antibody system involving platelets (30).

References


