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Complement-independent Ab-induced peroxide lysis of platelets requires 12-lipoxygenase and a platelet NADPH oxidase pathway

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AntiplATELET GPIIIa49–66 AB OF HIV-RELATED THROMBOCYTOPENIC PATIENTS INDUCES THROMBOCYTOPENIA AND PLATELET FRACTIONATION BY THE GENERATION OF Peroxide AND OTHER REACTIVE OXYGEN SPECIES (ROS). Here we report the presence of a functional platelet NADPH oxidase pathway that requires activation by the platelet 12-lipoxygenase (12-LO) pathway to fragment platelets. A new Ab-mediated mechanism is described in which the platelet 12-LO product, 12(S)-HETE activates the NADPH oxidase pathway to generate ROS.

Introduction

Patients with HIV-1 infection commonly develop immunologic thrombocytopenia (HIV-1-ITP) (1). Platelet survival studies indicate that early-onset HIV-1-ITP is often due to increased peripheral destruction of platelets, whereas patients with AIDS are more likely to develop decreased platelet production (2). Early-onset HIV-1-ITP is clinically indistinguishable from classic idiopathic autoimmune thrombocytopenia (AITP) and responsive to the same therapeutic agents used to treat classic AITP (1, 3–6). HIV-1-ITP is different from classic AITP, however, with respect to markedly elevated platelet-associated IgG, IgM, C3, C4, and the presence of circulating immune complexes (CICs) composed of the same (3, 4).

Past studies have revealed that these complexes also contain antiplatelet integrin GPIIIα (β3) Ab and its anti-idiotypic blocking Ab (8). Affinity purification of the Ab and epitope mapping of GPIIIα have revealed a high-affinity IgG1 Ab against GPIIIα residues 49–66 (7, 9). The presence of this Ab correlates inversely with platelet count (r = −0.71) and induces severe thrombocytopenia when introduced into mice (9) (mouse GPIIIα has 83% homology with human GPIIIα and macrophages with Fc receptors for human IgG1). Passively induced murine thrombocytopenia (9) as well as patient anti-idiotypic blocking Ab (8).

Circulating immune complexes from early-onset HIV-1-ITP patients contain platelet fragments (10). We have also demonstrated that affinity-purified anti-GPIIIα49–66, in the absence of complement, can induce platelet fragmentation in vitro and in vivo (10). This intriguing observation led to the discovery that Ab-induced platelet fragmentation was due to the generation of peroxide and other reactive oxygen species (ROS). The generation of ROS appeared to be due to the presence of an active NADPH oxidase pathway in platelets, since in vivo platelet fragmentation did not occur in mice deficient in the p47 phagocytic oxidase (p47phox) component of the NADPH (p47phox−/−) pathway and in vivo thrombocytopenia was approximately 60% less than that obtained with WT mice (10). The mechanism leading to this activation of ROS, however, was not defined nor was it clear whether the ROS activity was derived from the platelet or some other source.

The NADPH oxidase of granulocytes/macrophages is composed of five major components that coalesce onto the cell or vacuolar membrane to form an active electron donor in which superoxide, O2·−, is generated (11). Three cytoplasmic phox components, p47phox, p67phox, and p40phox, translocate to the cytoplasmic surface of the membrane (12–16) in independent association with activated Rac G protein. Rac binds to p67phox (17), and then binds to two membrane-localized components, gp91phox and p22phox, the α and β subunits of the cytochrome b complex (18, 19). This complex can bind NADPH and flavin adenine dinucleotide (FAD) (11, 20, 21). Several enzymes are required for the system to be activated after membrane perturbation by way of various physiologic ligands (fMLP, C5a, PAF, leukotriene B4, PAF, leukotriene B4, IL-8) that bind to pertussis toxin–sensitive (PTX-sensitive) G protein–coupled receptors (22). Activation is dependent upon lipid mediators such as phosphatidic and arachidonic acids and phosphatidylinositol (23–25). The enzymes involved include PI3K, which products form a scaffold for membrane attachment of p40phox and p47phox (26–28); protein kinase C, which phosphorylates p47phox, enabling its translocation to the membrane (29–31); cytosolic phospholipase A2 (cPLA2), which generates arachidonic acid from membrane phospholipids (32), serving to activate the association of p47phox with p22phox (23). Extracellular signal-regulated kinase (ERK) and p38...
mitogen–activated protein kinase (p38 MAPK) are required for the phosphorylation and activation of cPLA₂ (33–36). In both phagocytic and nonphagocytic cells, cPLA₂ participates in the generation of leukotriene B₄ (LTB₄), which appears to be required for ROS generation and chemotaxis (37). The relationship between LTB₄ and NADPH oxidase is unknown.

Here we (a) demonstrate that platelets have a functional, NADPH oxidase pathway; (b) indicate that the pathway is activated by membrane perturbation of GPIIIa₄₉–₆₆; and (c) describe a new ROS pathway in which 12(S)-HETE, a product of platelet 12-LOxygenase (12-LO), activates NADPH oxidase following Ab ligation of GPIIIa₄₉–₆₆. Thus, complement-independent Ab-induced platelet fragmentation through NADPH oxidase generation of ROS requires activation of platelet 12-LO.

Results

Absence of granulocytes in gel-filtered platelet preparations. Figure 1A demonstrates the absence of granulocytes/monocytes in a human gel-filtered platelet preparation as examined by flow cytometry with a granulocyte/macrophage–specific anti–CD11b Ab. It could be shown that greater than 99% of the platelets were free of CD11b reactivity. This was confirmed by a phase microscopy analysis of 2,000 platelets, in which no leukocytes were visualized. Thus, contamination of platelets with granulocytes/monocytes or cross talk between both cells was ruled out.

Presence of active NADPH oxidase in murine platelets and gp91phox in human platelets. Although our previous studies have indicated that ROS are necessary for Ab-mediated platelet fragmentation, the source of the ROS was not defined. In our previous report we demonstrated absence/impairment of anti–GPIIIa₄₉–₆₆–induced platelet fragmentation/thrombocytopenia in vivo with murine NADPH oxidase–deficient p47phox–/– mice, but did not examine the effect of Ab on platelet fragmentation in vitro. Therefore, a systemic effect or granulocyte/macrophage effect was not ruled out. We therefore incubated a different NADPH oxidase–deficient platelet preparation from gp91phox–/– mice with antiplatelet Ab. The gp91phox–/– platelets did not undergo platelet fragmentation in vitro when exposed to the anti–GPIIIa₄₉–₆₆ Ab (percentage fragmentation in control, 6.4% ± 0.06%, versus patient IgG, 5.9% ± 0.06%), whereas WT mouse platelets did fragment (percentage fragmentation in control, 5.7% ± 0.06%, versus patient IgG, 31.8% ± 5.4%), ruling out a systemic effect.

The presence of membrane gp91phox was also demonstrated in human gel-filtered platelets by flow cytometry with a specific mAb against gp91phox (Figure 1B). Platelets had significant reactivity that was similar to that seen in granulocytes.

Classical inhibitors of the NADPH oxidase pathway inhibit Ab-induced platelet fragmentation in vitro. To determine whether platelets had a functioning NADPH oxidase pathway responsive to membrane perturbation, we employed classical inhibitors of the system to determine whether or not they would inhibit Ab-induced platelet fragmentation. PTX, an inhibitor of G protein–coupled receptors, inhibited in vitro platelet fragmentation at 5 ng/ml (28% fragmentation in untreated versus 5.5% after PTX and 6% after control treatment, representative of two similar experiments). Similar results were obtained with other inhibitors (Figure 2) known to interfere with this pathway. IC₅₀ values for inhibition of several key enzymes were estimated to be PI3 kinase with LY294002 at 10 µM, protein kinase C with bisindolylmaleimide at 20 nM, p38 MAPK with SB203290 at 5 µM, and PLA₂ with bromophenacyl bromide at 50 µM.

Figure 1

platelets were exposed to varying concentrations of 12(S)-HPETE and 12(S)-HETE, its stable breakdown product, and the level of platelet fragmentation was assessed. Both products were as effective as anti–GPIIIa49–66 Ab in inducing fragmentation (Figure 4B). In contrast, the nonbiologically derived stereoisomer, 12(R)-HETE, was without effect in this assay (Figure 4B), as was the granulocyte 5-LO product, LTB4 (data not shown), indicating specificity. Second, confirming the connection between 12-LO and the NADPH oxidase system, we determined that, similar to Ab from HIV-1-ITP patients, 12(S)-HETE (200 nM) alone was sufficient to induce the translocation of p47phox and p67phox from cytosol to membrane fractions (Figure 3D). As predicted from Figure 2, 12(S)-HETE–induced translocation was also inhibited by the protein kinase C inhibitor, bisindolylmaleimide (Figure 3D). Furthermore, the time courses for platelet fragmentation induced by either patient anti–GPIIIa49–66 or by 12(S)-HETE were nearly identical (Figure 4C). Finally, confirming the central role of 12-LO in Ab-induced platelet fragmentation, we quantified the generation of 12(S)-HETE by platelets after exposure to Ab. These platelets generate substantial levels of the biologically active lipid over a time course that is consistent with the induction of fragmentation by Ab (Figure 4D).

Platelet fragmentation with 12(S)-HETE requires interaction of platelet 12-LO with the platelet NADPH oxidase pathway. If 12(S)-HETE induces platelet fragmentation by ROS generation, it follows that interventions that inhibit ROS action should also block the 12(S)-HETE activity. Thus, we tested the ability of 12(S)-HETE (100 nM) to fragment platelets in the presence of increasing concentrations of catalase and diphenylene iodonium (DPI). Both effectively blocked 12(S)-HETE–induced platelet fragmentation (Figure 5A) as they do Ab-induced platelet fragmentation (10). It also follows that 12(S)-HETE should induce the generation of measurable ROS, and, thus, we tested the ability of 12(S)-HETE treatment to induce platelet-dependent oxidation of the indicator dye, dichlorodihydrofluorescein (DCFH). 12(S)-HETE caused platelets to oxidize DCFH in a dose-dependent manner (Figure 5B). The concentration of H2O2 equivalents generated by platelets treated with 12(S)-HETE, as determined by DCFH oxidation, was sufficient to induce platelet fragmentation when oxidation was induced with equivalent amounts of H2O2 (17 nM, n = 2). Similar observations were noted with Ab (data not shown).

The role of platelet 12-LO, Rac2, and NADPH oxidase in Ab-induced platelet fragmentation was further demonstrated by the absence of Ab-induced thrombocytopenia in vivo in 12-LO+/–, gp91phox–/–, and Rac2–/– mice, as compared with WT mice (Figure 6A). Similar results were found in vitro where 12(S)-HETE was unable to induce platelet fragmentation in either gp91phox–/– or Rac2–/– mice (Figure 6B). Nevertheless, platelets from 12-LO–/– mice responded normally to 12(S)-HETE (Figure 7A), despite the absence of Ab-induced platelet fragmentation. Ab-induced platelet

**Figure 2**

Inhibition of Ab-induced platelet fragmentation with classic inhibitors of the NADPH oxidase pathway. Gel-filtered human platelets were preincubated with inhibitors at various concentrations for 15 minutes prior to the addition of 25 µg/ml affinity-purified anti–GPIIIa49–66 (AP) or control (C) IgG for 4 hours. Ci, control IgG plus inhibitor; BPB, boprophenacyl bromide. Platelet fragmentation was monitored by flow cytometry of platelets labeled with anti–CD41-FITC Ab. Platelet fragments were monitored by enumeration of fluorescent particles in gates established to detect smaller particles. A, n = 4; B, n = 4; C, n = 3; D, n = 6.

**Figure 3**

Functional evidence for NADPH oxidase activity. Gel-filtered human platelets were incubated with 25 µg/ml of control or patient Ab at 37°C, extracted, and immunoblotted. (A) Effect of Ab on Akt activation. C5, C15, C30, and P5, P15, and P30 refer to minutes of incubation with control or patient (P) Ab, respectively. Akt-P, phosphorylated Akt. (B) Effect of patient Ab on ERK-1/2 activation. Upper lanes refer to reactivity with Ab directed against phosphorylated ERK (ERK-P). Lower lanes refer to Ab directed against internal control ERK protein (representative of five experiments). (C) Effect of Ab on translocation of p67phox, p47phox, and PLA2 from platelet cytosol to membrane. Platelets were incubated with Ab for 30 minutes at 37°C, extracted, separated into cytosol and membrane components, and then immunoblotted with specific Ab’s. M, marker for p67phox and p47phox. Ab, P15 refers to platelets in buffer, prior to incubation. Cc, Crm, and Pc, Pm refer to Ab directly against cytoplasmic and membrane fractions, respectively. (D) Effect of 12(S)-HETE (200 nM) on translocation of p67phox and p47phox from platelet cytosol to membrane fractions. Hm, Hc, Cm, and Cc refer to cytoplasmic and membrane fractions from 12(S)-HETE and control incubations, respectively. PHm and PHc refer to incubation with 12(S)-HETE and protein kinase C inhibitor, bisindolylmaleimide at 20 nM. Representative of three experiments.
fragmentation in 5-LO–/– mice was no different from WT mice (Figure 7B), indicating that the change in 12-LO–/– was not a nonspecific effect related to the gene deletion.

**Discussion**

These data indicate that platelets have a functioning NADPH oxidase pathway capable of destroying platelets by the generation of ROS, following perturbation of platelet GPIIIa with a specific anti–GPIIIa49–66 Ab. Activation of NADPH oxidase requires prior activation of platelet 12-LO. It is therefore apparent that the 12-LO and NADPH oxidase pathways are linked, with 12-LO being upstream of NADPH oxidase. This is the first description of a role for 12-LO in platelets.

Suggestive evidence for the presence of an NADPH oxidase in platelets has been reported by us (10), as well as others (38, 39). None of these reports present rigorous evidence for a functioning granulocyte-like NADPH oxidase system, however. Furthermore, the activation of the NADPH oxidase and subsequent platelet fragmentation induced by Ab requires generation of the 12-LO product, 12(S)-HETE. This article presents such evidence based upon the following observations: (a) gel-filtered platelets are free of contaminating granulocytes/monocytes; (b) platelets from two different NADPH oxidase–deficient mice, p47 phox–/– (10) and gp91 phox–/–, failed to undergo either Ab or 12(S)-HETE–induced platelet fragmentation, nor does the Ab induce thrombocytopenia in vivo; (c) human platelets contain membrane-bound gp91phox; (d) platelet p47phox, p67phox, and cPLA2 translocate from cytoplasm to membrane following Ab-induced fragmentation; (e) a diverse group of classic inhibitors of NADPH oxidase activation also inhibit Ab-induced platelet fragmentation in vitro; (f) Ab-induced platelet fragmentation is associated with the activation of PI3K and ERK.

Indirect evidence for ligand-activated NADPH oxidase activity as well as components of the NADPH oxidase complex have been reported in nonphagocytic cells (smooth muscle) (40, 41), chondrocytes (42), kidney epithelium (43), and fibroblasts (44). Treatment with DPI, an inhibitor of the FAD component of neutrophil NADPH oxidase, can inhibit ROS production in these nonphagocytic cells when stimulated by ligands (40, 42, 44). On the other hand, p40phox and p47phox are dispensable for NADPH oxidase activity in a cell-free system, whereas p67phox is required (45). Indeed there has been little rigorous evidence supporting the existence of functional p47phox and p67phox in nonphagocytic cells. A recent suggestion has been made that other systems may regulate p67phox or that nonphagocytic cells do not require p67phox for ROS generation (46).

The role of ROS in cell messenger signaling is an emerging concept. Although high concentrations of ROS are used in host defense, there is evidence that lower concentrations (1–2% of granulocyte generation) are required for cell signaling. There have been

**Figure 4**

Role of platelet 12-LO in anti–GPIIIa49–66–induced platelet fragmentation. (A) Lipooxygenase inhibitors: 5-, 8-, 11-eicosatriynoic acid (ETI), nordihydroguaiaretic acid (NDGA) (n = 4), and cinnamyl-3-, 4-dihydrocycanoamine (CDC) (n = 8) were added to gel-filtered human platelets 15 minutes prior to Ab, and percentage of fragmentation measurements were made after a 4-hour incubation. Cg, control IgG (25 µg/ml); Ci, control IgG plus inhibitor (25 µg/ml); 0, patient IgG without inhibitor (25 µg/ml). (B) Effect of 12-LO products, 12(S)-HETE and 12(S)-HPETE (n = 8), on platelet fragmentation. 12(R)-HETE is a nonlipoxygenase-derived stereoisomer (n = 3). (C) Comparison of Ab versus 12(S)-HETE–induced platelet fragmentation (typical of three experiments). (D) Effect of patient IgG (Pg) vs. control IgG (Cg) on platelet 12(S)-HETE production (typical of three experiments).

**Figure 5**

Effect of 12(S)-HETE on platelet oxidation. (A) Gel-filtered human platelets were preincubated with various concentrations of catalase and DPI for 15 minutes prior to the addition of 12(S)-HETE (100 nM) for 4 hours. Cb, control buffer; H, 12(S)-HETE with no drug. Increasing numbers refer to doubling concentrations of catalase, starting at 125 µg/ml, and DPI, starting at 10 µM. (B) Gel-filtered platelets were loaded with DCFH and incubated with 12(S)-HETE at various concentrations for 4 hours at 37°C. Oxidation was measured by DCFH fluorescence.
several recent reports demonstrating a transient oxygen burst in cells stimulated by various cytokines: IL-1β, TGF-β1 (47), TNF-α (42, 48); growth factors: EGF (49, 50), PDGF (51, 52), VEGF (53), βFGF (54), thrombin (55); and apoptotic signals (56–58). All induce ROS, which is required for tyrosine kinase receptor phosphorylation, cell stimulation, and mitogenesis, since ROS scavengers (and/or antioxidants) inhibit these reactions.

Of particular interest is the observation that the nonphagocytic cell cytokine and growth factor ROS pathways also require arachidonic acid release by PLA2. This requires the Rac-1 G-coupled protein (49), as well as activation of PI3K, ERK, and p38 MAPK, as is the case with the granulocyte NADPH oxidase system. ERK and p38 MAPK are involved in arachidonic acid release induced by H2O2 and PDGF in mesangial cells (34). ERK is known to increase cPLA2 activity following cPLA2 phosphorylation by ERK. The p38 MAPK also phosphorylates and activates cPLA2 in thrombin or collagen-activated platelets (59, 60). It has been reported in mesangial cells that both PDGF as well as H2O2 induce the phosphorylation/activation of ERK (33, 34, 52). In human platelets, H2O2-induced phosphorylation/activation of cPLA2 with arachidonic acid release requires p38 MAPK (61).

The ROS burst of the cytokine/growth factor–induced pathways in nonphagocytic cells can be blocked by inhibitors of the lipooxygenase pathway. It is unlikely that lipooxygenase metabolism per se results in ROS generation (62), since arachidonic acid oxidation is independent of ROS production. It is more likely that 12-LO generates a downstream product that activates the NADPH oxidase pathway. In phagocytic cells, the pathway inhibited by lipooxygenase inhibitors can also be inhibited with DPI (antiflavoprotein oxidase), activated with LTB4, and blocked with antagonists of its receptor as well as by PTX (37). Yet, nonphagocytic cells do appear to express this NADPH oxidase pathway (40, 41, 43, 44). Thus, the presence, as well as relative roles of the various pathways, is unclear, and downstream events from receptor binding to induction of ROS are unknown.

Our current data provide an explanation for these observations by demonstrating that both the platelet 12-LO and NADPH oxidase pathways are interconnected with the platelet 12-LO pathway being upstream of the NADPH oxidase pathway. This is supported by our observations that the 12-LO product, 12(S)-HETE, induces platelet fragmentation even in platelets from 12-LO–deficient animals. That this is a specific pathway related to the generation of 12(S)-HETE is emphasized by the fact that Ab-induced platelet fragmentation is normal in platelets from 5-LO–deficient mice and by the observed inability of 12(R)-HETE, a synthetic stereoisomer of the biologically active lipid, to induce platelet fragmentation. It is likely that both the 12-LO and NADPH ROS-inducing pathways are involved in cell signaling, as with other nonphagocytic cells. In platelets, however, it can be pathophysiologically subverted to cell fragmentation and destruction following anti–GPIIIa49–66 ligation.

The physiological importance of the 12-LO and NADPH oxidase pathways in platelets has not yet been firmly established. Although platelets can engulf bacteria (63), there is no evidence that they can phagocytose and kill them. Platelets can generate ROS, particularly following activation with collagen, but not with ADP or thrombin; collagen-induced aggregation can be inhibited with catalase (64). It is suggested that Ab-induced ROS may have possible other effects on nonplatelet bystander cells such as megakaryocytes or endothelial cells, which could be contributing to the pathophysiology of the disease. Our data raise the possibility that 12-LO inhibition may be beneficial in reversing the thrombocytopenia in patients with AI TP. Indeed, preliminary data reveal that this is so, since the platelet 12-LO inhibitor CDC can significantly inhibit in vivo platelet particle formation and thrombocytopenia in mice. Unfortunately, no drugs are currently available for use in humans, so further testing of this hypothesis will have to wait for the development of effective inhibitors.

Methods

Human population. Patient sera were obtained from early-onset HIV-1–infected thrombocytopenic patients without AIDS and control subjects (healthy laboratory personnel). These studies

Figure 6

Effect of anti–GPIIIa49–66 and 12(S)-HETE on murine platelet count and fragmentation of WT, NADPH oxidase gp91phox–/–, Rac-2–/–, and 12-LO–/– platelets. (A) Platelet count. Mice were treated with 25 µg of control or patient anti–GPIIIa49–66, and platelet counts were monitored at various time intervals (n = 5 for each group). (B) Fragmentation. Percentage of platelet particles were enumerated after in vitro exposure to anti–GPIIIa49–66 (25 µg/ml) (Ga), control IgG (Cg), or 12(S)-HETE (100 nM) (H) for 4 hours at 37°C.

Figure 7

Effect of anti–GPIIIa49–66 and 12(S)-HETE on murine platelet fragmentation of (A) 12-LO–/– and (B) 5-LO–/– mice. Gel-filtered murine platelets were treated with control Ab (Ca), anti–GPIIIa49–66 (Ga), 12(S)-HETE, or its stereoisomer, 12(R)-HETE (H). Note absence of anti–GPIIIa49–66–induced platelet fragmentation with 12-LO–/– mouse platelets and its presence with 5-LO–/– mouse platelets. Note presence of 12(S)-HETE–induced platelet fragmentation with WT, 12-LO–/–, and 5-LO–/– platelets. n = 3.
were approved by the New York University Medical Center Institutional Review Board.

Mice. WT mice were C57BL/6. All KO mice were of this background. Mice (p91 phox−/−, p47 phox−/−, 12-LO−/−, and 5-LO−/−) were obtained from The Jackson Laboratory (Bar Harbor, Maine, USA). Rac-2−/− mice were kindly provided by David A. Williams (Indiana University School of Medicine, Indianapolis, Indiana, USA). Animal work was approved by the New York University School of Medicine Institutional Review Board.

Materials. All reagents were obtained from Sigma-Aldrich (St. Louis, Missouri, USA) unless otherwise designated. These reagents include inhibitors of G-coupled protein receptors, PTX; P13K, LY294002; protein kinase C, bisindolylmaleimide; phospholipase A2, bromophenacyl bromide; ROS, catalase and DPI. The p38 MAPK inhibitor, SB203290, was obtained from Calbiochem-Novabiochem Corp. (San Diego, California, USA). Lipoxigenase inhibitors, ETI, NDGA, and CDC, and eicosanoids, 12(S)-HETE, 12(S)-HPETE, and 12(R)-HETE, were obtained from Biosmol Research Laboratories (Plymouth Meeting, Pennsylvania, USA). DCFH diacetate (DCFH-DA) was obtained from Molecular Probes Inc. (Eugene, Oregon, USA). Goat polyclonal Ab’s against p47 phox and p67 phox were obtained from Santa Cruz Biotechnology Inc., Santa Cruz, California, USA. Mouse mAb against human or anti–CD41b-FITC for murine GPIIb was a gift of H.L. Malech (NIH, Bethesda, Maryland, USA).

Affinity purification of antiplatelet GPIIIa49–66. IgG was isolated by polyethylene glycol–precipitable immunocomplexes (8), followed by protein A affinity chromatography. G-200 gel filtration, affinity chromatography and affinity purification on 2% paraformaldehyde-fixed platelets as described (7). The antiplatelet Ab was eluted with 0.1 M glycine buffer, pH 2.9, and further purified on a GPIIIa49–66 peptide-affinity column, as described (7).

Preparation of leukocyte/platelet–enriched suspension. Three milliliters of heparinized blood were added to 9 ml of 1.25% dextran in isotonic saline, and red blood cells were allowed to settle to the bottom at room temperature for 90 minutes. The supernatant was first treated with 1 vol of distilled water for 60 seconds to remove contaminating red blood cells, followed by 4 vol of 4.25% NaCl to achieve isotonicity. The supernatant was then centrifuged and the cell pellet was washed and suspended in 1.5 volumes of Tyrode’s buffer.

Induction of platelet particles. Gel-filtered human platelets were prepared from blood collected in 5.2 mM EDTA. Murine blood was collected by cardiac puncture into heparin (1 unit/ml, final concentration). Gel-filtered platelets were prepared from platelet-rich plasma applied to a Sepharose 2B column preincubated with Tyrode’s buffer (pH 7.4). Platelets (106/ml) were either unlabeled or labeled with 10 µg/ml of an anti–CD61-FITC mAb or an anti–CD41b-FITC for murine GPIIb (PharMingen, San Diego, California, USA) for 30 minutes at 4°C. The cells were then centrifuged at 1,000 g for 6 minutes at room temperature and resuspended in Tyrode’s buffer. Ten microliters of platelet suspension (105 platelets) were then incubated with 10 µl of anti–GPIIIa49–66 Ab (250 µg/ml) and 80 µl of Tyrode’s buffer for 0–4 hours at 37°C and then stored in an ice bucket prior to measurements of the percentage of platelet particles by fluorescent flow cytometry.

Assay of platelet particle formation. Both fluorescent-labeled and unlabeled platelet particles were measured by flow cytometry, employing a FACScan (Becton Dickinson Immunocytometry Systems, Mountain View, California, USA). Similar results were obtained using both labeled and unlabeled platelets. Debris and dead cells were excluded using scatter gates. Only cells with low orthogonal light scattering were included in the sorting gates. Gates were adjusted for control platelets by exclusion of other blood cells. Fluorescent-labeled intact platelets were monitored in the right upper quadrant with the y axis measuring forward scatter and the x axis measuring fluorescence. For nonfluorescent particles, the y axis measured forward scatter, and the x axis measured side scatter. A shift in fluorescent or nonfluorescent particles from right upper quadrant to left upper quadrant and left lower quadrant reflected the percentage of platelet particle induction of 10,000 counted platelets/particles.

Assay of platelet oxidation. Gel-filtered platelets were loaded with 10 µM DCFH-DA for 30 minutes at 37°C as described previously (10) and challenged with platelet Ab. Oxidation was quantified by measuring the increase in mean fluorescence with a flow cytometer.

Assay of P13K and ERK-1/2 activity. Platelets (5 × 107) were incubated in 100 µl of buffer containing control or patient IgG for 4 hours and then extracted in an equal volume of ice-cold platelet lysate buffer containing 50 mM Tris, 5 mM EDTA, 1% Triton X-100, 0.1% sodium cacodylate, 150 mM NaCl, 1 mM PMSF, 1 mM Na2VO3, 1 mg/ml aprotinin, and 10 µg/ml leupeptin. Aliquots were then tested by immunoblot analysis.

P13K phosphorylation was detected by phosphorylation of its substrate, AKT, by Western blot analysis using an affinity-purified polyclonal rabbit anti-p8475AKT Ab, followed by a donkey anti-rabbit IgG-HRP (Promega Corp., Madison, Wisconsin, USA), and developed by chemiluminescence with substrate provided by Pierce Chemical Co. (Rockford, Illinois, USA). Actin was run as an internal control with a goat polyclonal IgG (c-11; Santa Cruz Biotechnology Inc.), followed by a monoclonal anti–goat IgG-HRP (Sigma-Aldrich).

ERK-1/2 phosphorylation was detected with a mouse monoclonal IgG2a against pERK (E-4), followed by a goat anti-mouse IgG2a-HRP (Santa Cruz Biotechnology Inc.), and developed by chemiluminescence. Nonphosphorylated ERK was run as an internal standard with a rabbit polyclonal IgG against ERK-1 (K-23; Santa Cruz Biotechnology Inc.), followed by protein A-peroxidase (Sigma-Aldrich), and developed with chemiluminescence.

Translocation of p67 phox, p47 phox, and cPLA2 to platelet membrane. Platelets (5 × 108) in 100 µl of buffer containing control or patient IgG were incubated for 4 hours at 37°C and extracted in relaxation buffer, with membrane and cytosol fractions prepared as described (36), with minor modifications. Cells were suspended in 50 mM HEPEs (instead of 10 mM PIPES) with protease inhibitors (1 mM PMSF, 5 mM EDTA, 10 µg/ml trasyol, and 10 µg/ml leupeptin or 1 tablet/50 ml of protease inhibitor cocktail, 1697498; Roche Diagnostics, Indianapolis, Indiana, USA) and frozen and thawed three times (instead of sonication). Insoluble cellular elements were removed by centrifugation at 30,000 g for 1 minute at 4°C, and the supernatant was adjusted to 5 mM EDTA, 1 mM Na2VO3, and 5 mM NaF, as described. The supernatant was then centrifuged at 30,000 g for 1 hour and the membrane pellet was resuspended in the original suspension volume of 0.34 M sucrose/half-strength relaxation buffer containing 1 mM DTT. Cytosol and membranes were stored at −70°C. The p47 phox, p67 phox, and cPLA2 were then tested by immunoblot analysis with a specific Ab, followed by an appro-
Assay of platelet 12-HETE production. Gel-purified platelets (10^9/100 μl) were incubated with control or patient Ab as described above. The interaction was stopped by addition of 1 vol of ice-cold acetone (giving a total volume of 200 μl) and the samples kept at −70 °C until extracted. Lipids were extracted into ether and analyzed by reverse-phase HPLC as previously described (65). The 12-HETE was quantified by UV absorbance at 237 nm.

Determination of mouse platelet counts. Twenty microliters of blood was drawn from the orbital sinus into unopettes (5854; Becton Dickinson Immunocytometry Systems) with optimum anticoagulant concentration and diluted for platelet enumeration by phase-contrast microscopy.