Role of molecular mimicry of hepatitis C virus protein with platelet GPIIIa in hepatitis C–related immunologic thrombocytopenia

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Patients with HIV-1 immune-related thrombocytopenia (HIV-1–ITP) have a unique Ab against platelet GPIIIa49-66 capable of inducing oxidative platelet fragmentation in the absence of complement. HIV–1-seropositive drug abusers are more prone to develop immune thrombocytopenia than non–drug abusers and have a higher coinfection with hepatitis C virus (HCV) than non–drug abusers (90% vs 30%). Molecular mimicry was sought by screening a phage peptide library with anti–GPIIIa49-66 antibody as bait for peptides sharing homology sequences with HCV. Several phage peptide clones had 70% homology with HCV protein. Sera from dually infected thrombocytopenic patients with HCV and HIV-ITP reacted strongly with 4 nonconserved peptides from HCV core envelope 1. Reactivity correlated inversely with platelet count (r² = 0.7, P < .01). Ab raised against peptide PHC09 in GPIIIa−/− mice induced thrombocytopenia in wild-type mice. Affinity-purified IgG against PHC09 induced oxidative platelet fragmentation in vitro. Drug abusers dually infected with HCV and HIV-1 had a greater incidence and severity of thrombocytopenia as well as titer of anti–GPIIIa49-66/PHC09 Ab.

In the present report, we demonstrate the following: (1) four HCV core-envelope peptides from a nonconservative region display molecular mimicry with GPIIIa49-66 by reactivity with anti-GPIIIa49-66 Ab. (2) The strongly reactive SAIHNRNSG peptide (PHC09) was examined more extensively. PHC09 injected into NZB/W F1 mice induced an Ab capable of inducing oxidative platelet fragmentation—induced by molecular mimicry with an HCV peptide in addition to HIV nef peptide in HIV-1-ITP.

Introduction

Thrombocytopenic patients with early HIV-1 infection have a shortened platelet survival due to an autoantibody against an epitope on platelet surface integrin GPIIIα, GPIIIa49-66 (CAPESIEFPVSEARVLED).1-3 Their sera have increased immune complexes that contain platelet fragments as well as anti–GPIIIa49-66 Ab. The presence of anti–GPIIIa49-66 Ab correlates inversely with platelet count (r = −0.71) and induces severe thrombocytopenia when injected into mice. This antibody is unique in that it induces complement-independent platelet fragmentation by oxidative platelet fragmentation due to the release of reactive oxygen species through activation of 12-lipoxygenase and NADPH oxidase.4,6

HIV-1 immune-related thrombocytopenia (HIV-1–ITP) is more frequent in drug abusers compared with non–drug abusers (37% vs 16% incidence, respectively), and more severe in HIV–1-seropositive drug abusers than non–drug abusers (platelet count < 10 × 10^9/L in 52% vs 9%, respectively).7,8 A striking feature of HIV-1 infection in drug abusers is the frequent coinfection with hepatitis C virus (HCV).9-13 The overall prevalence of HCV infection among HIV–1–infected individuals is 30% to 50%9 in non–drug abusers, with rates of coinfection as high as 90% in intravenous drug abusers.9,13 We asked whether coinfection with HCV facilitates ITP and, if so, what the mechanism would be. The presence of a relatively high-affinity immunodominant Ab against GPIIIa49-66 in HIV-1–ITP patients suggested antigen-driven B-cell clonal expansion. We therefore investigated whether coinfection of HCV in HIV–1–ITP patients enhances the likelihood of inducing anti–GPIIIa49-66 Ab due to molecular mimicry of hepatitis C with GPIIIa49-66, as we have shown for nef with HIV–1–ITP.14


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into NZB/W F1 mice induces thrombocytopenia that correlates with murine anti-PCH09 Ab level. (5) Thrombocytopenic drug abusers dually infected with HIV-1 and hepatitis C have a greater incidence and titer of anti–GPIIIa49-66 Ab as well as greater incidence and severity of thrombocytopenia.

Methods

Human population

Coded stored frozen sera (sent to the clinical laboratory for platelet-Ab testing) were randomly obtained from thrombocytopenic intravenous drug abusers with both HCV and HIV infection, non-drug abuser hepatitis C patients, non-drug abuser HIV-ITP patients, and healthy control subjects. Liver chemistries (albumin, alkaline, phosphatase) were comparable in all 3 groups. These studies were approved by the New York University Medical Center Institutional Review Board.

Mice

Female BALB/c and C57BL/6 mice were obtained from Taconic Farms (Germantown, NY). Integrin GPIIIa−/− knockout mice and NZB/W F1 mice were obtained from The Jackson Laboratory (Bar Harbor, ME). Animal work was approved by the New York University School of Medicine Animal Review Board.

Reagents

All reagents were obtained from Sigma-Aldrich (St Louis, MO) unless otherwise designated. 2′,7′-Dichlorodihydrofluorescein diacetate (DCFH-DA) was obtained from Molecular Probes (Eugene, OR).

Peptides

Peptide GPIIIa49-66 (CAPESIEFPVSEARVLED), PHC09 (SAIHIRRNASG), PHC07 (IFDPQAPPGARS), PHC09 preceded by cysteine (CSAIHIRNASG), PHC09-H1 (SAYQVRNSSG), PHC09-H2 (SAYQVRNSTG), PHC09-H3 (SAYEVRNVSG), and irrelevant 10-mer peptide (GIGALFLGFL) were synthesized by Biosynthesis (Lewisville, TX).

Antibodies

Human anti–GPIIIa49-66 Ab and rabbit anti–GPIIIa49-66 Ab were obtained or produced in our laboratory as described previously.1,2 Mouse Ab against HCV peptides (PHC09) or mouse Ab against rHCV core envelope 1 peptide. Positive sera were purified as described.

Assay of platelet particle formation

Gel-filtered murine or human platelets were isolated from EDTA-anticoagulated blood and labeled with anti–CD61-fluorescein isothiocyanate (FITC, human) or anti–CD41-FITC (murine) as previously described.3 Fluorescent-labeled platelets/particles were measured by flow cytometry using a FACSscan (Becton Dickinson Immunocytometry Systems, Mountain View, CA). Gates were adjusted for 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. A shift in fluorescence from right upper quadrant to left upper quadrant and left lower quadrant reflected the percentage of platelet particle induction in 10,000 enumerated events.

Assay of platelet oxidation

Gel-filtered platelets were loaded with 10 μM DCFH-DA for 30 minutes at 37°C as described4 and challenged with platelet Ab. Intracellular DCFH is converted to a fluorescent form by hydrogen peroxide generated in this reaction. Oxidation was quantified by measuring the increase in mean fluorescence by flow cytometry.

Induction of passive thrombocytopenia in mice

Six-week-old female BALB/c mice were randomly divided into 4 groups (n = 4 per group). Purified control IgG, patient IgG, or mouse IgG against peptide PHC09 (50 μg of each) was injected intraperitoneally into Balb/c mice, and platelet counts were followed for 24 hours.

Determination of mouse platelet counts

Platelet counts were determined from 20-μL blood draw into Unopettes (no. 365855; Becton Dickinson Immunocytometry Systems), containing optimal anticoagulant concentration and diluent for quantitating platelet counts by phase-contrast microscopy.

Expression of recombinant HCV core envelope 1 in E coli

HCV core envelope 1 C-histidine tag cDNA encoding 24 amino acids (167-191) of the core and 160 amino acids of the envelope (192-350) and containing the PHC09 sequence was cloned into expression vector PET29a (Novagen, Madison, WI). PHC09 is composed of amino acids 190 to 199 that contain the 2 C-terminal amino acids of the core and the 8 N-terminal amino acids of the envelope. Escherichia coli BL21 (DE3) PlysS cells were transformed with the PET29a-HCV core-envelope 1 plasmid cultured in LB

ELISA

Peptide (20 μg/mL) in 0.1 M sodium bicarbonate buffer (pH 9.6) was adsorbed to a 96-well ELISA plate at 4°C overnight, and blocked with blocking buffer (3% BSA in PBS-Tween [0.1%]) at room temperature for 2 hours. Primary antibody was added at room temperature for 1 hour, followed by horseradish peroxide (HRP)–conjugated secondary IgG for 1 hour at room temperature. Plates were washed 6 times with Tris-buffered saline 0.1% Tween 20 at each step in the procedure. The ABTS reagent used to develop the HRP color reaction at 405 nm was obtained from Pierce Chemical.

Production of anti-PHC09 Ab in GPIIIa−/− mice

GPIIIa−/− mice aged 6 to 8 weeks were immunized by intraperitoneal injection of 50 μg KLH-conjugated PHC09 with equal volume of mineral oil adjuvant for primary immunization. Booster injections were performed 3 times every 2 weeks. Immune sera were titered by ELISA on the PHC09 peptide. Positive sera were purified as described.

Screening of the phage display peptide library

Anti–GPIIIa49-66-66 Ab was incubated with 1012 to 1013 PhD-7 phage from a 7-mer linear peptide library obtained from New England Biolabs (Beverly, MA) in 2% nonfat milk in PBS at room temperature for 3 hours. Protein A– and alternating protein G–conjugated agarose beads were added overnight at 4°C. The beads were then centrifuged and washed 20 times with 0.1% Tween 20–PBS. Positive phages were eluted with trypsin–PBS by shaking for 10 minutes at room temperature, and eluted phage was titered and amplified as previously described.14 After the third round of panning, 30 clones were randomly selected for enzyme-linked immunosorbent assay (ELISA) and the positive clones sequenced. Phage peptide sequences were analyzed for sequence similarity to other proteins using the BLAST algorithm of the blast program and the database of the National Center for Biotechnology Information (NCBI).18
medium (Bacto-tryptone [1%; Difco Laboratories, Detroit, MI], Bacto-

yeast extract [0.5%; Difco Laboratories], NaCl [1%], pH 7.0) containing
carbenicillin (50 μg/mL) and chloramphenicol (34 μg/mL) with shaking at
37°C until the OD (600 nm) was 0.4 to 0.6. Expression was induced by the
addition of 1 mM IPTG at 37°C for 4 hours with shaking, and purified by
Ni-NTA agarose resin. The protein was dialyzed in PBS, and the purity
verified with 12% sodium dodecyl sulfide–polyacrylamide gel elec
trophoresis (SDS-PAGE).

Immunodeficiency animal model

Two animal models were used. (1) NZB/W F1 mouse model: NZB/W F1
mice were directly challenged with rHCV core envelope 1 protein
containing the corresponding PHC09 sequence, or scrambled peptide
(25 μg/mouse, every 10 days for 9 injections). Platelet counts and antibody
titer versus PHC09 were monitored after every injection. (2) BALB/c
mouse model: BALB/c mice were immunosuppressed, as described19
with an initial 6 mg cyclophosphamide/mouse (Mead Johnson, Princeton, NJ)
and maintained with additional doses of 2 mg/mouse every week. BALB/c
mice were challenged with control or rHCV core envelope 1 protein
(25 μg/mouse) every 10 days for 9 injections after the total leukocyte count
reached less than 2.5 × 10^9/L. Platelet counts and antibody titers versus
PHC09 were monitored as above.

Results

Identification and characterization of selected phage clones

To test the hypothesis that an HCV epitope contributes to the
generation of anti–GPIIIa49-66 Ab, we screened a filamentous
phage display 7-mer peptide library using anti–GPIIIa49-66 anti-
body as bait, and focused on peptides sharing homology sequences
with HCV protein. Thirty phage clones that reacted with anti–
GPIIIa49-66 Ab by ELISA were randomly selected. Clones with
nonspecific cross-reactivity to BSA were also screened by ELISA.
Twenty clones reacted positively with anti–GPIIIa49-66 Ab with
high OD value and low cross-reactivity with BSA (ratio > 3;
Figure 1A). Three positive clones (PHC09 [SIHIRNG], PHC07
[SFDPGLF], and PHC01 [VDWPRSE]) showed sequence homol-
gy with HCV protein. Clone PHC09 was in a nonconserved core
envelope 1 region of hepatitis C virus. We also searched the
database for other nonconserved clones with homology for PCH09.
Six such clones were found; PHC09-H1-H6, PHC09-H5, and H6
had the highest homology for PCH09 (Table 1). These clones were
synthesized as 10 mers and were compared for reactivity with
GPIIIa49-66, PHC09, conserved PHC01 and PHC07, and nef.
Figure 1B demonstrates the strongest Ab reactivity with PHC09
and PHC09-H5. Table 1 also records the HIV genotype. Note that
PHC09 is a relatively rare genotype (HCV6) in the United States
(1.5%), whereas PHC09-H1 to PHC09-H-6 are more common
HCVla or HCV1b genotypes (38% and 36%, respectively).

Table 1. HCV core envelope 1 isolates homologous with PHC09

<table>
<thead>
<tr>
<th>Clones</th>
<th>Homology, %</th>
<th>HCV sequence</th>
<th>HCV gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>PHC09</td>
<td></td>
<td>100% SIHIRNASG</td>
<td>Core envelope 1 (HCV,6)</td>
</tr>
<tr>
<td>PHC09-H1</td>
<td>60</td>
<td>SAYOVRNNSG</td>
<td>Core envelope 1 (HCV,1a)</td>
</tr>
<tr>
<td>PHC09-H2</td>
<td>50</td>
<td>SAYOVRNSTG</td>
<td>Core envelope 1 (HCV,1a)</td>
</tr>
<tr>
<td>PHC09-H3</td>
<td>60</td>
<td>SAYEVRNSG</td>
<td>Core envelope 1 (HCV,1b)</td>
</tr>
<tr>
<td>PHC09-H4</td>
<td>70</td>
<td>SAYHVNRNSSG</td>
<td>Core envelope 1 (HCV,1a)</td>
</tr>
<tr>
<td>PHC09-H5</td>
<td>70</td>
<td>SAIEVRNSG</td>
<td>Core envelope 1 (HCV,1b)</td>
</tr>
<tr>
<td>PHC09-H6</td>
<td>70</td>
<td>SAYEVRNASG</td>
<td>Core envelope 1 (HCV,1b)</td>
</tr>
</tbody>
</table>

Candidate clone PHC09 is located in a hepatitis C virus nonconserved region. H1-H6 represent homologous PHC09 isolates. The underlined regions denote an identical amino acid sequence between screened 7-mer peptide and PHC09. Bold letters represent identical amino acid sequence between 10-mer PHC09 and PHC09 isolates. Column 1 denotes the percentage homology of the isolates. Superscript numbers refer to amino acid position in HCV core envelope 1.
Reactivity of Ab with a modified epitope

Since the core envelope 1 protein is partially cleaved during virus secretion from the infected cell with removal of 2 amino acids from the core, we tested the remaining 8-mer peptide from 3 different HCV isolates. All 3 had similar or greater activity than the PHC09 10-mer peptide (Figure 1C).

Cross-reactivity of HCV-ITP, HIV-ITP, and HCV-HIV-ITP sera with GPIIIa49-66, PHC09, PHC09-H5, PHC07, and nef

Since HIV-1/HCV-ITP drug abusers have a higher incidence and severity of ITP, we reasoned that they might have a greater incidence of anti-GPIIIa49-66 Ab secondary to molecular mimicry with HCV peptides as well as recently reported HIV- nef. Figure 2A through C
demonstrate considerably increased Ab reactivity versus GPIIIa49-66 in 12 of 15 HIV/HCV-ITP drug abusers (Figure 2C) compared with 2 of 15 HCV-ITP patients (Figure 2A; \( P < .001 \)) and 6 of 15 HIV-ITP patients (Figure 2B; \( P < .05, \chi^2 \) analysis). Similar differences were noted for the other 4 molecular mimicry HCV and nef Ags in the 3 groups of patients (Table 2). Of further interest was the presence of Ab to all 4 Ags in 7 of 15 HIV/HCV-ITP patients compared with 1 HCV-ITP patient and 1 HIV-ITP patient (\( P = .019 \)). A greater severity of thrombocytopenia, less than 30,000 platelet counts \( \times 10^9 / L \) (Figure 2D), was also noted in 8 of 15 HIV/HCV-ITP drug abusers compared with 2 HCV-ITP (\( P < .025 \)) and 3 HIV-ITP (\( P = .06 \)) patients. Thus, the greater incidence and severity of thrombocytopenia in HIV-1/HCV-ITP drug abusers could be explained on the basis of molecular mimicry of GPIIIa49-66 with the 4 nonconserved HCV peptides and HIV-1 nef.

In this regard, it is of interest to cite the work of Yong and Wang\(^2\) who used an antihuman HCV core polyclonal Ab to pan a phage peptide library for hepatitis C epitope mapping. An analysis of their data reveals the presence of a dominant linear epitope at residues 19 to 25 of the N-terminal end. This peptide, PQX-V(I)XFP, has 86% homology with platelet GPIIIa51-57 (PESIEFP), which resides within the GPIIIa49-66 peptide, confirming our observations.

### Clinical association of HCV peptide–specific antibodies and thrombocytopenia

To determine the clinical association of HCV peptide and thrombocytopenia in patients, we obtained 15 serum samples from thrombocytopenic HIV-1/HCV-infected drug abusers and looked for a correlation between their platelet count and titer versus the 3 HCV peptides. Figure 3A and B demonstrates an inverse correlation between patient platelet count and Ab titer versus the PHC09 peptide (\( r^2 = 0.7, P < .01 \)) and the PHC09-H5 peptide (\( r^2 = 0.53, P < .05 \)). Since Ab versus PHC09 provided the best correlation (Figure 3A) and highest titers (Figure 2A-C), we focused on anti-PHC09 Ab located in HCV core envelope 1.

### Effect of the HCV peptide–specific antibodies on Ab-induced oxidative platelet fragmentation

To assess the potential effect of HCV peptide on Ab-induced platelet fragmentation, peptide PHC09 (SAIHIRNASG) was then tested for its ability to inhibit affinity-purified patient GPIIIa49-66 Ab-induced platelet fragmentation in vitro. PHC09 significantly inhibited Ab-induced human platelet fragmentation at a (IC\(_{50}\)) molar Ab/peptide ratio of 1:5 (similar to anti–GPIIIa49-66 peptide CAPESIEFPVSEARVLED), whereas irrelevant peptide had no effect at 1:100 (Figure 4A).

### Induction of oxidative platelet fragmentation with Ab

We next affinity-purified IgG with a PHC09 affinity column and confirmed that (similar to anti–GPIIIa49-66) anti-PHC09 IgG induces oxidative platelet fragmentation in vitro (Figure 4B,C). We therefore looked for platelet fragments in the immune complexes of 10 HCV-ITP patients by immunoblot. They were found in patient nos. 10 and 13 (Figure 4D). We next isolated IgG from the serum immune complexes of patients nos. 10 and 13 as well as patients nos. 2 and 6 who had undetectable platelet fragments. Both positive patients tested (nos. 10 and 13) had immune complex IgG capable of oxidatively fragmenting platelets, whereas the 2 control patients (nos. 2 and 6) were unreactive (Figure 4E,F).

### Effect of mouse Ab against HCV peptide on induction of thrombocytopenia in mice

The in vivo relevance of this peptide was further tested by raising an Ab against the KLH-conjugated peptide in GPIIIa\(^{−/−}\) KO mice, which do not recognize the peptide as self, avoiding the problem of mouse tolerance. The mouse serum Ab against PHC09 was shown to bind to GPIIIa49-66 (data not shown). Affinity-purified mouse anti-PHC09 IgG, mouse control IgG, and rabbit anti–GPIIIa49-66 IgG (50 \( \mu \)g/mouse, \( n = 4 \)/group) were then injected intraperitoneally into Balb/c mice, and platelet counts followed for 24 hours. Figure 5 demonstrates that whereas mouse control IgG had no effect, rabbit anti–GPIIIa49-66 IgG or mouse anti-PHC09 IgG induced thrombocytopenia.

### Pathological effect of rHCV core envelope 1 on mouse platelet counts

Since PHC09-SAIHINASG is a molecular mimic of GPIIIa49-66 and located in HCV core envelope 1, we investigated its in vivo pathological

### Table 2. Ab reactivity against various peptides in HCV-ITP, HIV-ITP, and HCV-HIV-ITP patients

<table>
<thead>
<tr>
<th></th>
<th>GPIIIa49-66</th>
<th>PHC09</th>
<th>PHC09-H5</th>
<th>PHC07</th>
<th>Nef</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCV-ITP</td>
<td>2</td>
<td>3</td>
<td>6</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>HIV-ITP</td>
<td>6</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>HCV-HIV-ITP</td>
<td>12</td>
<td>10</td>
<td>8</td>
<td>10</td>
<td>7</td>
</tr>
<tr>
<td>( P^* )</td>
<td>.046</td>
<td>.001</td>
<td>.007</td>
<td>.001</td>
<td>NS</td>
</tr>
</tbody>
</table>

\*\( \chi^2 \) analysis of positive Ab reactivity in the various groups. Upper \( P \) value row is difference between HCV-HIV-ITP and HIV-ITP. Lower \( P \) value row is difference between HCV-HIV-ITP and HCV-ITP.
effect on the mouse platelet count. Autoimmune SLE NZB/W F1 mice and immunosuppressed BALB/c mice were challenged with multiple rHCV core-envelope 1 injections. Platelet counts decreased in 8 of 8 NZB/W F1-treated mice (\( P < 0.001 \); mean platelet count of 1,003 ± 0.029 vs 0.87 ± 0.074 \( \times 10^9/\text{L} \); Figure 6A,B), whereas no decrease in platelet count was noted in the comparison groups immunized with control protein or rHCV core envelope 1 peptide in BALB/c immunosuppressed mice (data not shown). In addition, the serum titer to peptide PHC09 was elevated in NZB/W F1 mice immunized with rHCV core envelope 1, whereas the control group had no significant effect (Figure 6C). The platelet decrease correlated inversely with serum titer versus PHC09 (\( r^2 = 0.63, n = 16, P < 0.05 \)) (Figure 6D).

Since platelet counts represent an equilibrium between platelet production and destruction, we reasoned that the modest decrease in platelet count could have been dampened by a compensatory increase in platelet production by the challenged bone marrow. We therefore affinity-purified the serum IgG with PHC09, to increase its relative concentration. Figure 6E demonstrates a more pronounced thrombocytopenia (60% decrease in platelet count compared with control IgG).

**Discussion**

Our current data clearly demonstrate that the HCV core envelope 1 can induce functionally active anti–GPIIIa49-66 Ab by mimicking platelet GPIIIa49-66 as antigen and inducing immune thrombocytopenia in NZB/W F1 mice. Anti–HCV core envelope 1 Ab correlates inversely with platelet counts in HIV-1 drug abusers with HCV infection. To our knowledge, this is the first report that an HCV epitope can induce anti–GPIIIa49-66 Ab by molecular mimicry.

We now describe a second immune complex–associated platelet immunologic disorder (HCV-ITP) in which a unique Ab is developed against GPIIIa49-66 capable of inducing oxidative platelet fragmentation. It is now apparent that this new pathologic mechanism of platelet destruction is not unique for HIV-1–ITP and could possibly be associated with other immune complex–associated disorders in which serum complexes may contain platelet fragments and anti–GPIIIa49-66 Ab.

Attempts to induce thrombocytopenia by injection of HCV peptide PHC09 into immunosuppressed wild-type mice were unsuccessful—either due to immune tolerance (mouse GPIIIa has...
PHC09 mice. Percentage platelet decrease correlated inversely with serum titer versus percentage platelet count decrease and serum titer versus PHC09 in NZB/W F1 (HCV; C), whereas the control protein (control) had no effect. (D) Correlation between PHC09 was elevated in NZB/W F1 mice immunized with rHCV core envelope 1 in comparison group immunized with control protein (A). The serum titer to peptide obtained from virus isolates of genotypes 1a, 1b, 2b, and 3a.25

192 to 202 amino acids of E1 from 12 different E1 proteins donor infected with HCV genotype 1b reacts with the N-terminal envelope region, which is likely to retain the epitope specificity. Indeed, a human monoclonal Ab (H111) derived from a hepatitis C virus envelope to avoid host immune surveillance. Nevertheless, it is now apparent that at least some patients with HCV-ITP may develop their ITP due to molecular mimicry. This is supported by our ability to induce immunologic thrombocytopenia in NZB/WF1 mice with a core-envelope construct. It is well known that thrombocytopenia increases with severity of disease—and has been claimed that this may be due to liver disease rather than Ab. We consider this debatable because of patients whom we have studied with severe liver cirrhosis who had abundant IgG on their platelets as well as in their immune complexes.16

It is of interest that both HCV and HIV-1 each contain different structural elements that mimic peptide sequences in GPIIIa49-66 and that both viruses are capable of inducing GPIIIa49-66-specific antibodies capable of causing thrombocytopenia. A possible explanation is that both the HCV and HIV-1 mimicry epitopes are in nonconserved highly mutated regions of the 2 viruses, and that mimicry of host proteins serves to help viruses escape host immune surveillance. Therefore the mutations of these mimicry regions are likely to contribute to the autoimmunity in general, and in our study, thrombocytopenia more specifically. It is also likely that coinfection of HCV and HIV-1 increases the chances of molecular mimicry.

In summary, these findings demonstrate the potential of HCV or other pathogens to induce antplatelet Ab by molecular mimicry—and suggests that platelet membrane GPIIIa49-66 may be a common immunodominant target for these pathogens, particularly HIV-1/HCV for the induction of immunologic thrombocytopenia.

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Authorship

Contribution: W.Z. performed and analyzed most of the experiments; M.A.N. performed the platelet fragmentation experiment; W.B. sug-
gested the HCV genotype experiments; Z.L. designed the phage display experiments; and S.K. guided the project and wrote the paper.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

References


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