Spontaneous Control of Primary Hepatitis C Virus Infection and Immunity Against Persistent Reinfection

WILLIAM O. OSBURN,* BRIAN E. FISHER,* KIMBERLY A. DOWD,* GISELLE URBAN,* LIN LIU,§ STUART C. RAY,*‡ DAVID L. THOMAS,* and ANDREA L. COX*‡

*Departments of Medicine and‡Oncology, Johns Hopkins Medical Institutions, Baltimore, MD;§Department of Infectious Diseases, Southwest Hospital, Third Military Medical University, Chongqing, China

BACKGROUND & AIMS: We followed patients with ongoing hepatitis C virus (HCV) exposure following control of an initial HCV infection to determine whether primary control conferred protection against future persistent infections. METHODS: Twenty-two active injection drug users (IDU) who had cleared a primary hepatitis C viremia for at least 60 days were monitored monthly. Reinfection was defined as the detection of a new HCV infection. Protection was assessed based on the magnitude and duration of viremia following reinfection and generation of T-cell and neutralizing antibody (nAb) responses. RESULTS: Reinfection occurred in 11 IDU (50%) who previously spontaneously controlled primary HCV infection. Although viral clearance occurs in approximately 25% of patients with primary infections, spontaneous viral clearance was observed in 83% of reinfected patients. The duration and maximum level of viremia during subsequent episodes of reinfection were significantly decreased compared with those of the primary infection in the same subjects. In contrast to chronic infection, reinfection was associated with a significant increase in the breadth of T-cell responses. During acute infection, nAbs against heterologous viral pseudoparticles were detected in 60% of reinfected subjects; cross-reactive nAbs are rarely detected in patients who progress to chronic infection. CONCLUSIONS: HCV reinfection is associated with a reduction in the magnitude and duration of viremia (compared with the initial infection), broadened cellular immune responses, and generation of cross-reactive humoral responses. These findings are consistent with development of adaptive immunity that is not sterilizing but protects against chronic disease.

Abbreviations used in this paper: ALT, alanine aminotransferase; HCVpp, hepatitis C virus pseudoparticle; IDU, intravenous drug users; nAb, neutralizing antibodies; PBMC, peripheral blood mononuclear cells; RLU, relative light units.

Intravenous drug use is the most significant risk factor for infection with hepatitis C virus (HCV) in the United States, as most new infections occur in intravenous drug users (IDU). The incidence rate of HCV infection among IDU in metropolitan areas of the United States, Europe, and Australia has been shown to be 10% to 30% per year and a recent meta-analysis by Hagan and colleagues revealed that by the 10th year of drug use 75%—80% of IDU have been infected with HCV at least once. At the same time, spontaneous HCV clearance typically occurs in <30% of infected IDU.

The occurrence and outcomes of HCV reinfection have implications regarding vaccine development. Currently, no HCV vaccine has been licensed, despite the public health need. However, work in the chimpanzee model suggests that immunity against HCV can be generated by initial infection and vaccination. Clearance of multiple infections with homologous and heterologous virus has been observed in chimpanzees. Moreover, Landford and colleagues reported that clearance of both homologous and heterologous viral rechallenges was associated with decreased duration and magnitude of viremia. However, the existence of protective immunity in humans remains controversial.

Similar evidence for protective immunity has been reported in two studies in humans. Mehta and colleagues and Grebely and colleagues demonstrated a decreased risk for development of viremia in previously infected IDU compared to naive IDU, even after accounting for risk behavior. Conversely, a recent study by Aitken and colleagues reported a higher rate of infection in previously cleared young IDU compared to naive young IDU. There are major technical limitations to these human studies that would cause protective immunity to be underestimated. Instances of reinfection with rapid clearance of viremia would often be missed by infrequent sampling, and viral sequencing is necessary to distinguish new infection from recrudescence of pre-existing viremia. However, persons who recovered once might recover more often a second time, even without generation of protective immunity, due to host factors. More rapid and effective control of subsequent infections than the first infection would suggest a role for adaptive protective immunity rather than fixed genetic factors. Thus, it is
also important to assess the magnitude and duration of viremia in the same person with sequential infections.

In this study, we have longitudinally followed young IDU with ongoing exposure following control of their initial infection to assess the kinetics and immunological parameters of HCV reinfection. Using a strict definition of reinfection as viremia with heterologous virus with at least 60 days of aviremia following control of the primary infection, we were able to document a new infection in half of the individuals who had previously cleared infection. Previous clearance of a primary HCV infection was associated with high reinfection clearance rates and, compared to initial infection, with a decreased duration and magnitude of viremia during subsequent infections. Re-infection with a heterologous virus also resulted in generation of new T-cell responses not seen with the initial infection and cross-reactive anti-HCV neutralizing antibodies. Taken together, these data demonstrate that previous clearance of HCV alters the outcomes and kinetics of secondary infections, providing further evidence that generation of protective immunity against HCV is possible.

**Methods**

**Participants**

In a prospective study of young IDU in Baltimore, Maryland, the incidence, immunology, and virology of HCV infection were examined. Participants were invited to enroll in a prospective study of acute hepatitis C if they are at risk for HCV infection because of drug use but are anti-HCV antibody and HCV RNA negative. Those who consented are provided counseling to reduce the risks of drug use and monthly blood samples were collected as described previously. All participants with acute HCV infection were referred for evaluation for possible treatment. The study protocol was approved by the Institutional Review Board of the Johns Hopkins School of Medicine.

We identified people who developed antibodies to HCV between 1997 and 2007 with sufficient follow-up to evaluate outcomes of infection. HCV RNA testing was performed on serum or plasma separated from blood using a quantitative HCV RNA assay, described here later. HCV RNA testing was performed on samples collected before seroconversion until a negative result was obtained to determine the time of initial viremia and after seroconversion to evaluate the outcomes of infection. Peripheral blood mononuclear cells (PBMC; \(10^8\)–\(10^{10}\)) were collected from infected persons who were eligible for and consented to unit blood donation or apheresis.

**HCV RNA Assays**

**Determination of viral load.** Total RNA was extracted from serum using a Qiagen MinElute Virus column (Qiagen, Valencia, CA) according to manufacturer’s instructions. To determine the concentration of HCV RNA in blood samples, we used a quantitative reverse transcription polymerase chain reaction (PCR) assay (TaqMan HCV analyte-specific reagent, Roche Molecular Diagnostics, Indianapolis, IN) with the generation of DNA amplification products monitored on a Cobas TaqMan Analyzer (Roche Molecular Diagnostics). This assay has a lower limit of detection of 50 IU/mL.

**Phylogenetic analysis.** HCV Core-E1 sequences were obtained from the first and last viremic specimen of the initial period of viremia and from the first and last viremic specimen of any subsequent period of viremia. E2 sequences were obtained from viremic specimens within the first 100 days of an initial infection, following the detection of new genotype during initial infection, and during reinfections. Total RNA was extracted from serum using a QiAamp viral RNA mini column (Qiagen, Valencia, CA) according to manufacturer’s instructions. Direct sequencing of reverse transcription PCR products from the Core-E1 region was performed as described previously. Sequences were aligned using ClustalX (www.clustal.org) and trimmed to equal length using BioEdit (Ibis Technologies, Carlsbad, CA). The E2 region was reverse transcribed using genotype-specific primers and amplified using a nested PCR strategy with genotype-specific primers (Supplementary Table 1). Purified E2 PCR products were cloned into the entry vector, pDONR221, using Gateway technology (Invitrogen, Carlsbad, CA). Sequences were assembled into contigs using Aligner (CodonCode Corporation, Dedham, MA). Genetic divergence between Core-E1 sequences and E2 protein divergence from H77 was determined using the DNADist DNA matrix or ProtDist Phylip programs, respectively, included in the BioEdit software package. Viral sequences were identified as unique when the Core-E1 divergence between two sequences was \(\geq 0.05\). Genotypes were determined by comparing Core-E1 sequences to HCV genotype reference sequences using the Los Alamos National Laboratories HCV Phylogenetic Placement Service with pairwise distance analysis. All E2 sequences have been deposited into Genbank/EMBL/DDBJ with accession numbers (Genbank accession number GU055316-GU055395).

**Viral recovery and reinfection.** HCV controllers were defined as individuals with anti-HCV antibodies and HCV RNA undetectable by the COBAS Taqman reverse transcription PCR quantitative assay for a period of at least 60 days. Chronic infection was defined as continuous viremia or recurrent viremia with the same virus as determined by Core-E1 phylogenetic analysis. Reinfec-
tion was defined as the presence of new viremia, defined as a genetically unique HCV by Core-E1 phylogenetic analysis, in HCV controllers. The outcome of reinfection was analyzed in subjects who had follow-up for at least 120 days postreinfection. Control of reinfection was defined as undetectable HCV RNA for at least 60 days. Persistent reinfection was defined as continuous viremia...
or recurrent viremia with the same virus following the detection of reinfection. The midpoint between sample collection dates was used to determine duration. Subjects were excluded from analysis of HCV reinfection if they met any of the following criteria: (1) had gaps in follow-up >365 days, (2) had positive hepatitis B surface antigen at time of enrollment, or (3) could not successfully undergo Core-E1 sequencing of the initial HCV infection.

**Alanine Aminotransferase (ALT) Assay**
ALT levels in plasma samples were determined by the Johns Hopkins Hospital clinical laboratory.

**Interferon-γ ELISpot Assay**
HCV-specific CD8+ T-cell responses were quantified by interferon-γ ELISpot assay as previously described.7 Briefly, PBMC were screened for recognition of HCV-specific antigens using overlapping peptides and previously described optimal HCV epitopes. Pooled cytomegalovirus, Epstein-Barr virus, and influenza antigens and phytohemagglutinin were used as positive controls. A new T-cell response was defined as recognition of a peptide by PBMCs collected >200 days following the initial detection of viremia that was not recognized by PBMCs collected at earlier time points.

**H77 Pseudoparticle Production and Neutralization Assays**
HIV-H77 (HCV genotype 1a) and HIV-murine leukemia virus pseudoparticles containing the luciferase reporter gene were produced as described elsewhere.19,20 Neutralization assays were performed as described previously.21 Pseudoparticle infection, indicated by luciferase activity, was measured in terms of relative light units (RLUs) in the presence of subject serum (RLUtest) versus average infection in the presence of 2 HCV-negative serum specimens (RLUcontrol). Percent neutralization was calculated as 100 × [1 – (RLUtest/RLUcontrol)]. Results are reported as 50% inhibitory dilution values. As a control, all subject serum neutralized murine leukemia virus pseudoparticles <50%.

**Statistical Analysis**
Wilcoxon rank-sum test, paired \( t \) test, Kruskal-Wallis one-way analysis of variance by ranks, followed by Dunn’s multiple comparisons and Fisher’s exact test were used to evaluate statistically significant differences between groups. Differences were considered statistically significant when \( P < .05 \).

**Results**
From 1997 to 2007, we detected anti-HCV antibody seroconversion in 113 individuals (Figure 1). No period of aviremia >60 days was observed in 82 of these seroconverters, indicative of chronic infection. Of the 31 persons who controlled viremia, 9 were excluded from subsequent analysis for reinfection, permitting analysis of reinfection in 22 subjects.

**Analysis of Reinfection**
Reinfection with a heterologous virus was documented in 11 cleared subjects while no new viremia was detected in 11 subjects. Overall, the incident reinfection rate was 30.1 reinfections per 100 person-years. The median interval between HCV testing, total follow-up from seroconversion, follow-up postclearance, subject age at the time of seroconversion, and subject gender composition did not differ between the Reinfection and No-reinfection groups, respectively (Table 1). All subjects were Caucasian.

Phylogenetic analysis of viral sequences during initial infection and subsequent infections was performed to confirm reinfection. Seven subjects were reinfected with genotype 1a, 2 with genotype 1b, and 2 with genotype 3a (Table 2). Reinfection with a new virus of the same subtype occurred in 6 subjects, while reinfection with a new virus of a different subtype or genotype occurred in 3 and 5 subjects, respectively.

To examine the frequency of virus replacement in chronically infected subjects, we performed phylogenetic analysis of viral sequences obtained during various time points of chronic infection. This analysis revealed the appearance of a genetically distinct virus in 19 of 82 chronically infected subjects. Ten of these unique viruses were a different genotype, while 9 were the same genotype as the initial infecting virus (data not shown).
Outcome of Reinfection

The 11 reinfected subjects were monitored to assess the outcome of reinfection. One reinfected subject was excluded from analysis due to inadequate follow-up after initial detection of reinfection to determine outcome, while 2 subjects cleared 2 reinfections each. Therefore, we were able to analyze outcomes of a total of 12 reinfections in 10 subjects (Figure 1). Representative graphs are shown in Figure 2, illustrating HCV RNA and ALT levels throughout the history of infection. Various outcomes following control of initial infection are shown; no reinfection (Figure 2A), cleared reinfection (Figure 2B), and persistent reinfection (Figure 2C). Additional graphs are included in Supplementary Figure 1. Interestingly, increases in blood ALT levels of at least 2.5-fold over baseline were observed during 10 of 12 reinfections (Table 2). In total, 10 reinfections were controlled and 2 reinfections resulted in persistent viremia (follow-up during persistent reinfection: 1435 and 2268 days). The ratio of clearance to persistence in reinfection was nearly reversed and significantly different from that of primary infection (P = .001). The median total follow-up time following first detection of viremia and median testing interval in subjects with a primary infection and subjects with a reinfection were not different (P = .217 and P = .663, respectively; data not shown). These data suggest that spontaneous clearance of a reinfection is significantly more likely than spontaneous clearance of a primary infection.

Infection Kinetics During Primary Infection and Reinfection

Frequent monitoring of HCV infection status allowed for assessment of the kinetics of viremia during initial and subsequent infections within the same subjects, as has previously been done with experimental infection of chimpanzees. The maximum concentration of HCV RNA detected in blood during reinfection was

Table 1. Follow-Up Data and Demographics of Control Reinfection and Control No-Reinfection Subjects

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Follow-up from seroconversiona (d)</th>
<th>Follow-up postclearancea (d)</th>
<th>Testing intervala (d)</th>
<th>% Female</th>
<th>Ageb (y)</th>
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<tbody>
<tr>
<td>Control</td>
<td>11</td>
<td>1.269 (981–1,856)</td>
<td>702 (505–1,397)</td>
<td>32</td>
<td>55</td>
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<tr>
<td>Reinfection</td>
<td>11</td>
<td>984 (741–1,791)</td>
<td>815 (344–966)</td>
<td>33</td>
<td>55</td>
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</table>

P value

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<th>Subjects</th>
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<th>Follow-up postclearancea (d)</th>
<th>Testing intervala (d)</th>
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<td>33</td>
<td>55</td>
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P value

Table 2. Phylogenetic Analysis of Viral Sequences and Plasma ALT Levels During Primary Infection and Reinfection Within the Same Subject

<table>
<thead>
<tr>
<th>Subject ID</th>
<th>ALTa</th>
<th>Genotype</th>
<th>ALTb</th>
<th>Genotype</th>
<th>ALTb</th>
<th>Genotype</th>
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<th>Genotype</th>
<th>ALTb</th>
<th>Divergencec</th>
<th>Outcomed</th>
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<tr>
<td>18</td>
<td>13</td>
<td>1a</td>
<td>74</td>
<td>1a</td>
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<td>0.109</td>
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<td></td>
<td></td>
<td></td>
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<td>19</td>
<td>16</td>
<td>1a</td>
<td>411</td>
<td>1a</td>
<td>577</td>
<td>0.071</td>
<td>P</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>27</td>
<td>30</td>
<td>3a</td>
<td>559</td>
<td>1a</td>
<td>553</td>
<td>0.562</td>
<td>C</td>
<td></td>
<td></td>
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<td>48</td>
<td>16</td>
<td>3a/1a</td>
<td>258</td>
<td>1a</td>
<td>63</td>
<td>0.592, 0.595</td>
<td>C, C</td>
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<td>1a</td>
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<td>0.103</td>
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<td></td>
<td></td>
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<td>133</td>
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<td>1a</td>
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<td>1a</td>
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<td>0.116</td>
<td>C</td>
<td></td>
<td></td>
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<tr>
<td>152</td>
<td>17</td>
<td>1a</td>
<td>535</td>
<td>1b</td>
<td>347</td>
<td>0.375, 0.383</td>
<td>C, C</td>
<td></td>
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<tr>
<td>170</td>
<td>6</td>
<td>1a</td>
<td>38</td>
<td>3a</td>
<td>61</td>
<td>0.628</td>
<td>NA</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>172</td>
<td>16</td>
<td>1b</td>
<td>387</td>
<td>1b</td>
<td>64</td>
<td>0.163</td>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>176</td>
<td>22</td>
<td>1a</td>
<td>785</td>
<td>3a</td>
<td>62</td>
<td>0.625</td>
<td>C</td>
<td></td>
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<tr>
<td>180</td>
<td>18</td>
<td>3a/1a</td>
<td>475</td>
<td>1a</td>
<td>84</td>
<td>0.102</td>
<td>C</td>
<td></td>
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</table>

aBaseline alanine aminotransferase (ALT) value prior to infection; all subjects returned to baseline prior to reinfection.

bMaximum ALT value during infection.

cFirst value represents divergence between primary infection and reinfection A. Second value represents divergence between reinfection A and reinfection B.

dC, cleared; P, persistent; NA, excluded from analysis of outcome. The first and second letters indicate the outcome of the first and second reinfections, respectively.

eNA, subject never returned to baseline following reinfection A.
approximately 3 logs lower compared to initial infection even when persistent reinfections were included in the analysis (Figure 3A). Similarly, the duration of viremia during primary infection was nearly 4-fold longer than in subsequent infections in subjects who cleared reinfections (Figure 3B). These results suggest that immunologic events associated with clearance of an initial infection alter the kinetics of subsequent infections.

**Cellular Immune Responses Following Exposure to a New Virus**

Loss of breadth and specificity of T cells generated in the acute phase of HCV infection with a resultant paucity of cellular immune responses is one of the hallmarks of chronic HCV infection. Because reinfection is associated with altered infection kinetics, we investigated whether reinfection altered cellular immune responses to HCV, and specifically whether reinfection was associated with generation of new T-cell responses. Exposure to a genetically distinct virus following a period of aviremia was associated with acquisition of a significantly greater number of new T-cell responses than in persistent viremia (Figure 4), whether chronic infection was with the same virus or genetically distinct viruses (viral switch). In reinfected subjects, exposure to a new virus elicited new T-cell responses in all 4 subjects from whom sufficient PBMC were obtained to evaluate cellular immune responses. Three of 4 subjects with new responses were infected with only a genotype 1a virus during both initial and subsequent infections. These new T-cell responses were directed against epitopes evenly distributed across the HCV polyprotein (Supplementary Figure 3 and Supplementary Table 2). There was no significant correlation between the number of new T-cell responses and decreases in maximum HCV RNA concentrations during reinfection ($R^2 = .36; P = .40$). While the number of new T-cell responses appeared to be negatively correlated with increased ALT during reinfection ($R^2 = .89$), this correlation was not statistically significant due
to small sample size (P = .087). In contrast, evolution of the same viral sequence in subjects with chronic infection resulted in new cellular responses in only 1 of 11 subjects tested, and new cellular responses were detected in only 3 of 10 subjects in whom multiple dominant viral sequences were detected during initial infection. The number of new T-cell responses in these 2 groups was significantly less than in reinfected subjects. These results suggest that exposure to a new virus is not sufficient to elicit new cellular immune responses and that previous complete control of viremia is associated with enhanced generation of new cellular immune responses in response to a new HCV infection. However, 1 reinfection subject with new T-cell responses developed a persistent reinfection, suggesting that development of new T-cell responses does not provide absolute protection against persistence. This led us to investigate the humoral response to reinfection.

**Humoral Immune Responses During Reinfection**

Because the appearance of neutralizing antibodies (nAb) corresponds with clearance of initial infections in some individuals,24 we investigated the role of nAb in control of reinfection. nAbs against a genotype 1a pseudoparticle were detected during reinfections in subjects with and without nAb responses to the H77 E2 protein sequence. Sequences were not obtainable from 3 reinfection viruses due to low HCV RNA levels, but E2 divergence from H77 is shown for the remaining 9 viruses in Figure 5C. E2 sequence analysis revealed similar amino acid divergence from H77 of viruses obtained during reinfections with and without detectable nAbs. Interestingly, some viral sequences were highly divergent from H77 during reinfections in which nAb responses were detectable. Likewise, the sequence divergence of the initial infection stimulating virus from H77 within these groups was not different (Figure 5D). Taken together, these results suggest that, in some subjects, clearance of reinfection is associated with the generation of cross-reactive nAb independent of sequence similarity to stimulating viruses during both initial infection and reinfection.

**Discussion**

Results of our longitudinal analysis of IDU demonstrate that control of an initial HCV infection is associated with abbreviated subsequent viremia, reduced magnitude of viremia, generation of new T-cell responses, the appearance of cross-reactive anti-HCV nAbs, and a higher rate of clearance. Frequent monitoring of HCV infection status in IDU who clear a primary infection demonstrated that reinfection of IDU following control of a primary infection is common, with detection of 1 or more subsequent infections in 50% of cleared subjects. The incident reinfection rate in our study was 30.1 reinfections per 100 person-years. This reinfection rate is very similar to the incident primary infection rate previously reported in the same cohort (27.2 infections/100 person-years), strongly suggesting that prior clearance of HCV infection does not provide sterilizing immunity against reinfection.

Some previous studies demonstrated lower rates of reinfection, but these studies had long intervals between testing. Given the brief duration of viremia with reinfection, these longer testing intervals and therefore less frequent HCV RNA testing likely missed multiple episodes of recurrent viremia. Consistent with this, our reinfection rate is similar to that shown in a study by Aitken and colleagues in which shorter intervals between HCV RNA testing were employed and in which reinfection was documented in 46% of previously cleared IDU. Because approximately half of IDU are demonstrated to have been reinfected, studies with limited longitudinal
analysis of HCV infection status might incorrectly identify viremia as persistent infection following initial exposure when, in fact, it is a reinfection following control of the primary infection. This would lead to underestimation of the actual clearance rate of a single infection.

Studies in older, seropositive IDU have shown that prior clearance of HCV decreases the rate of persistent reinfection compared to initial infection. In contrast, a study in younger, seropositive IDU recently demonstrated that reinfection of previously cleared IDU occurred at a higher rate than primary infections, after accounting for risk factors. These conflicting results may in part be explained by the limitations of these previous studies. In these studies, the cohorts of reinfection subjects consisted primarily of individuals in whom seroconversion had occurred at an unknown time prior to assessment of reinfection. It is possible that the cohorts of older IDU represent a selection of subjects who have controlled multiple infections. The duration of reinfection in subjects who have cleared multiple infections may be progressively shorter with each reinfection, thereby decreasing the likelihood of detecting reinfection in these individuals.

In addition, the inclusion of seropositive, aviremic subjects precludes phylogenetic confirmation that any newly observed viremia is genetically distinct from previous infections. Longitudinal analysis of HCV infection has demonstrated that recurrent, short periods of aviremia followed by viral resurgence are common during acute primary infections. Therefore, reinfection criteria based solely on sequential HCV RNA negative blood samples prior to an HCV RNA positive sample, without sequencing, allow for remergence of the same virus during early primary infection to be misclassified as reinfection. In the current study, phylogenetic analysis of viral sequences coupled with the requirement of a temporal association between viral sequence divergence and aviremia providing complementary methods of confirming reinfection, thereby increasing confidence that reinfections were accurately classified.

While a reduced incidence of reinfection would indicate the presence of sterilizing immunity, protective immunity need not be sterilizing. In addition to a higher rate of control of subsequent infections, the duration and magnitude of secondary HCV viremia were significantly decreased relative to primary infection. The lower duration and magnitude of viremia we observed in subsequent infections suggest that prior exposure to HCV provides protective immunity against persistent reinfection because fixed genetic factors associated with control

Figure 5. Reinfection is associated with generation of cross-reactive neutralizing antibody (nAb). Representative graphs demonstrating nAb titers in (A) a subject who controls reinfection and (B) a persistently reinfected subject. Solid line represents reciprocal 50% inhibitory dilution nAb titer against a genotype 1a hepatitis C virus (HCV) pseudoparticle (H77). Undetectable nAb titers were assigned a value of 25. Black circles represent HCV RNA concentrations (IU/mL) detected in serum or plasma samples obtained at given time points from date of first detection of viremia. Dotted line denotes the HCV RNA limit of detection. Samples below the HCV RNA limit of detection were assigned a value of 25 IU/mL. Viral E2 protein sequence divergence from H77 E2 protein sequences of viruses obtained during (C) reinfection and (D) initial infection from subjects in whom nAb were and were not detected during reinfection. No significant difference in the E2 protein sequence divergence between groups was observed during reinfection and initial infection.
of infection would not be expected to adapt and become more efficient at viral control like the immune response. An effective vaccine must be able to evoke protective immunity against multiple HCV subtypes and genotypes due to the broad genetic diversity of HCV. Even though reinfection was common in our study, prior clearance was associated with protective immunity against subsequent persistent infection with viruses of similar and differing genotypes.

Adaptive immunity has been shown to play a vital role in clearance of primary HCV infection as well as reinfection. In humans, sustained and broad CD4+ and CD8+ T-cell responses are associated with viral control. Additionally, increased levels of serum ALT, which have been associated with T-cell activity in some studies, have been observed during acute infection. However, the magnitude and breadth of CD8+ T-cell responses declines with increasing duration of viremia in chronically infected individuals, suggesting that the effectiveness of cellular immunity in controlling infection declines with time. The role of adaptive immune cells in controlling reinfection has been investigated in chimpanzees, the only animal model for HCV infection. Control of secondary infections in chimpanzees was associated with robust CD8+ T-cell activity resulting in rapid clearance. Likewise, depletion of CD8+ T cells during reinfection resulted in viral persistence, while restoration of CD8+ T cells led to rapid clearance. However, serum ALT levels in chimpanzees remain stable during rechallenge, perhaps indicative of the differences in HCV infection parameters in chimpanzees compared to humans. Increased blood ALT levels were observed during reinfection and new T-cell responses were detected in reinfected individuals at a higher frequency than during chronic infection, suggesting cellular immune responses may contribute to the protective immunity afforded by previous clearance. Moreover, all reinfected subjects with detectable new T-cell responses had lower maximum viremia during reinfection. However, new T-cell responses and increased levels of ALT during reinfection were observed in a subject who developed persistent viremia, indicating that enhanced cellular immune responses provide incomplete protection against persistent reinfection. Our cellular immune response studies were limited by the fact that not all reinfected subjects were eligible for or consented to large-volume blood draws, thereby precluding analysis of T-cell responses in these subjects. Also, T-cell responses were screened using peptides derived from a genotype 1a virus, possibly underestimating T-cell responses during reinfection with non-genotype 1a virus. More detailed characterization of these responses in reinfected individuals may provide further insight into the role of cellular immunity in controlling reinfection and may illustrate immune mechanisms to target with a vaccine.

Although the appearance of neutralizing antibodies has been shown to drive E2 sequence evolution and correspond with control of viremia during initial infection, the role of neutralizing antibodies in clearance of reinfection in humans is unknown. In our study, 60% of reinfected subjects generated nAb against a heterologous hepatitis C virus pseudoparticle (HCVpp) that was highly divergent from the stimulating virus detected during both the initial infection and reinfection in these subjects. Previous experiments have demonstrated that during chronic infection nAbs fail to or weakly neutralize heterologous HCVpp during the acute phase of infection (<33 weeks), while neutralization titers against autologous HCVpp are significantly higher. These findings suggested that the reactivity of nAbs during initial infection is dependent upon the stimulating virus. In contrast, nAb with high titers against heterologous virus were detected in reinfected individuals independent of the sequence of the stimulating virus during initial infection and reinfection, suggesting the presence of cross-reactive nAbs able to neutralize heterologous HCVpp in these subjects. Detection of cross-reactive nAbs during reinfections of short duration suggests that clearance of an initial infection alters subsequent humoral responses to repeated HCV infection, thereby resulting in rapid generation of broadly neutralizing antibodies. Conversely, the ability to generate cross-reactive nAbs may be a marker of the ability to control infection rather than the result of repeated infection. The appearance of broadly neutralizing nAb before detectable reinfection supports this. However, nAb responses were detected in both individuals with persistent reinfection, demonstrating that, nAbs, like enhanced T-cell responses, provide incomplete protection against persistent reinfection.

Moreover, the high prevalence of cross-reactive nAbs observed in our study versus the absence of cross-reactive nAbs during chronic infection suggests that nAb responses generated during reinfection may be qualitatively different than in initial infection. Cross-reactive human monoclonal antibodies against conserved epitopes important in CD81-mediated viral entry have been described. It is possible that prior clearance of an HCV infection results in selection of antibodies targeting conserved epitopes, thereby allowing for neutralization of heterologous HCVpp. Further investigation of the epitopes targeted by nAb responses during reinfection may lead to identification of relevant immunogenic targets for inclusion in future vaccine development.

One limitation of our study is that a period of 60 days of aviremia may be unnecessarily conservative. We detected a genetically distinct virus in 19 subjects with constant viremia or aviremic periods of <60 days prior to detection of the new virus, precluding classification of these subjects as reinfected. It is possible that these individuals controlled a primary infection and were subsequently reinfected between clinical visits, resulting in the
absence of sufficient aviremia to meet our stringent criteria. This would lead to an underestimate of both the true rate of primary infection clearance and the frequency of reinfection. However, defining the appearance of a new virus as a reinfection without sufficient aviremia might potentially overestimate the incidence of reinfection. Sequence-specific PCR analysis of viral sequences in one transfusion recipient infected with multiple HCV strains by exposure to blood from different donors revealed a newly dominant viral sequence late in infection that was detected in the inoculum but not observed in earlier samples from the infected subject. Distinguishing new infection from infection with multiple strains with alternating dominance will require the development of highly sensitive assays for tracking specific viral sequences.

Collectively, the results of our study provide clear evidence supporting protective immunity associated with clearance of primary HCV infection. The fact that protective immunity following HCV infection is possible highlights the plausibility of a preventive vaccine. In the absence of sterilizing immunity, it may still be possible to target immune mechanisms with immunization that favor clearance in naive individuals following infection. Careful study of subjects who clear a primary infection and are subsequently reinfected will be important in determining the immunological correlates critical in the development of such vaccination strategy.

**Supplementary Data**

Note: To access the supplementary material accompanying this article, visit the online version of *Gastroenterology* at www.gastrojournal.org, and at doi: 10.1053/j.gastro.2009.09.017.

**References**


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Reprint requests
Address requests for reprints to: Andrea L. Cox, MD, PhD, Department of Medicine, Johns Hopkins University, Rangos Building, Suite 530, 855 N. Wolfe Street, Baltimore, Maryland 21205. e-mail: acox@jhmi.edu.

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