Naturally selected hepatitis C virus polymorphisms confer broad neutralizing antibody resistance

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Introduction

Despite recent advances in hepatitis C virus (HCV) treatment, a vaccine against the virus is still urgently needed (1–3). Vaccine design is challenging due to extensive world-wide genetic diversity of the virus and rapid viral evolution in infected individuals (4–7).

Most HCV-infected individuals develop neutralizing antibodies (nAbs) against the virus, but viral evolution at highly variable loci in HCV E2 (envelope) can lead to escape from isolate-specific nAbs (8–11). However, isolation of human mAbs capable of neutralizing multiple diverse HCV isolates has shown that nAbs may also target more conserved regions of the E1 and E2 proteins (12–25). Discovery of these broadly neutralizing mAbs has raised hope that a vaccine inducing similar nAbs could prevent HCV infection.

Due to limited availability of diverse strains of replication competent HCV (HCVcc) and previously limited availability of diverse E1E2 clones for use in HCV pseudoparticles (HCVpp), the breadth of neutralization of anti-HCV mAbs has generally been measured against small panels of HCV isolates (12, 16–29). In addition, epitopes of neutralizing mAbs have generally been mapped by alanine scanning mutagenesis and binding assays. While these studies provide useful information on key mAb-binding residues, they do not measure the neutralization resistance conferred by the vast array of HCV envelope polymorphisms that occur in nature.

Therefore, we developed a novel panel neutralization method that uses the natural variation of HCV E1E2, allowing measurement of neutralizing breadth, clustering of mAbs with similar resistance profiles, and identification of neutralization resistance polymorphisms anywhere in E1E2. Hundreds of unique E1E2 clones were isolated from individuals infected with genotype 1 HCV, and, from this library, 19 genotype 1a and 1b HCV E1E2 clones were selected to maximize sequence diversity among clones (30). The panel of 19 clones contains 94% of amino acid polymorphisms present at greater than 5% frequency in a reference panel of 643 genotype 1 HCV isolates from GenBank (31). These clones were used to produce a panel of HCVpp, which were tested for neutralization by 18 previously described broadly neutralizing anti-HCV human mAbs, including some of the most potent and broadly neutralizing mAbs described to date (refs. 12, 16–20, and Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI78794DS1). mAbs were grouped using hierarchical clustering analysis of pairwise neutralization resistance profile correlations and, using novel methods, E1E2 sequences were analyzed to identify polymorphisms associated with broad nAb resistance.

Results

Each neutralizing mAb produces a neutralization profile across the HCVpp panel. Eighteen mAbs were each tested for neutralization of each of 19 clonal genotype 1 HCVpp, the largest genotype 1 panel against which these mAbs have been tested. Neutralization
by representative mAbs is shown in Figure 1A, with neutralization results for all mAbs shown in Supplemental Figure 1. mAbs HC84.26 and AR4A showed the greatest neutralizing breadth, reducing by at least 50% the infectivity of 17 of 19 HCVpp and 18 of 19 HCVpp, respectively, at 10 μg/ml mAb. (Supplemental Figure 1). Notably, sensitivity to each mAb varied across the HCVpp panel, with some E1E2 clones highly sensitive (relative infection <0.1), some moderately resistant (relative infection 0.1–0.5), and some highly resistant (relative infection >0.5) to each mAb. For mAb HC84.26, neutralization sensitivity across the panel varied by more than 1,000-fold. To validate accuracy, IC₅₀ of 6 of the mAbs against full-length HCVcc bearing H77 E1E2 (32) were compared with the relative infection of H77 HCVpp measured using the same mAbs. The correlation between HCVcc IC₅₀ and HCVpp relative infection was significant (r = 0.93, P < 0.02, Supplemental Figure 2). In addition, full mAb dilution curves were performed and IC₅₀ were calculated for 24 HCVpp/mAb combinations. The IC₅₀ correlated significantly with the relative infection measured for the same mAb/HCVpp combinations (r = 0.91, P < 0.0001, Supplemental Figure 3).

**Ranking of resistance of library HCVpp reveals relationships between mAbs.** To further validate the unique pattern of HCVcc panel resistance to each mAb, rankings of HCVpp neutralization resistance to 2 closely related mAbs targeting the same linear epitope (HC33.4 and HC33.8) were compared and found to have extremely high correlation (r = 0.94, P < 0.0001, Figure 1B). Rankings of resistance to 2 unrelated mAbs (HC33.4 and HC84.22) were also compared and, as predicted, showed no correlation (r = 0.07, P = 0.75, Figure 1C). Surprisingly, a highly significant correlation was observed between the resistance profiles of mAbs HC84.22 and AR3C (r = 0.84, P < 0.0001, Figure 1D). Prior alanine scanning mutagenesis suggested that these mAbs bind to distinct epitopes (Supplemental Table 1 and refs. 12, 33). However, the recent publication by Kong and colleagues of the crystal structure of AR3C in complex with E2 core and crystal structures of 2 mAbs related to HC84.22, HC84.41 and HC84.27, with their peptide epitopes, suggest that AR3C and HC84.22 likely bind to overlapping E2 epitopes, supporting the correlation between resistance profiles observed here (21, 34).

Pairwise correlations among all mAb resistance profiles were calculated (Figure 2). Invariably, mAbs predicted to bind overlapping epitopes based on prior alanine scanning clustered together in this analysis (i.e., HC84.22 and HC84.26; AR3A, AR3B, AR3C, and AR3D; HC33.4 and HC33.8; AR4A and AR5A), but, surprisingly, many mAbs thought to have distinct epitopes based on prior mapping (i.e., HC84.22 and AR3C; HC33.4 and AR4A) also showed significant correlation in their neutralization resistance profiles. This suggests that polymorphisms, possibly outside of known binding epitopes, may confer resistance to multiple mAbs or that multiple epitopes may be genetically linked.

Hierarchical clustering analysis using pairwise correlations of neutralization resistance profiles placed all mAbs, except nonneutralizing mAb CBH-4B, into 3 strongly supported neutralization clusters (Figure 2). The 8 mAbs falling in neutralization cluster 1 (NC1), including CBH-2, HC84.26, CBH-5, HC84.22, AR3A, AR3B, AR3C, and AR3D, were chosen for further analysis.

Sequence analysis reveals polymorphisms associated with broad mAb resistance. A method was developed to identify polymorphisms in E1E2 associated with mAb resistance. A representative analysis for mAb HC84.22 is shown in Figure 3A, and the analyses for the remaining NC1 mAbs are shown in Supplemental Figure 4. The amino acid sequence of each of the 14 E1E2 clones most resistant to each mAb was compared with the sequences of the 5 most sensitive E1E2 clones to identify polymorphisms present in any resistant clone but absent in all sensitive clones. Sites with substitutions in at least 2 resistant clones were included in a summary panel for each mAb. The results for all mAbs were compiled, as shown in Figure 3B, to identify polymorphisms associated with resistance to multiple NC1 neutralizing mAbs.

Interestingly, the majority of candidate NC1 mAb resistance polymorphisms fell among positions 416–560, which span the front layer, CD81-binding loop, and the central β-sheet of E2. Polymorphisms in resistant clones relative to sensitive clones were 4-fold more likely in this region than in the remainder of the E1E2 protein (P < 0.0001) (Figure 3B). As expected, some of the resistance-associated polymorphisms were at positions previously identified as binding residues for NC1 mAbs (Figure 3C). For example, D431 and F442 were identified by alanine scanning as binding residues for CBH-2 (19) and HC84 mAbs (12), respectively, and polymorphisms at those sites were also associated with resistance to those mAbs in this analysis. In addition, polymorphisms at 4 sites identified recently as contact residues for mAb AR3C in the E2 core/AR3C crystal structure (residues 433, 438, 442, and 446; ref. 34) were also associated with resistance to mAb AR3C in this analysis.

Notably, most resistance-associated polymorphisms were not at previously identified NC1 mAb-binding residues, and many neutralization-resistant E1E2 clones did not have polymorphisms at any previously identified mAb-binding residues (Figure 3C and Supplemental Figure 4). Eleven resistance-associated polymorphisms, including D431E and F442I, were selected for further analysis.

Polymorphisms at residues 431, 442, and 560 confer relative resistance to multiple broadly neutralizing mAbs. The 11 mutations of interest were introduced individually or in combination by site-directed mutagenesis into an E1E2 clone, 1b09, which was sensitive to neutralization by each of the NC1 mAbs (Figure 4). Surprisingly, introduction of glutamic acid at D431, a known binding residue for CBH-2, conferred relative resistance to 6 of 7 NC1 mAbs, namely CBH-2, CBH-5, AR3A, AR3B, AR3C, and HC84.22. F560Y also conferred a small but significant increase in resistance to 6 of 7 mAbs. Introduction of F442I, individually or in combination with other mutations, conferred resistance to HC84.22 and HC84.26. Most of the remaining mutations tested, including N430D, L433I, L438I, K446E, and A475T, conferred small but statistically significant increases in resistance to mAbs HC84.22 and/or HC84.26. These same 11 mutations were also introduced into a second neutralization-sensitive E1E2 clone (1a53) with similar results (Supplemental Figure 5). When D431E and F442I were back mutated in resistant E1E2 clones, in which they occurred naturally, the predicted loss of mAb resistance was also observed. (Supplemental Figure 6).

Of note, while introduction of D431E into clone 1b09 by site-directed mutagenesis conferred relative resistance to 6 of 7 NC1
mAbs, some E1E2s with naturally occurring E431 were sensitive to most NC1 mAbs (Supplemental Figures 7 and 8), suggesting that E1E2 context, the combined effect of all polymorphisms in individual E1E2 clones, can influence the resistance phenotype of D431E. In addition, although F442 is an AR3C contact residue in the AR3C/E2 crystal structure, introduction of F442I into clone 1b09 did not confer detectable resistance to AR3C, and back mutation of isoleucine to phenylalanine at 442 in an E1E2 clone 1b09 did not confer detectable resistance to AR3C, and back mutation of isoleucine to phenylalanine at 442 in an E1E2 clone in which I442 occurred naturally did not increase sensitivity to AR3C (Figure 4, Supplemental Figure 6, and ref. 34). However, introduction of F442I or F442L into a different sensitive E1E2 clone, 1a53, did confer AR3C resistance (Supplemental Figure 5). This suggests that resistance conferred against some mAbs by F442I is also dependent on E1E2 context.

As shown in Supplemental Figure 9, introduction of D431E, F442I, or F560Y into clone 1b09 reduces HCVpp infectivity, with F442I reducing entry more than 10-fold. Interestingly, introduction of additional polymorphisms L438I, A475T, and P453S along with F442I partially restores infectivity. Most importantly, infectivity of some HCVpp produced with E1E2 clones containing naturally occurring D431E, F442I, or F560Y polymorphisms was greater than the median infectivity of the HCVpp panel (Supplemental Figure 9B), suggesting that compensatory mutations can reverse the fitness cost of these polymorphisms.

Polymorphisms in the central β-sheet confer broad resistance. While D431E, F442I, and F560Y polymorphisms may partially explain the correlation in resistance profiles among NC1 mAbs, none of these polymorphisms confer high level resistance to all NC1 mAbs. Therefore, we further investigated 2 NC1-resistant E1E2 clones, 1a129 and 1a142. Clone 1a129 was resistant despite lacking F442I, D431E, and F560Y (Supplemental Figure 8). Clone 1a142 carries the F442I polymorphism, explaining its resistance to HC84.22 and HC84.26, but it also showed significant resistance to the other NC1 mAbs (Figure 1 and Supplemental Figure 1), which cannot be attributed to F442I. To identify polymorphisms responsible for this resistance, the sequence encoding the E2 front layer, β-sheet, or front layer/CD81-binding loop/β-sheet from resistant E1E2 clones 1a129 and 1a142 was transferred into sensitive E1E2 clone 1b09. These E1E2 chimeras are referred to herein as 1b09/1a129-F or 1b09/1a142-FCβ and 1b09/1a129-FCβ or 1b09/1a142-FCβ, respectively. E1E2 chimeras 1b09/1a129-F and 1b09/1a142-FCβ were poorly functional in HCVpp so were not studied further. The remaining 4 chimeras were functional.

As shown in Figure 5, resistance to HC84.22, AR3A, and CBH-2 was conferred by transfer of the sequence encoding amino acids 526–569 (β-sheet) from resistant clone 1a129 into sensitive E1E2 clone 1b09 (15- to 55-fold increase in resistance). This was also true when the same region (β-sheet) was transferred from resistant clone 1a142 into sensitive E1E2 clone 1b09 (4- to 8-fold increase in resistance, Figure 6). Figure 6 also shows that amino acids 412-444 (front layer) from clone 1a142, including F442I, confer a 2- to 31-fold increase in clone 1b09 resistance to HC84.22, CBH-2, and HC84.26, while the F442I mutation alone confers resistance to HC84.22 and HC84.26 but not to CBH-2. Together, these results suggest that mutation at the F442 site is critical for high level HC84.26 resistance and adequate but not necessary for HC84.22 resistance. Polymorphisms in clones 1a129 and 1a142 between amino acids 562 and 569 (β-sheet) confer resistance to HC84.22, AR3A, and CBH-2.
Sequence analysis of amino acids 526–569 in resistant E1E2 clones 1a129 and 1a142 relative to sensitive clone 1b09 revealed 7 in-common variant sites in the resistant clones (Figure 7A). Each of these 7 polymorphisms in the 1b09/1a129-β chimera were back mutated individually or in combination to match clone 1b09 to identify the polymorphisms responsible for the broad resistance of the 1a129 and 1a142 clones. I538V, Q546L, and T563V in the central β-sheet were critical for maintaining the mAb resistance of the 1b09/1a129-β chimera (Figure 7B). Back mutation at any of these 3 positions resulted in a 2- to 3-fold increase in sensitivity to mAbs CBH-2, CBH-5, AR3A, and HC84.22 (Figure 7C). Notably, the effect of these mutations was additive, as back mutation of V538 to I, L546 to Q, and V563 to T together resulted in 7- to 11-fold increase in sensitivity. As shown in Figure 8A, introduction of I538V/Q546L/T563V into sensitive clone 1b09 conferred relative resistance to 6 NC1 mAbs (median 8.5-fold increase in resistance), and back mutation of the 3 polymorphisms in naturally resistant clone 1a129 conferred sensitivity to all NC1 mAbs (median 5-fold decrease in resistance). Introduction of I538V/Q546L/T563V into sensitive clone 1b09 reduced binding of NC1 mAbs CBH-2 and AR3C, while back mutation of the 3 polymorphisms in naturally resistant clone 1a129 increased mAb binding (Figure 8B). Together, these data suggest that I538V/Q546L/T563V polymorphisms confer resistance to NC1 mAbs by reducing mAb binding.

Residues 538, 546, and 563 fall in the central β-sheet of E2, distant from the binding site of AR3C (ref. 34 and Figure 8C). None of these 3 positions are contact residues for AR3C or known binding residues for any other NC1 mAb. Introduction of I538V/Q546L/T563V reduced entry of the 1b09 HCVpp, suggesting a potential fitness cost (Supplemental Figure 10). However, entry of HCVpp produced with E1E2 clones carrying naturally occurring I538V/Q546L/T563V, including resistant clone 1a129, was not reduced, suggesting that compensatory mutations can reverse the fitness cost of these polymorphisms. Interestingly, I538V/Q546L/T563V was present in the majority of the genotype 1a E1E2 clones in the panel, including some clones, like 1a154 (H77), that are NC1 mAb sensitive (Supplemental Figure 11). This suggests that, as with D431E and F442I, the nAb resistance conferred by I538V/Q546L/T563V can be E1E2 context dependent.

I538V/Q546L/T563V polymorphisms confer resistance to plasma neutralization. We also measured the effect of I538V/Q546L/T563V on neutralization by plasma from HCV-infected subjects (Figure 9). Strikingly, introduction of I538V/Q546L/T563V into sensitive E1E2 clone 1b09 conferred resistance to neutralization by each of 18 plasma samples tested (median relative infection 0.33 vs. 0.65 after I538V/Q546L/T563V, P = 1E–4). Back mutation of these polymorphisms in a naturally neutralization-resistant clone, 1a129, led to increased sensitivity to neutralization by all plasma samples tested (median relative infection 0.76 vs. 0.09 after V538I/L546Q/V563T, P = 8E–6). Notably, this effect on neutralization sensitivity was also observed with plasma from the source subject of resistant E1E2 clone 1a129. Together, these results suggest that this combination of polymorphisms likely confers resistance to neutralization in vivo.

Discussion

Using analysis of neutralization resistance profiles, we identified neutralization-resistant E1E2 clones and mAb neutralization clusters that were not predicted by prior epitope mapping, and we characterized novel polymorphisms conferring relative resistance to up to 6 mAbs in one of these neutralization clusters. Resistance polymorphisms I538V/Q546L/T563V are not at known contact or binding residues for any of the mAbs to which they confer resistance, and they fall within the central β-sheet of E2 rather than the front layer/CD81-binding loop that is the likely binding site for NC1 mAbs. However, these polymorphisms reduce mAb binding to E1E2, suggesting that they may induce structural changes in E2 that reduce mAb binding and neutralization.
Based on prior epitope mapping, the mAbs analyzed in this study could be divided into up to 10 distinct antigenic groups (antigenic domain A–D, antigenic region 1–5, epitope “1”). However, we show for the first time that all of these mAbs, except nonneutralizing mAb CBH-4B, segregate into only 3 distinct neutralization clusters. This suggests that resistance polymorphisms outside of known epitopes may confer resistance to many anti-HCV human mAbs, including some of the most broadly neutralizing mAbs described to date. The identification of clusters of mAbs with highly correlated resistance profiles could have significant implications for HCV treatment with mAb combinations or vaccination designed to induce mAbs against multiple epitopes. This method could be used to identify combinations of mAbs with complementary resistance profiles (i.e., NC1 and NC2 mAbs) that may be more likely to provide additive protection when used together.
neutralization clustering, as it showed synergistic neutralization by a combination of mAbs, HC84.26 and AR4A, which we have identified as NC1 and NC2 mAbs, respectively, in this study (35).

Increasing evidence suggests that mutations outside of known binding epitopes may confer resistance to anti-HCV mAbs. The Carlsen study found no correlation between variation in mapped mAb epitopes and mAb resistance (35). In addition, studies by Keck et al. (14) and Fofana et al. (36) described mutations arising in single HCV-infected subjects that fell outside of known mAb-binding epitopes and conferred relative resistance to multiple anti-HCV mAbs. We extend those findings by showing that broad resistance polymorphisms at noncontact residues are a common mechanism of nAb resistance, as they were repeatedly observed in a representative panel of E1E2 clones generated from 19 HCV-infected subjects. We also show that these resistance polymorphisms can have additive effects and may have phenotypes that are E1E2 context dependent.

These findings are also relevant for HCV persistence in vivo. The I538V/Q546L/T563V polymorphisms confer resistance to neutralization by plasma from HCV-infected subjects, suggesting that these polymorphisms may contribute to HCV persistence during chronic infection. Importantly, identification of these resistance polymorphisms in multiple circulating E1E2 clones from multiple subjects also suggests that any fitness cost of these polymorphisms can be adequately reversed by compensatory mutations elsewhere in the virus.

Multiple resistance-associated polymorphisms, including 4 at sites identified as contact residues for AR3C in the recently published crystal structure of E2 core/AR3C (34), did not confer detectable resistance when introduced into sensitive clone 1b09. This may be because most of the tested mutations are biochem-
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Physically conservative and may have subtle effects on neutralization resistance that are not detectable in vitro using current methods. The effect of some of these minor mutations could be additive, as is the case with IS38V/Q546L/T563V. Some of these mutations may also depend on an E1E2 sequence context not tested in this study. Work is ongoing to expand the HCVpp panel and better understand the influence of multiple mutations and E1E2 context on neutralization resistance.

In conclusion, we have developed a novel, rapid method, which uses the natural variation of HCV E1E2 to characterize patterns of resistance to neutralizing mAbs, identifying clusters of mAbs with shared resistance profiles that were not fully predicted by prior epitope mapping. Using this method, we identified naturally selected polymorphisms in E2 at both contact and noncontact residues that confer resistance to multiple broadly neutralizing mAbs and plasma samples. This neutralization clustering analysis suggests that broad resistance polymorphisms at noncontact residues may be a major mechanism of immune evasion by HCV, facilitating viral persistence and presenting a challenge for HCV vaccine development.

Methods

Source of mAbs. mAbs CBH-4B, CBH-2, CBH-5, CBH-7 (18), HC-1 (19), HC84.22, HC84.26 (12), HC33.4, and HC33.8 (20) were generated in the Fouvng laboratory. mAbs AR1A, AR2A, AR3A, AR3B, AR3C, AR3D (17), AR4A, AR5A (16) were a gift from Mansun Law (The Scripps Research Institute, La Jolla, California, USA).

Source of plasma. Plasma samples were obtained from the BBAASH (37) cohort (Andrea Cox, Johns Hopkins University School of Medicine). Samples from each of the 18 HCV-infected subjects who had previously shown at least 50% neutralization by 1:100 plasma dilution (37) cohort (Andrea Cox, Johns Hopkins University School of Medicine, La Jolla, California, USA).

Figure 6. Polymorphisms between amino acids 526 and 569 (β-sheet) of resistant E1E2 clone 1a142 confer resistance to multiple broadly neutralizing mAbs. Neutralization by the indicated mAb of HCVpp with E1E2 between resistant clone 1a142 and sensitive clone 1b09 (1b09/1a142-β and 1b09/1a142-F [F442I]), E1E2 with F442I introduced by site-directed mutagenesis (1b09/F442I), and with resistant E1E2 clone 1a142 (1a142 wild type), which carries F442I naturally. Chimeras with the front layer (1a142-F, amino acids 412-444) from resistant clone 1a142 are marked with cyan. Chimeras with [β6-I8] (1a142-β), amino acids 526-569) from resistant clone 1a142 are marked in red. Orange indicates clones with F442I polymorphisms. Dashed lines indicate 50% neutralization. Error bars indicate SD between duplicate wells.

Production of HCVpp library. Generation and validation of a diverse library of 19 genotype 1 HCVpp and its use for quantitation of neutralizing breadth has been previously described (30). The HCVpp panel used in this study is identical, except that 2 E1E2 clones, 1b20 and 1a114, were replaced by related clones 1b21 and 1a116, which gave more consistent HCVpp infectivity results. Sequences for E1E2s 1b21 and 1a114 have been deposited in GenBank, accession numbers KM660628-9. Plasma samples for production of the library were obtained from HCV-infected subjects in the BBAASH cohort (37), Irish Anti-D cohort (38), and Swan Project (39). An E1E2 alignment consisting of 634 genotype 1 clones previously selected for their representation of the worldwide diversity of genotype 1 sequences was obtained and cleaned using BIOedit (31). The package “seqinr” was used to load data into R (http://www.r-project.org). An amino acid polymorphism was considered represented by the 19-clone panel if it was observed in at least 5% of the worldwide sequences and was observed in the 19-clone panel.

HCVpp production and infectivity screening. HCVpp were produced by lipofectamine-mediated transfection of HCV E1E2 and pNL4-3. Luc.R-E plasmids into HEK293T cells as previously described (40, 41). Neutralization assays were performed as described previously (8). For neutralization at a single mAb concentration, HCVpp were incubated for 1 hour with mAb at a final concentration of 10 μg/ml and then added in duplicate to Hep3B target cells for 5 to 6 hours before media were changed. Infection was determined after 3 days by measurement of luciferase activity of cell lysates in RLUs. The majority of HCVpp used in neutralization assays produce RLUs greater than 2 × 10^5, which is more than 100-fold above background measured using mock pseudoparticles lacking E1E2. All HCVpp used in neutralization assays produced RLUs of at least 200,000, which is at least 10-fold above background. HCVpp with low infectivity were stored at 4°C until use to avoid freeze thawing, which reduces infectivity. With a few noted exceptions, HCVpp compared with each other were either all freeze thawed once or all stored at 4°C without freeze thawing until use.

Relative infection (neutralization) was calculated by dividing the production of HCVpp library. Generation and validation of a diverse library of 19 genotype 1 HCVpp and its use for quantitation of neutralizing breadth has been previously described (30). The HCVpp panel used in this study is identical, except that 2 E1E2 clones, 1b20 and 1a114, were replaced by related clones 1b21 and 1a116, which gave more consistent HCVpp infectivity results. Sequences for E1E2s 1b21 and 1a114 have been deposited in GenBank, accession numbers KM660628-9. Plasma samples for production of the library were obtained from HCV-infected subjects in the BBAASH cohort (37), Irish Anti-D cohort (38), and Swan Project (39). An E1E2 alignment consisting of 634 genotype 1 clones previously selected for their representation of the worldwide diversity of genotype 1 sequences was obtained and cleaned using BIOedit (31). The package “seqinr” was used to load data into R (http://www.r-project.org). An amino acid polymorphism was considered represented by the 19-clone panel if it was observed in at least 5% of the worldwide sequences and was observed in the 19-clone panel. HCVpp production and infectivity screening. HCVpp were produced by lipofectamine-mediated transfection of HCV E1E2 and pNL4-3. Luc.R-E plasmids into HEK293T cells as previously described (40, 41). Neutralization assays were performed as described previously (8). For neutralization at a single mAb concentration, HCVpp were incubated for 1 hour with mAb at a final concentration of 10 μg/ml and then added in duplicate to Hep3B target cells for 5 to 6 hours before media were changed. Infection was determined after 3 days by measurement of luciferase activity of cell lysates in RLUs. The majority of HCVpp used in neutralization assays produce RLUs greater than 2 × 10^5, which is more than 100-fold above background measured using mock pseudoparticles lacking E1E2. All HCVpp used in neutralization assays produced RLUs of at least 200,000, which is at least 10-fold above background. HCVpp with low infectivity were stored at 4°C until use to avoid freeze thawing, which reduces infectivity. With a few noted exceptions, HCVpp compared with each other were either all freeze thawed once or all stored at 4°C without freeze thawing until use.

Relative infection (neutralization) was calculated by dividing the RLUs produced in the presence of the test mAb (RLU_{test mAb}) at 10 μg/ml by the RLUs of the same HCVpp in the presence of the same concentration of normal human IgG (RLU_{control IgG}). For IC_{50} determination, HCVpp were incubated with serial 2.5-fold dilutions of mAb, starting from 50 or 10 μg/ml, and compared with infection with the same dilutions of normal human IgG.
Selection of resistance-associated polymorphisms. E1E2 sequences of neutralization-sensitive clones were compared with those of neutralization-resistant clones using VisSPA v1.6 (http://sray.med.som.jhmi.edu/SCRoftware/VisSPA/) to identify variants present in each resistant E1E2 clone but in none of 5 neutralization-sensitive clones. First, each E1E2 clone was ranked by increasing resistance to each mAb. The amino acid sequence of each of the 14 E1E2 clones most resistant to each mAb was compared with the sequences of the 5 E1E2 clones most sensitive to that mAb to identify polymorphisms present in each resistant clone but absent in all sensitive clones. Sites with polymorphisms in at least 2 resistant clones but in no sensitive clones were then coded according to whether polymorphisms were present more often in the 7 most highly resistant clones or the 7 more moderately resistant clones. These results were included in a summary panel for each mAb. The results for all NC1 mAbs were compiled to identify polymorphic sites that were associated with resistance NC1 neutralizing mAbs.

N430D was tested by site-directed mutagenesis along with other resistance-associated polymorphisms despite appearing in only one resistant clone, since asparagine 430 is a contact residue for mAb AR3C in the AR3C/E2 core crystal structure.

Generation of site-directed mutants. Mutations of interest were introduced into E1E2 clones by site-directed mutagenesis using QuikChange Lightning Multi (Stratagene). All clones were sequenced in their entirety after mutagenesis to confirm that no unintended additional mutations were introduced.

Generation of E1E2 chimeras. E1E2 chimeras were generated with E1E2 clone 1b09 and regions encoding amino acids 412–444, amino acids 526–569, and amino acids 412–569 from clones 1a142 and 1a129. 1b09 E1E2 lacking the nucleotides encoding these regions was amplified by PCR from the cloned 1b09 plasmid using Accuprime Pfx (Invitrogen) and the following primer sets: 5′ CD81P11b09, CATAAGTTCAACGCGTCCGGATGCCCAG; 3′ CD81Full1b09, GACTCTCTGGGACGCTCCGCTTGAG; 5′ CD81Full1b09, AGGGGGCGCCGAACCGATCGGTTGTCC; and 3′ CD81P21b09, AACATCGGGGGGTCGGTAACAACACCCTG. The regions encoding amino acids 412–444, 526–569, and 412–569 were amplified by PCR from 1a129 plasmids using the following primers: 5′ CD81BS1a129, GCGTCCCAGAGAGTCCAGCTGGTCAACACTAACGGTAGTTGGCAC; 3′ CD81BS1a129, GACCCCCCCGATGTTACAGGGGGGGCCTCCGCACACTTTGGTGAATCC; 3′ CD81P11a129, CGCGTTGAACTTATGGCGGTAGAAGAGCCCCGCCACCCAGCCG; and 5′ CD81P21a129, CGGTTCGGCGCCCCTACCTACAGCTGGGGTGAGAATGATACGGAC. Primers were also modified to match clone 1a142 in order to amplify the same regions from these clones. Primers have 15 base pair 5′ ends with homology to the 1b09 sequence. Chimeras were then constructed using In-Fusion cloning (Clontech).

Figure 7. V538I/L546Q/V563T mutations in E2 confer additive sensitivity to NC1 mAbs. (A) Alignment of amino acids 526–569 of sensitive E1E2 clone 1b09 and resistant clones 1a129 and 1a142. Homology to the 1b09 amino acid sequence is indicated by a dot. Positions that differ between the sensitive and both resistant clones are highlighted in yellow. Arrows indicate contact residues for AR3C in the E2 core/AR3C crystal structure. (B) The dashed line indicates relative infection of HCVpp with chimeric E1E2 1b09/1a129-β in the presence of the mab AR3A, adjusted to 1. Each subsequent bar indicates the fold change in neutralization resistance after the indicated mutation(s) were introduced. Error bars indicate SD between duplicate wells. (C) Neutralization by serial dilutions of the indicated mAb of HCVpp with chimeric E1E2 1b09/1a129-β (pp1b09/1a129-β) or pp1b09/1a129-β after introduction of the indicated mutation(s). Error bars indicate SD between duplicate wells.
HCV NS5A immunostaining. Huh7.5.1 cells plated at 10,000 cells per well on a 96-well plate the previous day were incubated with viral supernatants overnight. 72 hours after infection, cells were fixed with 4% formaldehyde for 20 minutes and then stained for HCV NS5A using primary anti-NS5A antibody 9E10 (a gift of Charles Rice, The Rockefeller University, New York City, New York, USA) at 1:20,000 dilution in PBS, 3% bovine serum albumin, and 0.3% Triton X-100 for 1 hour at room temperature. Cells were washed twice with PBS and stained using secondary antibody Alexa Daylight 488–conjugated goat anti-mouse IgG (Life Technologies) at 1:1,000 dilution in PBS, 3% bovine serum albumin, and 0.3% Triton X-100 for 1 hour at room temperature. Cells were washed twice with PBS and stained using secondary antibody Alexa Daylight 488–conjugated goat anti-mouse IgG (Life Technologies) at 1:1,000 dilution in PBS, 3% bovine serum albumin, and 0.3% Triton X-100 for 1 hour at room temperature. Cells were washed twice with PBS and then stored covered in 100 μl PBS at 4°C. To determine peak infectivity, wells were examined with a Zeiss invert microscope with an Olympus camera. Viral supernatants with peak infectivity were kept for further experiments.

HCVcc neutralization assays. Huh7.5.1 cells were plated at 10,000 cells per well onto 96-well plates. On the following day, antibodies were serially diluted 2.5-fold in growth medium, starting from 50 μg/ml mAb. Three to six replicates were also performed with 50 μg/ml nonspecific IgG. Merimeric E1E2 clones were sequenced in their entirety after construction to confirm that no unintended mutations were introduced.

Production and neutralization of HCVcc. Human hepatoma Huh7.5.1 cells (a gift of Jake Liang, NIH, Bethesda, Maryland, USA) were maintained in DMEM supplemented with 10% fetal bovine serum, 1% sodium pyruvate, and 1% l-glutamate. HCVcc chimera H77/JFH1 (32) was a gift of Jens Bukh (Copenhagen University Hospital, Copenhagen, Denmark). 30 μg plasmid DNA was linearized using XbaI (New England Biolabs) and then gel purified using the QIAEX II Kit (Qiagen). Purified linear DNA was used for in vitro RNA transcription using the T7 RiboMAX Express Large Scale RNA Production System (Promega). A phenol-chloroform extraction was performed to purify RNA products that were then stored at –80°C.

Transfection of Huh 7.5.1 cells with HCVcc RNA was performed as previously described (29). Transfection supernatants were collected every 3 to 4 days and stored at –80°C for titering by HCV NSSA immunostaining.

Figure 8. I538V/Q546L/T563V mutations in E2 confer resistance to 6 NC1 mAbs by reducing mAb binding to E1E2. (A) The first bar in each graph indicates relative infection of wild-type pp1b09 in the presence of the indicated mAb, adjusted to 1. Subsequent bars indicate fold change in neutralization resistance of the indicated HCVpp relative to pp1b09. Values are the means of 2 to 6 independent experiments performed in duplicate, and error bars indicate SD. *P < 0.05, **P < 0.005 by t test. (B) Binding of mAbs CBH-2 and AR3C to E1E2 protein in an ELISA. For each mAb, binding to each of the E1E2 variants was compared at a single mAb concentration selected to produce binding within the linear range of the assay. Values are normalized for relative binding of control NC2 mAb HC33.4. Error bars indicate SD between duplicate wells. (C) Crystal structure from Kong et al. of E2 core with AR3C (34), from the Protein Data Bank, accession 4MWF, with E2 colors modified as in Figure 5. Residue 431 is purple, residue 442 is orange, and residues 538 and 563 are blue. The likely position of residue 546 is indicated with a blue asterisk. AR3C Fab is tan.

Figure 9. I538V/Q546L/T563V mutations in E2 confer resistance to neutralization by plasma from HCV-infected subjects. Introduction of I538V/Q546L/T563V into sensitive E1E2 clone 1b09 or back mutation of V538 to I, S546 to Q, and V563 to T in resistant clone 1a129 confers increased resistance or sensitivity, respectively, to neutralization by plasma samples from 18 HCV-infected subjects. Each box represents relative infection of the indicated HCVpp in the presence of each of 18 plasma samples. Dots indicate neutralization by plasma from the source donor of resistant E1E2 clone 1a129. Plasma was tested for neutralizing activity at a 1:100 dilution. Black bars indicate medians. Whiskers indicate 10th and 90th percentiles. P values were calculated by Wilcoxon signed-rank test.
Viral supernatants diluted to fall in a linear range of infectivity were added to antibody dilutions in triplicate for 1 hour at 37°C. Medium was removed from cells, and the antibody-virus mix added onto cells and incubated overnight. Antibody-virus mix replaced with fresh medium the following day. After 72 hours, medium was removed and cells were fixed and stained. A composite image of each well was created using an LSM 510 Meta confocal microscope. Images were analyzed using ImageJ to quantify the total area of fluorescence in each well. Neutralization was calculated by comparing the ratio of infection at each concentration of mAb to infection in the presence of nonspecific IgG or the most dilute mAb concentration, whichever had a higher mean value.

**HCV E1E2 ELISA.** mAb binding to E1E2 was quantitated using an ELISA as previously described (14). Briefly, 293T cells were transfected with one of the following E1E2 expression constructs: wild-type b09, b09 with 1538V/Q546L/T563V introduced by site-directed mutagenesis, wild-type a1a29, or a1a29 with V538I/L546Q/V563I introduced by site-directed mutagenesis. Cell lysates were harvested at 48 hours. Plates were coated with 500 ng Galan-thus nivalis lectin (Sigma-Aldrich) and blocked with PBS containing 0.5% Tween 20, 1% nonfat dry milk, and 1% goat serum, and E1E2-containing cell lysates were added. mAbs were assayed in duplicate at 5-fold serial dilutions, starting at 10 μg/ml, and binding was detected with HRP-conjugated anti-human IgG secondary antibody (BD Pharmingen no. 555788). For each mAb, a dilution was chosen that produced binding within the linear range of the assay (OD 0.1–3.5) for all variants, and binding to each E1E2 variant was compared at that mAb concentration. Binding of each mAb was normalized for protein expression by correcting for any difference in binding of control NC2 mAb HC33.4 between E1E2 variants.

**Hierarchical clustering of mAb patterns.** Effects on infectivity of the panel of HCVpp by mAbs were compared pairwise for all mAbs using the Spearman correlation. Spearman ρ and P values were then used as input for hierarchical clustering, as implemented in the “pvclust” package for R (http://cran.r-project.org/web/packages/pvclust/index.html) (42). Approximately unbiased values greater than 95 are considered strongly supported by the data. This clustering, depicted as a tree, was also used to order a matrix of correlation values produced using the “corrplot” package for R (http://cran.r-project.org/web/packages/corrplot/index.html) (43).

**Statistics.** Neutralization curves for HCVpp and HCVcc were fit by nonlinear regression in SigmaPlot, and mAb IC₅₀ was calculated based on that curve. Significance of the difference in proportion of positions with polymorphisms between amino acids 416 and 560 relative to the rest of the E1E2 sequence (Figure 3) was calculated using a z test. Comparison of relative infection (neutralization) of wild-type clones and site-directed mutants by mAbs (Figures 4 and 8) was performed using 1-sided t tests, since prior evidence supported increased resistance of mutant clones. Comparison of relative infection (neutralization) of wild-type clones and site-directed mutants by donor plasma (Figure 9) was performed using Wilcoxon signed-rank test. Comparison of infectivity (fitness) of wild-type and mutant clones (Supplemental Figure 10) was performed using 1-sided t tests. In all cases, a P value of less than 0.05 was considered significant.

**Study approval.** All protocols were approved by the Institutional Review Board of the Johns Hopkins Hospital, and informed consent was obtained from all study participants.

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