A Diverse Panel of Hepatitis C Virus Glycoproteins for Use in Vaccine Research Reveals Extremes of Monoclonal Antibody Neutralization Resistance

Richard A. Urbanowicz,a,b C. Patrick McClure,a,b Richard J. P. Brown,a,b† Theocharis Tsoleridis,a,b Mats A. A. Persson,c Thomas Krey,d,e William L. Irving,a,b Jonathan K. Ball,a,b Alexander W. Tarr,a,b

School of Life Sciencesa and NIHR Nottingham Digestive Diseases Biomedical Research Unit,b The University of Nottingham, Nottingham University Hospitals NHS Trust, Nottingham, United Kingdom; Karolinska Institutet, Department of Clinical Neurosciences, Center for Molecular Medicine, Karolinska University Hospital Solna, Stockholm, Sweden; Unité de Virologie Structurale, Dérépartement de Virologie, Institut Pasteur,d and CNRS UMR 3569,e Paris, France

ABSTRACT

Despite significant advances in the treatment of hepatitis C virus (HCV) infection, the need to develop preventative vaccines remains. Identification of the best vaccine candidates and evaluation of their performance in preclinical and clinical development will require appropriate neutralization assays utilizing diverse HCV isolates. We aimed to generate and characterize a panel of HCV E1E2 glycoproteins suitable for subsequent use in vaccine and therapeutic antibody testing. Full-length E1E2 clones were PCR amplified from patient-derived serum samples, cloned into an expression vector, and used to generate viral pseudoparticles (HCVpp). In addition, some of these clones were used to generate cell culture infectious (HCVcc) clones. The infectivity and neutralization sensitivity of these viruses were then determined. Bioinformatic and HCVpp infectivity screening of approximately 900 E1E2 clones resulted in the assembly of a panel of 78 functional E1E2 proteins representing distinct HCV genotypes and different stages of infection. These HCV glycoproteins differed markedly in their sensitivity to neutralizing antibodies. We used this panel to predict antibody efficacy against circulating HCV strains, highlighting the likely reason why some monoclonal antibodies failed in previous clinical trials. This study provides the first objective categorization of cross-genotype patient-derived HCV E1E2 clones according to their sensitivity to antibody neutralization. It has shown that HCV isolates have clearly distinguishable neutralization-sensitive, -resistant, or -intermediate phenotypes, which are independent of genotype. The panel provides a systematic means for characterization of the neutralizing response elicited by candidate vaccines and for defining the therapeutic potential of monoclonal antibodies.

IMPORTANCE

Hepatitis C virus (HCV) has a global burden of more than 170 million people, many of whom cannot attain the new, expensive, direct-acting antiviral therapies. A safe and effective vaccine that generates both T cell responses and neutralizing antibodies is required to eradicate the disease. Regions within the HCV surface glycoproteins E1 and E2 are essential for virus entry and are targets for neutralizing antibodies. Screening of vaccine candidates requires suitable panels of glycoproteins that represent the breadth of neutralization resistance. Use of a standard reference panel for vaccine studies will ensure comparability of data sets, as has become routine for HIV-1. Here, we describe a large panel of patient-derived HCV glycoproteins with an assessment of their neutralization sensitivity to defined monoclonal antibodies, which has enabled us to predict their likely efficacy in the wider HCV-infected population. The panel could also be important for future selection of additional therapeutic antibodies and for vaccine design.

The recent development of direct-acting antiviral therapies (DAA) able to potently inhibit hepatitis C virus (HCV) replication is a major milestone toward limiting the burden of the disease, but these expensive therapies are likely to remain unattainable by the majority of the 170 million people with persistent HCV infection. Eradication of the global burden of liver disease caused by HCV infections will require the introduction of a safe, effective vaccine. While the immune correlates of vaccine-induced protection are not completely understood, generation of both effective T cell responses (1) and neutralizing antibodies (2–7) is likely to be essential. One of the major challenges in successful HCV vaccine design is the extreme genetic diversity of HCV populations (8), which results from immune-driven adaptation and escape (9, 10).

The HCV surface glycoproteins E1 and E2 are the major targets of neutralizing antibodies (reviewed in reference 11). Regions within these proteins are essential to facilitate interactions with host cell receptors during entry (12–14). This conservation and their functional importance make them highly desirable targets for therapeutic antibodies and vaccines. However, these regions are thought to be shielded by hypervariable regions, which act as immunological decoys (15, 16) and are highly glycosylated (17).

Many neutralizing monoclonal antibodies (MAbs) have been isolated from infected humans (18–22) and experimentally immunized animals (23–26). The vast majority of broadly neutralizing monoclonal antibodies target epitopes that overlap sites involved in the interaction of E2 with host CD81 (21, 27),
blockading the entry cascade. Antibodies targeting other regions appear to have restricted reactivity and low neutralizing potency. An exception to this is the MAb AR4A, which recognizes a conserved neutralization epitope outside the CD81 binding region (28).

Experimental HCV glycoprotein vaccines have achieved varied levels of success (26, 29–32). Similarly, the performance of neutralizing monoclonal antibodies in clinical trials has been extremely variable (33, 34). Rational screening of lead therapeutic antibodies and vaccine candidates requires access to suitable panels of viral glycoproteins that represent the breadth of neutralization resistance. However, this has been hampered by the limited number of viruses or glycoproteins available for screening (30, 35). Provision and use of standard reference panels for vaccine and antibody studies will ensure comparability of data sets, as has become routine practice for HIV-1 (36). This will ensure that efforts can be focused on the most promising candidates and will prevent advancement of vaccines and treatments that have a high risk of failing against viruses circulating in HCV-infected populations.

Here, we describe the generation of a large panel of patient-derived HCV envelope glycoproteins from individuals at different stages of disease and infected with different genotypes, together with an assessment of their relative infectivities and neutralization sensitivity to defined monoclonal antibodies, which has enabled us to predict their likely efficacy in the wider HCV-infected population. The panel will also be important for future selection of additional therapeutic antibodies and for vaccine design (36).

**MATERIALS AND METHODS**

**Glycoprotein cloning and phylogenetic analysis.** cDNA sequences encoding full-length E1E2 were amplified from RNA extracted from patient sera and cloned into the pcDNA3.1 V5his D-TOPO expression vector (Life Technologies), as previously described (37). As controls, the E1E2 genes from the widely used HCV strains H77F, JFH1, and J6 were cloned into the same vector. Nucleotide sequences were determined by Sanger sequencing and aligned using Clustal W. Phylogenetic relationships were determined using maximum-likelihood analyses, as previously described (10). Bootstrap analysis with 500 replicates was performed to provide statistical support to the sequence clusters.

**Cell lines.** The human embryonic kidney 293T and Huh-7 human hepatoma cell lines were purchased from ECACC, and the Huh-7.5 cell line was obtained from Apath, LLC. All were grown in Dulbecco’s modified essential medium (DMEM) (Invitrogen) supplemented with 10% fetal bovine serum (FBS) and 0.1 mM nonessential amino acids (Invitrogen).

**Antibodies.** Anti-E2 MAbs 1/7, L1, XTLab68, and AP33 have been previously described (23, 34, 38), as has the anti-E2 nanobody (nB) D03 (26).

**HCVpp generation and infectivity and neutralization assays.** Cloned E1E2 genes were used to generate HCV pseudoparticles (HCVpp), essentially as described previously (37). Pseudoparticles generated in the absence of the E1E2 plasmid were used as a negative control. Core expression, for a subset of HCVpp production runs, was analyzed by Western blotting (35), and there were no discernible differences in expression levels (not shown). Infectivity assays were conducted as previously described (35). For neutralization assays, pseudotype virus was mixed with defined concentrations of antibody, incubated for 1 h at 37°C, and then added to Huh-7 cells. Cultures were incubated and read as for the infectivity assay. All assays were done in triplicate. The 50% inhibitory concentration ($IC_{50}$) titer was calculated as the MAb concentration that caused a 50% reduction in relative light units (RLU) compared to the level in the virus control wells after subtraction of cell control RLU. All the data were fitted using nonlinear regression plots with no constraint on the Hill slope (GraphPad Prism version 6.05).

**HCVcc generation and neutralization assay.** Genotype 2 (Gt2) cell-culture-infectious (HCVcc) clones were generated essentially as described previously, using restriction digest cloning and PCR (39). Neutralization assays were performed as previously described (26). The percent infection was determined by comparison to the number of cells infected in the absence of inhibitors, as determined by NNSA staining with MAB 9E10 (40).

**Statistical analysis.** Viruses were compared with respect to overall neutralization sensitivity by rank ordering based on mean $log_{10} IC_{50}$ titers across the five antibodies. A heat map describing the clustering patterns was generated using the heat map tool of the Los Alamos database (http://www.hiv.lanl.gov/content/sequence/HEATMAP/heatmap.html). The robustness of membership within a given cluster was evaluated by bootstrap resampling of the data. Frequency distribution plots fit a normal distribution curve, as shown by the $D'$Agostino and Pearson omnibus normality test (GraphPad Prism version 6.0). Highlighter plots were generated using the tool in the Los Alamos database (http://www.hiv.lanl.gov/content/sequence/HIGHLIGHT/highlighter_top.html).

**Nucleotide sequence accession numbers.** The E1E2 sequences described in this study have been deposited in GenBank under accession numbers KU285151 to KU285228.

**RESULTS**

Production of the HCV E1E2 glycoprotein panel. A total of 883 E1E2 clones were generated from 3,909 patients enrolled in the Trent HCV cohort. Of these, 493 clones from 63 different subjects were sequenced and identified as containing complete E1E2 open reading frames. These were then screened for infectivity using a murine leukemia virus (MLV)-based pseudoparticle assay. One hundred eighteen clones were classified as infectious (yielding a relative luminescence value at least 10-fold greater than background). Of these, 78 clones from 36 different patients were selected for use in the neutralization panel, of which 5 (14%) were sampled during the acute early phase and 31 during chronic infection (86%) (Table 1). In addition, the standard reference strains H77, JFH-1, and J6 were included. The clones were chosen to ensure that they represented the major HCV genotypes (Fig. 1A) and a range of infectivities (Fig. 1B), although glycoproteins that conferred very low levels of infectivity were omitted from subsequent analyses because of unacceptably high intra- and interassay variability compared to an isolate with higher infectivity. Invariably, these low-infectivity clones were relatively easy to neutralize and therefore deemed less clinically relevant than neutralization-resistant strains.

Estimated inhibitory concentrations of antibody are consistent between experimental runs. Due to the large number of clones included in the neutralization assays and the inherent variability of RLU values that could be observed between experimental runs, we compared $IC_{50}$ between two independent neutralization assays to allow us to evaluate interassay reproducibility. HCVpp supplemented with six clones previously shown to exhibit different neutralization resistance phenotypes were tested in two separate neutralization assays using the same CD81 binding site MAs that were to be used for the subsequent analyses of the full panel. There was good correlation between the two different runs for all of the antibodies tested (Fig. 2), irrespective of the genotype or the magnitude of the $IC_{50}$. These
FIG 1 E1E2 glycoprotein clones, representing the major HCV genotypes, show varying degrees of infectivity. Shown are maximum-likelihood phylogenetic analysis (A) and relative infectivity (B) of the 81 E1E2 clones used for the antibody neutralization panel. (A) The genetic distance for each of the branch lengths shown in the phylogenetic tree is indicated by the scale bar, and the level of bootstrap support (for those branches supported by >95% of replicates) is indicated above each branch. (B) HCVpp generated without a glycoprotein envelope reproducibly gave RLU values of less than 20, and therefore, a cutoff 200 was used to determine if the clone was infectious; only those clones defined as infectious are shown. The data are mean values of triplicates ± standard deviations (SD).

UKNP, United Kingdom Nottingham Panel, followed by the genotype, patient number, and isolate number.
The estimated inhibitory concentrations of antibodies are consistent between different neutralization assay runs. IC$_{50}$s for MAb AP33, MAb 1:7, MAb L1, nAb D03, and XTL68, estimated from two independent neutralization assays using HCVpp supplemented with E1E2 proteins representing the six major genotypes, were compared to assess the impact of interassay variability. The IC$_{50}$ titer for each repeat assay was plotted for the entire data set (All IC$_{50}$s) or for each of the 6 HCVpp. The solid lines show the best fit of the experimental data. Pearson correlation coefficient $R$ values are shown, along with the associated $P$ values.

**TABLE 1** Numbers of clones analyzed from each of the six major HCV genotypes

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Total no. of clones isolated</th>
<th>No. of clones screened$^a$</th>
<th>No. infectious clones$^a$</th>
<th>No. of infectious clones in the panel$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gt1</td>
<td>504</td>
<td>280 (32)</td>
<td>92 (23)</td>
<td>58 (23)</td>
</tr>
<tr>
<td>Gt2</td>
<td>85</td>
<td>19 (9)</td>
<td>8 (5)</td>
<td>6 (5)</td>
</tr>
<tr>
<td>Gt3</td>
<td>184</td>
<td>130 (12)</td>
<td>7 (3)</td>
<td>5 (2)</td>
</tr>
<tr>
<td>Gt4</td>
<td>35</td>
<td>19 (3)</td>
<td>6 (3)</td>
<td>5 (3)</td>
</tr>
<tr>
<td>Gt5</td>
<td>51</td>
<td>36 (6)</td>
<td>3 (2)</td>
<td>2 (2)</td>
</tr>
<tr>
<td>Gt6</td>
<td>24</td>
<td>9 (1)</td>
<td>2 (1)</td>
<td>2 (1)</td>
</tr>
<tr>
<td>Total</td>
<td>883</td>
<td>493 (63)</td>
<td>118 (37)</td>
<td>78 (36)</td>
</tr>
</tbody>
</table>

$^a$ Total number (number of unique subjects).
results show that our experimental approach was subject to low run-to-run variability.

**Patient-derived glycoprotein clones of the same subtype can differ markedly in their sensitivity to neutralization.** From the larger panel of 81 infectious clones, a subset of 5 genotype 1 clones, together with the reference molecular clone H77, were tested against four MAbs (AP33 [41], 1:7 [38], L1 [38], and XTL68 [34]) and one nanobody (nB D03 [26]), all of which target discrete epitopes located in the CD81 binding site. The IC<sub>50</sub>s are presented in Table 2, and the corresponding neutralization curves are presented in Fig. 3. Sensitivity to neutralization varied across the six genotype 1 isolates, with some being highly sensitive to the antibodies (e.g., UKNP1.2.3) while others were highly resistant (e.g., UKNP1.10.1). These data strongly support the view that sensitivity to a neutralizing antibody is determined at the isolate rather than the genotype level.

**Extensive variability in neutralization sensitivity of patient-derived envelope glycoproteins.** Neutralization assays were performed to determine the sensitivity of each of the 81 clones in the larger glycoprotein panel to neutralization by each of the five antibodies, and the means of the resulting IC<sub>50</sub>s for E1E2 HCVpp clones were calculated and plotted, together with their ranges, in mean rank order (Fig. 4). These mean IC<sub>50</sub>s constituted a spectrum. The inset shows that when the isolates are split into chronic and acute early phase samples, there is no significant difference in the log<sub>10</sub> IC<sub>50</sub>s between the groups. The most easily neutralized E1E2 clone (UKNP1.21.2) had a mean IC<sub>50</sub> of 0.013 μg ml<sup>-1</sup>, whereas the most resistant clone (UKNP2.4.1) had a mean IC<sub>50</sub> of 49 μg ml<sup>-1</sup>. When the neutralization sensitivities of the clones were analyzed according to individual antibodies (Fig. 5), their relative positions in the neutralization spectrum were on the whole maintained, indicating an inherent sensitivity to neutralization by CD81 binding site-targeting antibodies. However, a few clones had greater sensitivity to some antibodies than their relative positions in the plot of mean IC<sub>50</sub>s would suggest. For example, UKNP1.16.2 was very susceptible to neutralization by MAb L1.

### Table 2 IC<sub>50</sub>s of neutralization curves in Fig. 3

<table>
<thead>
<tr>
<th>Clone</th>
<th>MAb AP33</th>
<th>MAb 1:7</th>
<th>MAb L1</th>
<th>nB D03</th>
<th>MAb XTL68</th>
</tr>
</thead>
<tbody>
<tr>
<td>H77.20</td>
<td>0.1012</td>
<td>0.5388</td>
<td>2.031</td>
<td>9.167</td>
<td>0.6316</td>
</tr>
<tr>
<td>UKNP1.2.3</td>
<td>0.01467</td>
<td>0.009883</td>
<td>0.03123</td>
<td>0.02079</td>
<td>0.03315</td>
</tr>
<tr>
<td>UKNP1.3.2</td>
<td>0.1294</td>
<td>0.0553</td>
<td>0.06801</td>
<td>0.1913</td>
<td>0.05287</td>
</tr>
<tr>
<td>UKNP1.4.1</td>
<td>0.2245</td>
<td>0.8219</td>
<td>0.1371</td>
<td>0.2545</td>
<td>1.27</td>
</tr>
<tr>
<td>UKNP1.10.1</td>
<td>2.118</td>
<td>11.88</td>
<td>15.47</td>
<td>5.091</td>
<td>44.02</td>
</tr>
<tr>
<td>UKNP1.11.7</td>
<td>0.1408</td>
<td>0.1022</td>
<td>0.99</td>
<td>0.3288</td>
<td>0.1833</td>
</tr>
</tbody>
</table>

*The IC<sub>50</sub> for each genotype 1 HCVpp/antibody curve was calculated using nonlinear regression plots with no constraint on the Hill slope (GraphPad Prism version 6.05).*

**FIG 3** Genotype 1 clones show both resistant and sensitive phenotypes. HCVpp supplemented with genotype 1 E1E2 proteins H77.20, UKNP1.2.3, UKNP1.3.2, UKNP1.4.1, UKNP1.10.1, and UKNP1.11.7 were neutralized by increasing concentrations of MAb AP33, MAb 1:7, MAb L1, nB D03, and MAb XTL68, and the resulting infectivity was plotted against the antibody concentration.
(Fig. 5), while UKNP1.12.1 was very susceptible to neutralization by MAb XTL68 (Fig. 5). No neutralization was observed when antibodies were tested against a negative control (vesicular stomatitis virus G [VSV-G]) or when pseudoparticles were assayed using anti-tetanus toxin MAb (Wyeth) (data not shown). The mean $\text{IC}_{50}$ (standard deviations) for MAbs AP33, 1:7, L1, D03, and XTL68 were 0.6872 (1.587), 2.105 (5.909), 4.581 (14.59), 2.289 (4.600), and 8.887 (24.69), respectively.

The neutralization phenotypes of patient E1E2 proteins are consistent in both HCVpp and HCVcc models of infection. To determine if the same pattern of neutralization sensitivity also applied to the cell culture model, two Gt2 E1E2 clones from the panel were transferred into the J6/JFH-1 cell-culture-infectious HCV backbone (Fig. 4, arrows). These clones and three reference strains, JFH-1, J6/JFH-1, and H77/JFH-1, represented a breadth of neutralization sensitivities, as predicted from the HCVpp panel, and were tested against MAb 1:7 (Fig. 6, left, and Table 3). Even with this small subset, there was a significant correlation ($r = 0.8938; P = 0.0152$) between the neutralization data obtained in the HCVpp and HCVcc systems (Fig. 6, right).

Increased infectivity conferred by E1E2 clones is associated with increased neutralization resistance. To investigate the relationship between infectivity and neutralization sensitivity, the mean $\log_{10} \text{IC}_{50}$ were plotted against the relative infectivity (as defined by the RLU value) of each of the E1E2 clones (Fig. 7). There was a direct correlation between increased neutralization resistance, as indicated by increasing $\text{IC}_{50}$, and increased infectivity, and this correlation was statistically significant ($r = 0.8513; P < 0.0001$). This trend was also maintained when the analysis was performed using individual MAb $\text{IC}_{50}$ data (not shown). E1E2 clone UKNP2.4.1 had the highest level of infectivity, and it was also the most resistant to antibody neutralization.

The glycoprotein panel can be used to predict the likely efficacy of monoclonal antibodies against circulating HCV strains. An important aim of this study was to determine whether the glycoprotein panel could be used to inform future vaccine and therapeutic antibody efficacy by predicting likely neutralizing potencies against a wider set of patient-derived isolates. Plots of $\log_{10} \text{IC}_{50}$ s for each antibody showed a normal distribution (Fig. 8), meaning that each antibody plot could be used to predict an antibody concentration that would exceed the $\text{IC}_{50}$ for 95% (equal to the mean of the neutralization distribution plus 2 standard deviations) or 99% (mean plus 3 standard deviations) of HCV isolates (Table 4). For MAb AP33, a concentration of 8 $\mu$g ml$^{-1}$ would be sufficient to meet the $\text{IC}_{50}$ of 95% of circulating HCV strains, but this would need to be increased to 59 $\mu$g ml$^{-1}$ in order to effectively neutralize 99% of circulating isolates. In comparison, concentrations of XTL68 of 117 $\mu$g ml$^{-1}$ and 1.51 mg ml$^{-1}$ would be needed to provide the same level of neutralization for 95% and 99% of circulating HCV isolates, respectively.

Patient-isolated E1E2 can be categorized into distinct neutralization phenotype clusters. In order to categorize the E1E2 panel into groups based on similar neutralization phenotypes, we next performed hierarchical cluster analysis of the $\text{IC}_{50}$ titers for
Pseudotype viruses incorporating different E1E2 clones derived from the major HCV genotypes display diverse susceptibilities to antibody neutralization. Eighty-one distinct patient-derived and reference E1E2 clones derived from genotypes 1, 2, 3, 4, 5, and 6 were assessed for their neutralization sensitivity to five monoclonal antibodies. The log_{10} IC_{50} for MAbs AP33, 1:7, L1, and XTL68 and nB D03 are plotted according to the E1E2 clone’s rank order presented in Fig. 4.
TABLE 3 Comparison of neutralization IC\textsubscript{50}s of patient-isolated clones in both the HCV\textsubscript{pp} and HCV\textsubscript{cc} infection models

<table>
<thead>
<tr>
<th>HCV\textsubscript{cc} clone</th>
<th>Neutralization rank\textsuperscript{a}</th>
<th>IC\textsubscript{50} (μg ml\textsuperscript{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HCV\textsubscript{pp} 1:7</td>
<td>HCV\textsubscript{cc} 1:7</td>
</tr>
<tr>
<td>JFH-1</td>
<td>32</td>
<td>0.06</td>
</tr>
<tr>
<td>H77.20</td>
<td>60</td>
<td>0.55</td>
</tr>
<tr>
<td>UKNP2.5.1</td>
<td>65</td>
<td>1.47</td>
</tr>
<tr>
<td>J6</td>
<td>71</td>
<td>0.82</td>
</tr>
<tr>
<td>UKNP2.4.1</td>
<td>81</td>
<td>19.11</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Position in the ranking of the mean IC\textsubscript{50} HCV\textsubscript{pp} neutralization values presented in Fig. 4.

DISCUSSION

Future HCV vaccines or antibody-based therapies will need to be effective against the majority, if not all, of the viruses present in infected individuals. This requires good preclinical screening of interventions to identify the best leads, which in turn relies upon the availability of reference panels of patient-derived HCV isolates. While some groups have reported successful culture of patient-derived HCV, these methods generally exhibit poor replication efficiency and therefore do not represent a tractable means of predicting neutralization sensitivity \textit{in vivo}; instead, surrogate models, such as the HCV\textsubscript{pp} system or chimeric HCV\textsubscript{cc}, have proven to be more robust tools (reviewed in reference 42). Previous studies on antibody neutralization have used the cell line Huh7.5 (40) or primary hepatocytes (43); we chose the former, as that cell line is more amenable and the HCV entry process is comparable to that observed in primary hepatocytes (44–46).

We had previously shown that patient-derived E1E2 clones were differentially susceptible to neutralization by human polyclonal sera, and this appeared to be unrelated to the genotype (35). Here, we extend those preliminary studies by generating a significantly larger panel of E1E2 HCV\textsubscript{pp} clones, drawn from different genotypes that are representative of individuals at different stages of infection. This marks a major advancement in the field, where existing knowledge of protective antibody determinants has been derived from much smaller or genotype-restricted HCV\textsubscript{pp} or HCV\textsubscript{cc} panels (28, 39, 47, 48). Importantly, our data highlight the fact that intra- or intergenotypic differences do not noticeably impact sensitivity to neutralization (at least not with respect to CD81 binding site-targeting MAbs) and that this property is determined at the isolate level. In our initial screening, we identified a number of clones that were not functional in the HCV\textsubscript{pp} system. This has been observed previously (12), and the reasons for this lack of functionality remain unclear.

One of the most striking observations was the extreme variability in neutralization sensitivity that patient-derived E1E2 clones demonstrated. While many E1E2 clones showed similar degrees of sensitivity to CD81 binding site-targeting MAbs, there were a number of isolates that were especially sensitive or resistant. Comparisons between resistant and sensitive strains failed to identify common sequence substitutions or motifs giving rise to this difference. Identifying mechanisms of resistance will be important in understanding neutralizing antibody responses, although it is likely that sequence differences underlying these different neutralization phenotypes will be highly isolate specific.

While our panel contained predominantly subtype 1a and
1b viruses, there was a sufficiently large number of clones representing the other major genotypes to be confident that the panel could provide a robust prediction of the neutralizing potency of individual MAbs against HCV strains present in infected individuals. For example, the panel has enabled us to estimate that 8 µg ml\(^{-1}\) of MAb AP33, which equals the mean IC\(_{50}\) plus 2 standard deviations, will provide at least 50% neutralization for \(\sim 95\%\) of patient-derived isolates, whereas for XTL68, this would increase to 117 µg ml\(^{-1}\). Importantly, this type of screening strategy provides insight into the likely potency of any given MAb when used against any patient-derived isolate. Given the observed correlation between in vitro and in vivo anti-HCV activities of MAbs (49, 50), this approach could be used to provide a quick indication of an antibody’s potential clinical value. In this respect, it is important to note that in a previous clinical trial, MAb XTL68 had only modest effect when administered to HCV-infected individuals (34). It is also important to note that other factors, such as the potential for MAb-driven escape (51), are also important parameters in determining the likely clinical value of any particular antibody or antibody combination. Finally, resistance to MAb neutralization was correlated with the degree of infectivity, as determined by the HCVpp assay. Such a relationship between increased antibody neutralization resistance and improved HCVpp infectivity has been reported previously in analyses of polyclonal sera obtained from a small cohort of liver transplant recipients (43).

Having established that isolates showed a wide range of neutralization sensitivities, we performed cluster analysis to define clusters of MAb sensitivity. The bulk of isolates showed variable patterns of neutralization—easily neutralized by some but not all MAbs tested. However, there were two distinct clusters that represented the two extremes, one highly sensitive and one highly resistant group. Both clusters contained viruses drawn from highly divergent genotypes. The mean IC\(_{50}\) for all five antibodies, of the resistant group was 290 times greater than that observed for the sensitive group. This was also true for the antibodies individually. The resistant viruses pose a major challenge for vaccine development and cannot be ignored; otherwise, deployment of a less than optimal vaccine would result in the selection and eventual dissemination of a virus population refractive to neutralization, as has been observed in the most promising HIV-1 vaccine trial to date (52).

Importantly, two strains (H77c and JFH-1) that have been used extensively to test the potency of candidate vaccine-induced sera fall outside the neutralization-resistant cluster. In contrast, the J6 reference strain is a more relevant reference strain to include in this type of analysis, which probably explains the variable neutralization observed for E1E2-elicited immune sera (29). This is consistent with the antibody neutralization resistance of this clone that was evident, but not commented upon, in previous studies (29, 53).

HCVpp represents a rapid, amenable way to test antibody neutralization against a large number of different isolates. Importantly, we have shown, at least for a subset of isolates, that the HCVpp-derived neutralization data accords with those obtained using chimeric HCVcc. While significant advances have been made in generating more diverse panels of infectious virus, it is still a very laborious process, so the range of isolates available is still very limited (28, 39). Although the HCVpp system provides greater flexibility with respect to the number of HCV isolates that can be tested easily, our data show that while there was good correlation between the two systems, HCVpp were more easily neutralized than the corresponding HCVcc chimeras. Therefore, while HCVpp is an easy system for rapid screening of MAbs and sera, the findings would need to be validated in the HCVcc system. Also, HCVcc chimeras utilizing more clinically relevant isolates are required for in vivo studies. Therefore, creation of chimeras of the clones representing resistant-cluster isolates is under way.

Our study focused on an initial characterization of the panel, using MAbs that targeted the CD81 binding site. These MAbs represent the majority of murine and human cross-neutralizing anti-HCV MAbs. Differences in sensitivity to antibodies and polyclonal sera targeting other neutralizing determinants will be important to assess in the future. However, it is important to reiterate...
that in our previous analysis of a much smaller panel using polyclonal sera, a similar spectrum of neutralization was observed (35).

In conclusion, this study provides the first objective categorization of cross-genotype patient-derived HCV E1E2 clones according to their sensitivity to antibody neutralization and showed that individual HCV isolates can be categorized as having a clearly distinguishable neutralization-sensitive, -resistant, or -intermediate phenotype. We were able to use the panel to predict the likely efficacies of a number of CD81 binding site MAbs against circulating HCV strains. This reference panel will be indispensable for future studies of the therapeutic potential of existing and newly-derived HCV isolates.

![Figure 8](image_url)

**TABLE 4** IC50s from frequency distribution plots in Fig. 5

<table>
<thead>
<tr>
<th>Antibody</th>
<th>IC50 (μg ml⁻¹) 95%</th>
<th>IC50 (μg ml⁻¹) 99%</th>
</tr>
</thead>
<tbody>
<tr>
<td>AP33</td>
<td>8</td>
<td>59</td>
</tr>
<tr>
<td>1:7</td>
<td>22</td>
<td>196</td>
</tr>
<tr>
<td>L1</td>
<td>54</td>
<td>662</td>
</tr>
<tr>
<td>D03</td>
<td>32</td>
<td>293</td>
</tr>
<tr>
<td>XTL68</td>
<td>117</td>
<td>1,517</td>
</tr>
</tbody>
</table>

IC50s were estimated from the frequency distribution plots based on the mean values plus 2 standard deviations (95%) or mean values plus 3 standard deviations (99%).
In hierarchical cluster analysis, the E1E2 clones can be grouped according to their sensitivity to monoclonal antibody neutralization. HCV E1E2 pseudoviruses \((n = 81)\) were assessed for neutralization sensitivities using five MAbs targeting discrete epitopes of the CD81 binding site. Individual clones are listed on the right, and antibodies are indicated at the base of the heat map. The magnitude of neutralization \((\log_{10} IC_{50} \text{ titer})\) is denoted by the color, where higher values of neutralization are represented by lighter colors (e.g., light yellows) and lower values are represented by more saturated, dark colors (e.g., dark red). Boxes are drawn around viral isolates that were resistant (blue) and sensitive (black) to neutralization that grouped with >90% probability according to bootstrap resampling (10,000 replicates) of the data set.

### TABLE 5 Mean IC\textsubscript{50}s for the resistant and sensitive neutralization clusters for all antibodies and individual antibodies\(^a\)

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Combined</th>
<th>AP33</th>
<th>I:7</th>
<th>L1</th>
<th>D03</th>
<th>XTL68</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitive</td>
<td>0.05468 (0.03714)</td>
<td>0.0136 (0.01475)</td>
<td>0.06553 (0.07598)</td>
<td>0.0675 (0.1175)</td>
<td>0.02213 (0.0177)</td>
<td>0.1046 (0.1257)</td>
</tr>
<tr>
<td>Resistant</td>
<td>15.9 (14.74)</td>
<td>2.541 (2.903)</td>
<td>8.545 (11.11)</td>
<td>20.39 (28.23)</td>
<td>7.501 (7.053)</td>
<td>40.52 (43.51)</td>
</tr>
</tbody>
</table>

\(^a\)Numbers in parentheses are standard deviations of the means.
discovered MAbs, used alone or in combination, as well as the potency of immune-generated sera.

ACKNOWLEDGMENTS

We thank Takaji Wakita for plasmid pJFH-1; Charles Rice for the Huh7.5 cell line, MAb 9E10, and the H77/JFH-1 chimera; and Francois Loïc Costet for plasmid phCMV.

R.A.U., A.W.T., and J.K.B. designed the research; R.A.U., C.P.M., R.J.P.B., T.T., and A.W.T. performed the research; M.A.A.P., T.K., and W.L.I. contributed reagents; R.A.U., A.W.T., and J.K.B. wrote the paper; and we all edited and approved the final paper.

The human antibodies L1 and 1:7 are protected in patents owned by Molecules of Man AB, a spinoff company based on discoveries made at Karolinska Institutet, Sweden; M.A.A.P. has a financial interest in the company as a shareholder. We have no other conflicts to declare.

FUNDING INFORMATION

Medical Research Council (MRC) provided funding to Richard A. Urbanowicz and Jonathan K. Ball under grant number G0801169. Seventh Framework Programme (FP7) provided funding to Richard A. Urbanowicz and Jonathan K. Ball under grant number 305600.

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

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