Predicting spontaneous clearance of acute hepatitis C virus in a large cohort of HIV-1-infected men

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ABSTRACT

Objective An epidemic of acute hepatitis C virus (HCV) infection in HIV-positive men-who-have-sex-with-men (MSM) is emerging in Europe, Australia and the USA. The aim of this study was to characterise the natural history of primary HCV in this setting and to assess host and viral factors which predict spontaneous clearance.

Methods This prospective longitudinal cohort study was carried out in 112 HIV-positive patients who were followed in a single centre (the St Mary's Acute HCV Cohort). Plasma and peripheral blood mononuclear cells (PBMCs) were obtained at monthly intervals for 3 months and at 3-monthly intervals thereafter for a median of 45 months (IQR=29–68 months). The primary end point was spontaneous clearance of HCV. Cox regression was used to assess the impact of clinical and virological variables on outcome, including liver function, CD4 count, rate of HCV RNA decline, T cell response and clonal sequence evolution within the HCV E2 envelope gene.

Results 15% of patients cleared HCV spontaneously, while 85% progressed towards chronicity. The latter group included a significant proportion of ‘fluctuating’ progressors (37.5%), in whom a fall followed by a rise in viral load measurements is useful in decisions regarding the necessity and timing of treatment. Exposure to the virus is usually unnoticed by patients until they develop cirrhosis of the liver or hepatocellular carcinoma following many years of silent infection. Thus, while large cohorts of patients with chronic hepatitis C virus (HCV) genome, the asymptomatic nature of early infection has meant that its natural history has only been partially described. Exposure to the virus is usually unnoticed by patients until they develop cirrhosis of the liver or hepatocellular carcinoma following many years of silent infection. Twenty-one years after the discovery of the hepatitis C virus (HCV) genome, the asymptomatic nature of early infection has meant that its natural history has only been partially described. Exposure to the virus is usually unnoticed by patients until they develop cirrhosis of the liver or hepatocellular carcinoma following many years of silent infection. Thus, while large cohorts of patients with chronic infection have been extensively described, insights into early host–virus dynamics have been less well characterised. Follow-up of point-source outbreaks—for example, nosocomial infection in hospitals and...
rhesus-negative women who received contaminated immunoglobulin during pregnancy—have provided valuable insights into transmission risks and the likelihood of spontaneous clearance.1,2 One large multicentre community-based study of patients mainly infected by iatrogenic exposure or intravenous drug use revealed that viral clearance usually occurs within 3 months.3 This was recently observed in another longitudinal study of intravenous drug users in whom female sex was also associated with spontaneous clearance.4

Very little is known about early HCV in HIV-infected individuals, in whom HCV is emerging as a major concern, with liver disease now a major cause of mortality in patients with access to highly active antiretroviral therapy.5 Globally, 4–5 million people are infected with both viruses, due to shared routes of transmission.6 An emerging epidemic of acute HCV infection in HIV-positive men in urban centres in Europe,8 9 Australia10 and the USA11 has provided a rare opportunity to study the progression of early disease prospectively in a large number of individuals. Patients are identified early in infection, despite lack of symptoms, due to the detection of abnormal liver function tests at routine clinic follow-up.

The majority of patients infected acutely with HCV progress to chronic infection (55–90%). A number of mechanisms may be responsible for the persistence of virus in these individuals. The virus has an extremely rapid generation time—producing on average, a trillion viral particles every day. Furthermore, the viral genome is copied imperfectly due to the lack of proof-reading function of the viral RNA polymerase.12 The result is the production of a swarm of related, but genetically different viral sequences that circulate at any given time; the quasispecies. The ability of the host immune system to recognise and eliminate each variant strain is highly likely to influence the final outcome. In a small, well-characterised cohort (n=12) of HIV-negative individuals, it has been shown that those who progress to chronicity have higher viral diversity than those who clear the virus spontaneously.13 Evolution of the hypervariable region-1 (HVR-1) of the viral envelope-2 (E2) protein, which contains both B and T cell epitopes, is associated with progressive disease, while a narrowed quasispecies repertoire occurs in patients who subsequently spontaneously clear infection. Spontaneous clearance (SC) of virus in HIV-negative individuals is accompanied by the emergence of strong multispecific CD8+ and CD4+ cell responses targeting a broad range of viral epitopes.14

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The majority of patients infected acutely with HCV progress to chronic infection (55–90%). A number of mechanisms may be responsible for the persistence of virus in these individuals. The virus has an extremely rapid generation time—producing on average, a trillion viral particles every day. Furthermore, the viral genome is copied imperfectly due to the lack of proof-reading function of the viral RNA polymerase.12 The result is the production of a swarm of related, but genetically different viral sequences that circulate at any given time; the quasispecies. The ability of the host immune system to recognise and eliminate each variant strain is highly likely to influence the final outcome. In a small, well-characterised cohort (n=12) of HIV-negative individuals, it has been shown that those who progress to chronicity have higher viral diversity than those who clear the virus spontaneously.13 Evolution of the hypervariable region-1 (HVR-1) of the viral envelope-2 (E2) protein, which contains both B and T cell epitopes, is associated with progressive disease, while a narrowed quasispecies repertoire occurs in patients who subsequently spontaneously clear infection. Spontaneous clearance (SC) of virus in HIV-negative individuals is accompanied by the emergence of strong multispecific CD8+ and CD4+ cell responses targeting a broad range of viral epitopes.14

In this study, a cohort of 112 prospectively followed HIV-positive patients infected with acute HCV was used to characterise clinical patterns of infection and identify parameters predictive of SC, based on host factors such as CD4 count, liver function and sequential viral load (VL) sampling. Longitudinal genetic sequence analysis and T cell function determination was carried out in 50 patients (10 clearers and 40 progressors) at multiple time points. We hypothesised that rapid diversification of virus would be expected in patients who progress to chronicity, while SC of disease would be predicted by limited quasispecies diversity accompanied by strong T cell responses.

MATERIALS AND METHODS

Patient cohort

Informed consent in writing was obtained from each patient, the study protocol conformed to the ethical guidelines of the 1975 Declaration of Helsinki and ethical approval was granted by the Riverside Research Ethics Committee.

A cohort of 112 patients with HIV and acute HCV (102 with chronic HIV/acute HCV and 10 with acute HIV/acute HCV co-infection) were prospectively recruited from 2005 to 2009 in a single centre (the St Mary’s Acute Hepatitis C Cohort). Plasma and peripheral blood mononuclear cell (PBMC) samples were obtained prospectively at monthly intervals for 3 months and at 3-monthly intervals thereafter for a median of 3 years. Stored frozen samples (HIV VL samples obtained every 3 months between 1994 and 2009) were also tested retrospectively by HCV reverse transcription–PCR (RT–PCR) in order to date the length of HCV infection. The date of infection was calculated as the midpoint between the last negative PCR or antibody test and first positive HCV PCR.

Patients were offered 48 weeks of treatment with pegylated interferon α and weight-based ribavirin (800–1200 mg) 12 weeks following HCV diagnosis unless this was contraindicated or SC occurred.

Case definition and end points

Acute HCV diagnosis was defined by a positive HCV RNA RT–PCR test within 6 months of a preceding negative RT–PCR or antibody test.

The primary end point of the study was SC of HCV, defined as two successive negative RT–PCR tests (<12 IU/ml) 3 months apart.

Sustained virological response (SVR) was defined as HCV RNA negativity (<12 IU/ml) 24 weeks after the end of 48 weeks of treatment with pegylated interferon α and ribavirin.

VL dynamics, liver function, viral diversity following clonal analysis and T cell responses were also assessed at multiple time points. VL observations were censored at the time of treatment.

Amplification of E2 HVR-1 and NS5B

Viral RNA was extracted using the QIAamp Viral RNA kit (Qiagen, Crawley, UK). Amplification of E2 HVR-1 and NS5B was performed by nested RT–PCR using a combination of geno-type-specific primers16 17 and a One-Step RT–PCR kit (Qiagen). The HVR-1 is the most heterogeneous section of the HCV genome and can therefore be used to determine intragenotype differences in HCV strains.20 Purified PCR products were cloned into TOPO-4 vector (Invitrogen, Paisley, UK) and, following bacterial transformation, 20 colonies per plate were selected and used for plasmid DNA preparation and sequencing.

Sequence analyses

Sequences were aligned using Clustal W. Phylogenetic trees were constructed with MEGA 4.0 using the Neighbor–Joining method and Kimura two-parameter distance for all substitutions, and the inferred phylogenies were tested with 1000 bootstrap replications. Genetic diversity was calculated using the Hamming distance, which is defined as the number of amino acid differences between two sequences using the formula (1–s)×100, where s is the fraction of shared sites in two aligned nucleotide sequences. Genetic distance was calculated as the mean percentage difference between all samples at any time point.

Unique strains were defined as those originating from a distinct branch on the phylogenetic tree with a bootstrap value of >70 and a >10% genetic distance when compared with other strains derived from the same patient. Superinfection was defined as evidence of unique strains not present at baseline sampling, but evident at a later time point. Mean intrastrain...
corrected genetic distance and Hamming distances were calculated for all patients.

Non-synonymous substitutions per non-synonymous site divided by synonymous substitutions per synonymous site (dN/dS) ratios were calculated using MEGA 4.0. dN/dS values >1 were considered as evidence of positive selection.

**Genetic sequence data**

All viral sequences have been submitted to EMBL (accession numbers: HQ616913-HQ619203).

**ELISpot assays**

PBMCs were separated from whole blood by density gradient centrifugation and stored in liquid nitrogen. Thawed PBMCs were added to 96-well plates coated with mouse antihuman interferon \( \gamma \) (IFN\( \gamma \)) antibody. Peptides were added and the cells were incubated overnight. These were then washed off and mouse antihuman-IFN\( \gamma \) monoclonal antibody and alkaline phosphatase-conjugated antibiotin monoclonal antibody were added followed by incubation with 5-bromo-4-chloro-indolyl-phosphate-nitroblue-tetrazolium. Spot-forming units (SFUs) were counted on an ELISpot reader. Further details are given in Appendix 1 available online.

**Error rates**

In order to exclude contamination error as the mechanism for superinfection events, multiple time points were assessed and amplification of NS5B was carried out at different time points. In order to distinguish superinfection from co-infection with varying dominance of virus, PCR amplification was carried out using strain-specific primers. Further methodology details, including measures taken to prevent error, are available in Appendix 2 online.

**Statistical analyses**

As appropriate, Cox regression survival analysis, Kaplan–Meier survival plots, Mann–Whitney, \( \chi^2 \) and Fisher exact tests were used to analyse the significance of observations using the statistical computer package STATA 10.0. VL observations were right-censored at the time of treatment or end of follow-up. HRs were calculated with 95% CIs. \( p \) Values of ≤0.05 were considered to be statistically significant.

**RESULTS**

**Cohort characteristics**

A total of 112 patients were recruited to the study and followed-up for a median of 45 months (IQR=29–69 months; figure 1). Sequential VL measurements revealed three patterns of infection (figure 2); progression to chronicity with an initial peak viraemia and chronic plateau viraemia (PV) phase (47.5%), fluctuating viraemia (FV) defined as a fall followed by a rise in VL of >1 log\(_{10} \) IU/ml (37.5%); and SC (15%). Demographic and clinical parameters are shown in table 1.

**Sequential bilirubin and alanine aminotransferase (ALT) measurements and outcome**

Early infection in progressors (PV and FV groups) and spontaneous clearers (SC) was associated with a rise in ALT and bilirubin. The peak SC bilirubin and ALT levels were significantly higher than in the progressor groups (table 1; figure 2).

**Viral dynamics and outcome**

SC was associated with an initial increase followed by a rapid fall in VL (2.2 log\(_{10} \) drop within 100 days and 3.46 log\(_{10} \) drop within 200 days; table 2) leading to clearance at a median of 108 days (95% CI 58 to 170 days). Thirteen FV patients became transiently PCR negative, but took on average 154 days to reach negativity (95% CI=113, 323 days; p=0.04). Of the 55 patients who achieved a 2.2 log\(_{10} \) drop in VL, 17 (49%) spontaneously cleared infection and 38 (51%) did not. Those who spontaneously cleared infection took a median of 45 versus 120 days to achieve a 2.2 log\(_{10} \) drop in VL (p=0.01).

**Genetic sequence and phylogenetic analyses**

Phylogenetic analysis of majority quasispecies compared with reference sequences revealed several monophyletic clusters of
infection within genotype 1a (77%), 4d (15%), 5a (7%) and 1b (4%), indicating several introductions of the virus into the MSM population (Appendix 3 online). Six per cent were infected with two genotypes and 2% had three genotypes (1a, 3a and 4d). SC occurred only in patients with genotype 1a infection, and this was significantly associated with outcome (p = 0.04).

In order to determine which phylogenetic factors predicted SC, Hamming distance (diversity), number of quasispecies and dN/dS ratios were calculated in 50 patients (40 progressors and 10 clearers) at multiple (median = 3) time points (table 3).

Initially, genetic diversity was noted to vary considerably over time in all groups (Appendix 4 online) but did not correlate with VL. In order to investigate if this variation was the result of superinfection events, Neighbor–Joining phylogenetic analyses of all longitudinally acquired E2 HVR-1 sequences (n = 2291) were next carried out, revealing three groups of patients (figure 5); those who spontaneously cleared infection (n = 10; 20%), those who progressed to chronicity with increasing quasispecies diversity but no evidence of superinfection (n = 20; 40%); and those who progressed to chronicity with evidence of superinfection events (n = 20; 40%; figure 5). A number of supplementary analyses were carried out to confirm these results (Appendix 2 online).

Mean intrastrain calculations of genetic diversity and dN/dS ratios were used for patients found to have multiple strains of infection. Following this correction, diversity and dN/dS ratios increased gradually with time in progressors, but decreased in clearers (table 3, figure 4).

**T cell responses**

IFNγ production in response to pooled peptides spanning the entire viral genome were measured by ELISpot at two time...
points in 40 patients with genotype 1a infection. Total responses at baseline were significantly higher in SC versus PV (415 versus 118 SFU/million cells; p = 0.006) and FV (415 versus 50 SFU/million cells; p = 0.0001) patients while responses were not significantly different by 3 months. In FV patients, transient viral control was associated with strong T cell responses. While viral control was associated with strong T cell responses, while viral 6 months. In FV patients, transient viral control was associated with strong T cell responses. While viral control was associated with strong T cell responses, while viral in VL.

Table 1 Clinical measures and outcome (spontaneous clearance vs progression)

<table>
<thead>
<tr>
<th>Clinical measure (units, normal range)</th>
<th>Spontaneous clearance (median, 95% CI)</th>
<th>Progression Plateau viraemia (median, 95% CI)</th>
<th>Fluctuating viraemia (median, 95% CI)</th>
<th>Range</th>
<th>HR (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>17 (15%)</td>
<td>53 (47.5%)</td>
<td>42 (37.5%)</td>
<td></td>
<td>1.00 (0.94 to 1.06)</td>
<td>1.00</td>
</tr>
<tr>
<td>Age (years)</td>
<td>37 (32 to 44)</td>
<td>39 (37 to 42)</td>
<td>39 (37 to 40)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline HCV VL log10 (IU/ml)</td>
<td>6.11 (1.79 to 7.35)</td>
<td>6.25 (5.71 to 6.44)</td>
<td>5.89 (5.06 to 6.13)</td>
<td></td>
<td>0.75 (0.55 to 1.01)</td>
<td>0.06</td>
</tr>
<tr>
<td>Peak HCV VL log10 (IU/ml)</td>
<td>6.11 (1.79 to 7.35)</td>
<td>6.52 (6.35 to 6.88)</td>
<td>6.21 (5.87 to 6.72)</td>
<td></td>
<td>0.61 (0.46 to 0.80)</td>
<td>0.0001**</td>
</tr>
<tr>
<td>HIV VL (copies/ml)</td>
<td>&lt;50 (&lt;50 to 7044)</td>
<td>&lt;50 (&lt;50 to 1185)</td>
<td>87 (&lt;50 to 6847)</td>
<td></td>
<td>1.00 (1.00 to 1.00)</td>
<td>0.40</td>
</tr>
<tr>
<td>Elevated bilirubin (0–17 μmol/l)</td>
<td>30 (12 to 60)</td>
<td>16 (14 to 19)</td>
<td>14 (12 to 17)</td>
<td>&lt;20</td>
<td>1†</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>20–40</td>
<td>1.64 (0.48 to 5.6)</td>
<td>0.43</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>≥40</td>
<td>5.04 (1.60 to 15.92)</td>
<td>0.006**</td>
</tr>
<tr>
<td>Peak ALT (0–40 IU/l)</td>
<td>347 (128 to 1792)</td>
<td>308 (204 to 420)</td>
<td>500 (316 to 659)</td>
<td></td>
<td>1†</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt;1000</td>
<td>2.62 (1.01 to 6.80)</td>
<td>0.048*</td>
</tr>
<tr>
<td>CD4 count (300–1400 × 10^3/μl)</td>
<td>650 (490 to 829)</td>
<td>510 (439 to 640)</td>
<td>520 (453 to 619)</td>
<td>&lt;650</td>
<td>1††</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>≥650</td>
<td>2.66 (1.02 to 6.91)</td>
<td>0.045*</td>
</tr>
<tr>
<td>Nadir CD4 (300–1400 × 10^3/μl)</td>
<td>340 (200 to 519)</td>
<td>310 (280 to 350)</td>
<td>315 (270 to 367)</td>
<td>≥200</td>
<td>1††</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt;200</td>
<td>1.51 (0.52 to 4.35)</td>
<td>0.45</td>
</tr>
</tbody>
</table>

Table 2 Median viral load (VL) drop in untreated spontaneous clearers and progressors

<table>
<thead>
<tr>
<th>Transmission</th>
<th>SC (95% CI)</th>
<th>PV (95% CI)</th>
<th>FV (95% CI)</th>
<th>HR* (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDU§</td>
<td>3/13 (23%)</td>
<td>13/44 (30%)</td>
<td>13/38 (36%)</td>
<td>1.63 (0.27 to 2.75)</td>
<td>0.36</td>
</tr>
<tr>
<td>INDU¶</td>
<td>9/12 (75%)</td>
<td>31/42 (74%)</td>
<td>27/34 (79%)</td>
<td>0.92 (0.25 to 3.46)</td>
<td>0.94</td>
</tr>
<tr>
<td>Sexual</td>
<td>14/100 (100%)</td>
<td>46/46 (100%)</td>
<td>39/39 (100%)</td>
<td>1.63 (0.57 to 4.70)</td>
<td>0.36</td>
</tr>
<tr>
<td>HAART</td>
<td>11/15 (73%)</td>
<td>31/53 (58%)</td>
<td>23/42 (55%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3 Viral sequence analyses and predictors of spontaneous clearance

<table>
<thead>
<tr>
<th>Clearers (95% CI)</th>
<th>Progressors (PV+FV) (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline quasispecies</td>
<td>3 (2 to 5)</td>
<td>0.21</td>
</tr>
<tr>
<td>Cumulative quasispecies (two samples taken &lt;150 days apart)</td>
<td>7 (7 to 8)</td>
<td>0.24</td>
</tr>
<tr>
<td>Baseline Hamming distance</td>
<td>0.00001†</td>
<td></td>
</tr>
<tr>
<td>Cumulative Hamming distance (two samples taken &lt;150 days apart)</td>
<td>0.0001†</td>
<td></td>
</tr>
<tr>
<td>Baseline HVR-1 dN/dS</td>
<td>0.004 (0.45 to 1.77)</td>
<td>1.68 (1.40 to 2.02)</td>
</tr>
<tr>
<td>Cumulative HVR-1 dN/dS (two samples taken &lt;150 days apart)</td>
<td>0.03*</td>
<td></td>
</tr>
</tbody>
</table>

Fluctuating viraemia

Patients with a fluctuating VL were more likely to have evidence of superinfecting strains than those with plateau persistence (63% vs 29%; p = 0.04). However, not all PV patients had evidence of superinfection. Among those with single strain infection, persons with recrudescent viraemia had similar dN/dS ratios at baseline but higher cumulative ratios following a second sample than plateau progressors (1.78 vs 0.76; p = 0.02). At baseline, those with single strain recrudescence viraemia had higher viral diversity (Hamming distance 2.84 vs 0.49; p = 0.04). Peak bilirubin, ALT and number of quasispecies did not differ between...
Responsive to treatment

Patients who did not achieve spontaneous clearance 12 weeks following diagnosis were offered treatment with 48 weeks of IFNα and ribavirin, and 66% achieved an SVR.

DISCUSSION

An emerging epidemic of acute HCV in HIV-positive MSM has for the first time allowed characterisation of the natural history of early HCV infection in a large number of HIV-positive patients. Unravelling the determinants of spontaneous clearance of HCV is crucial in order to increase our understanding of the biological mechanisms which define protective immunity and to improve the management of patients who face a lengthy course of treatment which is toxic, expensive and not always effective. HCV infection in HIV-positive individuals is a growing clinical problem associated with an increased rate of disease progression and reduced response to treatment.21 In this longitudinal study of 112 HIV-positive patients with acute HCV, the key predictors of spontaneous clearance were VL dynamics, liver function, viral diversity and T cell response.

Three courses of infection were observed, based on VL changes over time (figure 2). SC (15%) was associated with a rapid decrease in VL. Patients who progressed towards chronicity exhibited two patterns of viraemia; the majority (47.5%) had an initial rise in VL followed by a PV phase, analogous to that seen in other chronic viral infections when the rate of viral clearance...
drop versus a 0.03 log_{10} drop in FV and 0.58 log_{10} drop in FV within 100 days of infection (HR=1.78; p<0.0001; table 2), increasing to a 3.46 versus 0.17 (PV)/0.85 (FV) log_{10} drop by 200 days (HR=1.68; p<0.0001).

Cox regression analysis indicated that the best diagnostic tests for the prediction of SC were the maximum log_{10} drop in VL within 100 days, peak bilirubin (≥40 mmol/l), peak ALT (≥1000 IU/l) and baseline CD4 count (≥650×10^6/l) (tables 1, 2).

A difficulty encountered in predicting the outcome of infection is in distinguishing between those who will develop a FV and those who will go on to spontaneously clear infection, particularly those patients who become transiently RNA negative. Survival analysis was used to compare the time taken to a 2.2 log_{10} drop or a negative VL in FV and SC patients. The time to a 2.2 log_{10} drop was significantly shorter in SC versus FV patients (45 vs 120 days; p=0.01) as was the time to a negative VL in the FV patients who did not spontaneously clear infection (108 vs 154 days); thus the rate of decline of viraemia is key in predicting outcome.

A detailed phylogenetic and genetic sequence analysis of 2291 longitudinally acquired clonal E2 HVR-1 sequences allowed us to track the impact of superinfection, and to assess whether measurements of viral diversity and selection had predictive value. Several monophyletic clusters were present (Appendix 3 online), in keeping with multiple transmission events in the MSM population, as previously described. Within individuals, the presence of superinfection was suggested by marked non-linear variation in genetic distance over time (Appendix 3 online). Phylogenetic tree analyses (figure 3) revealed evidence of the introduction of new infecting strains in 40%, both of the same genotype and with new genotypes. This high rate of superinfection is in keeping with the likely ongoing high risk behaviour of this group of patients and the observation of reinfection events in other cohorts. The majority of patients with FV had evidence of reinfection with new strains of HCV (65%). Strain-specific PCR was carried out to reduce overestimation of superinfection due the limitations of clonal analysis.

In order to investigate the influence of positive selection on the diversification of viral strains, mean intrastrain Hamming distance and dN/dS ratios were calculated. Both were significantly higher in progressors by 150 days of infection, consistent with similar observations in HIV-negative individuals. In the main, a gradual increase in viral diversity and dN/dS ratio was observed over time in FV patients as a result of generation of positively selected variants associated with a significantly weaker T cell response. In contrast, viral diversity and the dN/dS ratio decreased gradually in SC patients (table 3, figure 4) in association with a strong, multispecific T cell response.

When patients with superinfection were excluded from the analysis, genetic diversity at baseline was greater in the FV group than in the FV group, possibly reflecting a more diverse infecting quasispecies.

FV patients as a group displayed a slow rise in diversity over time and a rise followed by a fall in the dN/dS ratio. However, T cell responses, viral diversity and VL were highly variable in individuals within this group. This variation in FV patients would be in keeping with a model of partial immune control and may reflect within some individuals a delayed partially effective immune response and in others the emergence of escape variants or an unsustained protective immune response due to B or T cell failure, for example as a consequence of PD-1 receptor upregulation (a marker of T cell exhaustion and disease progression both in HIV-positive individuals and in acute HCV infection).
Alternatively, it could represent reinfection with the same strain of virus, which has been experimentally demonstrated in the chimpanzee model. A third explanation would be the persistence and re-emergence of virus from sites outside the bloodstream, for example the liver, PBMCs or the central nervous system. Changes observed within the E2 HVR-1 region may reflect direct antibody-driven responses and indirectly reflect T cell responses to other parts of the genome such as the non-structural proteins. Protective immunity in HIV-negative individuals is known to be associated with broad and multispecific CD4 and CD8 T cell responses. A higher baseline CD4 count ($\geq 650 \times 10^6/l$) was noted in SC patients in this cohort (table 1). We therefore aimed to ascertain if decreased dN/dS ratios and viral diversity within E2 in SC and PV patients were related to a protective T-cell response, by measuring IFNγ responses to overlapping pooled peptides spanning the entire HCV genome. We found that SC patients exhibited significantly stronger multispecific IFNy responses to overlapping pooled peptides spanning the entire HCV genome. We found that SC patients exhibited significantly stronger multispecific IFNy responses to overlapping pooled peptides spanning the entire HCV genome. SC patients also had significantly increased responses to NS3 helicase and NS5B than PV progressors. Further studies to characterise specific epitopes within these regions are needed.

One unusual case within the cohort suggested that reactivation of HCV may rarely occur in immunosuppressed individuals. This patient had evidence of reactivation of HCV 2 years after successful treatment for acute disease (figure 5C). This coincided with a period of severe immunosupression (nadir CD4 count $\leq 10^6/l$ and Pneumocystis jerovecii infection) requiring hospitalisation and may have been the result of loss of T cell control of a virus present in the liver but not in the bloodstream, or extremely low level viraemia below the limit of detection. Such cases are rare but have been previously reported in the literature. This study is the first longitudinal and detailed analysis of clinical outcome of acute HCV in a large cohort of HIV-positive patients, incorporating viral dynamics, detailed viral sequence evolution and T cell responses. It has a number of limitations. First, it is restricted to an HIV-positive population, although patterns of evolution during early infection may also apply to those with HCV monoinfection. Steps were taken to minimise error rates attributable to PCR contamination and clonal analysis, as detailed in the Supplementary information online. In order to overcome the problem of delayed HCV seroconversion
in HIV-positive individuals, as previously reported in this cohort (to avoid the inclusion of chronically infected patients), we used retrospective PCR testing on stored HIV VL samples to date the time of infection. More work is needed to further define specific T cell targets and class I/II restriction of these responses, as well as the impact of viral escape.

Acute HCV infection is an emerging global epidemic in HIV-infected men and requires new public health planning and treatment strategies. These data suggest that the outcome of infection can be predicted by the measurement of VL dynamics as well as CD4 count and liver function. Lower genetic diversity and stronger T cell responses are also associated with SC, but were less useful than clinical measures in predicting outcome. Targeting of high risk patients—for example, with no evidence of rapid decline in VL within 100 days—may increase the efficacy of treatment while those more likely to clear infection spontaneously could be monitored for a longer period of time, thus avoiding a rigorous and expensive treatment.

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Predicting spontaneous clearance of acute hepatitis C virus in a large cohort of HIV-1-infected men

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