Characterization of anti-HIV-1 neutralizing and binding antibodies in chronic HIV-1 subtype C infection

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Characterization of anti-HIV-1 neutralizing and binding antibodies in chronic HIV-1 subtype C infection


**A R T I C L E   I N F O**

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**A B S T R A C T**

Neutralizing (nAbs) and high affinity binding antibodies may be critical for an efficacious HIV-1 vaccine. We characterized virus-specific nAbs and binding antibody responses over 21 months in eight HIV-1 subtype C chronically infected individuals with heterogeneous rates of disease progression. Autologous nAbs of study exit plasma against study entry viruses were significantly higher than contemporaneous responses at study entry (p = 0.002) and exit (p = 0.01). NAb breadth and potencies against subtype C viruses were significantly higher than for subtype A (p = 0.03 and p = 0.01) or B viruses (p = 0.03; p = 0.05) respectively. Gp41-IgG binding affinity was higher than gp120-IgG (p = 0.0002), IgG–FcγRIIa affinity was significantly higher than FcγRIIa (p < 0.005) at study entry and FcγRIIb (p < 0.05) or FcγRIIIa (p < 0.005) at study exit. Evolving IgG binding suggests alteration of immune function mediated by binding antibodies. Evolution of nAbs was a potential marker of HIV-1 disease progression.

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**Introduction**

In 2010 alone, there were an estimated 2.4–2.9 million new HIV-1 infections worldwide with 70% of these new infections occurring in sub-Saharan Africa (UNAIDS, 2011). This high HIV incidence makes the development of a protective vaccine a global public health priority. Such a vaccine will likely need to elicit antiviral antibodies, similar to the successful vaccines against other viral infections such as hepatitis B, measles, mumps and polio that are thought to mediate their effects primarily through antibody mechanisms (Plotkin, 2008). However, despite intense efforts in the study of HIV envelope structure and immunogen design, the development of an efficacious vaccine able to induce broadly effective antibody responses remains elusive.

During natural HIV infection, only a subset of individuals, less than 30%, develop broad, cross-neutralizing antibodies after many years (Gray et al., 2011; Li et al., 2009; Sather et al., 2009; Simek et al., 2009; Stamatazos et al., 2009). Such individuals have been an important source of new neutralizing monoclonal antibodies (mAbs) against the HIV envelope that have enhanced our understanding of HIV pathogenesis, envelope structure and provided clues for rational immunogen design (Pejchal et al., 2011; Walker et al., 2010; Wu et al., 2010). However, clinical benefit of anti-HIV antibodies has not yet been definitively demonstrated. Given that HIV-1C is the predominant circulating and most rapidly spreading subtype worldwide (Esparza, 2005; Hemelaar et al., 2011), screening, characterizing and understanding the types of nAbs produced by HIV-1C-infected individuals; and defining the potencies and breadth of these nAbs may contribute to the design of the next generation of envelope immunogens.

While only some people develop cross-neutralizing antibodies, autologous nAbs (AnAbs) appear in almost all HIV-infected individuals within the first year. A number of studies have shown that contemporaneous viruses are less sensitive to AnAbs than
earlier autologous viruses suggesting that viral evolution and escape occurs rapidly and this remains a significant obstacle to HIV vaccine development (Delwart et al., 1997; Moore et al., 2009; Richman et al., 2003; Rong et al., 2009; Wei et al., 2003). Several mechanisms of viral escape have been documented; these include insertions and deletions of amino acids, amino acid substitutions and shifting the position of N-linked glycans in Env (Frost et al., 2005; Lynch et al., 2011; Moore et al., 2009; Rong et al., 2009). Further understanding of HIV antibody escape patterns and mechanisms may help to inform better immunogen design to overcome Env diversity and immune escape.

Although the major focus of the HIV vaccine field is the development of immunogens able to induce broadly neutralizing antibodies, V1–V2 binding antibodies appear to have played some role against HIV-1 acquisition in the RV144 vaccine trial (Haynes et al., 2012). The role of binding or non-neutralizing antibodies in inhibiting virus replication through an Fc (fragment crystallizable) receptor (FcR)-dependent mechanism has been demonstrated (Peressin et al., 2011). FcRs are part of the immunoglobulin (Ig) superfamily and bind to the Fc portion of antibodies forming a bridge between the cell bearing the target antigens and the effector cell (Nimmerjahn and Ravetch, 2007). FcγRs have either an activating or inhibitory function with relative IgG binding affinities. FcγRI (high affinity), FcγRIIa and FcγRIIb (medium–low affinity)—all have activating functions; and FcγRIb has an inhibitory function (medium–low affinity) (Forthal and Moog, 2009; Siberil et al., 2007). The effector cell subsequently mediates virus killing through antibody-dependent cell mediated cytotoxicity (ADCC) or through antibody-dependent cell-mediated viral inhibition (ADCVI). Some studies have shown that the strength of binding interaction between the Fc region of the antibody and the FcγRs and/or FcγRIb potentially increases ADCC activity and Fc binding affinity could be altered through deglycosylation or site-specific mutagenesis, abrogating downstream ADCC (Jefferis, 2009; Lazar et al., 2003; Rong et al., 2009; Wei et al., 2003). Several mechanisms of viral escape have been documented; these include insertions and deletions of amino acids, amino acid substitutions and shifting the position of N-linked glycans in Env (Frost et al., 2005; Lynch et al., 2011; Moore et al., 2009; Rong et al., 2009). Further understanding of HIV antibody escape patterns and mechanisms may help to inform better immunogen design to overcome Env diversity and immune escape.

Table 1
Clinical description of study participants.

<table>
<thead>
<tr>
<th>Patient Identification</th>
<th>Sex</th>
<th>Age at baseline</th>
<th>Viral load study entry (copies/ml)</th>
<th>Viral load study exit (copies/ml)</th>
<th>CD4 T-cell count study entry (cells/µl)</th>
<th>CD4 T-cell count study exit (cells/µl)</th>
<th>CD4 T-cell decline rate (cells/µl per month)</th>
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<tbody>
<tr>
<td>SK010</td>
<td>F</td>
<td>31</td>
<td>6480</td>
<td>345,000</td>
<td>649</td>
<td>268</td>
<td>−10</td>
</tr>
<tr>
<td>SK035</td>
<td>F</td>
<td>31</td>
<td>5800</td>
<td>2950</td>
<td>680</td>
<td>322</td>
<td>−7</td>
</tr>
<tr>
<td>SK036</td>
<td>M</td>
<td>32</td>
<td>5100</td>
<td>10,600</td>
<td>936</td>
<td>575</td>
<td>−7</td>
</tr>
<tr>
<td>SK169</td>
<td>F</td>
<td>30</td>
<td>2210</td>
<td>2440</td>
<td>561</td>
<td>437</td>
<td>−4</td>
</tr>
<tr>
<td>SK200</td>
<td>F</td>
<td>40</td>
<td>14,360</td>
<td>24,900</td>
<td>595</td>
<td>416</td>
<td>−7</td>
</tr>
<tr>
<td>SK221</td>
<td>F</td>
<td>30</td>
<td>9740</td>
<td>23,700</td>
<td>503</td>
<td>297</td>
<td>−3</td>
</tr>
<tr>
<td>SK233</td>
<td>F</td>
<td>22</td>
<td>11,800</td>
<td>18,900</td>
<td>547</td>
<td>218</td>
<td>−8</td>
</tr>
<tr>
<td>SK312</td>
<td>F</td>
<td>59</td>
<td>3460</td>
<td>3630</td>
<td>545</td>
<td>881</td>
<td>−9</td>
</tr>
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</table>
exit (median nAb IC50 of 305; range: 210–1,076; \( p=0.01 \)). There were no correlations between the rates of CD4 T-cell decline or viral loads with for the nAb IC50 titers of study entry Envs against the study exit plasma or between contemporaneous nAb IC50 titers at study entry or study exit—Fig. 1(C) and (D).

**Autologous neutralization titers correlated with the length of the hypervariable regions V1–V2**

Previous studies have shown an inverse association between neutralizing antibody titers and length of variable loops (V1–V2) and numbers of potential N-linked glycosylation sites (PNGs) (Chackerian et al., 1997; Gray et al., 2007; Pinter et al., 2004; Rong et al., 2007; Sagar et al., 2006). To better understand the relationship between genotypic characteristics and neutralization sensitivity, the average amino acid length of the hypervariable loops of Env and PNGs were correlated to average autologous nAb IC50 titers (Fig. 2). Significant inverse correlation was found between the lengths of V1–V2 and nAb IC50 titers (\( r=-0.80; \ p=0.02 \)) for the entry viruses tested against the study exit plasma. There was a trend towards C2-V5 length correlation with nAb IC50 titers (\( r=-0.67; \ p=0.08 \)) and no significant correlations for V1–V5 (\( r=0.52; \ p=0.2 \)). The same analysis was extended for nAb IC50 titers and the numbers of potential N-linked glycosylation sites.

Fig. 1. (A) Schematic illustrating the autologous neutralizing antibody challenge assays in all participants over a median of 21 months from study entry to study exit. The average nAb IC50 titers for study entry and exit Envs used in the assays are shown. The study entry Envs were tested against nAbs from the study entry (contemporaneous) and study exit plasma samples. Likewise the study exit Envs were tested against nAbs from the study entry and study exit (contemporaneous) plasma samples. Fig. 1(B) depicts the autologous nAb IC50 titers in study participant entry and exit plasma samples for all participants. Fig. 1(C) and (D) shows the CD4 T-cell decline rate and the log viral load changes over time respectively. \( p \)-Values < 0.05 were considered significant. \( p \) Value was calculated using the two-tailed Mann-Whitney non-parametric test overall. All the \( p \) values (\( p<0.0125 \)) remained statistically significant after Bonferroni adjustment for multiple comparisons.
sites (PNGs) in the various hypervariable loops and there were no significant correlations observed.

Neutralization breadth scores and potency scores indicate subtype-specific responses in chronic infection

To assess neutralization breadth in this chronic infection cohort, we investigated changes in plasma neutralization activity over a median of 21 months. Fig. 3 depicts the profile of plasma neutralizing activity against a panel of heterologous Envs for the eight study participants at study entry and study exit. A total of 20 pseudoviruses from subtype A, B and C (as depicted in Table 2) were used to determine breadth and potency scores of neutralization according to the criteria previously described (Blish et al., 2007). Results are presented as a breadth score per subtype virus panel, and a total breadth score including all the subtype A, B and C viruses. Results are presented as potency scores per subtype, and total potency scores including all the subtype A, B and C viruses.

There was a wide range of variation in neutralization titers with most of the sera from the participants displaying preferential heterologous activity against the subtype C panel (Fig. 3). Significantly higher breadth scores were observed against subtype C viruses (median: 3; range: 1–7) compared to subtypes A (median: 2; range: 0–4; \( p=0.03 \)) or B (median: 2; range: 0–6; \( p=0.01 \))—Fig. 3. Likewise, significantly higher potency scores were observed against subtype C viruses (median: 7.6; range: 0–152.1) compared to subtypes A (median: 4; range: 0–13.8; \( p=0.03 \)) or B (median: 3.9; range: 0–26.9; \( p=0.05 \)). Three of the participants (SK200, SK221 and SK233) showed a consistent increase of neutralization breadth scores and potency scores from study entry to study exit for subtypes C, B and A respectively translating into overall increase in total breadth and potency scores as well. Analysis of these participants indicated significantly higher viral load at study entry compared to study exit (\( p=0.04 \)); suggesting that increasing breadth may be a result of higher antigenic stimulation.

SK200 displayed the highest breadth scores and also had potent nAb IC50 scores (Fig. 3) to six of the eight subtype C viruses tested including five of six tier 2 viruses. SK200 had a total neutralization breadth score of 16 (range: 2–16) and the overall potency score was 171.1 (range: 3–177.1) at study exit. SK200 neutralized subtype C – tier 2 viruses – CAP45.G3 and CAP239.G3 at study entry and exit plasmas, and CAP45.G3 in particular, was neutralized at nAb IC50 titers of >2000 (range: 45–5326). SK200 also displayed the highest titers >1000 at study entry (range: 45–1316) and study exit (range: 45–1829) to a subtype A tier 1 virus – Q23ENV17.

CD4 T-cell count and viral load are not correlated with neutralization breadth

A number of studies have shown an inverse relationship between neutralization breadth and CD4 T-cell counts and a positive correlation with viral loads (Euler et al., 2010; Gray et al., 2011; Plantadosi et al., 2009; Sather et al., 2009). To better understand the relationship between neutralization and clinical disease markers in this subtype C chronic cohort, we investigated the association of CD4 T-cell counts and viral loads with neutralization breadth or potency over a median of 21 months. However, there were no significant correlations between CD4 T-cell counts or viral loads and total breadth and potency scores (data not shown). A larger sample size may have been able to detect significant correlations between breadth or potency and markers of disease progression.

Mapping of potential epitopes targeted by cross-neutralizing antibodies

The putative targets of neutralization in participants displaying the highest breadth scores for neutralization at study exit (SK200, SK221 and SK233) against the reference panel viruses were further investigated (using study exit plasma samples). Single point mutations in V2 (N160A, K165E or L165A) and C (N332A) were introduced into CAP45.G3, ConC, Du156.12, TR0.11 and Q323 and tested for loss of sensitivity against the three plasmas.

As depicted in Table 3, 5.6-fold and 4-fold drops in neutralization titers (relative to the wild type neutralization titers) were seen when plasma samples from SK221 and SK233, respectively,
than for gp41 (median: 96.8; range: 71.9–112.1; gp120 (median: 137; range: 11.3–153.0) was significantly lower. Interestingly, at study entry, the median IgG binding affinities for p24, gp120 or gp41 or gp41.

There were no significant differences in the median EC50 binding affinities between study entry and study exit for p24, gp120 or gp41. Specifically and the binding affinity of IgGs to HIV-1 specific antigens. pants at study entry and study exit time points and measured the infection over 21 months. The neutralization titer is shown as reciprocal plasma dilution required to inhibit 50% of virus infectivity when the virus is challenged with the participant's plasma. The highest titer ( > 1,000) is shown in red, and the lowest in light orange, yellow depicts a titer of < 1:45 that is below detection as shown above.

Table 3
Effects of single point mutations on neutralization sensitivity and summary of antibody specificities.

<table>
<thead>
<tr>
<th>Plasma sample identity</th>
<th>Fold effect of mutationa</th>
<th>Antibody specificity conferring breadth</th>
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<tbody>
<tr>
<td></td>
<td>ConC N160A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ConC I165A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>ConC K169E</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAP45 N160A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAP45 I165A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAP45 K169E</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CAP45.G3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Du156.12</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N332A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Q23.17</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N332A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tro.11</td>
<td></td>
</tr>
<tr>
<td>SK200 study exit</td>
<td>1.1</td>
<td>Unknown</td>
</tr>
<tr>
<td>SK221 study exit</td>
<td>0.4</td>
<td>Quaternary, PG9/PG16 like N332</td>
</tr>
<tr>
<td>SK233 study exit</td>
<td>0.1</td>
<td>Quaternary, PG9/PG16 like</td>
</tr>
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* Calculated as wild type IC50/mutant IC50 for the plasma. Changes in titer of > 3 fold are shown in bold.

were tested against the K169E mutation in the CAP45 backbone. This suggested that the broadly neutralizing antibodies in these two participants targeted the V2 region and that a charge change from a lysine (K) to glutamic acid (E) at position 169 resulted in a disruption to the nAb epitope. In addition SK221 plasma also showed a 4.2 fold drop in neutralization titers to N332A made in the TRO.11 envelope indicating that the asparagine (N), or perhaps a potential N-linked glycosylation at this position, was essential for the antibody activity. Together, these results suggested that SK221 most likely had cross-neutralizing antibodies that target the V2 and C3 regions on Env. SK200, who displayed the highest neutralization breadth score, neutralized the wild-type and mutant viruses equivalently suggesting that this partic-

ular participant may have antibodies directed to other regions of Env, a profile that warrants further investigation.

HIV-specific IgG binding titers for p24, gp120 or gp41 in chronic infection over 21 months

We isolated total immunoglobulins (IgGs) from study partici-

pants at study entry and study exit time points and measured the specificity and the binding affinity of IgGs to HIV-1 specific antigens. There were no significant differences in the median EC50 binding affinities between study entry and study exit for p24, gp120 or gp41. Interestingly, at study entry, the median IgG binding affinities for gp120 (median: 137; range: 11.3–153.0) was significantly lower than for gp41 (median: 96.8; range: 71.9–112.1; p = 0.0002). Binding affinity for gp120 compared to p24 trended to be higher (median: 154.3; range: 57.9–293.9; p = 0.09)—Fig. 4. These findings may suggest that binding IgGs were selectively binding gp41 compared to gp120 during the chronic infection stage. In addition, there were no significant correlations between the CD4 T-cell counts or viral loads at study entry or exit with the EC50 of p24, gp120 or gp41 over the study period.

IgG binding affinities differ for the various FcγRI, IIA, IIB and IIIa

To determine if the binding affinities of the IgGs with various activating and inhibitory FcγRIs were different, we measured and compared the EC50 binding affinities over time between study entry and study exit time-points, and overall (Fig. 5A). IgG binding for FcγRI at study entry was significantly higher than for FcγRIIa (p < 0.0005). Likewise at study exit, the binding affinity of IgG for the FcγRI were consistently higher compared with FcγRI (p < 0.05) and FcγRIIa (p < 0.0005). Significantly higher binding affinities for FcγRIIa at study entry were noted when compared to FcγRIIa at study entry (p < 0.05). Likewise at study exit, the binding affinity of IgG for the FcγRIIa was higher compared to FcγRIIa (p < 0.05) in chronic HIV-1 C disease. Collectively, these results indicated that there was differential kinetics of binding affinities of the IgG during chronic infection, which may then impact downstream non-neutralizing activity through ADCC. Sustained high affinity IgG binding to FcγRI may be preserved during chronic infection. There were no significant correlations between viral loads or CD4 T-cell counts with the EC50 IgG binding titers to the various receptors (FcγRI I, IIA, IIB and IIIa)—Fig. 5(B) and (C).

Discussion

Neutralizing and binding antibodies may play a role in HIV disease progression but these parameters have rarely been investigated concurrently. We therefore undertook this study in individuals chronically infected with HIV-1 subtype C, the most abundant
subtype worldwide, to characterize neutralizing and binding antibody patterns in chronic disease, and to determine whether certain genotypic env characteristics are associated with autologous or heterologous nAb responses. Our data indicated that nAbs did not appear to protect against disease progression, rather, greater type-specific neutralization breadth and potency against subtype C Envs and increasing autologous nAb titers were associated with chronic disease progression. NAb IC50 titers were correlated with env genotypic characteristics, including increased amino acid length in hypervariable regions V1–V2 of gp120.

In this study, nAb potency or breadth did not predict disease progression rate in individuals with chronic HIV-1 subtype C infection. However, a number of interesting findings were apparent. Firstly, high-level neutralizing titers to contemporaneous autologous virus were not observed in most of the participants. Instead, we observed significantly higher autologous responses over time (when the study entry viruses were tested against the study exit plasma) compared to contemporaneous responses, which suggests that nAb are continuously evolving during chronic infection. These findings also argue that an increased nAb titer per se is not effective at attenuating disease progression but rather is a marker of disease progression. Secondly, autologous nAb IC50 titers correlated inversely with longer amino acid length for V1–V2 length indicating that increased length in this domain may be masking key neutralization epitopes. Indeed, evidence for V2 dependent epitopes was observed in SK221 and SK233. This data suggests that V1–V2 of Env may be persistently targeted by nAb in the chronic infection stage. Alternatively, there may be other intrinsic genetic differences of env that dictate neutralization potency and breadth that need to be further defined. Lastly, the dynamic nature of IgG binding affinity to various FcγRs and the selective high affinity binding to HIV-1 gp41 may be indicative of chronic disease progression.

The selective increase in neutralization breadth over time suggests that this parameter could be a marker of disease progression in chronic subtype C infection (Euler et al., 2010). Neutralization breadth did not significantly correlate with markers of disease progression (viral loads and CD4 T-cell counts) in this chronic HIV-1C infection cohort, although the small sample size and short time-frame may have severely limited our statistical power to address this issue. Gray et al. (2011) found significant correlations between CD4 T-cell count and viral load with neutralization breadth.
at six months post-infection only and not at later time points (Gray et al., 2011). These observations suggest that higher antigenic stimulation may dictate the breadth of heterologous antibody responses (Doria-Rose et al., 2010; Fraser et al., 2007; Goujard et al., 2006; Mellors et al., 1997; Piantadosi et al., 2009; Sather et al., 2009). Although increased neutralization antibody breadth may not be protective against disease progression, they may be effective against super-infection as has been suggested by some studies (Smith et al., 2006 and Deeks et al., 2006), although conflicting data exists (Blish et al., 2008). Together, these studies have implications for HIV vaccine design, as vaccine immunogens may need to be given over long periods of time to stimulate the B cell response, and to facilitate affinity maturation which appears to be necessary for antibodies to acquire cross-neutralizing activity (Pancera et al., 2010). For the participants who displayed potent cross-neutralizing antibody responses, their nAbs likely targeted quaternary V2 epitopes similar to PG9/PG16-like antibodies (Moore et al., 2011). In addition, nAbs that targeted the N332 glycan in C3 suggest that the asparagine in that position is essential for neutralization activity. In subtype C infection, the V1–V2 and C3 regions are the immunodominant regions commonly targeted by AnAbs particularly during the early stage of infection (Lynch et al., 2011; Moore et al., 2008, 2009; Rong et al., 2009). Our results indicate the V1–V2 and C3 regions remain immunodominant and the focus of the nAb response resulting in broadly cross-neutralizing antibodies even during chronic progressive HIV-1 disease.

A decline of p24 antibody titers as disease progresses has previously been reported (Allain et al., 1987; Binley et al., 1997; Forster et al., 1987; Lange et al., 1986), and it is therefore not surprising that the p24-specific IgGs may lose their affinity over time. Chargelegue and colleagues found that low p24 IgG affinity correlated with HIV-1 disease progression (Chargelegue et al., 1995). Low affinity IgG HIV-1 specific binding antibodies may be indicative of chronic disease progression. It is plausible that a relative decrease in the binding affinity for activating FcγRlla and FcγRlli (EC50) are shown. Only p-values < 0.05 are shown.

Fig. 5. (A) EC50 binding titers of IgGs for FcγRI, FcγRlla, FcγRllb and FcγRlli at study entry and exit time points. Intra- or inter-group comparisons were statistically different i.e. p < 0.05. Fig. 5(B) and (C). Correlation of CD4 T-cell counts at study entry and study exit with FcγRI, FcγRlla, FcγRllb and FcγRlli (EC50) are shown. Only p-values < 0.05 are shown.
affect FcγRI binding. Polymorphisms in the FcγRIa (Forthal and Moog, 2009), and the extent of glycosylation of the antibodies (Forthal et al., 2010) affect binding affinity. Additionally, a decrease in binding affinity of FcγRIa was associated with dysfunction in the expression of these receptors on the cells of the innate immune system (Dugast et al., 2011). Dugast and colleagues purport that HIV infection is associated with a number of changes in FcR expression on phagocytic cells that are associated with changes in their ability to respond to antibody-opsonized targets, leading to a failure in viral clearance in chronic infection (Dugast et al., 2011). We found a consistently high affinity binding of IgG to FcγRI over time which may suggest that the non-neutralizing pathway augmenting effector cell activity may be maintained during the chronic infection stage. However, in the absence of a functional assay, our data do not provide definitive answers on whether high affinity binding IgG to FcγRI is biologically significant in chronic infection and what the profiles of the various FcγRI–IgG binding were during the initial HIV-1 infection stages.

Some limitations of our study are worth noting. Firstly, we only tested a limited number of viral clones from each patient, and therefore may have biased the results of the study to the cloned variants only. Secondly, we do not know the exact time of infection for the study participants and relied on short-term median (21 months) follow-up immunological data, which may be an unrepresentative snap-shot of the entire natural history of disease progression for these participants. Thirdly, the small sample size meant that our study was underpowered to reach definitive conclusions regarding the significance of breadth of neutralization and autologous responses in attenuating disease. Fourthly, we used binding affinity assays as an indication of the strength of affinity of the Fc portion of IgG to the FcγRs which may affect downstream in vivo effector cell activity. Lastly, the dynamic nature of IgG binding affinity to various FcγRs and the selective high affinity binding to HIV-1 gp41 may be indicative of differences and evolution of the antibody mediated innate antiviral immune response in progressive HIV-1 subtype C infection, although this will require additional functional assays to prove.

Despite these caveats, the findings suggest that an ongoing de novo nAb response does not directly protect against disease progression during chronic HIV-1 subtype C infection, corroborating data from previous studies. Our results are also consistent with data from non-human primate studies showing that high titer neutralizing antibody titers are not found in sooty mangabeys (Li et al., 2010) suggesting that autologous nAbs are not part of the protection against disease progression in HIV and SIV infections. Furthermore, overall nAb breadth increased over time in most subjects, regardless of their disease status. The mechanism by which nAb breadth increased in these subjects is of interest. Thus, it will be important to determine which Env epitopes in chronically infected individuals elicit broadly neutralizing antibodies and whether these provide any clinical benefit to the patient. In addition, animal studies suggest that anti-HIV antibodies can protect against HIV-1 infection (Burton et al., 2011; Hessell et al., 2009, 2010) and vaccine-induced V1–V2 binding antibodies may be able to prevent infection (Haynes et al., 2012). Future studies will need to address whether neutralizing and binding antibodies from HIV infected individuals may be used to halt or reduce HIV acquisition.

**Materials and methods**

**Participants**

Participant samples were retrospectively identified from the Sinikithemba cohort, which is a prospective natural history study of HIV-1 infected individuals based at McCord Hospital, Durban, South Africa as previously reported (Kiepiela et al., 2004). Human subjects for this study were eight chronically-infected individuals with heterogenous rates of clinical disease progression as previously described (Archary et al., 2010). The median period of follow-up overall was 21 months. Ethical approval for this study was obtained from the University of KwaZulu-Natal Biomedical Research Ethics Committee and all participants gave written informed consent to participate in the study.

**Sample collection, CD4 T-cell counts and plasma viral load**

CD4 T-cell counts were performed at three-month intervals whereas viral loads were done at six-month intervals as previously described (Archary et al., 2010). Blood was drawn from each subject into EDTA tubes and plasma was separated by centrifugation and stored at −80 °C until use. Viral load was measured using the Amplicor Version 1.5 assay (Roche, Alameda CA, USA). CD4 T-cell counts were enumerated on a four color FACSCalibur flow cytometer (Becton Dickinson, Franklin Lakes, New Jersey, USA).

**Envelope clones**

Briefly, single genome PCR amplification derived env amplicons were directionally T/A cloned into the CMV-driven expression plasmid pcDNA3.1-V5 HisTOPO-TA and screened for biological function as pseudoviruses following co-transfection with an env-deficient subtype B proviral plasmid (SG3ΔEnv) into 293 T cells as described previously (Derdeyn et al., 2004; Haaland et al., 2009). A total of 20 standard reference envelope clones were obtained from the NIH AIDS Research and Reference Reagent Program and used in the heterologous neutralization assays. These included five subtype A, seven subtype B and eight subtype C envelope pseudoviruses as depicted in Table 2. Tier 1, 2 and 3 viruses were stratified previously based on a continuum of patterns of neutralization sensitivity—with tier 1A being the most sensitive, tier 1B above average, tier 2 displaying moderate to low sensitivity and tier 3 displaying the lowest sensitivity to neutralization (Seaman et al., 2010). The ConC plasmid based on the consensus of all the HIV-1 subtype C sequences from the Los Alamos database by 2001 was obtained from Dr Feng Gao (Kothe et al., 2006). The envelope plasmids containing single point mutations are described by Gray et al. (2011).

**Neutralization assays**

Patient plasma samples were evaluated for nAb activity against virions pseudotyped with autologous patient-derived viral Envs using a single reporter assay as described previously (Derdeyn et al., 2004; Gray et al., 2007; Li et al., 2005, 2006; Wei et al., 2003). Fig. 1(A) illustrates the schema used for autologous neutralizing antibody challenge assays where the study entry and exits Envs were tested for neutralization using the study entry and study exit plasma nAb samples for eight study participants. A total of 37 autologous Envs were tested against their study entry plasma and study exit plasma (range: 2–6 env clones per participant). Neutralization was measured as a reduction in luciferase gene expression after a single round of infection of TZM-βl cells (NIH AIDS Research and Reference Reagent Program). Two thousand infectious units of each pseudovirus was combined with five-fold dilutions of heat-inactivated participant plasma and incubated for 1 h at 37 °C as previously described (Rong et al., 2009). Subsequently, the virus and antibody reaction was added to the plated TZM-βl cells, and left to incubate at 37 °C for 48 h. The cells were then lysed and the luciferase activity was determined using a BioTek Synergy HT (BioTek, USA). The background
luminescence of the uninfected wells was subtracted from the test wells. The percentage of infectivity was calculated by dividing the number of luciferase units at each plasma dilution by the value in the well containing no test plasma. The dilution that yielded 50% inhibitory activity against the virus, known as the nAb IC50 titer was determined on Microsoft Excel. Each experiment was performed in duplicate and independently at least twice for replicability. The nAb IC50 titer was calculated as the reciprocal plasma dilution causing a 50% reduction of relative light units (IC50). Heterologous assays were done as previously reported (Montefiori, 2004). Briefly, positive neutralization was scored as an IC50 titer above 1:45. The nAb breadth and potency was determined as a score based on the median IC50 titer for each virus as reported by Blish et al. (2008).

**Plasma IgG isolation**

Total IgGs were purified from the participant’s plasma samples using the Melon Gel (Thermo Scientific, Surrey, United Kingdom) method according to the manufacturer’s instructions. Briefly, 500 μl of a 1:10 dilution of plasma was added to the column. The tubes were rotated for 5 min in order to ensure maximum capture of the IgGs and were then spun down for 1 min at 5000x g to elute the IgGs. IgG concentrations were measured on the NanoDrop 2000 Spectrophotometer (Thermo Scientific, Surrey, United Kingdom) and stored at 4 °C for downstream ELISA assays.

**Gp120, gp41, and p24 Binding ELISAs**

Binding affinities of the bulk population of IgGs purified from plasma were measured for the various HIV-1 specific antigens gp120, gp41 and p24. ELISA plates were coated with 80 μl/well of 250 ng/ml gp120 (Immune Technology, New Jersey, USA) or gp41 (Immune Technology, New Jersey, USA) and left for 1 h at RT. The plate was washed 3 times with 0.05% PBS–BSA at 4 °C and incubated O/N at 4 °C. The plates were washed three times with PBS Tween 0.05% and blocked with 5% PBS–BSA for 2 h at RT (100 μl/well). IgGs isolated previously were diluted in PBS and added to respective wells according to a ten times serial dilution to respective wells and incubated for 2 h at RT. The plates were washed three times with 0.05% PBS Tween. 100 μl anti-human IgG antibody, Fab fragment, peroxidase labeled was added to each well (KPL Protein Research Products, Maryland, USA) and left for 1 h at RT (1:500 diluted-per plate: 20 μl HRP in 10 ml PBS) and covered to protect it from light. The plate was washed three times with 0.05% PBS–Tween. The color was resolved with 100 μl/well of 0.05% phenylene diamine (OPD) (Sigma-Aldrich, St Louis, USA) – one OPD tablet was added to 11 ml of StrepHRP substrate), the reactions were stopped by adding 100 μl/well of 2.5 N sulfuric acid (Sigma-Aldrich, St Louis, USA). The plate was read at 490 nm using the Tecan Sunrise 16039400 plate reader (Tecan Group, Switzerland). The optical density data was exported to Microsoft Excel worksheet to calculate the 50% effective concentration (EC50) of each batch of total IgGs needed to bind HIV-specific gp41, gp120 and p24 antibodies and various Fc receptors (FcγRI, FcγRIIa – (R & D Systems, Minneapolis, USA) and FcγRIIIa – (R & D Systems, Minneapolis, USA) and left for 1 h at RT (1:500 diluted-per plate: 20 μl HRP in 10 ml PBS) and covered to protect it from light. The plate was washed three times with 0.05% PBS–Tween. The plate was washed three times with PBS Tween 0.05% and blocked with 250 μl/well 5% PBS–Bovine Serum Albumin (BSA) (Sigma-Aldrich, St Louis, USA) and then incubated O/N at 4 °C. The plates were washed three times with PBS Tween 0.05% and blocked with 5% PBS–BSA for 2 h at RT (100 μl/well). IgGs isolated previously were diluted in PBS and added to respective wells according to a ten times serial dilution to respective wells and incubated for 2 h at RT. The plates were washed three times with 0.05% PBS Tween. 100 μl anti-human IgG antibody, Fab fragment, peroxidase labeled was added to each well (KPL Protein Research Products, Maryland, USA) and left for 1 h at RT (1:500 diluted-per plate: 20 μl HRP in 10 ml PBS) and covered to protect it from light. The plate was washed 3 times with 0.05% PBS–Tween. One OPD tablet was added to 11 ml of phosphate citrate buffer + 4.4 μl hydrogen peroxide (H2O2); and 50 μl/well were added. The reaction was stopped by adding 50 μl 2.5 N H2SO4 per well. The plate was read at 490 nm on a microplate reader and the data analyzed using GraphPad Prism 5 software.

**FcGamma (γ) receptor binding ELISAs**

Binding affinities of the bulk population of IgGs purified from plasma were measured for the various activating – FcγRI, FcγRIIa and FcγRIIIa – (R & D Systems, Minneapolis, USA) and inhibitory receptors – FcγRIb.

Each ELISA plate (NI-NTA HisSorb™ Plate Qiagen, Dusseldorf, Germany) was coated with 100 μl/well of diluted FcγR (FcγRI, FcγRIIa, FcγRIIIa, and FcγRIib) at a concentration of 5 μg/ml using PBS. One receptor type per plate was added and incubated overnight (O/N) at room temperature (RT). The plate was washed three times with PBS–TWEEN (0.05%) and blocked with 250 μl/well 5% PBS–Bovine Serum Albumin (BSA) (Sigma-Aldrich, St Louis, USA) and then incubated for 1 h at RT. The plate was washed three times with 0.05% PBS–TWEEN. 100 μl/well of antibodies were added (the IgGs derived from the test plasma were diluted serially starting at 100 μg/μl diluted down to 1.6 μg/μl) to the respective wells and was left to incubate for 1 h at RT. The plate was washed three times with 0.05% PBS–TWEEN. 100 μl per well of anti-human IgG antibody, Fab fragment, peroxidase labeled was added (KPL Protein Research Products, Maryland, USA). 20 μl of HRP Fab (1 mg/ml) was added to 10 ml PBS 1X for one plate. The plate was incubated for 1 h at RT and covered to protect it from light. The plate was washed three times with 0.05% PBS–TWEEN. The color was resolved with 100 μl/well of O- phenylene diamine (OPD) (Sigma-Aldrich, St Louis, USA) – one OPD tablet was added to 11 ml of StrepHRP substrate), the reactions were stopped by adding 100 μl/well of 2.5 N sulfuric acid (Sigma-Aldrich, St Louis, USA). The plate was read at 490 nm using the Tecan Sunrise 16039400 plate reader (Tecan Group, Switzerland). The optical density data was exported to Microsoft Excel worksheet to calculate the 50% effective concentration (EC50) of each batch of total IgGs needed to bind HIV-specific gp41, gp120 and p24 antibodies and various Fc receptors (FcγRI, FcγRIIa – (R & D Systems, Minneapolis, USA) and FcγRIIIa – (R & D Systems, Minneapolis, USA) and left for 1 h at RT (1:500 diluted-per plate: 20 μl HRP in 10 ml PBS) and covered to protect it from light. The plate was washed three times with 0.05% PBS–TWEEN. One OPD tablet was added to 11 ml of phosphate citrate buffer + 4.4 μl hydrogen peroxide (H2O2); and 50 μl/well were added. The reaction was stopped by adding 50 μl 2.5 N H2SO4 per well. The plate was read at 490 nm on a microplate reader and the data analyzed using GraphPad Prism 5 software.

**Statistical analyses**

Comparisons of different parameters including nAb IC50 titers for both the autologous and heterologous responses, and binding affinities of the various HIV-specific antigens and FcγRs to IgGs was performed. Intra-group comparisons between study entry and study exit time-points were done using the Mann-Whitney U test and the Spearman rank correlation test. ANOVA was used to compare three or more groups and the Dunn’s post-test was used to perform pairwise comparisons. A general linear model was used to estimate the rate of change in viral load and CD4 T-cell count over time. Viral load was log-transformed to ensure normality. All reported p values are two-sided and are considered significant if less than 0.05. The statistical analysis was performed using GraphPad Prism 5 and SAS version 9.3 (SAS Institute Inc., Cary).

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