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Targeting the Late Stage of HIV-1 Entry for Antibody-dependent Cellular Cytotoxicity: Structural Basis for Env Epitopes in the C11 Region

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Summary

Antibodies can impact HIV-1 infection in multiple ways including antibody-dependent cellular cytotoxicity (ADCC), a correlate of protection observed in the RV144 vaccine trial. One of the most potent ADCC-inducing epitopes on HIV-1 Env is recognized by the C11 antibody. Here we present the crystal structure, at 2.9 Å resolution, of the C11-like antibody N12-i3, in a quaternary complex with the HIV-1 gp120, a CD4-mimicking peptide M48U1, and an A32-like antibody, N5i5. Antibody N12-i3 recognizes an epitope centered on the N-terminal "8th-strand" of a critical βsandwich, which our analysis indicates to be emblematic of a late entry state, after the gp120 detachment. In prior entry states, this sandwich comprises only 7-strands, with the 8th strand

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instead pairing with a portion of the gp120 C terminus. The conformational gymnastics of HIV-1 gp120 thus includes altered β -strand pairing – possibly to reduce immunogenicity – though nevertheless still recognized by the human immune system.

eTOC Blurb



Tolbert at al. describe a crystal structure of Fabs of A32- and C11-like antibody bound to single gp120 subunit. The C11-like antibody, N12-i3 recognizes new 8-stranded β -sandwich structure of gp120 that is formed at the late stage of HIV-1 entry.

Introduction

The native envelope (Env) glycoprotein trimer expressed at the surface of the viral particles or infected cells and its transition state structures induced on the cell surface after receptor/co-receptor binding constitute major epitope targets for potentially protective anti-HIV-1 antibody responses. The extreme genetic diversity of Env, its heavy glycosylation, and conformational masking are effective evasion mechanisms against antibodies targeting epitopes available on Env in its native, un-triggered state ((Moore et al., 2009; Wei et al., 2003). In contrast, Env transition state structures exposed after engagement of the Env trimer with host receptors during viral entry (Guan et al., 2013; Mengistu et al., 2015), or on infected cell surfaces after cis (Veillette et al., 2014) or trans (Finnegan et al., 2001) triggering by CD4 (CD4-induced [CD4i] epitopes) are attractive targets for antibodies with protective potential (reviewed in (DeVico, 2007; Lewis et al., 2017b). These transition state structures harbor the most conserved regions of Env and become temporarily available for antibody recognition, providing CD4i antibodies a limited window of opportunity to counter the infection. Theoretically, antibodies that target Env transition state structures could afford protection by virus neutralization but this is limited strictly to Tier-1 viruses that are not representative of transmitted founder viruses which typically exhibit Tier-2/3 neutralization resistance (Seaman et al., 2010). Recent analyses suggest that CD4i antibodies contribute to

protection via $Fc\gamma$ receptor ($Fc\gamma R$)-mediated effector functions such as antibody-dependent cellular cytotoxicity (ADCC) (Ferrari et al., 2011; Guan et al., 2013; Veillette et al., 2014). Previously, we defined Epitope Cluster A as a potent ADCC target recognized by CD4i mAbs (Guan et al., 2013). Epitope Cluster A maps to the inner domain and the extended N-, C-termini of gp120 ((Acharya et al., 2014; Gohain et al., 2015) and reviewed in (Lewis et al., 2014; Lewis et al., 2017a; Veillette et al., 2016)). Antibodies specific for Cluster A epitopes were shown to mediate the majority of the ADCC activity detected in chronicallyinfected individuals (Ferrari et al., 2011; Guan et al., 2013; Veillette et al., 2014) as well as being implicated in protection for the clinical RV144 vaccine trial (Bonsignori et al., 2012; Haynes et al., 2012; Tomaras et al., 2013) and our SHIV challenge studies in rhesus macaques (DeVico et al., 2007; Fouts et al., 2015) immunized with a gp120 transition state vaccine (Fouts et al., 2000). Epitope Cluster A is buried at the trimer axis on native Env and inaccessible for antibody recognition until interaction with cell surface CD4. We and others have shown that exposure of these epitopes on the virion spike and on the surface of infected cells strictly depends on the cell surface form of CD4, since triggering with soluble CD4 does not lead to exposure of these epitope targets (Acharva et al., 2014; Mengistu et al., 2015; Ray et al., 2014; Veillette et al., 2014). These data point toward the possibility of an additional energy component required for trimer unfolding and effective presentation of epitopes of the Cluster A region that is provided in vivo by conformational changes in the D3D4 region of cell-surface CD4 (Acharya et al., 2014; Gohain et al., 2015; Veillette et al., 2015). Recent data also indicate that small CD4 mimetic compounds and the CD4-peptide mimetic miniprotein M48U1 (Martin et al., 2003; Van Herrewege et al., 2008) are able to induce the CD4-bound conformation of Env at the surface of HIV-1 infected cells and thus sensitize HIV-1-infected cells to the ADCC killing mediated by Cluster A antibodies provided that anti-co-receptor binding site antibodies are present (Richard et al., 2016; Richard et al., 2015).

We reported previously that the Cluster A region consists of three epitope specificities as defined by ELISA competition for binding to CD4 triggered gp120 with the two prototype mAbs: A32 and C11 (Guan et al., 2013). One subgroup only competes with A32 (A32-like epitopes), the second only competes with C11 (C11-like epitopes), and the third competes with both A32 and C11 (hybrid A32-C11-like epitopes). The existence of these three subregions was confirmed using fluorescence correlation spectroscopy coupled with fluorescence resonance energy transfer (FRET-FCS) (Gohain et al., 2015). Recently, we defined the A32 and the mixed A32-C11 epitopes of Cluster A region at the atomic level (Acharya et al., 2014; Gohain et al., 2015; Tolbert et al., 2016). Both of these epitope regions mapped within the inner domain of gp120 including mobile layers 1 and 2 of constant regions 1 and 2 (C1-C2 region) in CD4-triggered gp120 (Acharya et al., 2014; Gohain et al., 2015; Tolbert et al., 2016). Further, the A32-C11 mixed epitope extended to a few residues in the 7-stranded β -sandwich of the gp120 inner domain (Gohain et al., 2015). Here, we continue with the mapping of the epitope subgroups within the Cluster A region and present the structural signatures of the C11 epitope subregion. We determined the crystal structure of the complex formed between the gp120 core_e that has extended N- and Ctermini along with the antigen binding fragments (Fab) of two Cluster A antibodies: the A32-like antibody N5-i5 (Guan et al., 2013) and the C11-like antibody N12-i3 (Guan et al.,

2013). This structure confirms that these two epitope regions are non-overlapping and defines new features of the gp120 N-terminus not seen previously. When bound to N12-i3, residues 31–42 of the gp120 N-terminus form an 8th β -strand to extend the 7-stranded β -sandwich formed by the gp120 inner domain seen in other Env structures including the PGT151-stabilized Env trimer (Lee et al., 2016), the CD4-trigered Env trimer (Ozorowski et al., 2017), and the gp120 core with N- and C-termini (Finzi et al., 2010; Pancera et al., 2010). These results indicate the importance of Epitope Cluster A as a major ADCC target for HIV-1 as well as identifying a new N-terminal structure that is implicated as an

Results

N12-i3 and C11 induce potent ADCC against CD4 positive HIV-1 infected cells

intermediate in a late entry state.

We previously derived mAb N12-i3 from memory B cells (BMem) of a Natural Viral Suppressor, an HIV-1 infected individual who suppressed HIV-1 replication to <400 copies/ml in the absence of antiretroviral therapy (Guan et al., 2009; Sajadi et al., 2009; Sajadi et al., 2007). As reported in (Guan et al., 2013) N12-i3 and C11 constitute the most potent ADCC antibodies of the Cluster A region against CEM-NKr-CCR5 target cells sensitized with gp120 (Gomez-Roman et al., 2006). This ADCC method is designed to detect effector functions of antibodies recognizing transitional epitopes exposed during the earliest stage of viral entry, the interaction of the Env trimer with the host cell receptor, CD4 (Guan et al., 2013). To further characterize whether N12-i3 is also capable of mediating ADCC against target cells productively infected by cell-free virus or cell-cell virus spread (Orlandi et al., 2016) we tested its activity in an RFADCC assay against EGFP-CEM-NKr-CCR5-SNAP cells infected for 5 days with a replication competent HIV-1_{BaL} molecular clone (Figure 1). As shown in Figure 1, N12-i3, similarly to C11, A32 and N5-i5, effectively recognizes and induces the killing of only infected or HIV-1-sensitized cells that have not downregulated cell surface CD4 (respectively $p24+_{high}/CD4+$ and $p24+_{low}/CD4+$) and is unable to recognize productively infected cells in which CD4 is downregulated (p24+/CD4cells). This is in agreement with recent reports indicating that epitopes within the Cluster A region are strictly CD4 inducible and become exposed on HIV-1 infected cells only if cell surface CD4 is still available on the same cell to trigger the Env trimer (Ding et al., 2015; Veillette et al., 2014).

N12-i3 binds to Env exclusively within the 8-stranded β -sandwich of CD4-triggered gp120 and recognizes a novel epitope formed at the gp120 N-terminus

We have shown previously that A32- and C11-like epitopes constitute two non-overlapping epitope subregions within the Cluster A region (Gohain et al., 2015; Guan et al., 2013). To describe the C11-like epitope for the first time at the atomic level and to precisely define the A32- and C11-like epitope subregions in context of a single Env antigen, we obtained the crystal structure of the complex formed between CD4-mimetic M48U1-triggered N/C-termini-gp120_{93TH057} core_e and antigen binding fragment (Fab) of representative antibody of each class: the A32-like mAb N5-i5 and the C11-like mAb N12-i3. The complex crystals belonged to space group C2 with two copies of the complex present in the asymmetric unit (Figure S1). The structure was solved at a resolution of 2.94 Å and refined to a final R/R_{free}

of 24.6/30.4%. The data collection and refinement statistics for the structure are summarized in Table 1.

The structure of N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120_{93TH057} core_e-M48U1 complex (Figure 2) confirms that N12-i3 and N5-i5 recognize non-overlapping areas of the CD4triggered gp120. Whereas the A32-like antibody N5-i5 binds to the gp120 Env utilizing the mode of binding and epitope footprint as described previously ((Acharya et al., 2014), Figure S2), the C11-like antibody N12-i3 recognizes a novel epitope, formed almost exclusively through residues of the 7-stranded β-sandwich of the inner domain and the extended N-terminus of gp120 of the C1 and C2 regions. Interestingly, in order to form the N12-i3 epitope, the extended N-terminus of gp120 (residues 31-42) undergoes a structural rearrangement to form a β 4-strand that docks parallel to the β 0-strand of the 7-stranded β sandwich (Figure 2, blow up). As a result of the rearrangement, a new 8-stranded β sandwich structure is formed consisting of the 7-strands contributed by residues of inner domain and the eight strand contributed by residues of the extended N-terminus of gp120. N12-i3 approaches CD4-triggered gp120 and recognizes its cognate epitope within the 8stranded β -sandwich at an approximately 90° angle as compared to the angle of approach of the A32-like antibody N5-i5. The N12-i3 and N5-i5 modes of binding allow both to bind simultaneously as predicted by competition ELISA, FRET-FCS, and SPR (Gohain et al., 2015; Guan et al., 2013).

Figure 3A shows the detailed map of N12-i3 and N5-i5 contacts within gp120. The N12-i3 epitope maps in close proximity to the N5-i5 epitope and is formed by residues, as measured by the buried surface area (BSA), of the newly formed 8-stranded β -sandwich structure (residues 31–40 of the extended N-terminus and 42–43, 45, 84–87 (C1 region), 224, 244–246 (C2 region) and 491 (C5 region) of the 7-stranded β -sandwich) with exception of one residue (Q⁸²) that maps to Layer 1 of inner domain (Figure 3). By binding the edge of the β -sheet, at the eighth strand, N12-i3 is able to grasp both sides burying 1693 Å² total surface area at the interface, the average calculated for two copies of the complex present in asymmetric unit of crystal (Table S1).

The antibody contacts to gp120 are mediated mostly by the heavy chain of N12-i3 (723 Å² of total BSA of the complex, Table S1, Figure S3) and its protruding CDR H2 that binds to the $\overline{\beta}4$ - and $\beta0$ -strands of the 8-stranded β -sandwich (Figure 2). The CDR H2 contacts to the β -strand are predominantly hydrophobic mediated by I^{52a}PIF and M⁵⁶P motifs with key interactions provided by the aliphatic ring of F⁵⁴ that stacks against Y³⁹, L⁴² and P⁴³ of the gp120 N-terminus stabilizing the Y³⁹YGVP turn (Figure 3B). The turn brings the $\overline{\beta}4$ -strand close to the $\beta0$ -strand and allows the formation of the 8-stranded β -sandwich structure. Although the contacts of N12-i3 to the 8-stranded β -sandwich are mediated mostly by its heavy chain and these interactions are strictly protein-protein contacts, the light chain provides possible contacts to the Env glycan shield. From the light chain, CDR L3 is involved in few stabilizing interactions to the $\overline{\beta}4$ - strand, and the CDR L1 is located in close proximity to the truncated sugars molecules, N-acetyl-D-glucosamine (GlcNAc), directly proximal to the asparagine residues at position 88 and 241 (Figure 2). However, in the complex, only GlcNAc attached to N²⁴¹ is involved in interactions (as accessed by 5Å cutoff) to the side chains of S³⁰ and S³¹ of CDR L1 (Figure 3B). Since the truncated sugar

molecules GlcNAc were introduced to help in co-crystallization studies and generated after the enzyme Endoglycosidase H (Endo H) treatment, we tested if N-linked branched wildtype glycans at positions 88 and 241 contribute to the N12-i3 binding. We tested binding of N12-i3 to the CD4-triggered gp120 expressed in wild-type HEK293 cells allowing production of the N-linked wild-type glycans and the CD4-triggered gp120 with sugars truncated to retain only the GlcNAc. As shown in Figure 3C, the affinity of N12-i3 for Env antigen remains unchanged for both CD4-triggered gp120 antigen preparations, indicating that branched glycans at positions 88 and 241 do not contribute to the N12-i3 binding.

The gp120 N-terminus exists in several conformations that change during viral entry

The gp120 subunit in the N12-i3 bound state assumes a unique conformation, not described previously for any of monomeric gp120 or Env trimers preparations studied in the unbound or the Fab-triggered states. Residues 35–39 of the gp120 N-terminus fold into a β -strand (strand β 4) that packs parallel to strand β 0 of the inner domain to form a stable 5-stranded β sheet (strands $\overline{\beta 4}$, $\beta 0$, $\beta 7$, $\beta 5$, and $\beta 25$) in the 8-stranded β -sandwich which also includes strands $\overline{\beta}3$, $\beta1$, and $\beta6$ (Figure 4A and B). The parallel $\overline{\beta}4$ - and $\beta0$ assembly is stabilized by a network of the main chain hydrogen bonds (Figure 4A, blow up). In the only other gp120 complex with full N- and C- termini studied in a context of the ternary Fab-CD4-triggerred gp120 complex, 48d Fab-N/C-termini-gp120_{Hxhc2}core_e- CD4 complex (PDB: 3JWD, (Pancera et al., 2010)), the N- and C- termini form a short antiparallel β-sheet consisting of strands $\overline{\beta}4$ and $\beta26$, respectively, that are separated from the 7-stranded β -sandwich (Figure 4A and B). A similar distribution of secondary structural elements with the $\overline{\beta}4$ - and $\beta26$ strands packing antiparallel and separated from the 7-stranded β -sandwich is also seen in the membrane associated Env pre-fusion soluble trimer, Fab PGT151-EnvJRFL (PDB: 5FUU, (Lee et al., 2016)) and other soluble SOSIP gp140 trimers. Of note, the β -sheet formed by the N-terminus in the N12-i3 complex (W³⁵VTVY³⁹) is nearly twice the length of that formed in the 48d complex (Y⁴⁰GV⁴²) and comparable in length to that seen between the Nand C- termini in the Env pre-fusion trimer (W35VTVYY40). In the N12-i3 Fab/N5-i5 Fab-N/C-termini-gp120_{93TH057} core_e complex the gp120 C-termini is disordered with no interpretable densities found for residues 493–511 of gp120. This indicates that N12-i3 epitope does not involve residues of gp120 C-terminus, and this epitope is likely formed only after the N- and C- termini are separated.

The N12-i3 epitope is highly conserved

We have shown previously based on co-crystal structures for A32 and other the A32-like monoclonal antibodies including N5-i5 that the A32-epitope region is highly conserved. Figure 4C and D shows both the N5-i5 and N12-i3 epitope footprints displayed on the surface of M48U1-triggered N/C-termini-gp120_{93TH057} core_e with surface colored according to protein sequence conservation determined using the Los Alamos HIV Sequence Database (www.hiv.lanl.gov). The coloring scheme ranges from most conserved resides shown in blue to the least conserved shown in red. The N12-i3 epitope maps to a highly conserved region and involves residues almost completely conserved such as W³⁵, which is mutated from the consensus in only about 0.34% of the sequences in the HIV Sequence Database Compendium (https://www.hiv.lanl.gov/content/sequence/HIV/COMPENDIUM/ compendium.html). Additionally, the most conserved residue at the gp120 N-terminus, G⁴¹

(mutated in less than 0.1% of sequences) contributes to the β -turn formation and the G⁴¹VP residues immediately following the 8th β -strand are all mutated in less than 0.2% of sequences. The gp120 N-terminal residues 35–45 which form the major anchoring point for N12-i3 are mutated from the consensus in less than 2% of gp120 sequences (less than 0.5% if one excludes residues 38, 39, and 44). The high conservation of this gp120 region most likely results from the fact that it contributes directly to the formation of the gp120/gp41 heterodimer of the Env trimer. As shown in Figure 4C (right panel), regions of the nascent N12-i3 and N5-i5 epitopes are buried at the heterodimer interface and are involved directly in interaction with the gp41 protomer, as previously suggested by mutagenesis (Finzi et al., 2010).

Variable levels of the N12-i3 and N5-i5 epitope become accessible for antibody recognition during Env trimer opening

The Cluster A region consisting of A32- and C11-epitope subregions is buried in native Env trimers on virions or on the surfaces of HIV-1 infected cells (Acharya et al., 2014; Gohain et al., 2015; Guan et al., 2013; Ray et al., 2014; Veillette et al., 2014). It was shown previously that this epitope region becomes exposed and is accessible for antibody recognition only after binding to cell surface CD4 (Mengistu et al., 2015) or on productively infected cells until CD4 is down-regulated by Nef and Vpu (Alsahafi et al., 2015; Ding et al., 2015; Richard et al., 2015; Veillette et al., 2015; Veillette et al., 2014; Veillette et al., 2016). Once exposed, Epitope Cluster A becomes a very potent ADCC target (Ferrari et al., 2011; Guan et al., 2013; Veillette et al., 2014). Recent data reveal also that the cell surface CD4-induced conformational rearrangements can be very effectively mimicked on the infected cell surface by small compound or peptide derived CD4 mimetics that alone or in presence of coreceptor binding site antibodies are capable of effective sensitization of infected cells to Cluster A antibody binding and ADCC killing (Richard et al., 2016; Richard et al., 2015). CD4 mimetics therefore constitute useful tools to the study of the structural rearrangements of the Env trimer which occur upon trimer triggering with cell surface CD4. On the other hand, our structural analysis indicates that the A32- and C11- epitope regions map to the non-overlapping gp120 regions that are buried at the pre-fusion trimer and may become available for antibody recognition at different levels during the CD4-induced opening of the Env trimer. The region of the nascent A32- epitope is nested inside the center of the prefusion trimer with multiple residues involved directly in contacts to the upper part of α 7 and a9-helix of gp41 ((Acharya et al., 2014) and Figure 4C, right panel). In contrast, residues of the nascent C11 epitope are localized at the bottom of the pre-fusion trimer (close to the viral membrane) with the N-terminus interacting with the gp120 C-terminus, disulfide loop region of the principal immunodominant domain and the β 27 strand the gp41 (Figure 4C, right panel). In the trimer, a 3-stranded antiparallel β -sheet is formed, assembled from the β 27 strand of gp41 and the $\overline{\beta}$ 4 and β 26 strands of gp120. In order to test if there is any difference in the accessibility of the A32- and C11-subregions during Env trimer opening we tested the ability of antibodies of the A32-subregion: A32 and N5-i5 and the C11-subregion: C11 and N12-i3 to recognize Env targets present at the surface of CEM-Nkr cells infected with a wild-type infectious molecular clone coding for functional Nef and Vpu proteins and ADA Env (48 hours post-infection as described and Experimental Procedures). Staining was performed in the presence or absence of the CD4 mimetic peptide M48U1 which was mixed

or not with the co-receptor binding site antibody 17b (Figure 5A and Figure S4). As we reported previously (Richard et al., 2015, Richard et al., 2016) the M48U1 mimetic combined with 17b is able to effectively trigger the conformational rearrangements of Env trimers present on infected cells to expose Cluster A epitopes. However, in the conditions tested (100nM M48U1 and 5 ug/mL 17b) we observed differences in the levels of antibody bound with A32 and the A32-like antibody N5-i5 binding more efficiently than C11 or the C11-like antibody N12-i3. This may indicate that the A32-like epitope region is exposed earlier during the CD4 mimetic-induced opening of Env trimer. Moreover, as shown in Figure 5B, we observed that while a Fab fragment of 17b was sufficient to expose A32 and N5-i5 epitopes within HIV-1_{JR-FL} Env trimers expressed at the surface of 293T cells, it failed to do so for C11 and N12-i3. A 17b Fab 2 or full 17b antibody were absolutely required to expose N12-i3 and C11 epitopes, implying that it is "easier" to expose A32 and N5-i5 epitopes than N12-i3 and C11 epitopes. This suggests that A32 exposure may precede C11 exposure (Figure 5B).

Finally, we tested if the CD4 mimetic M48U1 could be also effective in sensitizing the Env trimers present on pseudovirions for Cluster A antibody binding (Figure 5C). We showed previously (Ray et al., 2014) that triggering of the Env trimers present on pseudovirions with the soluble CD4 (sCD4) is insufficient to expose the Cluster A region. Figure 5C shows the binding of a panel of Cluster A antibodies to pseudovirions expressing the CCR5-tropic HIV-1_{BaL} envelope (a "tier 1" easily neutralized envelope, (Ray et al., 2014) in solution in the presence or absence of serial M48U1 concentrations ranging from 0 to 64 mM by FCS (Figure 5C). In the absence of M48U1, a small fraction of Cluster A mAbs (8 to 15% depending on the mAb) adopted the lower diffusion coefficient (8 μ m²/sec) indicative of virion binding. However, this fraction improved with increasing amounts of M48U1, reflecting higher mAb attachment to HIV-1_{BaL} virion particles. The binding signals for A32 and A32-like antibody N5-i5 increased from 16% to 37% and 15% to 31%, respectively, from 0 to 64 mM of M48U1. In contrast, the binding of C11 and N12-i3 only moderately improved reaching 17% to 27% in the presence of the highest amounts of M48U1. Overall, we observed increased binding of Cluster A mAbs to virions treated with M48U1 as compared to previous reports of Cluster A exposure with soluble CD4 (Ray et al., 2014). Our data is also consistent with the data from infected cells and Env trimers expressed at the surface of 293 cells, showing that the A32 subregion is more efficiently accessible for antibody recognition within HIV_{BaL} virions than the C11 subregion upon triggering with a CD4 mimetic. Of note, we also have tested the binding (data not shown) of the Cluster A antibody panel to HIV_{BaL} virions in presence of M48U1 ($25 \mu g/ml$) and $17b (10 \mu g/ml)$. Under these conditions we observed an increase in the binding of A32 (from 20% to 28% for virions treated with M48U1 alone or for virions treated with M48U1 and 17b in combination) but no noticeable differences in the binding of C11 or N12-i3.

Discussion

Fc γ R- mediated effector functions such as ADCC are thought to constitute an important component of protective immunity against HIV-1. They trigger clearing of virus particles or virus-infected cells through mechanisms involving interactions of the antibody constant (Fc) region with Fc γ R on the surface of effector cells including NK cells, monocytes,

macrophages, dendritic cells, and neutrophils (Sulica et al., 2001; Takai, 2002). ADCC responses appear relatively early during acute infection and are detectable as early as 21 and 48 days after SIVmac251 (Asmal et al., 2011; Sun et al., 2011) infection in macaques and acute HIV-1 (Pollara et al., 2010) infection in humans. These early ADCC responses, which in general are broadly reactive, precede the appearance of neutralizing antibody responses with similar breadth (Koup et al., 1991; Lyerly et al., 1987). ADCC is thought to eliminate virus-infected cells through a mechanism in which antibodies provide specificity for the release of cytotoxic mediators by linking the infected target to the effector cell.

The non-neutralizing, transitional Cluster A epitopes consisting of A32- and C11- and C11/A32 hybrid- epitopes (Guan et al., 2013) constitute major targets for ADCC responses in both acute and chronic HIV-1 infection as well as those induced by vaccination (Ampol et al., 2012; Bonsignori et al., 2012; Ferrari et al., 2011; Guan et al., 2013; Robinson et al., 2005). Early analyses of the antibody repertoire in patients on HAART during acute infection revealed high frequencies of B cells producing anti-Env antibodies directed at CD4i epitope targets that include the Cluster A region, with more than 47% of cells producing antibodies of this type (Robinson et al., 2005). These studies pointed toward CD4i epitopes as the most immunogenic during acute infection. Recent data are consistent with this early observation and confirm that chronically infected individuals also frequently elicit antibodies specific for this region. Although the majority of recent studies examine CD4i targets in context of the A32 epitope (Ferrari et al., 2011), growing evidence points toward the C11-epitope region as at least equally important, contributing significantly to plasmamediated ADCC activity of HIV-infected individuals. Our studies of natural virus suppressors who control HIV-1 without therapy have shown high frequencies of long-lived memory B cells for antibodies recognizing the conserved CD4i epitopes of this type (Guan et al., 2013; Guan et al., 2009). Half of a set of four Cluster A antibodies isolated from five HIV-1– infected individuals recognized the C11-epitope region (Guan et al., 2013). Recently Williams and colleagues (Williams et al., 2015) showed, using their novel strategy to isolate antibodies originated from unique B cell, that antibodies similar to C11 are common in HIVinfected individuals and antibodies targeting this epitope can be commonly produced. In this study, C11-like antibodies accounted for between 18-78% of ADCC activity in 9 chronically infected individuals and they were shown to develop from unique B-cell lineages in the same individual within the first 6 months of infection. The short time required for development as well as usual structural features of C11-like antibodies such as relatively short CDR H3s and low levels of somatic mutation of variable regions (Guan et al., 2013; Williams et al., 2015) support a role of the C11 epitope region as a potential vaccine target. It is possible that vaccination strategies designed to induce this type of antibody response will overcome the great challenges associated with the vaccines aimed to induce a broadly neutralizing antibody response. Interestingly, whereas the A32- epitopes were implicated in the protective effect of ADCC responses in humans induced in the RV144 vaccine trial (Bonsignori et al., 2012; Haynes et al., 2012; Tomaras et al., 2013), the vaccination studies in non-human primate (NHP) using the conformationally constrained gp120-CD4 immunogen (FLSC) (Fouts et al., 2000) directly identified C11-like antibodies as a correlate of protection (DeVico et al., 2007; Fouts et al., 2015).

The A32- and C11- epitopes constitute non-overlapping regions of the gp41-inteactive face of gp120. We describe here the crystal structure of the complex formed between Fab of antibody representing each of these two epitope subregion bound to the same antigen: CD4 mimetic-triggered gp120 core_e. Our structural analysis confirms that A32- and C11-like antibodies can simultaneously bind one CD4-triggered gp120 subunit occupying proximal epitope surfaces. Whereas our A32-like antibody, N5-i5, binds to discontinuous epitope exclusively within the inner domain mobile layers 1 and 2 (Gohain et al., 2015), the C11-like antibody, N12-i3, attaches at approximately 90° angle and recognizes residues of the newly formed 8-stranded β -sandwich of the inner domain of the C1-C2 region. N12-i3 binds at the top of the sandwich engaging the extended N-terminus (residues 31–42) to add an 8th strand to the 7-stranded β -sandwich structure known to be formed in pre-fusion Env trimers or monomeric gp120 studied in the context of other Fab-triggered states.

The structure of N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120_{93TH057} core_e-M48U1 complex provides the first look into molecular details of C11-epitope region in absence of structural information of the epitope of the prototype antibody C11. Although the exact epitope footprints of these two antibodies may not be exactly the same, they certainly recognize the same structural elements of CD4-triggered Env. We recently defined the putative contact region of C11 (Gohain et al., 2015) by ELISA, FCS-FRET, SPR competition assays and analyses of residues shown by mutagenesis to be involved in C11 binding (Finzi et al., 2010; Moore et al., 1994). Our studies mapped to the 7-stranded β -sandwich in the gp41interactive region, immediately adjacent to the A32-like epitope surface (Acharya et al., 2014; Gohain et al., 2015). Our predicted putative binding site overlaps well with N12-i3 epitope with most of residues defined by mutagenesis to form C11 epitope contributing also to the N12-i3-gp120_{93TH057} core_e interface. However, one structural signature of the C11 epitope region was missing at that time and could not be simply predicted based on available structural information: the formation of 8-stranded β -sandwich structure.

The N12-i3 epitope, similar to A32 subregion epitopes, maps to highly conserved regions of gp120 (only approximately 16% of sequences differ from Hxbc2 for residues 31–45 and only approximately 0.4% for residues 35–45). This high degree of conservation among HIV sequences predicts the broad reactivity of antibodies that target this region across HIV clades. Indeed, recently demonstrated strong cross-clade breadth with activity against 10 of 11 envelopes tested, including those from clades A, B, C, A/D and C/D was reported for two C11-like mAb isolated from HIV-specific memory B cells from a subtype A-infected Kenyan woman (Williams et al., 2015).

The 8-stranded β -sandwich structure as seen in the N12-i3 Fab/N5-i5 Fab- N/C-terminigp120_{93TH057} -core_e-M48U1 complex is unique, formed late during CD4-induced Env trimer opening most likely post-detachment of gp41 and gp120 subunits. We and others have shown that soluble CD4 induces a metastable conformation of the trimer in which gp120 protomers are displaced but still remain involved in the gp41 interaction thus incapable of exposure of the A32- and C11-epitope regions (Acharya et al., 2014; Ozorowski et al., 2017; Pancera et al., 2010; Wang et al., 2016). We are able, however, to induce the conformational changes of the viral trimer similar to those induced by cell surface CD4 with the CD4 miniprotein mimetic M48U1 alone or in combination with co-receptor binding site antibody.

M48U1 was initially developed to interfere with the initial steps of viral attachment to the CD4 cell receptor thus is capable of virus neutralization (Martin et al., 2003; Van Herrewege et al., 2008). Our earlier structural analysis confirm that M48U1 targets the Phe 43 cavity in a manner similar to that of CD4 but its cyclohexylmethoxyl group reaches deeper into the cavity. The increased surface area buried and improved shape complementarity of this group within the Phe43 pocket was found to contribute to M48U1 effectiveness in binding to HIV Env and CD4 antagonism (Stricher et al., 2008). The same feature most likely also contributes to the efficiency of M48U1 in trimer disassembly and exposure of Cluster A epitope region.

Our analyses of the binding of A32- and C11-like mAbs to the CD4-mimetic-M48U1 (or M48U1/17b mix)-triggered envelope trimers present on the infected cell surface, expressed at the surface of transfected cells or pseudo-viruses indicate that A32 epitope subregion is exposed more easily than the C11 subregion. At any given concentration of M48U1 (or M48U1/17b mix) we observed a better recognition by A32-like antibodies as compared to C11-like antibodies. This indicates the possibility that the A32-epitope is accessible earlier during CD4-induced opening of the HIV-1 Env trimer. Our recent data using confocal and super-resolution microscopy to track the exposure of Cluster A epitopes as they appear on single CCR5-tropic HIV-1 particles bound to target cell indicate that both the A32 and C11 epitope subregions become accessible for antibody recognition very rapidly upon target cell binding (Mengistu et al., 2015). The same level of accessibility of both epitopes was detected five minutes post-attachment of virus to the CD4+ target cells regardless of presence or absence of co-receptor. This indicates that structural rearrangements exposing both epitope structures occur very rapidly, most likely in the co-receptor independent manor.

Based on our data we propose the following model of the HIV-1 Env trimer rearrangements that lead to the exposure of the Cluster A epitopes during the entry process (Figure 6). In the pre-fusion Env trimer both the A32 and C11 subregion are occluded by the interaction between gp120 and gp41 and the N- and C- termini of gp120 clasped by gp41 (Figure 6, conformation from PDB: 5FUU, (Lee et al., 2016)). In this state, the gp120 inner domain is packed against a gp41 protomer preventing it from collapsing into a 6-helix bundle - the driving force behind membrane fusion. In the pre-fusion trimer, the N- and C- termini of gp120 are held by a triple-tryptophan clasp consisting of tryptophans 623, 628, and 623 of gp41 (Pancera et al., 2014). Upon binding of the Env trimer to the cell surface CD4, the inner domain of gp120 undergoes a series of conformation rearrangements with the formation of the bridging sheet and changes to the region of the nascent N5-i5 epitope (conformation from PDB: 5VN3, (Ozorowski et al., 2017)). In this state, the a0-helix and the $\overline{\beta}1$ -strand of the N5-i5 epitope are formed and the $\beta1$ helix is positioned to assume conformation as seen in the N5-i5-bound state. Although in this Env conformation the N5-i5 epitope is largely formed, it is not accessible for antibody recognition due to occlusion at the trimer interface. At this step, the 7-stranded β -sandwich remains intact with the N-, Ctermini occupying the gp41 tryptophan clasp, thus the N12-i3 epitope is not formed. Later in the entry process gp120 protomers are displaced from gp41 allowing the exposure of the A32 epitope subregions (PDB: 3JWD, (Pancera et al., 2010)). In this stage, the N5-i5 epitope is fully formed and exposed for antibody binding but not the N12-i3 epitope as the 8-stranded β -sandwich structure is yet to be assembled. Finally, the N- and C- termini are

released from gp41, and are most likely separated, permitting the addition of the 8th β -strand to the gp120 inner domain 7-stranded β -sandwich and formation of the N12-i3 epitope (the conformation of gp120 as observed in the N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120_{93TH057} -core_e-M48U1 complex).

STAR methods

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact: Marzena Pazgier, mpazgier@ihv.umaryland.edu

EXPERIMENTAL MODEL AND SUBJECT DETAILS

RFADCC and cell-surface staining was performed using EGFP-CEM-NKr-CCR5-SNAP cell line (Orlandi et al., 2016). EGFP-CEM-NKr-CCR5-SNAP cells were grown in R10 medium at 37°C.

METHOD DETAILS

Protein preparation and crystallization

N12-i3 and N5-i5 Fabs and gp120 protein for use in crystallization were prepared as previously described (Acharya et al., 2014; Gohain et al., 2015). N12-i3 only crystallized with gp120 in the presence of both N12-i3 Fab and fab of the A32-like antibody N5-i5. The dual Fab complex was made sequentially by combining gp120, M48U1, and N5-i5 Fab with a slight molar excess of N5-i5 Fab, purification by gel filtration on a Superdex 200 gel filtration column (GE Healthcare) (equilibrated with 5 mM Tris-HCI pH 7.2 and 150 mM sodium chloride), addition of N12-i3 Fab in slight molar excess to the purified complex, and purification again by gel filtration on the same gel filtration column.

Initial crystallization trials were performed in hanging drop crystallization trays with commercially available crystallization screens from Molecular Dimensions and Hampton Research. Crystals from screens were then optimized manually to produce crystals suitable for data collection. The N12-i3 Fab/N5-i5 Fab- N/C-termini-gp12093TH057 -coree-M48U1 complex crystals were grown from 15% PEG 2000 MME, 0.1 M Tris-HCl pH 8.0, and 0.1 M potassium chloride, and crystals were flash frozen in liquid nitrogen after a brief soak in crystallization buffer plus cryoprotectant, 15% MPD.

Data collection and structure solution

Data were collected on a PILATUS detector at SSRL beam line 12–2. The N12-i3/N5-i5 Fab complex crystallized in monoclinic space group C2 with cell dimensions a = 311.0 Å, b = 53.3 Å, c = 223.6 Å, and $\beta = 128.9$. The N12-i3/N5-i5 complex was solved by molecular replacement using Fab and gp120 coordinates from the N5-i5-gp12093TH057coree-CD4 structure (PDB: 4H8W) with PHASER (McCoy et al., 2007) from the CCP4 suite (Winn et al., 2011). Refinement was done with REFMAC (Murshudov et al., 2011) and PHENIX (Adams et al., 2002). Model building was done using COOT (Emsley and Cowtan, 2004). Data collection and refinement statistics are shown in Table 1.

RFADCC and cell-surface staining

Infection, staining and RFADCC were performed as described previously (Gohain et al., 2015; Orlandi et al., 2016). In summary, EGFP-CEM-NKr-CCR5-SNAP cells were spinoculated for 2 h at 2000 RPM at 12 °C in 96-well U-bottom plate (5 × 105 cells/well) with 240 ng of IMC BaL virus (control cells were incubated without virus), as measured by HIV-1 p24 antigen capture ELISA. As previously described (Gohain et al., 2015) the peak of infection was reached culturing the cells in the presence of Bal IMC at 37°C for 5 days, when EGFP-CEM-NKr-CCR5-SNAP cel ls were harvested, washed twice with R10 medium and divided in two groups: The first group was stained with N12-i3, A32, N5-i5, C11 and Synagis (MedImmune) mAbs labeled with human IgG Zenon-APC kit (Invitrogen), Live/ Dead Fixable Near-IR Dead Cell Stain (Molecular Probes) and (eFluor 450)-conjugated mouse anti-CD4 OKT4 mAb (eBioscience) for 30 min at RT. After washing and permeabilizing (Perm-Fix, BD-PharMingen,) the cells were stained with (PE)-conjugated mouse anti-p24 mAb (KC57-RD1; Beckman Coulter, Inc.). After two washes, HIV-1-infected or mock cells were fixed in 2% PFA.

The second group of Mock and Bal-infected EGFP-CEM-NKr-CCR5-SNAP cells was stained with SNAP-Surface Alexa Fluor 647 dye and utilized as targets for the RFADCC (Orlandi et al., 2016). Serial dilutions of N12-i3, A32, N5-i5, C11 and Synagis were incubated with mock and Bal-infected targets for 15 mins at RT and, subsequently with PBMC as effectors. The cells were allowed to interact for 3 hours at 37°C, at the end of which the cells were washed once with PBS and fixed with 2% PFA. Binding and RFADCC assays were then analyzed with an LSRII Fortessa flow cytometer (BDBiosciences). Data analysis was performed with FlowJo software (Tree Star, Inc., San Carlos, Calif.).

Infection and staining of CEM-NKr cells

The infectious molecular clone NL4.3 coding for GFP, the ADA envelope and functional Nef and Vpu proteins was used as previously described (Veillette et al., 2014). In order to achieve 20% of infection of CEM-NKr cells, the proviral vector and a VSVG-encoding plasmid were co-transfected in 293T cells by standard calcium phosphate transfection. Concentrated virus (over a 20% sucrose cushion) used to infect CEM-NKr cells by spin infection, as described (Veillette et al., 2014). Forty-eight hours post infection, cell-surface staining was performed as previously described and MFI histograms shows signal on live infected populations (Richard et al., 2015; Veillette et al., 2015). Binding of HIV-1-infected cells by Alexa-Fluor 647-conjugated Cluster A mAbs A32, N5-i5, N12-13 and C11 or coreceptor binding site 17b Ab was performed in presence of M48U1 (100 nM) or with equivalent volume of vehicle (PBS). Binding of HIV-1-infected cells with Alexa-Fluor 647conjugated Abs A32, N5-i5, N12-i3, C11 and 17b was performed alone or in combination with unconjugated 17b (5µg/ml). Detection of GFP+ infected cells was performed as described (Richard et al., 2015). The percentage of infected cells (GFP+) was determined by gating the living cell population based on the viability dye staining (Aqua Vivid, Thermo Fisher Scientific). Samples were analyzed on a LSRII cytometer (BD Biosciences, Mississauga, ON, Canada) and data analysis was performed using FlowJo vX.0.7 (Tree Star, Ashland, OR, USA).

To express the wild-type HIV-1JRFL Env for flow cytometric analysis, a stop codon was introduced into the expression plasmid for the codon-optimized Env, replacing the codon for Gly 711. This modification results in the expression of Env with a deletion of the cytoplasmic tail, which increases the level of Env on the cell surface. Env expression plasmids were transfected into 3×105 293T cells along with a pIRES-GFP vector at a ratio of 2 µg Env to 0.5 µg green fluorescent protein (GFP) expression plasmid. 16 h after transfection, cells were washed with fresh media and Env exposure was evaluated 24 h later. A32, N5-i5, N12-i3, and C11 were used to detect CD4-induced exposure and each of the antibodies was conjugated to an Alexa-Fluor 647 (AF647) probe (Invitrogen). Transfected 293T cells were incubated with either the 2G12 antibody (1 μ g/ml), which recognizes the outer domain of gp120, or AF647-A32, AF647-N12-i3, AF647-C11 (4 µg/ml) in the absence or presence of M48U1 (100 nM) or 17b, 17bFab1, and 17bFab2 (5 µg/ml) for 1-hour at room temperature. The 2G12 and 17b antibodies were detected using a goat anti-human antibody coupled to Alexa Fluor 647 (Invitrogen). Antibody binding was detected by gating on GFP-positive cells on an LSRII cytometer (BD Biosciences, Mississauga, ON, Canada). Data analysis was performed using FlowJo vX.0.7 (Tree Star, Ashland, OR, USA).

Fluorescence correlation spectroscopy measurements

FCS analyses were performed similar to previously reported (Gohain et al., 2015; Ray et al., 2014). mAbs A32, N5-i5, C11, N12-i3 and 17b were labeled with Alexa 647 probe with a dye-to-protein ratios of 3 for FCS experiments. HIV-1BaL pseudoviruses were produced by co-transfection of HEK293T cells with an Env-deficient HIV-1 backbone plasmid, pNL4-3-E-EGFP, along with Env expression plasmid pHIV-1-BaL 0.1. All FCS experiments used virus preparations diluted to a 10 µg/ml p24 equivalent. gp120 to p24 antigen ratios were typically 1:10 to 1:50. The 10 µg/ml p24 equivalent value typically corresponded to TCID50/ml values in the range of 200,000 to 650,000 (HIV-1BaL). Pseudoviruses with 10µg/ml p24 equivalent concentrations in 25µl reaction buffer were first treated for 90 min at 37°C with different concentrations (0 to 64 mM) of M48U1 to open up the cluster A epitopes and 100 µg/ml of nonspecific (Calbiochem) IgG1 to block nonspecific binding. This was followed by addition of 5 μ g/ml of Alexa Fluor 647-conjugated mAbs. MAbs were allowed to interact with virions for 90 min at 37°C. FCS measurements were performed in a confocal microscope (ISS Q2) with a high-numerical-aperture (NA) water objective (60x; NA 1.2). The excitation source was a Fianium SC-400 super-continuum laser with a NKT super-select AOTF filter. The fluorescence was collected by avalanche photodiodes through a band-pass (650 – 720 nm; Chroma) filter and a B&H SPC-150 card operated in a photon time-tag time-resolved (TTTR) mode. ISS VistaVision software was used to analyze the FCS data to assess the in vitro binding of mAbs to HIV-1 BaL pseudovirus particles by determining the translational diffusion coefficients. The percentage of total mAb (at given test concentrations) that shifts into the more slowly diffusing species which is the virionbound fraction reflects the relative magnitude of cognate epitope exposure in the target population of virions (Ray et al., 2014).

Surface plasmon resonance (SPR) measurements

Binding affinities of N12-i3 to full length single chain (FLSC) were assessed by surface plasmon resonance on a Biacore T-100 (GE Healthcare) as previously described (Tolbert et al., 2016). Varying concentrations of single-chain gp120BaL-sCD4 (FLSC) (Fouts et al., 2000) either expressed in 293 cells (FLSC with wild type glycosylation) or in GnT1- cells and deglycosylated with Endo Hf (New England Biolabs) (deglycosylated FLSC) were passed over N12-i3 IgG bound to a protein A coupled sensor chip. Binding affinities were calculated using a 1:1 binding model with the BIAevaluation software.

QUANTIFICATION AND STATISTICAL ANALYSIS

Antibody binding analyses

Data are presented as mean and standard deviation of the mean fluorescence intensity over mock (MFI/Mock). Data are the averages from four independent experiments. Statistical significant was evaluated using paired student t test, * P < 0.05, ** P < 0.01, ***P < 0.001, **** P < 0.00001; ns, not significant.

DATA AND SOFTWARE AVAILABILITY

Accession Numbers

The atomic coordinates for the structures presented in this study have been deposited in the Protein Data Bank under ID code: 5W4L

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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• the C11-like ADCC epitope footprint is defined at the molecular level

- the C11 subregion maps to a newly formed gp120 8-stranded β -sandwich
- the C11 and A32 epitope subregions are adjacent and non-overlapping
- the C11 epitope is formed upon release of the gp120 N-terminus from the Env trimer

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Figure 1. Binding of N12-i3 to EGFP-CEM-NKr-CCR5-SNAP cells infected with BaL infectious molecular clone (IMC)

EGFP-CEM-NKr-CCR5-SNAP were infected with IMC BaL virus and evaluated 5-days post-infection for (**A**) intracellular p24, cell surface levels of CD4 and (**B**) the binding of: N12-i3, Cluster A mAbs A32, N5-i5 and C11, and Synagis. Panel A shows the gating strategy for EGFP-CEM-NKr-CCR5-SNAP Bal-infected (Blue gates) and Mock (Red gate) cells, previously defined as GFP+ and L/D dye- cells. For Bal-infected cells two gates were delineated: p24+/CD4- and p24+_{low}/CD4+; one gate was delineated for Mock p24-/CD4+. (**C**) ADCC using the RFADCC method. The curves shown are normalized for plateau cytotoxicity values using C11.



Figure 2. Crystal structure of the N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120_{93TH057} -core_e-M48U1 complex

Only complementary determining regions (CDRs) involved in N12-i3 Fab binding are colored and shown. The N-terminus of gp120 (black) forms an eighth strand to the 7-strand β -sandwich (magenta) of the inner domain to create the N12-i3 epitope. See also Table S1 and Figure S1 and S2.

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Figure 3. N12-i3 epitope footprint

(**A**)N12-i3 (green) and N5-i5 (blue) gp120 contact residues mapped as spheres on the gp120 ribbon diagram and epitope footprints plotted on the gp120 surface with layers colored as in Figure 1. gp120 side chain (+) and main chain (–) contact residues colored green for hydrophobic, blue for hydrophilic and black for both as determined by a 5 Å cut off value over the corresponding sequence. Buried surface residues as determined by PISA are shaded. See also Figure S3. (**B**) Blow up view of the N12-i3 interaction with sugars at positions 88 and 241 (top) and the 8-stranded β -sandwich formation (bottom). CDR L1 and CDR H2 of the N12-i3 Fab are shown with the side chains of CDR L1 potentially contributing to contacts with the sugar at position 241 and the side chains of CDR H2

contributing to stabilization of the $Y^{39}YGVP$ turn of the 8-stranded β -sandwich shown as sticks. (C) Binding of N12-i3 to glycosylated and deglycosylated single-chain gp120BaL-sCD4 complex (FLSC). Surface plasmon resonance sensorgrams and binding kinetics are shown with wild type glycosylated FLSC – red curves and GnT1 cell grown deglycosylated FLSC – blue curves. Program fits to each sensogram are shown with black lines.

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^{466 471 476 481 486 491 496} NNESEIFRPGGGDMRDNWRSELYKYKVVKIEPLGVAPTK

Figure 4. Properties of the N12-i3 epitope

(A) Comparison of the gp120 conformation in the N12-i3 Fab/N5-i5 Fab- N/C-terminigp120_{93TH057} -core_e-M48U1 complex to the gp120 conformations in the Fab 48d-N/Ctermini-gp120_{Hxbc2}core_e-CD4 structure (PDB: 3JWD) and in the Env pre-fusion trimer, Fab PGT151-Env_{JRFL} (PDB: 5FUU). A blow-up view shows a network of hydrogen bonds stabilizing the interaction of $\overline{\beta}4$ - and β 0-strands. (**B**) The topology of the inner domain and N/C-termini of gp120 from the available crystals structures of gp120. The N-terminus as seen in the N12-i3 Fab/N5-i5 Fab complex structure is shown in black and the point of the C-terminus truncation is indicated by an arrow over the topology of the inner domain and

N/C-termini from the Fab 48d-N/C-termini-gp120_{Hxbc2}core_e-CD4 structure (PDB: 3JWD). (**C**) and (**D**) Sequence conservation among residues forming the N12-i3 and N5-i5 epitope. Residues colored dark blue differ from the Hxbc2 sequence in a range of 0.2–7% and residues colored red differ in a range of 87–99.9%. Only unique sequences in the database having an equivalent residue at each position were included in the calculated percentage representing approximately 32,000 sequences on average. (**C**) Ribbon diagram and molecular surface of the gp120 core_e from the N12-i3 Fab/N5-i5 Fab complex (left) and gp120/gp41 heterodimer of the Env pre-fusion trimer, Fab PGT151-Env_{JRFL} (PDB: 5FUU) (right). N12-i3 and N5-i5 epitope footprints and regions of the nascent N12-i3 and N5-i5 epitopes are shown over the molecular surfaces of gp120 core_e (solid line) and gp120/gp41 heterodimer, respectively (broken line). (**D**) Sequence, including variable regions, of gp120_{JRFL}. The N5-i5 and N12-i3 epitope footprints are highlighted with a solid line for N12-i3 and broken line for N5-i5.

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Figure 5. Exposure of A32-like and C11-like epitopes at the surface of HIV-1 infected cells and HIV-1 virions

(A) CEM.NKr cells were infected with the infectious HIV-1NL-4.3 ADA/GFP molecular clone; 48h post-infection cells were stained with Alexa Fluor 647-conjugated mAbs: 17b, A32, N5-i5, N12-i3 and C11 alone, in the presence of the CD4 mimetic M48U1 (100nM) or together with M48U1 and co-receptor binding site 17b (5µg/ml). Data are the averages of four independent experiments. (**B**) Binding to HIV-1_{JR-FL} Env trimers expressed at the surface of 293T cells. At 48-hours post-transfection, the exposure of Cluster A epitopes was assessed with Alexa Fluor 647 (AF647)-conjugated A32, N12-i3, or C11 antibodies in the presence or absence of 17b, the 17b Fab-fragment, or the Fab-2 fragment with or without

CD4mimetic-M48U1 (100nM). Data are the averages from four independent experiments. Statistical significant was evaluated using paired student t test, * P < 0.05, ** P < 0.01, ***P < 0.001; *** P < 0.0001; ns, not significant. See also Figure S4 and S5. (C) Effect of M48U1 on the binding of Cluster A mAbs to BaL pseudovirus particles in solution as determined by the FCS assay. The relative fraction of mAb that adopts a lower diffusion coefficient as a result of virion binding is shown. All measurements were performed in triplicate. Average values are shown; error bars indicate standard deviations.



Figure 6. Model of the exposure of the Cluster A epitopes during the entry process

In the pre-fusion Env trimer both epitopes are occluded by the interaction between gp120 and gp41 with the gp120 N and C termini clasped by gp41 (left panel). Binding of CD4 to the outer domain of gp120 induces formation of the co-receptor binding site which includes conformational changes to the gp120 inner domain (panel two from left). These changes to mobile layers 1 and 2 of the inner domain create the N5-i5 epitope and disrupt the gp120-gp41 interaction (panel three from left). Binding of co-receptor or further opening up of the trimer disengage the N-terminus of gp120 from the gp41 clasp. The N-terminus is then free to form the eighth strand of the inner domain β -sandwich creating the N12-i3 epitope (rightmost panel). Figures were assembled using available structures of the Env pre-fusion trimer (Cry-EM structure of cleaved wild type JR-FL Env trimer in complex with PGT151 Fab, PDB: 5FUU), CD4-triggered Env trimer (Cryo-EM structure of a B41 SOSIP Env trimers in complex with CD4 and antibody 17b, PDB: 5VN3) and 48d Fab-N/C-termini-gp120_{Hxbc2}core_e-CD4 complex (PDB: 3JWD) on which N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120_{93TH057} -core_e-M48U1 complex was superimposed. The bottom panels show the gp120 subunit only.

Table 1

Data collection and refinement statistics

	N12-i3 Fab/N5-i5 Fab- N/C- termini-gp120 _{93TH057} -core _e - M48U1
Data collection	
Wavelength, Å	0.9795
Space group	C2
Cell parameters	
a, b, c, Å	311.0, 53.3, 223.6
$\alpha, \beta, \gamma, ^{\circ}$	90, 128.9, 90
Complex/a.u.	2
Resolution, (Å)	50-2.94 (3.0-2.94)
# of reflections	
Total	229,966
Unique	62,153
R_{merg}^{b} , %	18.4 (92.3)
I/σ	6.0 (1.1)
Completeness, %	99.8 (99.9)
Redundancy	3.7 (3.8)
Refinement Statistics	
Resolution, Å	50.0-2.94
R ^{<i>C</i>} , %	23.1
R _{free} ^d , %	29.1
# of atoms	
Protein	18,731
Ligand/Ion	360
Water	20
Overall B value $(\text{\AA})^2$	
Protein	52.4
Ligand/Ion	63.3
Water	27.5
Root mean square deviation	
Bond lengths, Å	0.005
Bond angles, $^{\circ}$	1.14
Ramachandran ^e	
favored, %	88.8
allowed, %	8.0
outliers, %	3.2

Values in parentheses are for highest-resolution shell

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 $b_{Rmerge} = \Sigma | I - \langle I \rangle | \Sigma I$, where *I* is the observed intensity and $\langle I \rangle$ is the average intensity obtained from multiple observations of symmetry-related reflections after rejections

 $C_{R=\Sigma \parallel F_{O} \mid - \mid F_{C} \parallel / \Sigma \mid F_{O} \mid}$, where F_{O} and F_{C} are the observed and calculated structure factors, respectively

 d_{Rfree} = as defined by (Brunger, 1997)

^eCalculated with MolProbity (Chen et al., 2010)

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER	
Antibodies			
Human monoclonal N12-i3	Isolated from patient (Guan et al., 2013)	HIV Database: record number 2973	
Human monoclonal N5-i5	Isolated from patient (Guan et al., 2013)	HIV Database: record number 2971	
Human monoclonal A32	Pazgier lab	HIV Database: record number 660	
Human monoclonal C11	Pazgier lab	HIV Database: record number 707	
Human monoclonal 17b	Pazgier lab	HIV Database: record number 658	
Human monoclonal 2G12	Finzi lab	N/A	
Synagis (Palivizumab)	Medmiune	N/A	
(eFluor 450)-conjugated mouse anti-CD4 OKT4 mAb	eBioscience	Cat# 48-0048-42	
(PE)-conjugated mouse anti-p24 mAb (KC57-RD1)	Beckman Coulter, Inc	Cat# 6604667	
Bacterial and Virus Strains			
infectious molecular clone BaL virus	(Orlandi et al., 2016)	IMC Bal virus	
infectious molecular clone NL4.3 coding for GFP, the ADA envelope and functional Nef and Vpu	(Veillette et al., 2014)	N/A	
Chemicals, Peptides, and Recombinant Proteins			
full length single chain protein	(Fouts et al., 2000)	FLSC	
M48U1	(Martin et al., 2003)	CD4-peptide mimetic	
Endo H _f	New England BioLabs	Cat#P0703S	
Critical Commercial Assays			
human IgG Zenon-APC kit	Invitrogen	Cat# Z25451	
Live/Dead Fixable Near-IR Dead Cell Stain	Molecular Probes	Cat# L10119	
Fixation/Permeabilization Solution Kit	BD-Bioscience	Cat# 554714	
SNAP-Surface Alexa Fluor 647	New England BioLabs	Cat# S9136S	
Aqua Vivid	Thermo Fisher Scientific	Cat# L43957	
Deposited Data			
N12-i3 Fab/N5-i5 Fab- N/C-termini-gp120 _{93TH057} - core _e -M48U1 complex	This work	5W4L	
Fab 48d-N/C-termini-gp120 _{Hxbc2} coree-CD4 structure	(Pancera et al., 2010)	3JWD	
Fab PGT151-Env _{JRFL} structure	(Lee et al., 2016)	5FUU	
Cryo-EM structure of a B41 SOSIP Env trimers in complex with CD4 and antibody 17b	(Ozorowski et al., 2017)	5VN3	
N5-i5 Fab- gp12093TH057 -coree-d1d2CD4 structure	(Acharya et al., 2014)	4H8W	
Experimental Models: Cell Lines	1		
EGFP-CEM-NKr-CCR5-SNAP cells	(Orlandi et al., 2016)	N/A	
CEM-NKr cells	The following reagent was obtained through the AIDS Research and Reference Reagent Program, Division of	N/A	

REAGENT or RESOURCE	SOURCE	IDENTIFIER	
	AIDS, NIAID, NIH: CEM.NKR-CCR5 from Dr. Alexandra Trkola		
HEK293T cells	ATCC	ATCC® CRL-3216 TM	
Recombinant DNA			
pIRES-GFP vector	Clontech	Cat#6029–1	
pNL4-3- E-EGFP plasmid	(Ray et al., 2014)	N/A	
pHIV-1-BaL 0.1 plasmid	AIDS Research and Reference Reagent Program, Division of AIDS, NIAID	N/A	
Software and Algorithms			
FlowJo software	Tree Star,Inc., San Carlos, Calif.	https://www.flowio.com	
FlowJo vX.0.7 software	Tree Star, Ashland, OR, USA	http://docs.flowio.com/vx/faq/qeneral-faq/tree-star-flowio	
ISS Vista Vision software	ISS, Inc., Champaign, IL, USA	http://www.iss.com/microscopv/software/vistavision.html	
PHASER	(McCoy et al., 2007)	$http://www.phaser.cimr.cam.ac.uk/index.php/Phaser_Crystallographic_Software$	
CCP4	(Winn et al., 2011)	http://www.ccp4.ac.uk	
REFMAC	(Murshudov et al., 2011)	http://www.ccp4.ac.uk/html/refmac5/description	
PHENIX	(Adams et al., 2002)	https://www.phenix-online.orq	
Coot	(Emsley and Cowtan, 2004)	http://www2.mrc-lmb.cam.ac.uk/Personal/pemslev/coot/	
MolProbity	(Chen et al., 2010)	http://molprobitv.biochem.duke.edu/	