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Co-evolution of a broadly neutralizing HIV-1 antibody and founder virus

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Supplementary information includes Supplementary Tables 1–15 and Supplementary Figures 1–11.

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The GenBank accession numbers for 292 CH505 Envs are KC247375-KC247667, for 459 V_HDJ_H are 174 V_LJ_L sequences of antibody members in CH103 clonal lineage are KC575845-KC576303 and KC576304-KC576477, respectively. Coordinates and structure factors for unbound CH103 Fab as well as CH103 Fab in complex with ZM176.66 outer domain have been deposited with the Protein Data Bank under accession codes, 4JAM for CH103 Fab and 4JAN for CH103-gp120 complex.

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Abstract

Current HIV-1 vaccines elicit strain-specific neutralizing antibodies. However, cross-reactive neutralizing antibodies arise in ~20% of HIV-1-infected individuals, and details of their generation could provide a roadmap for effective vaccination. Here we report the isolation, evolution and structure of a broadly neutralizing antibody from an African donor followed from time of infection. The mature antibody, CH103, neutralized ~55% of HIV-1 isolates, and its co-crystal structure with gp120 revealed a novel loop-based mechanism of CD4-binding site recognition. Virus and antibody gene sequencing revealed concomitant virus evolution and antibody maturation. Notably, the CH103-lineage unmutated common ancestor avidly bound the transmitted/founder HIV-1 envelope glycoprotein, and evolution of antibody neutralization breadth was preceded by extensive viral diversification in and near the CH103 epitope. These data elucidate the viral and antibody evolution leading to induction of a lineage of HIV-1 broadly neutralizing antibodies and provide insights into strategies to elicit similar antibodies via vaccination.

Induction of HIV-1 envelope (Env) broadly neutralizing antibodies (BnAbs) is a key goal of HIV-1 vaccine development. BnAbs can target conserved regions that include conformational glycans, the gp41 membrane proximal region, the V1/V2 region, glycans-associated C3/V3 on gp120, and the CD4 binding site (CD4bs)^{1–9}. Most mature BnAbs have one or more unusual features (long heavy chain third complementarity determining regions [HCDR3s], polyreactivity for non-HIV-1 antigens, and high levels of somatic mutations) suggesting substantial barriers to their elicitation^{4,10–13}. In particular, CD4bs BnAbs have extremely high levels of somatic mutation suggesting complex or prolonged maturation pathways^{4–7}. Moreover, it has been difficult to find Envs that bind with high affinity to BnAb germline or unmutated common ancestors (UCAs), a trait that would be desirable for candidate immunogens for induction of BnAbs^{7,14–18}. Whereas it has been found that Envs bind to UCAs of BnAbs targeting gp41 membrane proximal region^{16,19}, and to UCAs of some V1/V2 BnAb²⁰, to date, heterologous Envs have not been identified that bind the UCAs of CD4bs BnAb lineages^{7,18,21–23}, although Envs that bind CD4bs BnAb UCAs should exist²¹.

Eighty percent of heterosexual HIV-1 infections are established by one transmitted/founder (T/F) virus²⁴. The initial neutralizing antibody response to this virus arises approximately 3

months after transmission and is strain-specific^{25,26}. This antibody response to the T/F virus drives viral escape, such that virus mutants become resistant to neutralization by autologous plasma^{25,26}. This antibody-virus race leads to poor or restricted specificities of neutralizing antibodies in ~80% of patients; however in ~20% of patients, evolved variants of the T/F virus induce antibodies with considerable neutralization breadth, e.g. BnAbs^{2,20,27–33}.

There are a number of potential molecular routes by which antibodies to HIV-1 may evolve, and indeed, types of antibodies with different neutralizing specificities may follow different routes^{6,11,15,34}. Because the initial autologous neutralizing antibody response is specific for the T/F virus³¹, some T/F Envs might be predisposed to binding the germline or unmutated common ancestor (UCA) of the observed BnAb in those rare patients that make BnAbs. Thus, although neutralizing breadth generally is not observed until chronic infection, a precise understanding of the interplay between virus evolution and maturing BnAb lineages in early infection may provide insight into events that ultimately lead to BnAb development. BnAbs studied to date have only been isolated from individuals who were sampled during chronic infection^{1,3–7,20,27,29}. Thus, the evolutionary trajectories of virus and antibody from the time of virus transmission through the development of broad neutralization remain unknown.

We and others have proposed vaccine strategies that begin by targeting unmutated common ancestors (UCAs), the putative naïve B cell receptors of BnAbs with relevant Env immunogens to trigger antibody lineages with potential ultimately to develop breadth^{6,11,13–16,18,19,21}. This would be followed by vaccination with Envs specifically selected to stimulate somatic mutation pathways that give rise to BnAbs. Both aspects of this strategy have proved challenging due to lack of knowledge of specific Envs capable of interacting with UCAs and early intermediate (I) antibodies of BnAbs.

Here we report the isolation of the CH103 CD4bs BnAb clonal lineage from an African patient, CH505, who was followed from acute HIV-1 infection through BnAb development. We show that the CH103 BnAb lineage is less mutated than most other CD4 binding site BnAbs, and may be first detectable by as early as 14 weeks after HIV-1 infection. Early autologous neutralization by antibodies in this lineage triggered virus escape, but rapid and extensive Env evolution in and near the epitope region preceded the acquisition of plasma antibody neutralization breadth defined as neutralization of heterologous viruses. Analysis of the cocrystal structure of the CH103 Fab and a gp120-core demonstrated a novel loop binding mode of antibody neutralization.

Isolation of the CH103 BnAb lineage

The CH505 donor was enrolled in the CHAVI001 acute HIV-1 infection cohort³⁵ approximately 4 weeks after HIV-1 infection (Supplementary Fig. 1) and followed for more than 3 years. Single genome amplification of 53 plasma viral Env gp160 RNAs (5) from 4 weeks after transmission identified a single clade C transmitted/founder (T/F) virus. Serologic analysis demonstrated the development of autologous neutralizing antibodies at 14 weeks, CD4 binding site (CD4bs) antibodies that bound to a recombinant Env protein (resurfaced core, RSC3)⁵ at 53 weeks, and evolution of plasma cross-reactive neutralizing

activity from 41–92 weeks after transmission³⁰ (Fig. 1, Supplementary Table 1, Supplementary Fig. 2). The natural variable regions of heavy- (V_HDJ_H) and light-chain (V_LJ_L) gene pairs of antibodies CH103, CH104, CH106 were isolated from peripheral blood mononuclear cells (PBMC) at 136 weeks after transmission by flow sorting of memory B cells that bound RSC3 Env protein^{36,5,13} (Fig. 1b). The V_HDJ_H gene of antibody CH105 was similarly isolated, but no V_LJ_L gene was identified from the same cell. Analysis of characteristics of V_HDJ_H (V_H4 –59 [posterior probability, PP = 0.99), D3–16 (PP=0.74), J_H4 [PP= 1.00]) and V_LJ_L ($V\lambda3$ –1 [PP=1.00], J\lambda1 [PP=1.00]) rearrangements in mAbs CH103, CH104, CH105 and CH106 demonstrated that these antibodies were representatives of a single clonal lineage we designated as the CH103 clonal lineage (Fig. 2, Supplementary Table 2).

Neutralization assays using a previously described^{5,37} panel of 196 geographically and genetically diverse Env-pseudoviruses representing the major circulated genetic subtypes and circulating recombinant forms demonstrated that CH103 neutralized 55% of viral isolates with a geometric mean IC₅₀ of 4.54 ug/ml among sensitive isolates (Fig. 1c, Supplementary Table 3). ELISA cross-competition analysis demonstrated that CH103 binding to gp120 was competed by known CD4bs ligands such as mAb VRC01 and the chimeric protein CD4-Ig (Fig. 1d); CH103 binding to RSC3 Env was also substantially diminished by gp120 with P363N and 371I mutations known to reduce binding of most CD4bs mAbs (Supplementary Fig. 3)^{5,30}.

Molecular characterization of the CH103 BnAb lineage

The RSC3 probe isolated CH103, CH104, CH105, and CH106 BnAbs by single cell flow sorting. The CH103 clonal lineage was enriched by V_HDJ_H and V_LJ_L sequences identified by pyrosequencing PBMC DNA^{34,38} obtained 66 and 140 weeks after transmission and cDNA antibody transcripts⁶ obtained 6, 14, 53, 92 and 144 weeks after transmission. From pyrosequencing of antibody gene transcripts, we found 457 unique heavy and 171 unique light chain clonal members (Figs. 2a, 2b). For comprehensive study, a representative 14 member BnAb pathway was reconstructed from V_HDJ_H sequences (1AH92U, 1AZCET and 1A102R) recovered by pyrosequencing, and V_HDJ_H genes of the inferred intermediate (I) antibodies (I1–I4, I7, I8)^{11,16,34} (Kepler, TB, Submitted, 2012) that were paired and expressed with either the UCA or I2 V_LJ_L depending on the genetic distance of the V_HDJ_H to either the UCA or mature antibodies (Fig. 2c, Supplementary Table 2). The mature CH103, CH104 and CH106 antibodies were paired with their natural V_LJ_L . The CH105 natural $V_HD_HJ_H$ isolated from RSC3 memory B cell sorting was paired with the V_LJ_L of I2.

Whereas the V_HDJ_H mutation frequencies calculated by using the method as described in the Online Methods of the published CD4bs BnAbs VRC01, CH31 and NIH45–46 V_HDJ_H are $30-36\%^{5-7,22,39}$, the V_HDJ_H frequencies of CH103 lineage CH103, CH104, CH105 and CH106 are 13–17% (Fig. 2c). Additionally, antibodies in CH103 clonal lineage do not contain the large (>3 nt) insertion or deletion mutations common in VRC01-class of BnAbs (1–3) with the exception of the V_LJ_L of CH103 which contained a 3 aa LCDR1 deletion.

It has been proposed that one reason CD4bs BnAbs are difficult to induce is heterologous HIV-1 Envs do not bind their UCAs^{7,18,22}. We wondered, however, whether the CH505 T/F Env, the initial driving antigen for the CH103 BnAb lineage, would preferentially bind to early CH103 clonal lineage members and the UCA compared to heterologous Envs. Indeed, a heterologous gp120 T/F Env, B.63521, did not bind to the CH103 UCA (Fig. 2d) but did bind to later members of the clonal lineage. Affinity for this heterologous Env increased four orders of magnitude during somatic evolution of the CH103 lineage, with maximal K_d values of 2.4 to 7.0 nM in the mature CH103–CH106 mAbs (Fig. 2d). The CH103 UCA mAb also did not bind other heterologous T/F Envs AE.427299, B.9021 and C.1086 (Supplementary Table 4), confirming lack of heterologous Env binding to CD4bs UCAs. Moreover, the gp120 Env RSC3 protein was also not bound by the CH103 UCA and earlier members of the clonal lineage (Supplementary Fig. 3a) and no binding was seen with RSC3 mutant proteins known to disrupt CD4bs BnAb binding (Supplementary Fig. 3b).

In contrast to heterologous Envs, the CH505 T/F Env gp140 bound well to all of the candidate UCAs (Supplementary Table 5) with the highest UCA affinity of $K_d = 37.5$ nM. In addition, the CH505 T/F Env gp140 was recognized by all members of the CH103 clonal lineage (Fig. 2d). Whereas affinity to the heterologous T/F Env B.63521 increased by over four orders of magnitude as the CH103 lineage matured, affinity for the CH505 T/F Env increased by no more than ten fold (Fig. 2d). To directly demonstrate Env escape from CH103 lineage members, autologous recombinant gp140 Envs isolated at weeks 30, 53 and 78 post-infection were expressed and compared with the CH505 T/F Env for binding to the BnAb arm of the CH103 clonal lineage (Supplementary Table 6, Supplementary Fig. 4). Escape mutant Envs could be isolated that were progressively less reactive with the CH103 clonal lineage members. Envs isolated from weeks 30, 53 and 78 lost UCA reactivity and only bound intermediate antibodies 3, 2 and 1 as well as BnAbs CH103, CH104, CH105 and CH106 (Supplementary Table 6). In addition, two Env escape mutants from week-78 viruses also lost either strong reactivity to all intermediate antibodies or to all lineage members (Supplementary Table 6).

To quantify CH103 clonal variants from initial generation to induction of broad and potent neutralization, we used pyrosequencing of antibody cDNA transcripts from five time points, weeks 6, 14, 53, 92 and 144 weeks after transmission (Supplementary Table 7). We found two V_HDJ_H chains closely related to, and possibly members of, the CH103 clonal lineage (Fig. 2a, Supplementary Table 7). Moreover, one of these V_HDJ_H when reconstituted in a full IgG1 backbone and expressed with the UCA V_LJ_L weakly bound the CH505 T/F Env gp140 at endpoint titer of 11 ug/ml (Fig. 2a). These reconstructed antibodies were present concomitant with CH505 plasma autologous neutralizing activity at 14 weeks after transmission (Supplementary Fig. 2). Antibodies that bound the CH505 T/F Env were present in plasma as early as 4 weeks after transmission (data not shown). Both CH103 lineage V_HDJ_H and V_LJ_L sequences peaked at week 53 with 230 and 83 unique transcripts, respectively. V_HDJ_H clonal members fell to 46 at week 144, and V_LJ_L members were 76 at week 144.

Polyreactivity is a common trait of BnAbs, suggesting that the generation of some BnAbs may be controlled by tolerance mechanisms^{10,21,40}. Conversely, polyreactivity can arise

during the somatic evolution of B cells in germinal centers as a normal component of B-cell development⁴¹. The CH103 clonal lineage was evaluated for polyreactivity as measured by HEp-2 cell reactivity and binding to a panel of autoantigens¹⁰. While earlier members of the CH103 clonal lineage were not polyreactive by these measures, polyreactivity was acquired in concert with BnAb activity by the intermediate antibody I2, I1, and clonal members, CH103, CH104 and CH106 (Supplementary Figs. 5a, 5b). The BnAbs CH106 and intermediate antibody I1 also demonstrated polyreactivity in protein arrays with specific reactivity to several human autoantigens, including elongation factor-2 kinase and ubiquitin-protein ligase E3A (Supplementary Figs. 5c and 5d).

Structure of CH103 in complex with HIV-1 gp120

Crystals of the complex between Fab CH103 and the ZM176.66 strain of HIV diffracted to 3.25-Å resolution, and molecular replacement identified solutions for Fab CH103 and for the outer domain of gp120 (Fig. 3a). Inspection of the CH103-gp120 crystal lattice (Supplementary Fig. 6) indicated the absence of the gp120 inner domain was likely related to proteolytic degradation of the extended gp120 core to an outer domain fragment. Refinement to R_{work}/R_{free} of 19.6%/25.6% (Supplementary Table 8) confirmed a lack of electron density for gp120 residues N terminal to residue Val 255_{gp120} or C terminal to Gly472_{gp120} (gp120 residues are numbered according to standard HXB2 nomenclature), and no electron density was observed for residues $301-324_{gp120}$ (V3), $398-411_{gp120}$ (V4) and $421-439_{gp120}$ (β 20–21). Superposition of the ordered portions of gp120 (gp120 residues are numbered according to standard HXB2 nomenclature) in complex with CH103 with the fully extended core gp120 bound by antibody VRC01⁷ indicated a highly similar structure (C α -rmsd 1.16 Å) (Fig. 3b). Despite missing portions of core gp120, the entire CH103 epitope appeared to be present in the electron density for the experimentally observed gp120 outer domain.

The surface bound by CH103 formed an elongated patch with dimensions of ~40 × 10 Å, which stretched across the site of initial CD4 contact on the outer domain of gp120 (Fig. 3c). The gp120 surface recognized by CH103 correlated well with the initial site of CD4 contact; of the residues contacted by CH103, only eight of these residues were not predicted to interact with CD4. CH103 interacted with these residues through side-chain contact with Ser256_{gp120} in loop D, main- and side-chain contacts with His364_{gp120} and Leu369_{gp120} in the CD4-binding loop, and main- and side-chain contacts with Asn463_{gp120} and Asp464_{gp120} in the V5 loop (Fig. 3d). Notably, residue 463 is a predicted site of *N*-linked glycosylation in strain ZM176.66 as well as in the autologous CH505 virus, but electron density for an N-linked glycan was not observed. Overall, of the 22 residues that mAb CH103 was observed to contact on gp120, 14 were expected to interact with CD4 (16 of these residues with antibody VRC01), providing a structural basis for the CD4-epitope specificity of CH103 and its broad recognition (Supplementary Table 9).

Residues $1-215_{\text{HC}}$ on the antibody heavy chain and $1-209_{\text{LC}}$ showed well defined backbone densities. Overall, CH103 utilizes a CDR H3 dominated mode of interaction, although all six of the complementarity-determining regions (CDRs) interacted with gp120 as well as the light chain framework region 3 (FWR3) (Supplementary Figs. 7a,b, Supplementary Tables

10 and 11). It is important to note that ~40% of the antibody contact surface was altered by somatic mutation, in two regions, in the CDR H2 and in the CDR L1, L2 and FWR3. In particular, residues 56_{HC} , 50_{LC} , 51_{LC} and 66_{LC} are altered by somatic mutation to form hydrogen bonds with the CD4-binding loop, loop D and loop V5 of gp120. Nevertheless, 88% of the CH103 V_HD_HJ_H and 44% of the V λ J λ contact areas were with amino acids unmutated in the CH103 germline, potentially providing an explanation for the robust binding of the T/F Env to the CH103 UCA (Supplementary Figs. 7c, 7d, and Supplementary Table 12).

Evolution of transmitted/founder Env sequences tracks acquisition of BnAb activity

Using single genome amplification and sequencing²⁴ we tracked the evolution of CH505 env genes longitudinally from the T/F virus through 160 weeks post-transmission (Fig. 4, Supplementary Fig. 8). The earliest recurrent mutation in Env, N279K (HIV-1 HXB2 numbering), was found at 4 weeks post-infection, and was in Env loop D in a CH103 contact residue. By week 14 additional mutations in loop D appeared, followed by mutations and insertions in V1 at week 20. Insertions and mutations in the V5 loop began to accumulate by week 30 (Fig. 4). Thus, the T/F virus began to diversify in key CD4 contact regions starting within 3 months of infection (Supplementary Figs. 8, 9). Loop D and V5 mutations were directly in or adjacent to CH103/Env contact residues. Although the V1 region was not included in the CH103-Env co-crystal, the observed V1 CH505 Env mutations were adjacent to contact residues for CD4 and VRC01 so are likely to be relevant. It is also possible that early V1 insertions (Fig. 4) were selected by inhibiting access to the CD4bs in the trimer or that they arose in response to early T cell pressure. CD4 binding-loop mutations were present by week 78. Once regions that could directly impact CH103-lineage binding began to evolve (loop D, V5, the CD4 binding, loop, and possibly V1), they were under sustained positive selective pressure throughout the study period (Fig. 4, Supplementary Figs. 8, 9, Supplementary Table 13).

Considerable within-sample virus variability was evident in Env regions that could impact CH103-linage antibody binding, and diversification within these regions preceded neutralization breadth. Expanding diversification early in viral evolution (4–22 weeks after transmission) (Supplementary Figs. 8, 9) coincided with autologous NAbs development, consistent with autologous NAb escape mutations. Mutations that accumulated from weeks 41–78 in CH505 Env contact regions immediately preceded development of NAb breadth (Fig 4, Supplementary Figs. 8, 9). By weeks 30–53, extensive within-sample diversity resulted from both point mutations in and around CH103 contact residues, and to multiple insertions and deletions in V1 and V5 (Supplementary Fig. 9). A strong selective pressure seems to have come into play between weeks 30 and 53, perhaps due to autologous neutralization escape, and neutralization breadth developed after this point (Fig. 4, Supplementary Figs 8, 9). Importantly, due to apparent strong positive selective pressure between week 30 and week 53, there was a dramatic shift in the viral population that is evident in the phylogenetic tree, such that only viruses carrying multiple mutations relative to the T/F, particularly in CH103 contact regions, persisted after week 30. This was followed

by extreme and increasing within time-point diversification in key epitope regions, beginning at week 53 (Supplementary Fig. 9). Emergence of antibodies with neutralization breadth occurred during this time (Supplementary Fig. 2, Supplementary Table 1). Thus, plasma breadth evolved in the presence of highly diverse forms of the CH103 epitope contact regions (Fig. 4, Supplementary Fig. 2).

To evaluate and compare the immune pressure on amino acids in the region of CH103 and CD4 contacts, we compared the frequency of mutations in evolving T/F sequences of patient CH505 during the first year of infection and in 16 other acutely infected subjects followed over time (Supplementary Fig. 10). The accumulation of mutations in the CH505 virus population was concentrated in regions likely to be associated with escape from the CH103 lineage (Supplementary Fig. 10a), and diversification of these regions was far more extensive during the first six months of infection in CH505 than in other subjects (Supplementary Fig. 10b). However, by one year into their infections, viruses from the other subjects had also begun to acquire mutations in these regions. Thus, the early and continuing accumulation of mutations in CH103 contact regions may have potentiated the early development of neutralizing antibody breadth in patient CH505.

Neutralization of autologous and heterologous viruses and the CH103 lineage

Heterologous BnAb activity was confined to the later members (I3 and later) of the BnAb arm of the CH103 lineage as manifested by their neutralization capacity of pseudoviruses carrying tier 2 Envs A.Q842 and B.BG1168 (Fig. 5a). Similar results were seen with Envs A.Q168, B.JRFL, B.SF162 and C.ZM106 (Supplementary Tables 14 and 15). In contrast, neutralizing activity of clonal lineage members against the autologous T/F Env pseudovirus appeared earlier with measurable neutralization of the CH505 T/F virus by all members of the lineage after the UCA except mAb 1AH92U (Fig. 5a). Thus, within the CH103 lineage, early intermediate antibodies only neutralized the T/F virus, while later intermediate antibodies gained neutralization breadth, indicating evolution of neutralization breadth with affinity maturation, and CH103-CH106 BnAbs evolved from an early autologous neutralizing antibody response. Moreover, the clonal lineage was heterogeneous, with an arm of the lineage represented in Fig. 5a evolving neutralization breadth and another antibody arm capable of mediating only autologous T/F virus neutralization. While some escape viruses are clearly emerging over time (Supplementary Table 4), it is important to point out that, whereas escape mutant viruses are driving BnAb evolution, the BnAbs remained capable of neutralizing the CH505 T/F virus (Fig. 5a). Of note, the earliest mutations in the heavy chain lineage clustered near the contact points with gp120, and these remained fixed throughout the period of study, while mutations that accumulated later tended to be further from the binding site and may be impacting binding less directly (Supplementary Fig. 11). Thus, stimulation of the CH103 BnAbs occurs in a manner to retain reactivity with the core CD4bs epitope present on the T/F Env. One possibility that might explain this is that the footprint of UCA binding contracts to the central core binding site of the CH103 mature antibody. Obtaining a crystal structure of the UCA with the T/F Env should inform this notion. Another possibility is that because affinity maturation is

occurring in the presence of highly diverse forms of the CD4bs epitope, antibodies that favor tolerance of variation in and near the epitope are selected instead of those antibodies that acquire increased affinity for particular escape Envs. In both scenarios, persistence of activity to the T/F form and early viral variants would be expected. Fig. 5b and Supplementary Fig. 11 show views of accumulations of mutations or entropy during the parallel evolution of the antibody paratope and the Env epitope bound by mAb CH103.

Vaccine Implications

In this study, we demonstrate that the binding of a T/F Env to an UCA B cell receptor of a BnAb lineage was responsible for the induction of broad neutralizing antibodies, thus providing a logical starting place for vaccine-induced CD4bs BnAb clonal activation and expansion. Importantly, the number of mutations required to achieve neutralization breadth was reduced in the CH103 lineage compared to most CD4bs BnAbs, although the CH103 lineage had reduced neutralization breadth compared to more mutated CD4bs BnAbs. Thus, this type of BnAb lineage may be less challenging to attempt to recapitulate by vaccination. By tracking viral evolution through early infection we found that intense selection and epitope diversification in the T/F virus preceded the acquisition of NAb breadth in this individual -- thus demonstrating the viral variants associated with development of BnAbs directly from autologous neutralizing antibodies and illuminating a pathway for induction of similar B cell lineages.

These data have implications for understanding the B cell maturation pathways of the CH103 lineage and for replicating similar pathways in a vaccine setting. First, we demonstrate in CH505 that BnAbs were driven by sequential Env evolution beginning as early as 14 weeks after transmission, a time period compatible with induction of this type of BnAb lineage with a vaccine given the correct set of immunogens. Second, whereas heterologous Envs did not bind with UCAs or early intermediate antibodies of this lineage, the CH505 T/F Env bound remarkably well to the CH103 UCA, and subsequent Envs bound with increased affinity to later clonal lineage members. This suggests that immunizations with similar sequences of Env or Env subunits may drive similar lineages. Third, the CH103 lineage is less complicated than those of the VRC01-class of antibodies because antibodies in this lineage has fewer somatic mutations, and no indels, except CH103 V_L has a deletion of 3 amino acid residues in the LCDR1 region. It should be noted as well that our study is in one patient. Nonetheless, in each BnAb patient, analysis of viral evolution should elucidate a similar pathway of evolved Envs that induce BnAb breadth. The observation that rhesus macaques infected with the CCR5-tropic SHIV-AD8 virus frequently develop neutralization breadth⁴² suggests the notion that certain envelopes may be more likely to induce breadth and potency than others.

Polyreactivity to host molecules in the CH103-lineage arose during affinity maturation in the periphery coincident with BnAb activity. This finding is compatible with the hypothesis that BnAbs may be derived from an inherently polyreactive pool of B cells, with polyreactivity providing a neutralization advantage via heteroligation of Env and host molecules^{21,43}. Alternatively, as CH103 affinity maturation involves adapting to the simultaneous presence of diverse co-circulating forms of the epitope⁴⁴, the selection of

antibodies that can interact with extensive escape-generated epitope diversification may be an evolutionary force that also drives incidental acquisition of polyreactivity.

Thus, a candidate vaccine concept could be to use the CH505 T/F Env or Env subunits (to avoid dominant Env non-neutralizing epitopes) to initially activate an appropriate naïve B cell response, followed by boosting with subsequently evolved CH505 Env variants either given in combination to mimic the high diversity observed in vivo during affinity maturation, or in series, using vaccine immunogens specifically selected to trigger the appropriate maturation pathway by high affinity binding to UCA and antibody intermediates¹¹. These data demonstrate the power of studying subjects followed from the transmission event through the development of plasma BnAb activity for concomitant isolation of both T/F viruses and their evolved quasispecies along with the clonal lineage of induced BnAbs. The finding that the T/F Env can be the stimulator of a potent BnAb and bind optimally to that BnAb UCA is a critical insight for vaccine design, and could allow the induction of BnAbs by targeting UCAs and IAs of BnAb clonal lineage trees¹¹.

Methods

Study subject

Plasma and peripheral blood mononuclear cells (PBMC) were isolated from serial blood samples that were collected from a HIV-1 infected subject CH505 starting 6 weeks after infection up to 236 weeks after infection (Supplementary Table 1) and frozen at -80° c and liquid nitrogen tanks, respectively. During this time, no anti-retroviral therapy was administered. All work related to human subjects was in compliance with Institutional Review Board protocols approved by the Duke University Health System Institutional Review Board.

Inference of unmutated common ancestor (UCA) and identification of clone members

The inference of the UCA from a set of clonally related genes is described in detail in manuscript by Kepler, T.B., "Reconstructing a B cell clonal lineage: I. Statistical Inference of Unobserved Ancestors (http://arxiv.org/submit/665968)". Briefly, we parameterize the VDJ rearrangement process in terms of its gene segments, recombination points, and nregions. Given any multiple sequence alignment A for the set of clonally related genes and any tree T describing a purported history, we can compute the likelihood for all parameter values, and thus the posterior probabilities on the rearrangement parameters conditional on A and T. We can then find the UA with the greatest posterior probability and compute the maximum likelihood alignment A* and tree T* given this UA, and then recompute the posterior probabilities on rearrangement parameters conditional on A* and T*. We iterate the alternating conditional maximizations until convergence is reached. We use ClustalW⁴⁷ for the multiple sequence alignment, dnaml (PHYLIP) to infer the maximum likelihood tree, and our own software for the computation of the likelihood over the rearrangement parameters. The variable regions of heavy- and light chain $(V_H DJ_H \text{ and } V_L J_L)$ gene segments were inferred from the natural pairs themselves. The posterior probabilities for these two gene segments are 0.999 and 0.993, respectively. We first inferred the UA from the natural pairs as described above. We identified additional clonally related variable region

sequences from deep sequencing and refine the estimate of the UCA iteratively. We identified all variable region sequences inferred to have been rearranged to the same V_HDJ_H and J_H , and to have the correct CDR3 length. For each sequence, we counted the number of mismatches between the sequence and the presumed V_HDJ_H gene up to the codon for the second invariant cysteine. Each iteration was based on the CDR3 of the current posterior modal UA. For each candidate sequence, we computed the number of nucleotide mismatches between its CDR3 and the UA CDR3. The sequence was rejected as a potential clone member if the z-statistic in a test for difference between proportion is greater than two⁴⁸. Once the set of candidates has been thus filtered by CDR3 distance, the UA was inferred on that larger set of sequences as described above. If the new posterior modal UA differed from the previous one, the process was repeated until convergence was reached. Due to the inherent uncertainty in UA inference, we inferred the 6 most likely VH UCA sequences resulting in 4 unique amino acid sequences that were all produced and assayed for reactivity with the transmitted/founder envelope gp140 (Supplementary Table 5).

Phylogenetic Trees

Maximum-likelihood phylograms were generated using the dnaml program of the PHYLIP package (version 3.69) using the inferred ancestor as the outgroup root, "speedy/rough" disabled, and default values for the remaining parameters. For the large antibody datasets, neighbor-joining phylogenetic trees were generated using the EBI bioinformatics server at http://www.ebi.ac.uk/Tools/phylogeny/ using default parameter values. All NJ trees were generated subsequent to the inference of the UAs.

Isolation of V_HDJ_H and V_L genes and expression of V_HDJ_H and V_LJ_L genes as full-length IgG1 recombinant mAbs

The V_HDJ_H and V_LJ_L gene segment pairs of the observed CH103, CH104 and CH106 and the V_HDJ_H gene segment of CH105 were amplified by RT/PCR of flow sorted HIV-1 Env RSC3 (re-surface core3) -specific memory B cells using the methods as described previously^{36,5–7,22}. To compare V_H mutation frequency of CH103, CH104, CH105 and CH106 with that of previously published of CD4bs BnAbs VRC01, CH31 and NIH45–46, V_H sequences of these antibodies were aligned to the closest V_H gene segment from the IMGT reference sequence set, and counted differences between the target sequence and the V_H gene segment up to and including the second invariant cysteine. The comparison 3' of Cys2 is omitted since the unmutated form of the ancestral sequence is not as well known.

Additional $V_H DJ_H$ and $V_L J_L$ and $V_L J_L$ genes were identified by 454 pyrosequencing. Clonally related $V_H DJ_H$ and $V_L J_L$ sequences derived from either sorted single B cells or 454 pyrosequencing were combined and used to generate neighbor-joining phylogenetic trees (Figs 2a and 2b). Antibodies that were recovered from single memory B cells are noted in the figure in red, and bolded lines show the inferred evolutionary paths from theUCA to mature BnAbs. For clarity, related V_H variants that grouped within monophyletic clades from the same time-point were collapsed to single branches, condensing 457 $V_H DJ_H$ and 174 V $V_L J_L$ variants to 119 and 46 branches, respectively, via the "nw_condense" function from the Newick Utilities package (v. 1.6)⁴⁹. The frequencies of $V_H DJ_H$ variants in each B cell sample are shown to the right of the $V_H DJ_H$ tree in Fig. 2a, and were computed from

sample sizes of 188,793, 186,626, and 211,901 sequences from weeks 53, 92, and 144, respectively. Two V_HDJ_H genes (IZ95W and 02IV4) were found at 14 weeks after transmission and paired with UCA V_LJ_L for expression as IgG1 mAbs. IZ95W mAb weakly bound the CH505 T/F Env gp140 with end-point titer of 11 ug/ml. Among heavy chain sequences in the tree, the mean distance of each to its nearest neighbor to was calculated to be 8.1 nt. The cumulative distribution function shows that, while there are pairs that are very close together (nearly 30% of sequences are 1nt from its neighbor), 45% of all sequences differ by 6nt or more from its nearest neighbor. The probability of generating a sequence that differs by 6 or more nucleotides from the starting sequence by PCR and sequencing is very small. The numbers of sequences obtained from a total of 100 million PBMC were within the expected range of 50–500 antigen-specific B cells.

Regarding the number of unique V_HDJ_H and V_LJ_L genes that we have isolated, we have analyzed this issue in a number of ways. First, we have clarified the calculations for the possible number of antigen-specific CD4bs memory B cells that could have been isolated from the samples studied. We studied 5 patient CH505 time points with pyrosequencing with ~20 million PBMC per time point for a total of 100 million PBMC studied. In chronic HIV, there is a mean of 145 total B cells per ul of blood, and a mean of 60 memory B cells per ul of blood⁵⁰. This high percent of memory B cells of ~40% of the total B cells in chronic HIV infection is due to selective loss of naïve B cells in HIV infection. Thus, in 100 ml (100,000 ul) of blood, there will be approximately 6 million memory B cells. If 0.1 to 1.0% are antigen specific, that that would be 6,000 to 60,000 antigen-specific B cells sampled, and if, of these, 5% were CD4bs antibodies, then from 300 to 3000 CD4 bs B cells would have been sampled in 100 million PBMC studied. This is completely compatible and within the range of the calculations of the reviewer above (50 CD4 bs B cells per 5 million PBMC), since we studied 100 million PBMC, there should, by these calculations, 1000 CD4bs B cells sampled. Either calculation therefore yields estimates that are completely compatible with the 474 V_HDJ_H genes amplified.

To further study the plausibility of sequences isolated, the second method of analysis we used was as follows. Among heavy chain sequences in the tree, one can compute the distance of each to its nearest neighbor. The mean distance to the nearest neighbor is 8.1 nt. The cumulative distribution function shows that, while there are pairs that are very close together (nearly 30% of sequences are 1nt from its neighbor), 45% of all sequences differ by 6nt or more from its nearest neighbor. The probability of generating a sequence that differs by 6 or more nucleotides from the starting sequence by PCR and sequencing is very small. We believe the number of genes represented in our sample is closer to 200 than to 50, and most likely is larger than 200.

The third analysis we performed was to compute the distance of each heavy chain sequences in the tree to its nearest neighbor. The mean distance to the nearest neighbor is 8.1 nt. We used agglomerative clustering to prune the sequence alignment. At the stage where no pairs of sequences were 3 nucleotides apart or closer, there were 335 of 452 sequences remaining; when no pairs are 6nt apart or closer, there are still 288 sequences remaining. Therefore with this analysis, we believe the number of genes represented in our sample is closer to 300 than

to 50, and may be larger. Thus, by the sum of these re-analyses, we believe that the number of genes in the trees in Figure 2 are quite plausible.

The isolated Ig V_HDJ_H and V_LJ_L gene pairs, the inferred UCA and intermediate V_HDJ_H and V_LJ_L sequences, and select V_HDJ_H gene sequences identified by pyrosequencing were studied experimentally (Supplementary Table 2) and used to generate a phylogenetic tree showing percentage of mutated V_H sites and time of appearance after transmission (Figs 2c) and binding affinity (Fig 2d). The isolated four mature antibodies are indicated in red, antibodies derived from 454 pyrosequencing are indicated in black, and inferredintermediate antibodies (I1–I4, I7, I8) are indicated by circles at ancestral nodes. The deep clades in this tree had modest bootstrap support, and the branching order and UCA inference were somewhat altered when more sequences were added to the phylogenetic analysis (compare the branching order of Fig. 2c and Fig. 2a). The tree depicted in Figs. 2c and 2d was used to derive the ancestral intermediates of the representative lineage early in our study, and marked an important step in our analysis of antibody affinity maturation. The V_HDJ_H and V_LJ_L genes were synthesized (GenScript, NJ) and cloned into pcDNA3.1 plasmid (Invitrogen, Grand Island, NY) for production of purified recombinant IgG1 antibodies as described previously^{51,52}. The V_HDJ_H genes of I1–I4, I7 and I8 as well as the V_HDJ_H of CH105 were paired with either the V_L gene of the inferred UCA or I2 depending on the genetic distance of the V_HDJ_H to either the UCA or mature antibodies for expressing as full-length IgG1 antibodies as described⁵¹ (Supplementary Table 2).

Recombinant HIV-1 Proteins

HIV-1 Env genes for subtype B, 63521, subtype C, 1086, and subtype CRF_01, 427299, as well as subtype C, CH505 autologous transmitted/founder Env were obtained from acutely infected HIV-1 subjects by single genome amplification²⁴ codon-optimized by employing the codon usage of highly expressed human housekeeping genes⁵³, *de novo* synthesized (GeneScript, NJ) as gp140 or gp120 (AE.427299) and cloned into a mammalian expression plasmid pcDNA3.1/hygromycin (Invitrogen, Grand Island, NY). Recombinant Env glycoproteins were produced in 293F cells cultured in serum-free medium and transfected with the HIV-1 gp140- or gp120-expressing pcDNA3.1 plasmids, purified from the supernatants of transfected 293F cells by using *Galanthus nivalis* lectin-agarose (Vector Labs, Burlingame, CA) column chromatography^{16,54,52}, and stored at –80°C. Select Envs made as CH505 T/F Env were further purified by superose 6 column chromatography to trimeric forms, and used in binding assays that showed similar results as with the lectin-purified oligomers.

Enzyme-Linked Immunoassay (ELISA)

Binding of patient plasma antibodies and CH103 clonal lineage antibodies to autologous and heterologous HIV-1 Envs was measured by ELISA as described previously^{34,52}. Plasma samples in serial 3-fold dilutions starting at 1:30 to 1:521,4470 or purified mAbs in serial 3-fold dilutions starting at 100ug/ml to 0.000ug/ml diluted in PBS were assayed for binding to autologous and heterologous HIV-1 Envs. Binding of biotin-labeled CH103 at the subsaturating concentration was assayed for cross competition by unlabeled HIV-1 antibodies and soluble CD4-Ig in serial 4-fold dilutions starting at 10ug/ml. The half

maximal effective concentration (EC_{50}) of plasma samples and mAbs to HIV-1 Envs were determined and expressed as either the reciprocal dilution of the plasma samples or concentration of mAbs.

Surface plasmon resonance (SPR) affinity and kinetics measurements

Binding K_d and rate constant (association rate k_a , dissociation rate k_d) measurements of mAbs and all candidate UCAs to the autologous Env C. CH05 gp140 and/or the heterologous Env B.63521 gp120 were carried out on BIAcore 3000 instruments as described previously^{19,43,45}. Anti-human IgG Fc antibody (Sigma Chemicals) was immobilized on a CM5 sensor chip to about 15000 Response Unit (RU) and each antibody was captured to about 50-200 RU on three individual flow cells for replicate analysis, in addition to having one flow cell captured with the control Synagis (anti-RSV) mAb on the same sensor chip. Double referencing for each mAb-HIV-1 Env binding interactions was used to subtract non-specific binding and signal drift of the Env proteins to the control surface and blank buffer flow respectively. Antibody capture level on the sensor surface was optimized for each mAb to minimize rebinding and any associated avidity effects. C.CH505 Env gp140 protein was injected at concentrations ranging from 2 to 25µg/mL and B.63521 gp120 was injected at 50 - 400 µg/mL for UCA and early intermediates (IA8, IA4), 10-100 μ g/mL (IA3), and 1–25 μ g/mL for the distal and mature mAbs. All curve fitting analysis were performed using global fit of to the 1:1 Langmuir model and are representative of at least three measurements. All data analysis was performed using the BIAevaluation 4.1 analysis software (GE Healthcare).

Neutralization assays

Neutralizing antibody assays in TZM-bl cells were performed as described previously⁵⁵. Neutralizing activity of plasma samples in 8 serial 3-fold dilutions starting at 1:20 dilution and for recombinant mAbs in 8 serial 3-fold dilutions starting at 50ug/ml were tested against autologous and herologous HIV-1 Env-pseudotyped viruses in TZM-bl-based neutralization assays using the methods as described^{5,37,55}. Neutralization breadth of CH103 was determined using previous described^{5,37} a panel of 196 of geographically and genetically diverse Env-pseudoviruses representing the major circulated genetic subtypes and circulating recombinant forms. The subtypes shown in Figure 1c are consistent with prior publications^{5,56}, and the clades described in Los Alamos database (www.hiv.lanl.gov). HIV-1 Subtype robustness is derived from the analysis of HIV-1 clades over time⁵⁷. The data were calculated as a reduction in luminescence units compared with control wells and reported as IC₅₀ in either reciprocal dilution for plasma samples or in μ g/ml for mAbs.

Crystallization of antibody CH103 and its gp120 complex

The antigen binding fragment (Fab) of CH103 was generated by LyS-C (Roche) digestion of IgG1 CH103 and purified as previously described⁷. The extended core gp120 of HIV-1 clade C ZM176.66 was used to form a complex with Fab CH103 by <u>using previously</u> <u>described methods⁵⁸</u>. Briefly, deglycosylated ZM176.66, constructed as an extended gp120 core⁵⁹, that was produced using the method as described previously⁷ and Fab CH103 were mixed at a 1:1.2 molar ratio at room temperature and purified by size exclusion chromatography (Hiload 26/60 Superdex S200 prep grade, GE Healthcare) with buffer

containing 0.35 M NaCl, 2.5 mM Tris pH 7.0, 0.02% NaN₃. Fractions of the Fab or gp120:CH103 complex were concentrated to ~10 mg/ml, flash frozen with liquid nitrogen before storing at -80° C and used for crystallization screening experiments.

Commercially available screens, Hampton Crystal Screen (Hampton Research), Precipitant Synergy Screen (Emerald BioSystems), Wizard Screen (Emerald BioSystems), PACT Suite and JCSG+ (Qiagen) were used for initial crystallization screening of both Fab CH103 and its gp120 complex. Vapor-diffusion sitting drops were set up robotically by mixing 0.2 µl of protein with an equal volume of precipitant solutions (Honeybee 963, DigiLab). The screen plates were stored at 20°C and imaged at scheduled times with RockImager (Formulatrix.). The Fab CH103 crystals appeared in a condition from the JCSG+ kit containing 170 mM ammonium sulfate, 15% glycerol and 25.5 % PEG 4000. For the gp120:CH103 complex (Supplementary Table 8), crystals were obtained after 21 days of incubation in a fungi-contaminated^{60,61} droplet of the PACT suite that contained 200 mM sodium formate, 20% PEG 3350 and 100 mM Bistrispropane, pH 7.5.

X-ray data collection, structure determination and refinement for the gp120:CH103 complex

Diffraction data were collected under cryogenic conditions. Optimal cryo-protectant conditions were obtained by screening several commonly used cryo-protectants as described previously⁷. X-ray diffraction data were collected at beam-line ID-22 (SER-CAT) at the Advanced Photon Source, Argonne National Laboratory, with 1.0000 Å radiation, processed and reduced with HKL2000⁶². For the Fab CH103 crystal, a data set at 1.65 Å resolution was collected with a cryo-solution containing 20% ethylene glycol, 300 mM ammonium sulfate, 15% glycerol and 25 % PEG 4000 (Supplementary Table 8). For the gp120:CH103 crystals, a data set at 3.20 Å resolution was collected using a cryo-solution containing 30% glycerol, 200 mM sodium formate, 30% PEG 3350 and 100 mM Bistrispropane, pH 7.5 (Supplementary Table 8).

The Fab CH103 crystal was in the P2₁ space group with cell dimensions at a=43.0, b=146.4, c=66.3, α =90.0, β =97.7, γ =90.0 and contained two Fab molecules per asymmetric unit (Supplementary Table 8). The crystal structures of Fab CH103 were solved by molecular replacement using Phaser⁶³ in the CCP4 Program Suite⁶⁴ with published antibody structures as searching models. The gp120:CH103 crystal also belonged to the P21 space group with cell dimensions at a=48.9, b=208.7, c=69.4, α =90, β =107.2, γ =90.0, and contained two gp120:CH103 complexes per asymmetric unit (Supplementary Table 8). The high resolution Fab CH103 structure was used as an initial model to place the Fab CH103 component in the complex. With the Fab CH103 position fixed, searching with the extended core gp120 of ZM176.66 in the VRC01-bound form as an initial model failed to place the gp120 component in the complex. After trimming the inner domain and bridging sheet regions from the gp120 search model, Phaser was able to correctly place the remaining outer domain of gp120 into the complex without significant clashes. Analysis of the packing of the crystallographic lattice indicated a lack of space to accommodate the inner domain of gp120, suggesting possible protease cleavage of the gp120 by the containing fungi during crystallization^{60,61}.

Structural refinements were carried out with PHENIX⁶⁵. Starting with torsion-angle simulated annealing with slow cooling, iterative manual model building was carried out on $COOT^{66}$ with maps generated from combinations of standard positional, individual *B*-factor, TLS refinement algorithms and non-crystallographic symmetry (NCS) restraints. Ordered solvents were added during each macro cycle. Throughout the refinement processes, a cross validation (R_{free}) test set consisting of 5% of the data was used and hydrogen atoms were included in the refinement model. Structure validations were performed periodically during the model building/refinement process with MolProbity⁶⁷ and pdb-care⁶⁸. X-ray crystallographic data and refinement statistics are summarized in Supplementary Table 8. The Kabat nomenclature⁶⁹ was used for numbering of amino acid residues in amino acid sequences in antibodies.

Protein structure analysis and graphical representations

PISA⁷⁰ was used to perform protein-protein interfaces analysis. CCP4⁶⁶ was used for structural alignments. All graphical representation with protein crystal structures were made with Pymol⁷¹.

Polyreactivity analysis of antibodies by HEp-2 cell staining, ANA assays and protein array microchip

All antibodies in CH103 clonal lineage were assayed at 50 ug/ml for autoreactivity to HEp-2 cells (Inverness Medical Professional Diagnostics, Princeton, NJ) by indirect immunofluorescence staining and a panel of autogens by ANA assays using the methods as reported previously¹⁰. The intermediate antibody (IA1) and CH106 were identified as reactive with HEp-2 cells and then selected for further testing for reactivity with human host cellular antigens using ProtoArray 5 microchip (Invitrogen, Grand Island, NY) according to the instructions of the microchip manufacturer. Briefly, ProtoArray 5 microchips were blocked and exposed to 2 µg/ml IA1, CH106 or an isotype-matched (IgG1, k) human myeloma protein, 151K (Southern Biotech) for 90 min at 4 °C. Protein-Ab interactions were detected by 1 µg/mL Alexa Fluor 647-conjugated anti-human IgG. The arrays were scanned at 635 nm with 10µm resolution using 100% power and 600 gain (GenePix 4000B scanner, Molecular Devices). Fluorescence intensities were quantified using GenePix Pro 5.0 (Molecular Devices). Lot-specific protein spot definitions were provided by the microchip manufacturer and aligned to the image.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Structure factors and coordinates for Fab CH103, both unbound and bound to HIV-1 gp120, are in the process of being deposited with the Protein Data Bank. We are also in the process of depositing next-generation sequencing data used in this study to NCBI Short Reads Archives. The opinions herein are those of the authors and should not be construed as official or representing the views of the U.S. Department of Health and Human Services, National Institute for Allergy and Infectious Diseases. H.X.L., R.L., T.Z., F.G., S.D.B., B.H.H., T.B.K., J.R.M., P.D.K. and B.F.H. have filed patent applications on mAbs and/or CH505 Envs used in this study.

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Figure 1. Development of neutralization breadth in donor CH505 and isolation of antibodies a, Shown are HIV-1 viral RNA copies and reactivity of longitudinal plasmas samples with HIV1-1 YU2 core gp120, RSC3 and negative control RSC3 3711(RSC3) proteins. **b**, PBMCs from week 136 was used for sorting CD19⁺, CD20+, IgG⁺, RSC3⁺ and RSC3⁻ memory B cells (0.198%). Individual cells indicated as orange, blue and green dots yielded mAbs CH103, CH104 and CH106, respectively, as identified by index sorting. **c**, The neutralization potency and breadth of the CH103 antibody are displayed using a neighbor joining tree created with PHYLIP package. The individual tree branches for 196 HIV-1 Envs representing major circulating clades are colored according to the neutralization IC50 values as indicated. **d**, Cross competition of CH103 binding to YU2 gp120 by the indicated HIV-1 antibodies, and soluble CD4-Ig was determined by ELISA.



Figure 2. CH103-clonal family with time of appearance, $\rm V_HDJ_H$ mutations, and HIV-1 Env reactivity

Phylogenies of $V_H DJ_H$ (**a**) and $V_L J_L$ (**b**) sequences from sorted single memory B cells and pyrosequencing. The ancestral reconstructions for each were done using the methods described in the Online Methods. The phylogenetic trees themselves were subsequently computed using neighbor joining on the complete set of DNA sequences (see Online Methods) to illustrate the correspondence of sampling date and read abundance in the context of the clonal history. Within time-point V_H monophyletic clades are collapsed to single branches; variant frequencies are indicated on the right. Isolated mature antibodies are red, pyrosequencing-derived sequences are black. The inferred evolutionary paths to observed matured antibodies are bold. **c**, <u>Maximum-likelihood phylogram showing the</u> CH103 lineage with the inferred intermediates (circles, I1–4, I7 and I8), and percentage mutated V_H sites and timing (blue), indicated. **d**, Binding affinity (Kd, nM) of antibodies to autologous CH505 (left box) and heterologous B.63521 were measured by surface plasmon reasonance (SPR) (right box).



Figure 3. Structure of antibody CH103 in complex with the outer domain of HIV-1 gp120 (OD) **a**, Overall structure of complex with gp120 polypeptide depicted in red ribbon and CH103 shown as a molecular surface (heavy chain in green and light chain in blue). Major CH103binding regions on gp120 are colored orange for Loop D, yellow for the CD4-binding site and purple for Loop V5. **b**, Superposition of OD bound by CH103 (red) and core gp120 bound by VRC01 (gray) with polypeptide shown in ribbon representation. **c**, CH103 epitope (green) on OD (red) with the initial CD4-binding site superposed (yellow boundaries) in surface representation. **d**, Sequence alignment of outer domains of the crystallized gp120 shown on the first line and diverse HIV-1 Envs recognized by CH103. Secondary structure elements are labeled above the alignment with gray dashed lines indicating disordered regions. Symbols in yellow or green denote gp120 OD contacts for CD4 and CH103, respectively, with open circles representing main-chain contacts, open circles with rays representing side-chain contact, and filled circles representing both main-chain and sidechain contacts.

	Loop D		V1 Loop	β23 - V5 Loop - β24	CD4 Binding Loop
	Weeks Post-Infection		PLOTLICTAT ASISS	ETERPOCONNEL ETERPOCONNEL AND A SECONDARY AND A S	TEOPSSOCOLETTT
	Autologous Neutralization ———		PLOYTLICTIAT ASISS	LTRDGGKNVT ETFRPGGGNWKDNVRS	TEOPSSOCOL ELTT
	Heterologous Neutralization (Tier 1)	H ^{201.0}	ECTLOTIF _{as} ,, _{nat} 83 85	LTRDCCKINT ETERPCCCNIKONIRS	TEOPSSOOL ETTH
		22 ^{1.0} 0.5 0.0	PLOTLICTNEL	LTRDGGKINT ETFRPGGGINKDWRS	TFOPSSGG <mark>dle</mark> itth
	Expanding Heterologous Neutralization ——— (Tier 2)	30 ^{1.0} 0.5	PLCVTLICTNAT _{As}	LTRDCCKINT DTETERPCCCI IN DAVRS	TEOPSSCOLETT
			PLOVILICI BATAS NATAS NS STATAS ISS	LTRDGGKNNT	TEOPSSOCOLEITT
	Full Breadth		PLOVILICIDATASARYASAATA, SAATASASSAT	LTRDCCENRT	TF@PSSCGDLE*TT
	Neutralization KEY		PLOVILIC <mark>IB</mark> AVATASNIMATAS NINATAS	LTRDGGENGPoggwißt Ettrpeggnikdwirs	TFOPSSGGD ee itth
GS NG PA # +	STYC polar IQ neutral IRH basic IE acidic AWFLIMV hydrophobic Lor indel (gap)		PLOTLICERIT _{e tannen s} itte sittes	LTRDCC INFILLEDTETERPECGINKOWRS	TERPSSGGDEETTH
			PLOVILICENT	LTRDCGINNTILLIEDTETFRPVCGNNKDNVSS	TEOPSSOCOLETTH .
	incomplete codo	n Env site	Env site	Env site	Env site
	CD4 binding-site contact				
×	x CH103 contact residue				
	CONTRACTS CD/4 & CHI103				

Figure 4. Sequence Logo displaying variation in key regions of CH505 Envs

The frequency of each amino acid variant per site is indicated by its height, deletions are indicated by grey bars. The first recurring mutation, N279K, appears at week 4 (open arrow). The timing of BnAb activity development (from Supplementary Fig. 2 and Supplementary Table 1) is on the left. Viral diversification, which precedes acquisition of breadth, is highlighted by vertical arrows to the right of each region. CD4 and CH103 contact residues, and amino acid position numbers based on HIV-1 HXB2, are shown along the base of each Logo column.



Figure 5. Development of neutralization breadth in the CH103-clonal lineage

a, Phylogenetic CH103 clonal lineage tree showing the IC50 (μg/ml) of neutralization of either the autologous T/F (C.CH505), heterologous tier clades A (A.Q842) and B (B.BG1168) viruses as indicated. **b**, Interplay between evolving virus and developing clonal lineage mapped on to models of CH103-developmental variants and contemporaneous virus. The outer domain of HIV gp120 is depicted in worm representation, with worm thickness and color (white to red) mapping the degree of per-site sequence diversity at each time point. Models of antibody intermediates are shown in cartoon diagram with somatic mutations at each time point highlighted in spheres and colored red for mutations carried over from I8 to mature antibody, cyan for mutations carried over from I4 to mature antibody, green for mutations carried over from I3 to mature antibody, blue for mutations carried over from I1 to mature antibody, magenta for CH103 mutations from I1. Transient mutations that did not carry all the way to mature antibody are colored in deep olive. The antibody (paratope) residues are shown in surface representation and colored by their chemical types as indicated.