Sequence and Structural Convergence of Broad and Potent HIV Antibodies That Mimic CD4 Binding

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Passive transfer of broadly neutralizing HIV antibodies can prevent infection, which suggests that vaccines that elicit such antibodies would be protective. Thus far, however, few broadly neutralizing HIV antibodies that occur naturally have been characterized. To determine whether these antibodies are part of a larger group of related molecules, we cloned 576 new HIV antibodies from four unrelated individuals. All four individuals produced expanded clones of potent broadly neutralizing CD4-binding-site antibodies that mimic binding to CD4. Despite extensive hypermutation, the new antibodies shared a consensus sequence of 68 immunoglobulin H (IgH) chain amino acids and arise independently from two related *IgH* genes. Comparison of the crystal structure of one of the antibodies to the broadly neutralizing antibody VRC01 revealed conservation of the contacts to the HIV spike.

T wo to three years after infection, some HIV-infected patients develop serum antibodies that can neutralize a broad spectrum of HIV viruses (1-4). Among the naturally occurring monoclonal antibodies, VRC01, an antibody cloned from memory B cells that targets the CD4-binding site (CD4bs) on the HIV envelope spike, is unusual in its potency and breadth (5, 6). As do other HIV antibodies obtained by the single-cell antigen-capture method (7), VRC01 shows high levels of somatic mutations that are essential for its activity (6-8). This high frequency of mutation is a potential impediment to antibody identification, because the mutated sequences

Fig. 1. The 2CC core captures CD4bs antibodies. (A and B) Top line indicates bait used for sorting, and below is the patient number. The number in the center of the pies denotes the number of antibodies; slices are unique clones and are proportional to clone size. Membership of an antibody in a B cell clone is determined by sequence analysis, in particular CDR3s and shared V and J genes of paired heavy- and light-chain genes (table S3). (A) Pie charts show a summary of detected B cell clones irrespective of their binding epitopes. (\bigstar) Star indicates clones found in both YU2-gp140 and 2CC core-sorted cells. (B) Pie charts show the distribution of antibodies binding to CD4bs, CD4i, CD4ind. (equally affected by D368R and I420R), and core (not affected by either D368R or I420R). Dashed lines indicate antibodies that were cloned but could not be produced. (C) Representative ELISAs on YU2gp120, mutants, and 2CC core.

may no longer be complementary to the primers used for immunoglobulin gene amplification (9). To avert this potential problem, we developed a new primer set specifically designed to address this problem (the 5' primer is set farther upstream to avoid the mutated region) (fig. S1A and table S1) (10). The new strategy was tested by sorting single B cells from a patient with high titers of broadly neutralizing antibodies (Pt 8) (table S2) that bind to an HIV gp120 core glycoprotein stabilized in the CD4-bound conformation and lacking the variable (V) loops 1 to 3 (2CC core) (fig. S1B) (11, 12). In contrast to the resurfaced protein used to clone VRC01, which was designed to focus on antibodies to the CD4bs, the 2CC core should capture additional antibodies including those specific to the CD4-induced co-receptorbinding site (CD4i) (11, 12).

In side-by-side comparisons, the new primer set increased recovery of immunoglobulin H (IgH) chains when compared with the original primer set (fig. S2, A and B) (9). As expected, the antibodies obtained with the new primer set were more mutated (fig. S2, A and C) (average 35.7 versus 19.8, P=0.0013, and maximum 85 versus 50 for *IgH*) and included clones not found with the original primer set (fig. S2, A and B). Moreover, the new primers also recovered VRC01related antibodies from cDNA samples isolated from single cells that had been sorted with the original YU2-gp140 trimer probe used to develop the single-cell antibody-cloning method (7, 13) (fig. S3, A and B).

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Four unrelated HIV-infected individuals showing high titers of broadly neutralizing antibodies were examined by using the 2CC core (table S2 and fig. S4, A and B). Two of these individuals, Pt 1 and Pt 3, had been studied previously, but their cloned antibodies could not account for their serologic activity (7). From single sorted B cells representing 200 different B cell clones that were diversified by mutation in germinal centers, 576 antibodies were obtained from a starting population of 1.5×10^{5} IgG⁺ memory B cells (Fig. 1A and table S3). Representative members of each expanded B cell clone were tested for binding to gp120 and were all positive (Fig. 1, B and C; fig. S5; and table S3). The site of antibody binding on the envelope spike was mapped by using mutant proteins that interfere with either the CD4bs [gp120(D368R)] (14-16), or the CD4i site [gp120(I420R)] (17). [These mutants have substitutions, respectively, Arg for Asp at position 368 and Arg for Ile at position 420.] NIH45-46, which is a VRC01 variant, and antibodies 3BNC60, 8ANC131, and 12A12 (antibodies selected on the basis of neutralizing activity, see below) (Fig. 1C, fig. S5, and table S3) showed binding patterns similar to VRC01's. Others, including 1B2530 and 8ANC195, could not be classified precisely solely on the basis of enzyme-linked immunosorbent assay (ELISA). As expected from earlier

Fig. 2. Antibodies captured by 2CC core have broad and potent HIV-neutralizing activity. (A) HIVneutralizing activity assayed on a limited panel of viruses. Top line indicates the donor number (table S2). Antibody clones are grouped, and individual clonal relatives are represented by a column grouped as in table S4. Each row represents one virus as indicated on the left. Colors indicate concentration of clones at the median inhibitory concentration (IC₅₀): red, \leq 0.1 µg/ml; orange, 0.1 to 1 µg/ml; yellow, 1 to 10 μ g/ml; green, \geq 10 μ g/ml; and white, not neutralized at any concentration tested. (\bigstar) Star indicates the representatives selected for the extended virus panel (table S5). (B) HIV-neutralizing activity assayed on an extended panel of viruses (tables S5 and S6). (C) Neutralization summary graph comparing the published IC₅₀ values of VRC01 (6) with NIH45-46 and 3BNC117 (tables S5 and S7). Each color represents a different HIV clade (black corresponds to CRF01AE, blue to clade B, green to clade G, pink to clade D, gray to clade CD, emerald to clade C, brown to clade ACD, magenta to clade AC, red to clade A, and orange to CRF02AG). Length of lines and size of circles are inversely proportional to IC₅₀. The distance between the outer and inner circle, as well as from the inner circle to the center of the spiders, in each span is two logs in IC_{50} concentration.

studies on HIV envelope–specific antibodies (8), 65% of the antibodies isolated by using the 2CC core were polyreactive (fig. S6) compared with 22.7% polyreactivity in healthy control memory B cells (18) and 17.3% in gp140-negative B cells from HIV-positive controls (8). Somatic hypermutation was likely required for development of high-affinity antigen binding and polyreactivity because reversion of four representative antibodies to the corresponding germ line led to complete loss of binding to YU2-gp140 (13) (fig. S6B and fig. S7, A to C).

HIV-neutralizing activity was measured in vitro by using an initial panel of eight viruses including three tier 1 clade A, B, and C, and five tier 2 clade B envelope (Env) pseudovirus variants (19, 20). The neutralizing activity of the antibodies was compared with VRC01 and purified serum IgG from the donors (Fig. 2A, fig. S4, and table S4). A selection of 11 representative antibodies showing high levels of neutralizing activity (Fig. 2A and table S5) were further tested on a panel of 15 additional tier 2 Env pseudovirus variants (Fig. 2B and table S6), including five viruses that are resistant to VRC01 (Fig. 2B and table S6). Of all of the antibodies tested, 88% showed some neutralizing activity, and six clones (RU01, 08, 10, 12, 16, and NIH45-46) contained antibodies that were highly potent and broad

(Fig. 2 and tables S4 and S6). These clones were also the most abundant among those captured by 2CC core in each of the four patients studied (Fig. 1A and table S3). Five antibodies representing four different broadly neutralizing founder B cell clones [clone RU01 (3BNC117 and 3BNC55), clone RU16 (12A12), clone RU12 (8ANC195), and NIH45-46] (table S5), were tested against an expanded panel of 118 tier 2 viral isolates from all known clades, including 32 transmitted founder viruses (Fig. 2C and table S7). [VRC01 had previously been tested on 82 of these (6).] The most impressive of the new antibodies, 3BNC117, belonging to a clone with 85 members (RU01), showed an arithmetic mean 80% inhibitory concentration (IC₈₀) on a combined group of 95 tier 2 viruses of 1.4 µg/ml (median and geometric mean IC₈₀ values 0.3 µg/ml, table S7B). When compared with the previously published VRC01 neutralization data, only 17 of the viruses tested were more sensitive to VRC01 than 3BNC117, and 3BNC117 showed greater breadth (Fig. 2, B and C, and tables S4, S6, and S7) (6). NIH45-46, a new variant of VRC01, was more potent than VRC01 on 62 of the viruses tested but still less potent than 3BNC117 (Fig. 2, B and C, and tables S4, S6, and S7) (6). Together, the new antibodies neutralized 96% of the 118 viruses tested (table S7A). Finally, the best antibodies were highly hypermutated,



and this was essential for their breadth and potency (fig. S7D and tables S4, S6, and S7).

Our cloning strategy captures antibodies produced by antigen-binding memory B cells in the blood, but circulating antibodies are not produced by these cells; they originate instead from plasma cells in the bone marrow (21). To determine whether the antibodies cloned from memory B cells are also found in the bone marrow plasma cell compartment, we purified plasma cells from paired bone marrow samples from patients 3 and 8 (table S2 and fig. S8A) and used polymerase chain reaction (PCR) to specifically amplify IgV_H genes from the clones RU01 and RU10 from memory B cells in these individuals (Fig. 2A, fig. S8A, and table S3). Members of these clones and large numbers of additional variants were readily identified in the respective plasma cell samples (fig. S8, B and C). We also verified that antibodies from clones RU01, RU08, and RU10 (Fig. 2A and table S3) are found in serum by mass spectrometry (fig. S9 and table S8).

To determine whether antibody affinity to gp120 is related to neutralizing activity, we compared the binding of the highly active antibodies, selected clonal relatives, and germ line-reverted progenitors by using surface plasmon resonance (SPR) (Fig. 3, A and B; figs. S7B and S10; and table S9). The top neutralizing antibodies (tables S4 to S7) showed affinities (K_A) ranging from $\simeq 10^7$ to 10^{12} (M⁻¹) to YU2-gp140 trimers (13) and $\simeq 10^7$ to 10^{11} (M⁻¹) to the 2CC core (11) (Fig. 3, A and B, and tables S4, S6 and S9). Consistent with their decreased neutralizing potency and breadth, 3BNC66, 3BNC156, and 3BNC55 displayed lower affinities to the YU2-gp140 trimer than 3BNC117, but, surprisingly, the affinities of these antibodies to 2CC core did not correlate with their neutralizing activity (Fig. 2, fig. S10, and tables S4 and S9). Finally, we were unable to detect binding by any of the germ line-reverted antibodies tested (Fig. 3B, fig. S7B, and table

S9). We conclude that the antibodies captured by the 2CC core tend to show higher affinity to the YU2-gp140 trimer than to the 2CC core.

When VRC01 binds to the HIV spike, it produces large conformational changes that mimic CD4 binding and expose the CD4i site (6). By contrast, the broadly neutralizing antibody b12 (22) and many other known CD4bs antibodies do not (23). To determine whether the ability to mimic CD4 is a shared feature of the most potent antibodies (tables S4 to S7), we expressed two different HIV spikes (24) on the surface of human embryonic kidney-293 (HEK 293T) cells and measured CD4i-antibody binding in the presence or absence of CD4 or CD4bs antibodies (Fig. 3C). With one exception, all of the antibodies tested resemble CD4 and VRC01 in that they facilitate CD4i-antibody binding to one or both viral spikes (Fig. 3C and fig. S11). The only antibody tested that did not share this characteristic, 8ANC195, was not a traditional CD4bs antibody in that it was equally sensitive to the D368R and I420R mutations (Fig. 1C and table S3), and it differed from the others in its neutralization pattern (Fig. 2, A and B; and tables S4, S6, and S7).

To determine whether highly active CD4bs antibodies share common sequence features, we aligned the 10 best antibodies (table S5): two variants each from independently derived antibody clones arising in each of the four patients studied and from NIH45 (Fig. 4, A and B; table S2; and fig. S3) (6). Comparison of the IgV_H regions revealed a conserved consensus sequence covering 68 IgV_H residues (Fig. 4A). It is noteworthy that the IgV_H consensus contains seven VRC01-gp120 contact residues, including Arg 71_{VRC01} , which mimics the key interaction of Arg59_{CD4} and Asp368_{gp120} (Fig. 4A) (5, 25). All 10 antibodies arise from only two closely related germline IgV_H genes that conserve six contact residues (Fig. 4A, fig. S12A, and table S3). The

codons of the consensus residues are highly somatically mutated in the 10 selected antibodies; however, the ratio of replacement to silent mutations in the consensus residues ranges from 0.7 to 1.7, whereas it is 3.5 to 22 in the nonconsensus residues, which together indicate that conservation of the consensus is strongly selected (fig. S13 and table S10). The light chain of VRC01 makes fewer contacts with gp120 (5). Consistent with its more limited role, comparison of the light-chain sequences of the same antibodies uncovers a less-extensive consensus covering 53 residues from the variable region of immunoglobulin light chain (VL residues), which includes three VRC01-gp120 contact residues (Fig. 4B and fig. S3B). Finally, like the heavy chains, the light chains arise from a limited set of germline genes: Two are derived from IgK1D-33, two from IgK3-11, and one from IgL1-47 (Fig. 4B, fig. S12B, and table S3) (5). Antibody 8ANC195, which differed from the others in several important respects (Figs. 1C, 2B, and 3C; fig. S11; and tables S4, S6, and S7), did not entirely conform to the consensus and did not arise from related heavy or light chains (Fig. 4, A and B; fig. S14; and table S3). Thus, there is significant sequence convergence among highly active agonistic CD4bs antibodies (HAADs).

To determine whether the structure of the antibodies in different patients is also conserved, we solved the crystal structure of the 3BNC60 Fab to 2.65 Å resolution and compared it with VRC01s (26) (Fig. 4, C to E). Superimposition of domains in the variable region of immunoglobulin heavy chain (V_H domains) from 3BNC60 and VRC01 in the VRC01-gp120 cocrystal structure (5) yields a root mean square deviation (RMSD) of 1.3 Å (calculated for 111 Ca atoms) with major differences confined to CDR2 residues 58 to 65 (3BNC60 numbering). Superimposing the structures suggests conservation of the recognition interface with gp120 (Fig. 4D). For example,

293T^{YU.2}



and table S5), and 12A germ line (GL)-reverted antibodies. (B) Graph shows K_A for representative antibodies (tables S5 and S9). (C) Graph shows mean fluorescence intensity of CD4i antibody 3-67 (7) binding to Bal-expressing 293T cells after incubation with the indicated antibodies. Table indicates whether or not an antibody induces CD4i site accessibility (fig. S11).

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Arg723BNC60 adopts a conformation similar to that of Arg71_{VRC01}, which mimics an important salt bridge normally formed between Arg59_{CD4} and Asp368_{gp120} (5, 25). Gln65_{3BNC60}, which corresponds to Gln64_{VRC01}, is within the residue segment (residues 58 to 65) that differs in structure from VRC01. The conformation of this region of 3BNC60, which is involved in a lattice contact in the crystals, is likely to change upon binding gp120, as it would clash with the CD4-binding loop on gp120. Superimposing the 3BNC60 and VRC01 V_L domains yields a RMSD of 0.9Å (calculated for 95 C α atoms) and shows that some of gp120-contacting residues are structurally conserved (Fig. 4E); Tyr913BNC60 and Glu903BNC60 adopt similar conformations as Tyr91_{VRC01} and Glu96_{VRC01}, which engage loop D of gp120 via polar interactions (Fig. 4E). The V_H and V_L domains of the two Fabs show similar interdomain interactions and overall orientation, as superimposition of the V_H and V_L domains of 3BNC60 and VRC01 yields an RMSD of 1.4 Å (calculated for 206 C α atoms). Overall, these structural comparisons suggest that 3BNC60 binds gp120 with the same architecture as observed for the binding of VRC01 (5).

Our experiments define a class of agonistic CD4bs antibodies, HAADs, that shares *IgVH* and *IgVL* consensus sequences including 10 of the contact residues between VRC01 and the HIV spike (Fig. 4, A and B). In five different donors, these antibodies originate from only two closely related *IgVH* and three *IgVL* genes (Fig 4, A and B). Although we cannot exclude the possibility that by using the 2CC core as bait for single-cell sorting might introduce a selection bias, gp140 sorts performed on two of the donors did not reveal additional broadly neutralizing antibodies (tables S3, E and F, and S4), which suggests that a significant fraction of the CD4bs-directed rep-

ertoire was captured in our experiments. Thus, the repertoire of antibodies that are available to solve the problem of broad and potent HIV neutralization by binding to the CD4bs appears to be somewhat restricted. Despite this restriction, HAADs account for a significant fraction of the memory B cell, and plasma cell compartments, and the circulating IgGs in the patients tested. Therefore, HAADs are not rare, and, once elicited, they contribute to the circulating antibody pool in patients that neutralizes the virus broadly. However, HAADs are not found in all patients with high titers of broad neutralizing activity (1, 27, 28). Moreover, broad neutralization can be achieved by targeting epitopes other than the CD4bs (29-34) or by combinations of antibodies to different epitopes (7, 35), or even combination of different CD4bs antibodies (table S7).

The high levels of neutralizing activity displayed by HAADs appear to be correlated with



Fig. 4. Sequence and structural conservation of HAADs. (**A**) Amino acid alignment of 10 selected HAADs (table S5), their germline genes, and 8ANC195. Residues are numbered according to the 3BNC60 structure. Framework (FR) and CDR regions are indicated. Red shading shows amino acid identity; yellow shows biochemical similarity. The consensus is defined by 70% similarity between the 10 selected HAADs. The consensus sequence is shown above; dashes in this sequence indicate nonconserved residues. Contact residues between VRC01 and gp120 are shown above the consensus as closed circles for main-and side-chain interactions, open circles main chain only, and stars side chains

only (5). (**B**) As in (A) for light chains. (**C**, **D**, and **E**) Crystal structure of 3BNC60 Fab. (C) Superimposed C α traces of the two Fab molecules in the 3BNC60 asymmetric unit are shown in green and red. Semitransparent surfaces are used to outline the heavy (cyan) and light (gray) chains. (D) Superimposition of the 3BNC60 V_H (red, C α trace) and VRC01 V_H (cyan, C α trace) shown with a ribbon representation of the CD4-binding loop. The salt bridge between Arg71_{VRC01} and Asp368_{gp120} is shown as dashed lines. (E) Superimposition of the 3BNC60 VL (red, C α trace). Hydrogen bonds between VRC01 and gp120 are shown as dashed lines. binding to a specific surface, which is similar, but not identical, to that contacted by CD4 (5, 36). Why a specific type of binding that mimics CD4 appears to be required is not known (6); however, it has been suggested that binding of soluble CD4, or CD4 mimetic compounds, to the HIV envelope trimer leads to the formation of a metastable intermediate that decays rapidly and loses the ability to fuse (37). We speculate that HAAD binding mimics soluble CD4 and destabilizes the trimer. Irrespective of their mechanism of action, HAADs might contribute to viremic control in a subset of HIV-infected individuals and may be useful in HIV prevention, or possibly even therapy, because of the low concentrations required for viral neutralization.

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- tabulated in the supporting online material, and the coordinates for the 3BNC60 Fab as well as the antibody sequences have been deposited in the Protein Data Bank and the European Molecular Biology Laboratory-European Bioinformatics Institute database [accession codes: 3RPI (3BNC60 coordinates), HE584535 to HE584554 (antibody sequences)]. We thank the HIV-positive donors for their support. We thank M. Jankovic for discussions and help with cloning of HIV envelope constructs; R. Wyatt for the plasmid of YU2-gp140;]. Mascola for the plasmid of 2CC-core and YU2-gp120 mutants; and M. Connors for access to patient NIH45; C. Gaebler, M. Warncke, P. M. Marcovecchio, and H. Gao for help with protein expression; K. Moss for patient coordination. We thank F. McCutchan, G. Shaw, B. Hahn, J. Baalwa, D. Montefiori, F. Gao, M. Thomson, J. Overbaugh, R. Swanstrom, L. Morris, J. Kim, L. Zhang, D. Ellenberger,

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Supporting Online Material

www.sciencemag.org/cgi/content/full/science.1207227/DC1 Materials and Methods Figs. S1 to S14 Tables S1 to S11 References (38–51)

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Yeast Rrn7 and Human TAF1B Are TFIIB-Related RNA Polymerase I General Transcription Factors

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Eukaryotic and archaeal multisubunit RNA polymerases (Pols) are structurally related and require several similar components for transcription initiation. However, none of the Pol I factors were known to share homology with transcription factor IIB (TFIIB) or TFIIB-related proteins, key factors in the initiation mechanisms of the other Pols. Here we show that Rrn7, a subunit of the yeast Pol I core factor, and its human ortholog TAF1B are TFIIB-like factors. Although distantly related, Rrn7 shares many activities associated with TFIIB-like factors. Domain swaps between TFIIB-related factors show that Rrn7 is most closely related to the Pol III general factor Brf1. Our results point to the conservation of initiation mechanisms among multisubunit Pols and reveal a key function of yeast core factor/human SL1 in Pol I transcription.

Transcription initiation by most eukaryotic and archaeal multisubunit polymerases (Pols) requires transcription factor IIB (TFIIB) or a TFIIB-related general transcription factor. TFIIB and TFIIB-like proteins (archaeal Tfb and the Pol III factor Brf) operate at multiple key steps during transcription initiation such as Pol binding, preinitiation complex formation, start site selection, promoter opening, and abortive initiation (1–4). Because of the fundamental and conserved role of TFIIB-like factors for other Pols, it was surprising that none of the Pol I– specific subunits or general initiation factors were known to share TFIIB homology.

Transcription of the *Saccharomyces cerevisiae* ribosomal DNA (rDNA) locus by Pol I requires four general transcription factors: UAS-binding upstream activity factor (UAF), TATA box– binding protein (TBP), core factor (CF) and the regulatory factor Rm3 (5). CF contains subunits Rm6, 7, and 11 (6, 7) and interacts with UAF, TBP, Rm3, and Pol I (5). CF is analogous to the human Pol I factor SL1, composed of four TBP-associated factors (TAFs), TAF1A, TAF1B, TAF1C, and TAF1D ((8, 9)), where TAF1A, B, and C are orthologous to yeast CF subunits Rm6, 7, and 11 ((10)). Both CF and SL1 recruit Pol I to its promoter, but the function of the CF/SL1 subunits, apart from TBP and Pol binding, is unknown ((5, 11, 12)).

We guessed that a Pol I TFIIB-like factor would have diverged considerably from related factors (13) because Pol I subunits share relatively low protein sequence conservation with their Pol II and Pol III counterparts (14). Using the homology detection program HHpred, which uses pairwise hidden Markov model profile comparisons that are more sensitive than traditional Web-based approaches (15), we detected highprobability matches between the Rm7 N-terminal 320 residues and the TFIIB family, indicating that

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