Near-Pan-neutralizing, Plasma Deconvoluted Antibody N49P6 Mimics Host Receptor CD4 in Its Quaternary Interactions with the HIV-1 Envelope Trimer

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ABSTRACT The first step in HIV-1 entry is the attachment of the envelope (Env) trimer to target cell CD4. As such, the CD4-binding site (CD4bs) remains one of the few universally accessible sites for antibodies (Abs). We recently described a method of isolating Abs directly from the circulating plasma and described a panel of broadly neutralizing Abs (bnAbs) from an HIV-1 "elite neutralizer" referred to as patient N49 (N49 Ab lineage [M. M. Sajadi, A. Dashti, Z. R. Tehrani, W. D. Tolbert, et al., Cell 173:1783–1795.e14, 2018, https://doi.org/10.1016/j.cell.2018.03.061]). Here, we describe the molecular details of antigen recognition by N49P6, an Ab of the N49 lineage that recapitulates most of the neutralization breadth and potency of the donor’s plasma IgG. Our studies done in the context of monomeric and trimeric antigens indicate that N49P6 combines many characteristics of known CD4bs-specific bnAbs with features that are unique to the N49 Ab lineage to achieve its remarkable neutralization breadth. These include the omission of the CD4 Phe43 cavity and dependence instead on interactions with highly conserved gp120 inner domain layer 3. Interestingly, when bound to BG505 SOSIP, N49P6 closely mimics the initial contact of host receptor CD4 to the adjacent promoter of the HIV-1 Env trimer to lock the trimer in the closed conformation. Altogether, N49P6 defines a new class of near-pan-neutralizing, plasma deconvoluted CD4bs Abs that we refer to as the N49P series. The details of the mechanisms of action of this new Ab class pave the way for the next generation of HIV-1 bnAbs that can be used as vaccine components of therapeutics.

IMPORTANCE Binding to target cell CD4 is the first crucial step required for HIV-1 infection. Thus, the CD4-binding site (CD4bs) is one of the most accessible sites for antibodies (Abs). However, due to steric constraints, only a few Abs are capable of targeting this site. Here, we show that the exceptional neutralization breadth and potency of N49P6, a near-pan-neutralizing Ab targeting the CD4bs isolated from the plasma of an HIV-1 "elite neutralizer," patient N49, are due to its signature combination of more typical CD4bs Ab-binding characteristics with unique interactions with the highly conserved gp120 inner domain. In addition, we also present a structural analysis of N49P6 in complex with the BG505 SOSIP trimer to show that N49P6 exhibits remarkable breadth in part by mimicking CD4’s quaternary interaction with the neighboring gp120 protomer. In its mode of antigen interaction, N49P6 is unique and represents a new class of CD4bs-specific bnAbs.

KEYWORDS CD4-binding site, HIV, N49P lineage, near-pan-neutralizing, neutralizing antibodies

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Broadly neutralization antibodies (bnAbs) capable of neutralizing diverse circulating HIV-1 strains are considered the key for a successful vaccine or passive prophylaxis against HIV-1. High mutation rates, polymorphisms, altered glycosylation patterns forming a protective “glycan shield,” and conformational heterogeneity of the envelope glycoprotein (Env) trimer all contribute to HIV-1 variability both within a patient and, to a greater extent, within a population driving viral escape from the immune system (1). Despite these obstacles, potent bnAbs have been isolated that target various distinct vulnerable epitopes within Env, including the CD4-binding site (CD4bs) (2), the V1/V2 loop, the V3 glycan patch (3), the membrane-proximal external region (MPER) (4), and the gp120-gp41 interface (5). Of particular interest is the CD4bs because the infectivity of HIV-1 largely relies on the successful engagement of Env with the host CD4 receptor. Due to this critical functional constraint, conservation of the CD4bs remains relatively high among circulating viruses, rendering them susceptible to neutralizing antibodies. Indeed, members of a group of antibodies targeting CD4bs exhibit both cross-clade neutralization and good potency. Many of them share similar characteristics in heavy chain gene usage and gp120 recognition mode and thus have been categorized as “VRC01 class” antibodies, after the first isolated member (6). The bnAbs within this class typically take longer to develop in natural infection than other bnAb specificities and contain, on average, a high degree of somatic hypermutation (SHM) and a short 5-residue CDR L3 (complementarity-determining region light chain 3) (7, 8). The emergence of single B cell sorting and monoclonal antibody (mAb) isolation enabled the discovery of many potent VRC01 class bnAbs from HIV-infected donors. Of note, some of the most potent CD4bs bnAbs isolated from B cells included N6, 1-18, 3BNC117, and VRC07 (9–12). However, memory B cell repertoires have been shown to differ from those from circulating plasma (13, 14), and there is often a discordance between memory B cell pools and anti-Env circulating antibody responses, which shows that the HIV neutralization profiles of memory B cell-derived mAbs do not always match those found in plasma (15–17).

Recently, by using proteomic and genomic analyses, we described a panel of near-pan-neutralizing antibodies, including N49P7, from the plasma of an HIV-1 “elite neutralizer,” donor N49. In addition to being an elite neutralizer, donor N49 belonged to a cohort of viremic controllers within a natural viral suppressor (NVS) cohort of HIV subtype B-infected donors (18, 19). N49P7 was distinct from other bnAbs in the VRC01 class as it made substantially more contact with conserved residues within the gp120 inner domain, contributing to its near-pan-neutralizing ability, with a median 50% inhibitory concentration (IC50) of 0.10 μg/ml (20). Also isolated from donor N49, mAb N49P6 exhibits slightly less potency with a similar remarkable neutralization breadth. A unique feature of both bnAbs N49P6 and N49P7 is that they completely recapitulate the neutralization breadth of the donor N49 anti-Env polyclonal plasma IgG, exhibiting broad and high potency with the ability to neutralize all 117 pseudoviruses in an HIV “global panel.” In this study, we characterize the unique mechanisms of the neutralizing activity of bnAb N49P6 by describing the molecular details of N49P6’s interaction with monomeric and trimeric Env antigens. The antigen-binding features of N49P6 are discussed in the broad context of antigen recognition of known CD4bs-specific bnAbs, including the VRC01-like class bnAbs, to describe the unique characteristics of Abs of the N49 lineage that allow them to achieve high potency and near-pan-neutralization breadth.

RESULTS
N49P6 shows extraordinary neutralization potency and breadth. Using a multi-clade, 117-global-pseudovirus panel, we examined the neutralization activity of monoclonal antibody (mAb) N49P6. Figure 1 shows the neutralization profile of N49P6 compared to N49P7, a near-pan-neutralizing mAb of the same lineage isolated from donor N49 and previously characterized by us (20). N49P6 showed high potency, with a median 50% inhibitory concentration (IC50) and 80% inhibitory concentration (IC80) of 0.31 μg/ml and 0.71 μg/ml,
respectively. N49P6, similar to N49P7, exhibits extraordinary neutralizing activity, capable of neutralizing 71.8% of all viruses at an IC50 of 1 mg/ml and 49.6% of all viruses at an IC80 of 1 mg/ml, compared to 86.3% and 77.8% for N49P7, respectively. Of note, both are among the broadest mAbs discovered thus far and show greater combined breadth and potency than VRC01, VRC07, 3BNC117, NIH45-46, PG9, PG16, PGDM1400, PGT121, PGT128, PGT145, PGT151, 8ANC195, and 10-1074 (20).

N49P6 recognizes highly conserved residues within gp12093TH057coree.

In order to better understand the features of N49P6 that give it its extraordinary breadth and potency, we determined the crystal structure of N49P6 Fab-gp12093TH057coree to a 2.55-Å resolution (Table 1). As shown in Fig. 2, N49P6 primarily binds through its VH1-2 heavy chain by mimicking many of the gp120-binding characteristics of CD4. Heavy chain CDR H2 and CDR H3 contribute the most to binding, representing 39% and 27% of the total buried surface area (BSA) of the Fab, respectively (see Table S1 in the supplemental material). CDR H1, in contrast, contributes only 5 Å² (0.7%) to the heavy chain BSA. Of these two major anchoring elements, CDR H2 contributes the most to

FIG 1 Neutralization profiles of mAbs N49P6 and N49P7. (A) A multiclade, 117-HIV-1-pseudovirus panel was tested against mAbs N49P6 and N49P7. IC50 and IC80 values are color-coded according to their potencies. Both mAbs N49P6 and N49P7 exhibited 100% breadth, with all pseudoviruses having an IC50 of <50 μg/ml. (B) Geometric mean IC50 and IC80 values of mAbs N49P6 and N49P7 against each clade and the whole panel.

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the BSA of the interface (373 Å² of the total 733-Å² heavy chain BSA), which includes a short antiparallel β-sheet (N49P6 residues 55 to 57) that interacts with residues in the gp120 CD4-binding loop (gp120 residues 365 to 368) (Fig. 2B). These contacts are “cha- peroned” by a salt bridge between Asp368 of gp120 and a heavy chain framework region 3 (FWR3) residue of N49P6, Arg71, found in many VRC01-like bnAbs (Fig. 2C). In addition, CDR H2 also interacts with gp120 loop D (through a hydrogen bond formed between Trp50 [CDR H2] and Asn280 [loop D]) and gp120 loop V5 (through contacts with gp120 residues 455 to 459). Interestingly, although N49P6 mimics many of CD4’s binding characteristics, Gly54 in CDR H2 of N49P6 bypasses the Phe43 (CD4)-binding cavity, a hydrophobic pocket used by CD4 and many CD4 mimetics to bind to Env (Fig. 2B). N49P6 shares this feature with N49P7, which indicates that this Ab lineage

| TABLE 1 Data collection and refinement statisticsa |  |
|--------------------------------------------------|---|
| Parameter | Value(s) for N49P6 Fab in complex with: |
| HIV-1 clade A/E strain | HIV-1 BG505 SOSIP.664 |
| 93TH057 gp120core | Env trimer ectodomain |
| Data collection |  |
| Wavelength (Å) | 0.979 | 0.979 |
| Space group | P2₁,2,2₁ | P2₁ |
| a, b, c (Å) | 65.4, 80.8, 195.2 | 164.7, 164.7, 164.7 |
| α, β, γ(°) | 90, 90, 90 | 90, 90, 90 |
| Complexes (ASU) | 1 | 1 |
| Resolution (Å) | 50–2.55 (2.59–2.55) | 50–4.05 (4.12–4.05) |
| No. of reflections | 111,144 | 68,987 |
| Unique | 31,853 (1,524) | 12,543 (608) |
| Rmerge b (%) | 27.6 (72.7) | 11.3 (100) |
| Rapim c (%) | 16.6 (55.6) | 5.3 (75.7) |
| CC1/2 d | 0.99 (0.56) | 0.99 (0.48) |
| Wilson B factor (1/Å²) e | 41 | 176 |
| l/σ | 6.4 (1.1) | 22.8 (1.0) |
| Completeness (%) | 92.0 (89.3) | 99.6 (99.8) |
| Redundancy | 3.1 (2.2) | 5.5 (4.6) |
| Refinement statistics |  |
| Resolution (Å) | 50.0–2.55 | 50.0–4.05 |
| R (%) | 21.9 | 25.5 |
| Rfree (%) | 27.3 | 31.5 |
| No. of atoms |  |  |
| Protein | 5,872 | 7,725 |
| Water | 39 |  |
| Ligand/ion | 154 | 544 |
| Overall B factor (Å²) | 63 | 219 |
| Protein | 55 |  |
| Water | 90 | 322 |
| Ligand/ion |  |  |
| Root mean square deviation | 0.011 | 0.008 |
| Bond lengths (Å) | 1.3 | 1.2 |
| Ramachandran plot (%) | 91.4 | 81.2 |
| Favored | 6.5 | 13.1 |
| Allowed | 2.1 | 5.7 |
| Outliers |  |  |
| PDB accession no. | 6OZ2 | 6OZ4 |

aValues in parentheses are for the highest-resolution shell. ASU, asymmetric unit.

bRmerge = \sum_j |J − \langle J\rangle|/\sum_j \langle J\rangle, where I is the observed intensity and \langle J\rangle is the average intensity obtained from multiple observations of symmetry-related reflections after rejections.

cRapim, as defined in reference 53.

dCC1/2, as defined by Karplus and Diederichs (54).

eWilson B factor, as calculated in reference 55.
developed an alternate strategy to compensate for the loss of binding energy by using Gly instead of an amino acid with a bulky hydrophobic side chain at this position (20); some, but not all, VRC01-like Abs use the Phe43 cavity for binding Env, although this may decrease the breadth in the presence of cavity-filling mutations such as Ile375 in the N6-resistant pseudovirus CT565_C7_48. An important anchoring element that may

![Crystal structure of the N49P6 Fab-gp120 complex.](image)

FIG 2 Crystal structure of the N49P6 Fab-gp120core complex. (A) Overall structure of the complex shown as a ribbon diagram. The complementarity-determining regions (CDRs) of N49P6 Fab are colored as follows: CDR H1 is black, CDR H2 is yellow, CDR H3 is red, CDR L1 is dark green, and CDR L3 is blue. The outer and inner domains of gp120 are dark and light gray, respectively. The outer domain D and E, CD4 binding, and V5 loops are shown in cyan, purple-blue, magenta, and green, respectively. Carbohydrates at positions N276 (loop D) and N355 (loop E) are shown as sticks. (B) N49P6 Fab-gp120core interface with coloring as described above for panel A. (Top) N49P6 Fab is shown as a molecular surface, and the gp120 contact residues are shown as sticks. (Bottom) A 180° view reveals the detailed interaction of N49P6 Fab and the gp120 surface. Contact residues of N49P6 Fab are shown as sticks, and all Fab contact residues are listed. gp120 is shown as a molecular surface and colored according to its electrostatic potential, with red, blue, and white representing negative, positive, and neutral electrostatic potentials, respectively. (C) Blowup view of the hydrogen bond network of the interaction between N49P6 Fab and the gp120 surface. Three hydrogen bonds (CDR H2 W50 and loop D N280, CDR H3 W100J and loop D N279, and CDR L3 E96 and loop D N280) and a salt bridge (heavy chain residue R71 and CD4-binding loop D368) are formed at the interface. Residues contributing to the interaction are shown as sticks, and the hydrogen bond network is shown with red dashed lines.
compensate for this loss in N49P6 is the 20-amino-acid (aa)-long CDR H3 (Fig. 2B and C). N49P6’s CDR H3 contacts gp120 loop D (gp120 residues 275 to 276 and 279 to 283), establishing a Trp100J (CDR H3)-Asp279 (loop D) hydrogen bond, and contacts the conserved gp120 inner domain layer 3 (gp120 residues 474 and 476). As we discuss below, the inner domain contacts are unique to the N49 lineage.

The contribution of the N49P6 lambda light chain to the gp120 core complex interface is minimal, representing only 25% of the total Fab BSA (18% if excluding the contribution from the Asn276 glycan). N49P6’s light chain contains both deletions in its 8-amino-acid-long CDR L1 and a shortened CDR L3 to minimize light chain-mediated viral escape. The truncated CDR L1 contacts only the N-acetylglucosamine linked to Asn276 on gp120 loop D. The short CDR L3 also contacts the Asn276 glycan as well as neighboring residues in loop D (gp120 residues 276 and 278 to 280) and loop V5 (gp120 residues 458 to 459), with CDR L3 Phe91 making van der Waals contacts with gp120 Asn279 and Thr278 and CDR L3 Glu96 forming a hydrogen bond with the gp120 Gly459 main chain. Steric clashes between the light chain and the Asn276 glycan are a major limiting factor in breadth for CD4bs bnAbs, with most having deletions in CDR L1 in addition to a 5-residue CDR L3; N49P6 has one of the shortest CDR L1s, 1 amino acid shorter than VRC01 and 3 amino acids shorter than N6 (Fig. S1). In addition, N49P6 and N49P7 have a Cys at framework residue 36 of the light chain, normally Tyr, which allows the light chain to rotate further away from gp120 and better accommodate the Asn276 glycan (20).

N49P6 closely resembles N49P7 in binding to gp120 93TH057core. N49P6 is clonally related to N49P7, another antibody from the same donor. N49P6 and N49P7 share similar breadths, with N49P7 being slightly more potent (Fig. 1). The available high-resolution crystal structure of the complex of the N49P7 Fab with gp120core, of the same clade A/E 93TH057 strain determined by us previously (20) allowed us to compare their mechanisms of attachment and determine what contributes to the breadth and potency of this antibody class. Figure 3 shows a structural comparison of both complexes and a detailed analysis of specific antibody-antigen contacts. Superimposition of the N49P6 and N49P7 complexes based upon gp120 93TH057core (Fig. 3A) revealed no significant structural differences (Cα carbon root mean square deviations of 1.53 Å for the complex and 0.89 Å for the complex minus the constant part of the Fab), although N49P6 has a 1-amino-acid-longer CDR H3 than N49P7 (Arg100C represents an insertion in N49P6 relative to N49P7). In the N49P6 gp120 complex, Arg100C points away from gp120 and does not contribute to binding. However, we noticed a slight difference in the relative orientations of VL and VH in the N49P6 structure, with VL tilting further away from loop D, which results in a slightly smaller light chain footprint on gp120, as shown in Fig. 3B. Both N49P6 and N49P7 recognize a mixed inner domain/CD4-binding-site epitope referred to as the iCD4bs epitope, which is composed of highly conserved regions in gp120 (Fig. 3D). As shown in other studies, the gp120 inner domain, consisting of three mobile layers and a 7- or 8-stranded β-sandwich, contributes to Env integrity as well as to conformational transitions during viral fusion and therefore harbors some of its most highly conserved residues (21, 22). Specifically, CDR H2 and CDR H3 of N49P6 interact with residues 97, 102, and 124 of layer 2 and residues 473, 474, 476, and 480 of layer 3.

Interestingly, structural analyses indicate that the slightly lower neutralization potency of N49P6 than of N49P7 can be attributed to a few sequence changes in CDR H2 and CDR H3 (lower BSA due to Ser53-versus-Met, Gly56-versus-Gln, and Thr57-versus-Val differences). In addition, the insertion of an extra Arg in CDR H3 of N49P6 (Arg100C) relative to N49P7 results in a slightly lower BSA for Glu100F and Val100H, even though their side chain positions are largely superimposable (Fig. 3C and D). The only other significant difference in BSAs between the two epitopes resides on the light chain, with N49P7 having a slightly higher BSA in its CDR L3 due to a Phe91-versus-Tyr difference in their sequences.

N49P6 and N49P7 are unique among VRC01-like CD4bs Abs in how they extend their gp120 footprint and increase their neutralization breadth. The CD4-binding site is recessed in the HIV-1 trimer and provides only limited access to binding partners. Only a few classes of Abs that target this site can meet these requirements and effectively neutralize HIV-1, i.e., those derived from the heavy chain germ line allele Vα1-2 (VRC01 class) or Vα1-46 (8ANC131 class) and those that predominately bind with their
CDR H3 (6). To better understand how N49P6 and also N49P7 fit into these categories, we compared their structures to those of bnAbs of this class whose gp120core structures are available, specifically, VRC01, VRC03, VRC07, NIH45-46, and N6 (Fig. 4 and Fig. S1). As can be expected due to their shared germ line VH1-2 allele, there is a high degree of similarity in how CDR H2 of each of these Abs binds gp120. The Thr57-to-Val difference between N49P6 and N49P7 that results in an approximately 12-Å² increase in the BSA for N49P7 seems common for all other members examined here (Fig. S1); Thr57 in N49P6 forms a weak hydrogen bond with gp120 Gly366 and may not pack as well against Ser365 as does Val57 in N49P7, which forms van der Waals contacts with both residues. Similarly, the choice of a larger hydrophobic residue than serine at position 53 seems the norm, with Met for N49P7 and Leu or Arg for all others except N6 with Gln.

A common strategy used by VRC01-like Abs to increase binding energy is the use of the CD4 Pheε1-binding cavity. Both VRC03 and N6 do this by having a Trp or Tyr at heavy chain position 54, respectively. N49P6 and N49P7 but also VRC01, VRC07, and NIH45-46 have a
glycine at position 54 and leave this cavity empty (Fig. S1). Phe43 cavity-filling mutations such as those found in many clade A/E HIV-1 strains, e.g., His375, may interfere with a bulky side chain at this position, as is seen for many small-molecule CD4-mimetic compounds that rely heavily on the Phe43 cavity. Having Gly at position 54 may therefore contribute to increased breadth at the expense of potency, although for CD4bs Abs, this explanation may not be as straightforward since His375, at least in the case of N6, does not prevent binding and neutralization (9).

**FIG 4** Structural comparison of binding modes between N49P6 and other broadly neutralizing CD4-binding-site antibodies. (A) Pie charts showing the buried surface area (BSA) contributions to gp120 binding. The outer domain loops: D, CD4 binding, and V5 are colored as shown. The other outer domain and layers 2 and 3 of inner domain are also colored as shown. (B) Comparison of interactions between layers 2 and 3 of the gp120 inner domain and CDR H3 of N49P6 with other bnAbs (N49P7, VRC01, VRC03, VRC07, NIH45-46, and N6). The molecular surface is displayed over the gp120core, and the CDRs are shown as a ribbon diagram, with coloring as described in the legend of Fig. 2. Layers 2 and 3 of the gp120 inner domain are shown in darker and lighter shades of gray, respectively. Layer 3-contacting residues are labeled on the surface of gp120. (C) BSA contributions to layer 2 and 3 binding for a range of broadly neutralizing CD4bs antibodies (N49P6, N49P7, VRC01, VRC03, VRC07, NIH45-46, and N6). Residues of layers 2 and 3 are color-coded based on their sequence conservation as described in the legend of Fig. 3. BSA values for N49P6, N49P7, VRC01, VRC03, VRC07, NIH45-46, and N6 contact residues are shown. (D) Contact residues of gp120 mapped onto the gp12093TH057core sequences. Contact residues are defined by a 5-Å cutoff and marked above the sequence with 1 for the side chain and 2 for the main chain to indicate the type of contact. Buried surface residues were determined by PISA and are shaded blue.
N49P6 and N49P7 are unique in their use of long 20- and 19-aa CDR H3s, respectively. These are the longest found among VRC01 class Abs; the next longest, found in VRC07 and NIH45-46, have 16-aa CDR H3s (Fig. S1). As shown in Fig. 4, the elongated CDR H3s allow N49P6 and N49P7 to increase their binding footprint to include the gp120 inner domain, specifically gp120 inner domain layer 3. N49P6 and N49P7 have the largest BSA contribution with gp120 inner domain layer 3 to the total BSA of the complex among all VRC01 class Abs (113 and 157 Å² for the N49P6 and N49P7 complexes, respectively, compared to a range of 34 to 99 Å² for the other CD4bs bnAb complexes) (Fig. 4A). These include contacts with conserved and highly conserved residues at positions 474, 476, and 480 of gp120 layer 3 mediated by Ser100, Lys/Arg100B, and Arg100D of the extended CDR H3 of N49P6/P7 (Fig. 4B and C); VRC07, NIH45-46, and N6 but not VRC01 and VRC03, reach residues in gp120 inner domain layer 3 through CDR H3 contacts mediated by Ala100/Ala99A and Arg100A/Arg99B of VRC07/NIH45-46 and Tyr98 of N6. VRC07 and NIH45-46 also rely on gp120 inner domain layer 2 contacts in an area also recognized by N49P6 and N49P7 but with noticeably lower BSAs. In contrast, the more typical VRC01-like bnAbs, including VRC01, VRC03, and, to a lesser extent, N6, use their CDR H3 to almost contact loop D residues exclusively, with limited contact with the layers in the gp120 inner domain (Fig. 4). The use of the conserved gp120 inner domain residues may be an important component contributing to the breadth of Abs targeting the CD4bs. It is worth noting that VRC07 and NIH45-46 are among the broader and more potent bnAbs in the VRC01-like class. N49P6 and N49P7 neutralize 98/117 (83.8%) and 115/117 (98.3%) members of the 117-pseudovirus panel with IC80 values of 50 µg/ml and 86/117 (73.5%) and 102/117 (87.9%) members with IC80 values of <50 µg/ml, respectively. VRC07 and NIH45-46 neutralize 106/116 (91.4%) and 94/110 (85.5%) members of the panel with IC80 values of <50 µg/ml and 102/116 (87.9%) and 88/115 (76.5%) members with IC80 values of <10 µg/ml. To put things into perspective, the MPER bnAb 10E8 neutralizes 88/114 (77.2%) members of the panel with IC80 values of <10 µg/ml.

N49P6 utilizes interprotomer contacts for binding to the HIV-1 trimer. To further investigate the interaction of N49P6 with HIV-1 Env antigen, we determined the crystal structure of N49P6 Fab in complex with BG505 SOSIP.664 at a 4.05-Å resolution (Table 1). Figure 5A shows the structure of the full complex generated by crystallographic symmetry, which consists of three gp120s, three gp41s, and three N49P6 Fab s. Each CD4bs of the trimer is occupied by an N49P6 Fab, and the primary contact for the Fab is the CD4bs, which largely mirrors the interaction of the Fab and gp120 in the monomer complex structure (Fig. 5B). The total CD4-binding loop BSAs are 188 Å² and 184 Å² for gp120 and trimer complexes, respectively; the loop D BSAs are 297 Å² and 310 Å²; and the loop V5 BSAs are 180 Å² and 240 Å² (Table S1). Differences in BSAs for the CD4bs between the monomer and trimer structures can largely be attributed to differences in sequence between clade A/E 93TH057 and clade A BG505 gp120 and differences in conformation due to deletions in the V1/V2 and V3 loops in the gp120 core structure, which can be seen in the BSA plot versus residue positions (Fig. 5C). The difference in sequence in loop V5 (residues 460 and 461) and layer 3 (residues 474 and 476) influences the BSAs for these residues and also likely neighboring residues, i.e., Gly459 and Gly473. A major conformational change at Gln428 represents the largest single-residue difference in BSAs between the two structures. Gln428 sits on the β20-β21 turn on the outer domain half of the bridging sheet, which is formed in the monomer but not the trimer complex. The inner domain half of the bridging sheet, the V1V2 loop, extends to the top of the trimer, which disrupts the bridging sheet in the trimer, while a deletion in the V1V2 loop facilitates its formation in the monomer. The better packing of N49P6 around Gln428 in the trimer in the absence of the bridging sheet is likely also an explanation for the slightly higher BSA for the trimer residue E102 than for the monomer due to their proximity in the two structures. Conversely, the higher BSA for the monomer in loop D residues around Asn276 may reflect differences in the glycans between the two structures. The gp120 complex was made
with HEK 293 GnT1-grown protein with all but the initial N-acetylglucosamine removed by Endoglycosidase H (EndoH) (New England BioLabs). The glycan on Asn276 in the trimer structure is present but disordered past the initial N-acetylglucosamine; the trimer complex was also made from GnT1-grown protein, but the
high-mannose N-glycans were left intact in the protein used for crystallization. The truncated glycan may permit N49P6 to have a more stable light chain interaction in the monomer than in the trimer, although the total light chain contributions to the BSA are largely identical between the two, 245 Å² for the monomer and 258 Å² for the trimer; the light chain’s contribution to binding is almost exclusively to the glycan at Asn276 and nearby residues in both structures.

Although the primary CD4bs for both complexes are similar, contacts with the adjacent protomer increase the BSA of the complex formed by each N49P6 Fab with the trimer to make it significantly larger (Table S1). These interprotomer interactions are identical for each Fab due to crystal symmetry and are made mostly by the framework region immediately preceding CDR H1 and one CDR H1 residue (Gly26, Tyr27, Asp28, and Tyr32). CDR H3 adds two arginines to the interface (Arg96 and Arg100C), including the one (Arg100C) inserted in the CDR H3 relative to N49P7 (Fig. 5A and B). Contacts within the adjacent promoter map to mobile layer 1 on the gp120 inner domain and include Tyr61, Glu64, Lys65, and His66 with Fab-gp120 hydrogen bonds (the carbonyl oxygen of Gly26 [FWR1] and the side chain of Lys65, the side chains of Asp28 [FWR1] and His66, the side chains of Tyr27 [CDR H1] and Glu64, the side chains of Arg96 [CDR H3] and Glu64, and the side chains of Arg100C [CDR H3] and Tyr61) and hydrophobic van der Waals interactions (formed between the side chains of Tyr27 [FWR1] and Lys65). The total BSA of the interface is 429 Å², 207 Å² from the trimer and 222 Å² from the heavy chain of N49P6, which adds approximately 15% more BSA to the trimer and 17% more BSA to the Fab (Fig. S1).

N49P6 mimics CD4 in its initial quaternary interaction with BG505 SOSIP.664. It has been shown previously that the first CD4 that binds a BG505 SOSIP trimer has both a primary and a secondary binding site (23). The primary site is essentially identical to that used in CD4 structures with monomeric gp120 (24). The secondary CD4bs resides on the adjacent gp120 within inner domain layer 1 (23). Interestingly, a structural alignment of the complexes formed between the BG505 SOSIP.664 trimer and three N49P6 Fabs or a single CD4 molecule (Protein Data Bank [PDB] accession number 5U1F) (23) reveals that the layer 1 gp120 residues used by N49P6 to contact the adjacent protomer are largely identical to the ones that CD4 uses in its initial quaternary interaction with the trimer (Fig. 6A through C). Although the resolution of the monovalent CD4-BG505 SOSIP.664 trimer complex was low, which introduces some ambiguity into the details of the interface, the CD4 contacts with the adjacent protomer were mapped to residues 62 to 66 of layer 1 and within the coreceptor-binding site, Lys207. (Fig. 6B) (23). Interestingly, N49P6’s interprotomer contacts involve three (Glu64, Lys65, and His66) of five residues utilized by CD4 within layer 1. Of note, residues Glu64, His66, and Lys207 are almost universally conserved across HIV-1 isolates, and mutational studies confirm their importance in proper spike function and viral infectivity (23). N49P6 capitalizes on the functional importance and the sequence conservation of these residues to increase its affinity and breadth.

It has been shown previously that the one CD4-bound BG505 SOSIP trimer transitions very quickly to a more open conformation with three CD4s bound that fully exposes the coreceptor-binding site (25). In this state, the contacts of CD4 with the adjacent protomer were lost (26) (Fig. 7). The binding of coreceptor or coreceptor-binding-site antibody stabilizes this open conformation with a few additional changes (26, 27). Charge inversion within the quaternary CD4-binding site in HIV-1 Env of His66 to Glu or Glu62 to Lys and Glu64 to Lys abrogates viral infectivity, highlighting the importance of this region (23). It is interesting to note that N49P6 interacts with these residues much like a charge-inverted virus to CD4, with Asp28 (N49P6) forming a hydrogen bond to His66 of the adjacent gp120 in the trimer and Arg96 (N49P6) forming a hydrogen bond to Glu64 (gp120) (Fig. 6C). The inferred binding based on distance in the CD4 complex is between Lys21 (or Lys22) (CD4) and His66 (gp120) and between Gin20 (CD4) and Glu64 (gp120), which would seem less stabilizing. This may reflect CD4’s role in infection, which is to drive the transition to an open conformation and expose the coreceptor-binding
FIG 6 Molecular details of the interprotomer contacts of N49P6. (A) Structural alignment of the N49P6 Fab-BG505 SOSIP.664 trimer complex and the BG505 SOSIP.664 trimer with one CD4 molecule bound (PDB accession number 5U1F). N49P6 Fab (orange) and CD4 (green) are shown as a ribbon diagram. The zoomed-in views depict the contact between N49P6 Fab or CD4 and the adjacent BG505 protomer. All contacting residues are shown as sticks. (B) Networks of interactions formed between N49P6 Fab/CD4 and the adjacent gp120 BG505 protomer as defined by either a 5-Å distance criterion cutoff for N49P6 or a 7.5-Å distance criterion cutoff for CD4 are shown as solid lines. Layer 1 residues are color-coded based on their sequence conservation as described in the legend of Fig. 3. (C) Binding footprints of N49P6 and CD4 on the adjacent gp120 BG505 protomer (surface representation) are colored in yellow. Contacting residues of N49P6 and CD4 are shown in a stick representation (framework region [FWR] in orange, CDR H1 in black, CDR H3 in red, and CD4 in green). (D) Binding footprints of other CD4bs Fabs (VRC01 [PDB accession number 5FYJ], VRC03 [PDB accession number 6CDI], 3BNC117, 1-18, CH31, NIH 45-46) on the adjacent gp120 BG505 protomer are shown in yellow. Contacting residues of the Fabs are shown in a stick representation (FWR in orange, CDRs in black, and CD4 in green). (Continued on next page)
site. In contrast, N49P6’s role in neutralization is to bind and lock the trimer in a closed state.

The N49P6 quaternary interaction is different from the quaternary interactions of other potent CD4bs Abs. N49P6 is not the only CD4bs bnAb to extend its footprint onto the adjacent protomer. Many of the broader and more potent CD4bs bnAbs use different interprotomer contact strategies to lock the trimer in a more closed conformation; some target elements of the gp120 inner domain, while others target the coreceptor-binding site at the base of the V3 loop. Figure 6C and D show the interprotomer contact details of many of these bnAbs compared to N49P6 and CD4. The most like N49P6 is the prototypic VRC01 class Ab VRC01, which binds to inner domain layer 1 of the adjacent gp120 with a total BSA of 102 Å², compared to 206 Å² for N49P6, 199 Å² from layer 1 and 7 Å² from layer 2 (2, 28). VRC03, another VRC01 class Ab from the same donor, uses heavy chain framework residues and a unique insertion in the heavy chain framework region to contact gp120 inner domain layer 1, 42 Å², but relies more heavily on inner domain layer 2, 90 Å², and the base of the V3 loop, 47 Å², for a total gp120 BSA of 179 Å² (29, 30). NIH45-46 also contacts inner domain layer 1 on the adjacent trimer, 38 Å², but without the heavy chain framework insertion contacts only layer 1. 3BNC117 contacts the adjacent gp120 inner domain layer 1, 40 Å², and the base of the V3 loop, 125 Å², for a total of 165 Å² (31). Similarly, CH31 uses both heavy chain framework and CDR H1 residues to bind the adjacent gp120 inner domain layer 1, 100 Å²; layer 2, 140 Å²; and the base of the V3 loop, 92 Å²; for a total of 332 Å² (29, 30). And finally, 1-18 uses its heavy chain framework and CDR H1 residues to contact inner domain layer 2, 104 Å², and the base of the V3 loop, 175 Å², for a total BSA of 279 Å² (Table S2) (10).

An important consideration when comparing quaternary contacts in the trimer is the degree to which the trimer “opens” upon Ab binding. The HIV-1 BG505 SOSIP trimer goes from a “closed” trimer with distances between protomers as measured by the α-carbon of residue 375 at the base of the Pheα cavity of 54.5 Å to an average of 69.6 Å in the fully open trimer with three CD4s bound (Fig. 7). The degree to which a trimer opens determines the degree to which quaternary contacts are possible. It is worth noting that all the CD4bs bnAbs that make contacts with the adjacent gp120 have BG505 SOSIP trimers in a more closed conformation, more closed than even the BG505 SOSIP trimer with only one CD4 bound, which has an average distance of 57.4 Å. The average distance between protomers ranges from 54.5 to 55.9 Å, with N49P6 having an average interprotomer distance identical to that of the unbound BG505 SOSIP trimer, 54.5 Å. Some BG505 SOSIP trimers are asymmetric, like NIH45-46, which has quaternary contacts for only two of the three possible adjacent protomers in the trimer, but most are largely symmetric. NIH45-46 also has the smallest adjacent protomer footprint, which may explain its inability to lock its BG505 SOSIP trimer in a closed conformation.

N49P6’s total adjacent protomer footprint is larger than those of VRC01, VRC03, NIH45-46, and 3BNC117 but smaller than those of 1-18 and CH31. However, N49P6 has the largest BSA for the inner domain layer 1 region that overlaps CD4’s interprotomer footprint, twice the layer 1 footprints of VRC01 and CH31 and roughly four times those of VRC03 and NIH45-46. VRC03, 3BNC117, 1-18, and CH31 focus their epitopes more toward inner domain layer 2 and the base of the V3 loop of the adjacent trimer. This difference in strategy may ultimately determine neutralization breadth since breadth is a function of residue conservation as well as affinity.
DISCUSSION

The primary CD4-binding site (CD4bs), the major contact interface between the HIV-1 Env spike and host receptor CD4, can be broken down into three interacting regions, the CD4-binding loop, loop D, and loop V5 (32). Host receptor CD4 makes a number of important contacts with each of these regions that are highly conserved across HIV-1 strains (24). Broadly neutralizing CD4bs antibodies utilize many, if not all,
of these residues to achieve their neutralization breadth (6, 29, 32, 33). Increasing neutralization potency, on the other hand, also requires the use of residues outside the CD4bs for effective competition with CD4. Which residues and their conservation across HIV-1 strains are often limiting factors in an antibody’s neutralization breadth, while the number and strength of the interactions outside the CD4bs are limiting factors in determining their neutralization potency.

HIV-1 has developed several strategies to prevent antibodies from binding the CD4bs and to pivot away from those that do bind to prevent neutralization. The CD4bs is recessed and shielded by heavy glycosylation on Env, which prevents all but a few antibodies from binding (34, 35). Broadly neutralizing antibodies that bind within the CD4bs have had to develop strategies to overcome this hurdle. It has been shown for the prototypical VRC01 class of CD4bs-specific bnAbs that changes, alone or in combination, in the lengths of gp120 loops D and V5 and the glycosylation pattern in loop D constitute major viral escape mechanisms through steric clashes with antibody CDR H2, the N terminus of the light chain, and/or antibody CDR L3 (36–38). VRC01-like antibodies adopt a short 5-amino-acid (aa)-long CDR L3 and often deletions in CDR L1 to accommodate changes in these regions (6, 29, 33). N49P6 together with N49P7 constitute a new branch of the VRC01-like class of CD4bs-specific bnAbs that, in addition to the strategy of shortening CDR L1 and L3, also utilize the light chain’s rotation/tilt to provide additional space and prevent steric clashes with loops D and/or V5. The latter is possible due to a mutation in light chain framework region 2 (FWR2), Tyr36 to Cys, which allows greater mobility of the heavy and light chains relative to one other. N49P6 also contains a deletion in its CDR L1 (8 aa) in addition to the extremely short CDR L3 (5 aa) found in VRC01-like bnAbs. The combination results in a minimal light chain footprint, which can be seen by the low buried surface area (BSA), 245 Å² (157 Å² without the contribution from the Asn276 glycan), compared to those of most other Abs in its class (see Table S2 in the supplemental material). In addition, N49P6’s CDR H2 also contains a Pro60ArgPro62 motif, which preferentially adopts a left-handed polyproline helical conformation (with phi and psi angles for Pro60 of −59° and 145°, Arg61 of −47° and −44°, and Pro62 of −58° and −11°, respectively) that points away from loop V5, further minimizing steric clashes with V5. Although N49P6 utilizes Asn276 and its attached glycan in its epitope, this interaction seems to be mainly limited to the initial N-acetylglucosamine. This is in contrast to VRC01 and many other VRC01-like bnAbs, which utilize more of the glycan in their epitope (39). A change in the glycan position from position 276 to position 279 in gp120, which keeps steric pressure on the CD4bs but removes many of the beneficial glycan interactions, results in neutralization resistance to many of these other CD4bs bnAbs. This change may be less of a problem for N49P6 due to its limited dependence on the Asn276 glycan.

The glycan at Asn276 prevents the binding of all but a few germ line antibody alleles to the CD4bs. Those that do either bind by utilizing a long CDR H3 to bypass light chain-mediated steric clashes or bind with one of two germ line alleles, V\textsubscript{\textgamma}1-2 (VRC01 class) or V\textsubscript{\textgamma}1-46 (8ANC131 class), in combination with light chains with deletions in CDR L1 and L3 (6, 29, 33). N49P6, as a member of the VRC01 class, utilizes V\textsubscript{\textgamma}1-2 with a relatively long CDR H2 and CDR H3, composed of 17 and 20 aa, respectively. The V\textsubscript{\textgamma}1-2 germ line gene has several characteristics of its CDR H2 that permit it to mimic CD4. For example, both the typical V\textsubscript{\textgamma}1-2 CDR H2 and CD4 form a short antiparallel β-sheet with residues in the gp120 CD4-binding loop. In addition, V\textsubscript{\textgamma}1-2 germ line-derived bnAbs also often mimic the salt bridge between Arg59 (CD4) and Asp368 (gp120) and utilize the hydrophobic cavity that binds Phe63 in CD4. The Phe63-binding cavity constitutes an important anchoring point for CD4bs-specific antibodies and small-molecule CD4-mimetic compounds that rely on hydrophobic contacts with residues lining the cavity (40). N49P6 preserves many of these features along with the Arg-Asp salt bridge mediated by heavy chain framework residue Arg71; however, N49P6, like N49P7, has a glycine at heavy chain position 54, which leaves the Phe63 cavity empty.

The Phe63 cavity is a key feature used by many small-molecule CD4-mimetic compounds to modulate the Env conformation and in many cases to drive Env to the CD4-bound, open conformation (40–42). To compensate for its loss, N49P6 establishes an
extensive network of interactions with loop D, loop V5, and, most importantly, the highly conserved residues within the gp120 inner domain that include inner domain layers 2 and 3. In addition, N49P6 also contacts inner domain layer 1 on the adjacent gp120 in the trimer, largely mimicking the interprotomer contact of host receptor CD4. The inner domain residues involved in the N49P6 epitope are highly conserved among HIV-1 clades, and N49P6 takes advantage of this. The mobile layers of the gp120 inner domain are involved in structural rearrangements of the HIV-1 trimer after CD4 attachment and coreceptor binding. They form the gp41-interacting face of gp120, and their conformational change serves as a driving force in trimer disassembly and cellular fusion. Their mutation comes at a fitness cost to the virus, which enables N49P6 breadth.

Finally, one of the most interesting features of how N49P6 engages HIV-1 Env is in its contact with the adjacent protomer. The mode of binding to the trimer and the scope of interprotomer contact residues largely resemble those of the interaction of host receptor CD4 in its initial binding to the HIV-1 Env trimer. Structural alignment of the N49P6-BG505 SOSIP trimer complex to the complex of the BG505 SOSIP trimer with one CD4 molecule bound (23) reveals a large overlap of the Fab/CD4 contact surface on the adjacent protomer. Other CD4bs-specific bnAbs use the adjacent protomer to increase their epitope footprint but tend to focus instead on the coreceptor-binding site at the base of the V3 loop and less on layer 1 at the site used by CD4. In this regard, N49P6 is unique in sharing many of the same contact residues used by CD4 in its initial binding to the trimer. Interestingly, it is known that following the initial contacts of one CD4 molecule to the HIV-1 BG505 SOSIP trimer, the engagement of a second and a third quickly follows, which leads to the full opening of the trimer (26). This is in contrast to the case of N49P6, where the BG505 SOSIP trimer is "locked" in the closed conformation with interprotomer distances almost identical to those of the unbound trimer.

Neutralization of the global panel of HIV-1 pseudoviruses by N49P6 is broad and potent but still leaves room for improvement, especially when IC80 values are used for comparison. Are there any clues from the more resistant sequences that could suggest other means to improve neutralization? Sequences from the panel that are better neutralized by N49P7 are largely conserved across N49P6 contact residues, suggesting that resistance is more subtle than those due to single point mutations (Fig. S2). Indeed, the generally high degree of conservation among epitope residues may imply that there is a large fitness cost for their mutation. Instead, one common theme among more resistant sequences in the panel and the sequences from donor N49, which are resistant to N49P6/7, is a larger V5 loop. Although most of the N49P6 epitope residues in the V5 loop reside at its base, insertions in this region may result in steric hindrance or conformational rearrangement that may lower Ab affinity. Examination of viral sequences from donor N49 shows exceptionally long V5 loops with the possible addition of a glycan, which may be a mechanism of resistance. The only other obvious difference in N49 donor sequences is the exceptionally long V1 region in the V1V2 loop (Fig. S2). To compensate, N49P7 has increased its binding to the trimer within the primary CD4-binding site. N49P6 has opted instead to increase its binding to the secondary CD4-binding site on the adjacent protomer in the trimer. Both approaches seem to have merit when applied to the global pseudovirus panel.

In conclusion, the N49 lineage of near-pan-neutralizing antibodies deconvoluted from the plasma of an HIV-1-infected individual represents a new class of CD4-binding-site-specific antibodies that we refer to as the N49P series. These antibodies omit the Phe43 cavity and rely strongly on the gp120 inner domain for binding to the primary gp120 protomer, and they mimic the initial contact of host receptor CD4 with the adjacent promoter of the HIV-1 Env trimer. When bound to the HIV Env trimer, N49P series antibodies lock the trimer in the closed conformation using residues from the initial contact of the CD4 receptor with the adjacent protomer in the trimer.

MATERIALS AND METHODS

Protein production and purification. N49P6 Fab was produced by papain digestion of N49P6 IgG as described previously (43, 44). BG505 SOSIP.664 was a kind gift from John Moore (Weill Cornell...
Medical College, New York, NY). Complexes were made with an excess of N49P6 Fab at ratios of 1:2:1 for the gp120 complex and 3.5:1 for the SOSIP complex and then purified by gel filtration chromatography on a Superdex 200 16/60 column (GE Healthcare) equilibrated with a solution containing 10 mM Tris-HCl (pH 7.2) and 100 mM ammonium acetate. Purified complexes were concentrated to approximately 5 to 10 mg/ml for use in crystallization trials.

**Neutralization assay.** An HIV-1 neutralization assay was performed by measuring the reduction in luciferase expression following a single round of virus infection in TZM-bl cells as previously described (20). Briefly, mAbs were tested against murine leukemia virus (MuLV) as a negative control and a panel of pseudoviruses. Threefold serial dilutions of mAbs were tested in duplicate in 10% Dulbecco’s modified Eagle growth medium (DMEM) (100 μl/well). A total of 200 50% tissue culture infective doses (TCID₅₀) of pseudoviruses (50 μl) were added to each well, and the plates were incubated for 1 h at 37°C. TZM-bl cells (1 x 10⁵ cells/well in 100 μl) were then added in 10% DMEM in the presence of DEAE-dextran (Sigma, St. Louis, MO) at a final concentration of 11 μg/ml. Each well contained a final volume of 250 μl. Assay controls included replicate wells of TZM-bl cells alone (cell control), TZM-bl cells with virus (virus control), and the MuLV control. After a 48-h incubation at 37°C, 150 μl of assay medium was removed from each well, and 100 μl of Bright-Glo luciferase reagent (Promega, Madison, WI) was added to each well. The cells were allowed to lyse for 2 min, the cell lysate (150 μl) was then transferred to a 96-well black solid plate, and the luminescence intensity was measured using a Victor 3 luminometer (PerkinElmer, Waltham, MA).

**Crystallization.** Initial crystal screens were carried out with commercially available sparse matrix crystallization screens from Molecular Dimensions (Proplex and MacroSol) using the hanging-drop vapor diffusion method with drops of 0.5 μl protein and 0.5 μl precipitant solution. Conditions that produced crystals were optimized to produce crystals suitable for data collection. N49P6 Fab-gp120_TOTALcore complex crystals were grown from a solution containing 15% polyethylene glycol 3350 (PEG 3350) and 0.1 M Tris-HCl (pH 8.0). Each well contained a final volume of 250 μl. Assay controls included replicate wells of TZM-bl cells alone (cell control), TZM-bl cells with virus (virus control), and the MuLV control. After a 48-h incubation at 37°C, 150 μl of assay medium was removed from each well, and 100 μl of Bright-Glo luciferase reagent (Promega, Madison, WI) was added to each well. The cells were allowed to lyse for 2 min, the cell lysate (150 μl) was then transferred to a 96-well black solid plate, and the luminescence intensity was measured using a Victor 3 luminometer (PerkinElmer, Waltham, MA).

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**Data collection and structure solution and refinement.** Diffraction data for both N49P6 complexes were collected at the Stanford Synchrotron Radiation Light Source (SSRL) at beamline BL12-2 equipped with a Dectris Pilatus area detector. N49P6 Fab-gp120_TOTALcore crystals belong to orthorhombic space group P2₁ ₂₁ ₂ with unit cell parameters of a = 65.4, b = 80.8, and c = 195.2 Å and one N49P6 Fab-gp120_TOTALcore complex present in the asymmetric unit (ASU). N49P6 Fab-BG505 SOSIP.664 complex crystals belong to cubic space group P₂₁ 3 with unit cell parameters of a = b = c = 164.7 Å and one-third of the N49P6 Fab-BG505 SOSIP.664 trimer present in the ASU. Data were processed and reduced with HKL2000 (45). The N49P6 Fab-gp120_TOTALcore complex structure was solved by molecular replacement with Phaser (46) from the CCP4 suite (47) based on the coordinates of the N49P6 Fab-gp120_TOTALcore complex (PDB accession number 6BCK). The N49P6 Fab-BG505 SOSIP.664 complex was solved using coordinates of gp120 and gp41 from BG505 SOSIP.664 in complex with PGT122 and 35O22 Fabs (PDB accession number 4TVP) and the refined N49P6 Fab coordinates from the N49P6 gp120 complex. Refinement was done with Refmac (48) and/or Phenix (49). Refinement was coupled with manual rebuilding and in COOT (50). The N49P6 Fab-gp120_TOTALcore complex diffraction to 2.55 Å and was refined to an R-factor of 0.219 and an R̄₉₀ of 0.273. The N49P6 Fab-BG505 SOSIP.664 complex diffraction to 4.05 Å and was refined to an R-factor of 0.255 and an R̄₉₀ of 0.315. Data collection/refinement statistics are shown in Table 1.

**Structure validation and analysis.** The quality of the final refined model was monitored using the program MolProbity (51). Structural alignments were performed using the program lsqkab from the CCP4 suite. The PISA Web server (52) was used to determine contact surfaces and interface residues. All illustrations were prepared with the PyMOL Molecular Graphics suite (http://pymol.org) (DeLano Scientific, San Carlos, CA, USA).

**Availability of data.** The data sets generated and/or analyzed during the current study are available in the Protein Data Bank (PDB) repository under accession numbers 6OZ2 and 6OZ4.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 1.5 MB.

**FIG S2**, TIF file, 1.1 MB.

**TABLE S1**, DOCX file, 0.02 MB.

**TABLE S2**, DOCX file, 0.02 MB.

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We declare that we have no competing interests.

W.D.T., D.N.N., and M.P. designed, performed research, and analyzed the data; Z.R.T. and M.M.S. isolated N49P6 antibody and performed neutralization assays; W.D.T., D.N.N., and M.P. wrote the manuscript; and all authors provided comments or revisions.
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