Structural basis of coreceptor recognition by HIV–1 envelope spike

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HIV–1 envelope glycoprotein (Env), which consists of trimeric (gp160)3 cleaved to (gp120 and gp41)3, interacts with the primary receptor CD4 and a coreceptor (such as chemokine receptor CCR5) to fuse viral and target–cell membranes. The gp120–coreceptor interaction has previously been proposed as the most crucial trigger for unleashing the fusogenic potential of gp41. Here we report a cryo–electron microscopy structure of a full–length gp120 in complex with soluble CD4 and unmodified human CCR5, at 3.9 Å resolution. The V3 loop of gp120 inserts into the chemokine–binding pocket formed by seven transmembrane helices of CCR5, and the N terminus of CCR5 contacts the CD4–induced bridging sheet of gp120. CCR5 induces no obvious allosteric changes in gp120 that can propagate to gp41; it does bring the Env trimer close to the target membrane. The N terminus of gp120, which is gripped by gp41 in the pre–fusion or CD4–bound Env, flips back in the CCR5–bound conformation and may irreversibly destabilize gp41 to initiate fusion. The coreceptor probably functions by stabilizing and anchoring the CD4–induced conformation of Env near the cell membrane. These results advance our understanding of HIV–1 entry into host cells and may guide the development of vaccines and therapeutic agents.
The atomic structures of four-domain CD4 and the gp120 core in complex with CD4 were excluded.

Overall structure of the CD4–gp120–CCR5 complex

The absence of any G proteins in the purified complex is consistent with the notion that gp120, unlike chemokines, does not require G-protein mediated signalling pathways in these cells (Extended Data Fig. 3a, b). To purify the wild-type CCR5 in its Env-bound conformation, we isolated the CD4–gp120–CCR5 complex from the CCR5-expressing cells (Extended Data Fig. 3a, b). The purified CD4–gp120–CCR5 complex eluted from a size-exclusion column as a single sharp peak at the expected volume, which confirmed its stability and conformational homogeneity (Extended Data Fig. 3c). Coomassie-stained sodium dodecyl sulfate–polyacrylamide gel electrophoresis and negative-stain electron microscopy of the peak fractions showed that all three components were in a stoichiometry of 1:1:1 (Extended Data Figs. 3c, 4a–c). The absence of any G proteins in the purified complex is consistent with the notion that gp120, unlike chemokines, does not require G-protein coupling for high-affinity binding to CCR5.

We acquired cryo-EM images on a Titan Krios electron microscope with a K2 Summit direct detector (Extended Data Fig. 4d) and used RELION for image processing. Two-dimensional (2D) class averages of the particle images showed secondary structural features for both gp120 and CCR5 (Extended Data Fig. 4e). Three-dimensional (3D) classification of the particles was performed (Extended Data Fig. 5a), using the low-pass-filtered 3D reconstruction of the complex in negative stain as an initial model (Extended Data Fig. 4c). The particles from the classes with defined structural features were combined and refined to generate a map at 4.6 Å resolution, showing all three components (Extended Data Fig. 5a). The structure was determined by rounds of 3D classification, 3D classification with signal subtraction and masked refinement, as described in Methods and Extended Data Fig. 5a–d. The final resolution was 3.9 Å, when the last two domains of CD4 were excluded.

Overall structure of the CD4–gp120–CCR5 complex

The atomic structures of four-domain CD4, the gp120 core in complex with CD4, and the modified CCR5—none of which was used in image processing—all fit as rigid bodies notably well to the electron microscopy density of the CD4–gp120–CCR5 at a 4.5 Å resolution (Fig. 1a, b and Extended Data Fig. 5b). The gp120 core, which contains inner and outer domains and the bridging sheet, occupied excellent density—this indicates that this region of gp120 has the same rigid structure in the ternary complex as it does in the CD4-induced conformation. Several N-linked glycans were marked by protruding densities from the protein surfaces, as expected for a fully glycosylated gp120. All four domains of the soluble CD4 were visible, with density for first two (D1–D2) slightly stronger than for last two (D3–D4) domains. There was also density for an N-linked glycan at Asn271 in the palmitoylated cytoplasmic tail of CCR5, which presumably interacts with membrane, was disordered in this detergent-solubilized complex. Extra densities between CCR5 and gp120 that cannot be explained by the existing structures define the details of the interaction between CCR5 and gp120.

Interfaces between gp120 and CCR5

The map from the masked refinement showed good density for the gp120 and CCR5 interfaces (Fig. 1c and Extended Data Fig. 6). As previously postulated, there are two major contacting interfaces between gp120 and CCR5 (Fig. 1d). The V3 loop of gp120 inserts into the CRS2 of CCR5 and makes contact with all the helices of the seven transmembrane helices. The map from the masked refinement showed good density for first two (D1–D2) slightly stronger than for last two (D3–D4) domains. There was also density for an N-linked glycan at Asn271 in the palmitoylated cytoplasmic tail of CCR5, which presumably interacts with membrane, was disordered in this detergent-solubilized complex. Extra densities between CCR5 and gp120 that cannot be explained by the existing structures define the details of the interaction between CCR5 and gp120.

V3 loop of gp120 and CRS2 of CCR5

The conserved 310-Gly-Pro-Gly-Arg(Gln)-313 motif at the tip of the V3 loop penetrates by approximately one third of the thickness of the lipid bilayer into the CRS2 pocket; the Pro311 residue reaches most of the V3 loop.

Fig. 1 | Cryo-EM structure of the CD4–gp120–CCR5 complex.

a, Cryo-EM map of the complex that contains HIV-1 gp120 (cyan), CCR5 (red), four-domain CD4 (green; D1–D4, domain 1–domain 4) and detergent micelle (grey).

b, Fit of structures of gp120 (RCSB Protein Data Bank code (PDB ID): 5VN3), CCR5 (PDB ID: SU1W) and four-domain CD4 (PDB ID: 1WIO) into the electron microscopy map shown in a. N271 of CD4 (green), N234, N262 and N362 of gp120 in cyan are N-linked glycosylation sites. c, The structure of the CD4–gp120–CCR5 complex was modelled on the basis of a 3.9 Å density map. d, Overall structure of the four-domain CD4–gp120–CCR5 complex shown in ribbon diagram. N, N terminus; C, C terminus; ECL2, extracellular loop 2; I, II, III, IV, V, VI, VII, transmembrane helices 1–7.

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The gp120 bridging sheet and CCR5 N terminus

The second major interface between Env and its co-receptor is formed by the N terminus of CCR5 and the bridging sheet of gp120. Helices I and VII of CCR5 are locked by a disulfide (Cys20–Cys269) of CCR5 in complex with gp120. Helices I and VII of CCR5 are locked by a disulfide (Cys20–Cys269); the transmembrane helices of CCR5 is limited, at least in the liganded state. The gp120 CCR5-binding site to gp41, as previously hypothesized. Whereas CD4 and/or CD4i antibody-induced conformational changes in the SOSIP-CCR5 with stabilizing mutations bound to the antagonist [5P7]CCL5 also adopts a conformation very similar to that of the gp120-bound wild-type CCR5, which suggests that the conformational freedom of the transmembrane helices of CCR5 is limited, at least in the liganded inactive forms (Fig. 3c).

Differences between CD4- and CCR5-bound gp120

The most unexpected aspect of the CD4–gp120–CCR5 structure is the presence of obvious allosteric changes that can propagate from the CCR5-binding site to gp41, as previously hypothesized. Whereas CD4 and/or CD4i antibody-induced conformational changes in the SOSIP-based Env trimer have recently been described, a comparison of the CD4- and CCR5-bound gp120 with the CD4-bound gp120 shows no major differences in the core region of gp120 (Fig. 4a and Extended Data Fig. 8a). In particular, an approximately 50 Å zone between the CCR5-binding site and the gp120–gp41 interface (including the inner...
and outer domains and the bridging sheet) remains almost invariant, which suggests that CCR5 does not induce any major structural changes that affect gp41. There are, however, some differences between the two structures. First, when CCR5 binds the V3 loop reconfigures to fit into the CR2S pocket (Fig. 3a). This conformation, which partially mimics that of the N terminus of [5P]CCL5, has not been seen for either unliganded or antibody-bound V3 loops. Second, a more substantial difference is seen in the region that includes the N and C termini of gp120, near its interface with gp41. In the pre-fusion SOSIP trimmer structure, the gp120 termini are surrounded by the so-called ‘four-helix collar’ of gp41, which closes by insertion of the side chain of Met530—located on one of the helices (α6)—into a hydrophobic clasp formed by three tryptophan residues on two other helices (α8 and α9). CD4 binding induces a shift of α6, which enables the fusion peptide to pack directly against the termini of gp120. In the CD4–gp120–CCR5 complex, the N and C termini bend back at pivot regions that contain the highly conserved 40-Gly-Val-Pro-42 and 489-Pro-Leu-Gly-491 sequences, respectively. In particular, the N terminus rotates almost by about 180° to pack against the surface of gp120 and occupies the space of the fusion peptide in the Env trimer (Fig. 4b). There is no obvious density for the rest of the C terminus, which is probably disordered—although the histidine tag could influence its conformation.

### Model for how CCR5 functions as an HIV-1 coreceptor

Because CCR5 binding does not appear to induce any allosteric changes that can unleash gp41 to fuse membranes, it is intriguing how CCR5 might function as an essential coreceptor. On the basis of the known structures of the HIV-1 Env trimers, the pre-fusion gp41— which wraps around the N and C termini of gp120—is no longer stable and is likely to enter an irreversible refolding process once gp120 dissociates. Thus, gp120 dissociation may be the crucial trigger that initiates gp41 refolding events, including the insertion of the fusion peptide of gp41 into the target membrane and the formation of the post-fusion conformation. In the pre-fusion conformation, the N and C termini of gp120 are gripped by the four-helix collar of gp41. CD4 binding leads to a large shift of the C terminus of helix α6 away from the gp120 termini; this creates a pocket, which is filled by the fusion peptide that packs against the pivot region (40Gly-Val-Pro) of the gp120 N terminus (Fig. 4b). When intrinsic conformational dynamics cause the fusion peptide to dissociate from the pocket, this opens up one side of the gp41 grip, and the gp120 N terminus can then bend back to adopt the conformation observed in the CCR5-bound structure (Extended Data Fig. 9). The rearrangements of the termini of gp120—which are probably independent of CCR5 binding—can prevent the fusion peptide from reoccupying the pocket and effectively weaken gp120–gp41 interactions, which possibly leads to complete dissociation. Indeed, spontaneous or CD4-induced gp120 shedding from Env trimers are well-documented for many HIV-1 isolates, which indicates that gp120 is prone to dissociation from gp41 even in the absence of a coreceptor. We note that the effect of the membranes and Env trimer organization remains unknown and will require further investigation.

If the rearrangement of gp120 termini to activate gp41 does not depend on CCR5 binding, the question arises as to why a coreceptor would be needed at all. First, premature gp120 dissociation would—in absence of a coreceptor—be non-productive; for a virion attached to the target-cell surface only through an Env trimer–CD4 contact, the distance between the fusion peptide and membrane surface can be about 160 Å (Fig. 4b). If gp120 dissociates, the fusion peptide would be too far away to reach the target membrane. The binding of gp120 to CCR5 can bring the fusion peptide of gp41 to within 70 Å of the membrane surface (Fig. 4b), which is consistent with the distance needed for the fusion peptide to translocate and reach the target membrane. Second, the gp120–CD4 association—measured by single-molecule force spectroscopy with infectious virions and live host cells—is unstable, and rapidly reversible unless CCR5 binding follows immediately. CCR5 is therefore needed to stabilize the CD4-induced conformational changes, which are already competent for promoting fusion. In particular, the tucking away of the V3 loop by CCR5 would prevent the Env trimer from moving back to the pre-fusion conformation, and help shift the equilibrium towards the irreversible step—that is, gp41 refolding. Third, membrane fusion may require more than one Env trimer to induce fusion pore formation, as shown for other viral fusion proteins. Because the number of Env trimers on the virion is low—about 14 trimers per virion—a long lifetime for the Env–receptor complex would be important for recruiting additional CD4- and coreceptor-primed trimers. Thus, our structure shows how a coreceptor can be essential for membrane fusion, despite the fact that it does not actively induce gp41 refolding.

### Coreceptor switch

The switch from CCR5 to CXCR4 is often associated with an accelerated increase in viral load and decrease in CD4+ T cells, as well as with faster disease progression. Our structure supports a model for how simple mutations in Env can achieve this seemingly complicated transition. First, the coreceptor is required only to stabilize the CD4-induced conformation, and not to trigger additional allosteric changes in Env through specific interactions. The switch can thus be accomplished if the V3 loop gains sufficient affinity for CXCR4 and does not release the coreceptor, as no specific mutations in Env are needed to make it able to be ‘triggered’ by CXCR4. Second, the overall dimensions of the CR2S pocket in the gp120-bound CCR5 and the liganded CXCR4 are very similar, which suggests that changes of the surface-exposed residues in the V3 loop to make it compatible with CXCR4 binding would be sufficient. Use of both coreceptors by R5X4 isolates further underscores the similarities between the two coreceptors. Third, the main contacts between gp120 and the CCR5 N terminus are electrostatic. CXCR4 has seven acidic residues in its N terminus (before the first disulfide-forming Cys residue) and CCR5 has four, in addition to two or three sulfotyrosines. No additional mutations in the bridging sheet region would be needed if the extra acidic residues in CXCR4 can replace the two critical sulfated tyrosines in CCR5. Finally, X4 V3 loops generally have more positive charges than those of R5 viruses, consistent with a more negatively charged CR2S in CCR5 than CXCR4. Evolution from CCR5 to CXCR4 use can indeed be achieved by multiple mutational pathways, but often gain net positive charges in the V3 region.
Therapeutic agents based on CCR5 antagonists
The V3 binding-site only partially overlaps with the minor subpocket of the maraviroc binding site (Fig. 3a). Maraviroc-resistant viruses can emerge either in infected individuals under the treatment, or by in vitro selection. Major changes in the escaped viruses map to the V3 region—some of these changes have a three-residue deletion, but in general they show no consistent patterns. Some resistant viruses can infect cells by recognizing the drug-bound CCR5. In addition, replication-competent HIV-2 viruses have been selected that lack the entire V3 loop. These data indicate that no specific structural determinants in the V3 region are required for the coreceptor to function, which is fully consistent with our conclusion that CCR5 does not actively trigger gp41 through specific interactions with gp120.

Our structure also suggests a general strategy for how to improve maraviroc-like therapeutic agents. Because the V3 loop mainly overlaps with maraviroc in the minor subpocket (which is primarily occupied by the triazole group of the compound), additional groups may be added to the triazole ring to enhance its competing power with the V3 loop and increase the barrier to drug resistance.

Online content
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METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and investigators were not blinded to allocation during experiments and outcome assessment.

Constructs and stable cell lines. The gene of the intact human C-C chemokine receptor type 5 (CCR5; NCBI reference sequence: NP_000570.1) was cloned into pcMV-IRE6-puro vector (Codex Biosolutions). Genes of HIV-1 gp120 (residues 1–507) from the isolate 92BR020 with a C-terminal hexahistidine tag, and of four-domain CD4 (residues 1–388) with a C-terminal twin strep tag [(GGGGS)2WSHPQFEK(GGGGS)2WSHPQFEK] were synthesized by GenScript and cloned into pcMV-IRE6-puro vector. HEK293T cell lines (Thermo Fisher Scientific) that were stably transfect (HEK293T-CCR5) with these constructs were generated either in-house or at Codex Biosolutions. In brief, 8 × 10⁶ HEK293T cells in 2 ml of DMEM containing 10% FBS and no antibiotics were seeded on a 6-well plate and incubated overnight. The cells were then transfected with the expression constructs using DNA-In 293 Transfection Reagent (MRT-GlobalStem), following a protocol recommended by the manufacturer. Twenty-four hours after transfection, the cells were transferred to a medium containing DMEM, 10% FBS and 1 μg/ml puromycin for selection. Single colonies were picked after 2–3 weeks, and transferred to 24-well plates in the same selective medium.

Protein expression was confirmed by both western blot and a fluorescence-activated cell sorting assay (see below). Positive clones were expanded, frozen and stored in liquid nitrogen. To grow cells in large-scale in suspension, we also generated stable cell lines expressing CCR5 with Exp293F (Exp293F-CCR5) cells (Thermo Fisher Scientific). Hybridoma cells for production of an anti-V3 antibody 447-52D were obtained from Dr. Zolla-Pazner (New York University). Cell line was authenticated for protein expression by western blot and/or flow cytometry, and other functional assays, such as cell-cell fusion and chemokine receptor assays. Our cell culture is routinely tested for mycoplasma contamination.

Purification of recombinant proteins. HIV-1 gp120 of the isolate 92BR020. Cells that express C-terminal His-tagged 92BR020 gp120 were grown in 250-ml roller bottles with DMEM containing 10% FBS and 1 μg/ml puromycin. The protein was purified by affinity chromatography using Ni-NTA agarose (Qiagen) followed by gel filtration chromatography, as previously described56,57. The peak fractions were pooled and concentrated to 10 mg/ml using a 10-kDa MWCO Millipore filter (MilliporeSigma).

Soluble CD4. Cells that express strep-tagged CD4 were grown in 250-ml roller bottles with DMEM containing 10% FBS and 1 μg/ml puromycin. Once the cells reached ~70% confluence, the medium was replaced with HEK293T serum-free expression medium. After 5 days, cell supernatants were collected and loaded onto a Strep-Tactin Sepharose (JBA Lifesciences) column. The column was then washed with 100 mM Tris-HCl, pH 8.0 and 150 mM NaCl. The protein was eluted with 100 mM Tris-HCl, pH 8.0, 150 mM NaCl and 2.5 mM dithiothreitol (IBA). Eluted fractions were concentrated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE), and peak fractions containing CD4 were pooled and dialysed against 100 mM Tris-HCl, pH 8.0 and 150 mM NaCl using a dialysis tubing with a Gatan US4000 CCD camera and operated at a voltage of 200 kV. Particles were recorded on a Philips Tecnai F20 electron microscope (Thermo Fisher Scientific) equipped with a Gatan US4000 CCD camera and operated at a voltage of 200 kV. Particles were

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picked manually and 2D class averages were generated by using EMAN2 software package\textsuperscript{53}. These 2D averages were used to generate a 3D initial model starting from a density of isotropic Gaussian distribution\textsuperscript{64}.

**Cryo-EM sample preparation and data collection.** To prepare cryo grids, 3 μl of the freshly purified CD4–gp120–CCR5 complex at 1.0 mg/ml was applied to a 1.2/1.3 Quantifoil grid (Quantifoil Micro Tools GmbH), which had been glow-discharged for 90 s at 20 mA. Grids were immediately plunge-frozen in liquid ethane using a Vitrobot (Thermo Fisher Scientific) with a blottting time of 4 s. The grids were first screened for ice thickness and particle distribution using a Talos Arctica transmission electron microscope (Thermo Fisher Scientific) operated at 200 kV and equipped with a K2 Summit direct detector (Gatan). For data collection, images were acquired with selected grids using a Titan Krios transmission electron microscope (Thermo Fisher Scientific) operated at 300 kV with a K2 detector. Automated data collection was carried out using SerialEM\textsuperscript{65} at a magnification of 130,000× and the K2 detector in super-resolution mode (pixel size, 0.529 Å) at a dose rate of ~6 electrons per physical pixels per second. Each movie had a total accumulated exposure of ~46 e/Å\textsuperscript{2} fractionated in 35 frames of 200 ms. Total of four datasets were acquired in different sessions using a defocus range of 1.0–2.8 μm.

**Image processing and 3D reconstructions.** For cryo-EM data, drift correction was performed using MotionCor2\textsuperscript{66} and images were binned 2 × 2 by Fourier cropping to a pixel size of 1.059 Å. The contrast transfer function was estimated using CTFIND4\textsuperscript{67} using motion-corrected sums without dose-weighting. Motion-corrected sums with dose-weighting were used for all other image processing. RELION 2.1\textsuperscript{68} was used for particle picking, 2D classification, 3D classification and refinement procedures. Around 2,000 particles were manually picked and classified by 2D classification to generate the templates for automatic particle picking. After the manual inspection of auto-picked particles, a total of 1,707,675 particles were extracted from 9,776 selected images (out of 10,530 movie stacks in total collected in four sessions). These particles were subjected to 2D classification in three groups, giving a total of 1,546,032 particles. The low-resolution negative-stain reconstruction of the complex was low-pass-filtered to a 60 Å resolution, and used as the initial model for 3D classification. A total of 7,600 particles were manually picked and classified by 2D classification to generate the templates for automatic particle picking. The atomic structure coordinates are deposited in the RCSB Protein Data Bank (PDB) under the accession numbers 6MEO and 6MET; and the electron microscopy maps have been deposited in the Electron Microscopy Data Bank (EMDB) under the accession numbers EMD-9108 and EMD-9109. All other related data generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

**Model building.** The initial model of gp120 was a homology model calculated by I-TASSER\textsuperscript{68}, using the cryo-EM structure of gp120 from the CD4-bound SOSIP trimer (PDB ID: 5VN3) as a template. The crystal structures of CD4 (PDB ID: 1WIO) and CCR5 (PDB ID: 5WU) were also used as initial templates for model building. Several rounds of manual building were performed in Coot\textsuperscript{69}. The model was finalized by refinement in Phenix\textsuperscript{70} against the 3.9 Å cryo-EM map. The refinement statistics are summarized in Extended Data Table 1.

**Data availability**

The atomic structure coordinates are deposited in the RCSB Protein Data Bank (PDB) under the accession numbers 6MEO and 6MET; and the electron microscopy maps have been deposited in the Electron Microscopy Data Bank (EMDB) under the accession numbers EMD-9108 and EMD-9109. All other related data generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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Extended Data Fig. 1 | Previously known structures of CCR5 and CXCR4. CCR5 and CXCR4 were identified as the coreceptors for HIV-1 entry in 1996.71–77. **a, b,** Crystal structures of a modified CCR5 (C224–N226 deleted and replaced with rubredoxin; ΔF320–L352; and the point mutations C58Y, G163N, A233D and K303E) in complex with the HIV entry-inhibitor maraviroc (PDB ID: 4MBS) (a) and a modified chemokine [5P7]CCL5 (an antagonist; PDB ID: 5UIW) (b). CCR5 is shown in ribbon diagram in blue, with the internally fused rubredoxin in magenta and the ligands in yellow. **c–e,** Crystal structures of an engineered CXCR4 in complex with a viral chemokine antagonist vMIP-II (PDB ID: 4RWS) (c), a small molecule antagonist IT1t (PDB ID: 3ODU) (d) and a cyclic peptide antagonist CVX15 (PDB ID: 3OE0) (e). CXCR4 is shown in green, the fused T4 lysozyme in magenta and the ligands in yellow.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Characterization of stable cell lines (HEK293T and Expi293F) expressing wild-type human CCR5. a, Chemokine receptor assay. HEK293T and HEK293T-CCR5 (stable) cells were treated with different concentrations of CCL5. \( F/F_0 \) is a fluorescence-signal ratio proportional to that of intracellular cAMP concentration at 40 min after CCL5 activation and at time 0. The dose–response curves were plotted for both HEK293T (black) and HEK293T-CCR5 (red) cells. The experiment was carried out in quadruplicate, and repeated at least three times with similar results. Error bars indicate the standard deviation calculated by the STDEV function in Excel.

b, Flow cytometry histograms of HIV-1 gp120 binding to CCR5 expressed on the cell surfaces in the absence (orange) or presence (red) of soluble CD4. HEK293T cells (black), CCR5-expressing cells only (grey) and CCR5-expressing cells with soluble CD4 only (blue) were negative controls. The experiment was repeated independently at least twice with similar results.

c, HIV-1 Env-mediated cell–cell fusion. HEK293T cells stably transfected with CCR5 were mixed with HIV-1 Env (gp160)-expressing cells in the absence or presence of soluble CD4. The CCR5 cells fuse with CD4-triggered Env cells very efficiently, and form large syncytia that cover almost the entire well. The experiment was repeated independently twice with similar results.

d, Chemokine receptor assay by various ligands. As in a, Expi293F and Expi293F-CCR5 (stable) cells were treated with CCL5, gp120, CD4 or the complex of gp120 and CD4. The dose–response curves were plotted for both Expi293F as a control (left) and Expi293F-CCR5 (right) cells, with different ligands as indicated. The experiment was carried out in quadruplicate and repeated at least three times with similar results. Error bars indicate the standard deviation calculated by the STDEV function in Excel.

e, Left, kinetic curves of 5 representative wells of HEK293T-CCR5 cells treated with 5 different ligands as indicated. ATP activates the endogenous \( G_q \)-coupled G-protein-coupled receptor (P2Y receptor), as a positive control. The ratio represents fluorescence intensity divided by baseline intensity. Right, dose–response curve of each ligand. The y axis is a background-subtracted ratio (peak fluorescent intensity ratio − 1). We conclude that our gp120 and gp120–CD4 do not activate G-protein-mediated calcium flux at the concentrations tested here. The experiment was carried out in quadruplicate and repeated twice with similar results. Error bars indicate the standard deviation calculated by the STDEV function in Excel.

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Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Purification of the CD4-gp120-CCR5 complex.

a, Schematic of expression constructs for HIV-1 gp120, human CCR5 and CD4. Segments of gp120 are designated as follows: C1–C5, conserved regions 1–5; V1–V5, variable regions 1–5; and His-tag, a six-histidine tag. Tree-like symbols represent glycans. Abbreviations used for segments of CCR5 are: N, N terminus; TM1–TM7, transmembrane helices 1–7; ECL1–ECL3, extracellular loops 1–3; ICL1–ICL3, intracellular loops 1–3; and CT, cytoplasmic tail. For CD4, the following abbreviations are used: D1–D4, immunoglobulin (Ig) domains 1–4; and strep tag, a purification tag. The transmembrane segment (TM) and cytoplasmic tail (CT) in grey are truncated in the expression construct.

b, Unmodified human CCR5 in complex with HIV-1 gp120 and four-domain CD4 was purified by the following steps. (1) Complex formation: HIV-1 gp120 (light blue) and strep-tagged, four-domain CD4 (green) were incubated with CCR5 (magenta)-expressing cells to allow formation of the CD4–gp120–CCR5 complex on cell surfaces. (2) Strep-tag purification: the CCR5 complex and some of the CD4–gp120 complex were captured to strep-tactin resin via the strep-tagged CD4 (strep tag in purple). They were eluted by d-desthiobiotin under mild conditions. (3) Negative selection by an anti-V3 antibody to remove the CD4–gp120 complex. The CCR5 complex was further purified by size-exclusion chromatography. The purified CD4–gp120–CCR5 complex was resolved by gel-filtration chromatography on a Superose 6 column in the presence of the detergent LMNG. The molecular-mass standards include thyoglobulin (670 kDa), ferritin (440 kDa), γ-globulin (158 kDa) and ovalbumin (44 kDa). The expected size of the CCR5 complex is ~310 kDa (120 kDa for gp120, 50 kDa for four-domain CD4, 40 kDa for CCR5 and ~100 kDa for LMNG micelle). Peak fractions were analysed by Coomassie-stained SDS–PAGE (lanes 1–3). Labelled bands were confirmed by western blot and protein sequencing. The experiment was repeated independently at least 15 times with similar results.
Extended Data Fig. 4 | Characterization of the CD4–gp120–CCR5 complex by electron microscopy. a, Representative image of the CD4–gp120–CCR5 complex in negative stain. The experiment was repeated independently at least 4 times with similar results. b, 2D averages of the negatively stained CD4–gp120–CCR5 complex. The box size of 2D averages is ~330 Å. c, 3D reconstruction of the negatively stained CD4–gp120–CCR5 complex, fitted with a gp120 structure containing an extended V3 loop (PDB ID: 2QAD), four-domain CD4 (PDB ID: 1WIO) and CCR5 (PDB ID: 4MBS). d, A representative cryo-EM image of the four-domain-CD4–gp120–CCR5 complex. Scale bar, 25 nm. Five independent large datasets were collected with similar results. e, 2D averages of the cryo-EM particle images show secondary structural features for both gp120 and CCR5.
Extended Data Fig. 5 | Single-particle cryo-EM analysis of the CD4–gp120–CCR5 complex. a, Data-processing workflow for the CD4–gp120–CCR5 complex. b, 3D reconstructions of the CD4–gp120–CCR5 complex refined with no mask at an overall resolution of 4.5 Å (left), and with a mask to exclude the last two domains of CD4 at a resolution of 3.9 Å (right), are coloured according to local resolution estimated by RELION. c, The angular distribution of the cryo-EM particles used in the reconstruction is also shown in respect to both the side and top views of the electron microscopy map. d, Gold standard Fourier shell correlation curves of the unmasked and masked electron microscopy reconstructions shown in b.
Extended Data Fig. 6 | Gallery of representative density for the CD4–gp120–CCR5 complex. Representative density in grey mesh from the 3.9 Å resolution electron microscopy map is shown for transmembrane helices TM1–TM7, the N terminus of CCR5, extracellular loop 3 (ELC3) near TM6, Tys10, Tys14 and Tyr15 (red model); two V3 regions; and for helix α1, N terminus, V3 loop, the bridging sheet and N-linked glycan at N262 of gp120 (cyan model).
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Comparison of the conformations of the V3 loop and [5P7]CCL5 in complex with CCR5, as well as of gp120-bound CCR5 and G-protein-bound β2 adrenergic receptor. 

a, The structures of the CD4–gp120–CCR5 and [5P7]CCL5–CCR5 complexes are superposed on CCR5 (red). The V3 loop of gp120 with its Pro311 in stick model is in cyan and [5P7]CCL5 with its Pro3 in stick model in yellow. Residues 309–316 of the V3 loop and residues 1–8 of [5P7]CCL5 adopt a very similar structure, and are highlighted in a rectangular box.

b, Superposition of the structures of the N terminus of the gp120-bound CCR5 (red) and the complementarity-determining region H3 loop of antibody 412d in complex with gp120 core (green). The electron microscopy density of the CD4–gp120–CCR5 complex is shown in grey. The positions of the sulfated tyrosine ('Tys') residues, including Tys10 and Tys14 (from CCR5) and Tys100 and Tys100c (from 412d), are indicated.

c, A model for interactions of three CD4 receptors and three CCR5 coreceptors with the SOSIP Env trimer. The side and bottom views of a composite structure of the CD4–CCR5–SOSIP Env trimer complex are shown. The model was generated using the CD4-bound SOSIP trimer (PDB ID: 5VN3) and the structure of the CD4–gp120–CCR5 complex from this study. All the structures were aligned on the basis of the core region of gp120. CCR5 is shown in red, CD4 in green, gp120 in blue, the gp120 of SOSIP in dark blue and the gp41 of SOSIP in grey. The crystallographic dimer of CCR5 (PDB ID: 4MBS) is also shown, on the left only, in a rectangular box. The observed crystallographic dimer of CCR5 or the transmembrane helix 5-mediated dimer by modelling does not seem to be relevant to binding to either monomeric or trimeric gp1207,78.

d, Superposition of the structures of the gp120-bound CCR5 (red) and the G-protein-bound β2 adrenergic receptor (blue). The position of TM6, which is critical for the activation of G-protein-coupled receptors, is indicated.
Extended Data Fig. 8 | Comparison of conformations of different structures of monomeric gp120 and various V3 loops. a, Comparison of structures of an unliganded gp120 core (PDB ID: 4OLV; purple), a CD4-bound monomeric gp120 core with the V3 loop (PDB ID: 2QAD; blue) and gp120 in complex with CD4 and CCR5 from this study (cyan). The gp120 core region is marked by a circle with a diameter of 50 Å. The N and C termini, V1V2 stem, V3 stem or loop and bridging sheet are indicated. b, Representative conformations that an HIV-1 V3 loop can adopt. From left to right, V3 loop in the unliganded SOSIP BG505 Env trimer (PDB ID: 4ZMJ); the first-V3-containing gp120 core in complex with CD4 and antibody X5 (PDB ID: 2B4C29); CD4- and 412d-bound monomeric gp120 core with V3 (PDB ID: 2QAD); CCR5-bound intact gp120 (this study); and V3 peptide in complex with antibody 447-52D (PDB ID: 3GHB36); antibody 268-D (PDB ID: 3GO1 37); antibody 2557 (PDB ID: 3MLV37); and antibody 10A37 (PDB ID: 5V6L 38). The root-mean-square deviation of each structure (except for 5V6L), relative to the CCR5-bound gp120 monomer, is shown at the bottom in parentheses.
Extended Data Fig. 9 | Model of HIV-1 Env activation to induce membrane fusion. A hypothesis of how the cellular receptors CD4 and CCR5 trigger the HIV-1 Env trimer to induce membrane fusion and viral entry. Left, virus attaches to the target cell by gp120 (cyan) binding to CD4 (green). Helix collar (gp41), the four-helix collar gripping the N- and C termini of gp120. Right, immediate binding by CCR5 (red) prevents rapid dissociation between gp120 and CD4, stabilizes the CD4-induced conformational changes within the Env trimer and brings the trimer close to the cell membrane. Simultaneous binding of gp120 to both CD4 and CCR5 may require bending in the cell membrane. The fusion peptide (magenta) of gp41 (grey) flips out owing to intrinsic conformational dynamics, which enables the bending back of the N and C termini of gp120. This bending blocks the fusion peptide from resuming its original position in the trimer. The movements of the fusion peptide and gp120 termini effectively weaken the non-covalent association between the two subunits and may lead to partial or complete dissociation of gp120, as well as a series of refolding events in gp41 to adopt the pre-hairpin intermediate conformation (with the fusion peptides inserting into the target-cell membrane). Extended helix (gp41), three helices in the fusion-intermediate conformation of gp41.
## Extended Data Table 1 | Cryo-EM data collection, refinement and validation statistics

|                     | CD4-gp120-CCR5 complex masked (EMDB-9108) (PDB 6MEO) | CD4-gp120-CCR5 complex overall (EMDB-9109) (PDB 6MET) |
|---------------------|------------------------------------------------------|-------------------------------------------------------|
| **Data collection and processing** |                                                      |                                                      |
| Magnification       | 130000                                               | 130000                                                |
| Voltage (kV)        | 300                                                  | 300                                                   |
| Electron exposure (e-/Å²) | ~46                                                  | ~46                                                   |
| Defocus range (µm)  | 1.2.8                                                | 1.2.8                                                  |
| Pixel size (Å)      | 0.529                                                | 0.529                                                  |
| Symmetry imposed    | C1                                                   | C1                                                    |
| Initial particle images (no.) | 1,707,575                                           | 1,707,575                                             |
| Final particle images (no.) | 307346                                              | 307346                                                |
| Map resolution (Å)  | 3.9                                                  | 4.5                                                   |
| FSC threshold       | 0.143                                                | 0.143                                                  |
| **Refinement**      |                                                      |                                                      |
| Initial model used (PDB code) | 5UIW, 1WIO, 2QAD and 5VN3                           | 5UIW, 1WIO, 2QAD and 5VN3                            |
| Map sharpening B factor (Å²) | -190                                                | -190                                                  |
| Model composition   |                                                      |                                                      |
| Non-hydrogen atoms  | 7462                                                 | 8911                                                  |
| Protein residues    | 887                                                  | 1074                                                   |
| Ligands             | 32                                                   | 32                                                    |
| R.m.s. deviations   |                                                      |                                                      |
| Bond lengths (Å)    | 0.004                                                | 0.009                                                  |
| Bond angles (°)     | 0.954                                                | 1.386                                                  |
| Validation          |                                                      |                                                      |
| MolProbity score    | 1.60                                                 | 1.97                                                   |
| Clashscore          | 3.51                                                 | 5.84                                                   |
| Poor rotamers (%)   | 0.25                                                 | 1.48                                                   |
| Ramachandran plot   |                                                      |                                                      |
| Favored (%)         | 92.77                                                | 90.83                                                  |
| Allowed (%)         | 7.23                                                 | 8.51                                                   |
| Disallowed (%)      | 0                                                    | 0.66                                                   |
**Reporting Summary**

Nature Research wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Research policies, see Authors & Referees and the Editorial Policy Checklist.

### Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

| n/a | Confirmed |
|-----|-----------|
| ☑  | n/a       |

- **The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement**
- **An indication of whether measurements were taken from distinct samples or whether the same sample was measured repeatedly**
- **The statistical test(s) used AND whether they are one- or two-sided**
  *Only common tests should be described solely by name; describe more complex techniques in the Methods section.*
- **A description of all covariates tested**
- **A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons**
- **A full description of the statistics including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)**
- **For null hypothesis testing, the test statistic (e.g. \( F, t, r \)) with confidence intervals, effect sizes, degrees of freedom and \( P \) value noted**
  *Give \( P \) values as exact values whenever suitable.*
- **For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings**
- **For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes**
- **Estimates of effect sizes (e.g. Cohen's \( d \), Pearson's \( r \)), indicating how they were calculated**
- **Clearly defined error bars**
  *State explicitly what error bars represent (e.g. SD, SE, CI)*

Our web collection on [statistics for biologists](#) may be useful.

### Software and code

**Policy information about availability of computer code**

| Data collection | SerialEM (3.5) |
|-----------------|---------------|

| Data analysis | Flowlo (10.5.3), EMAN2 (2.2), MotionCor2 (1.0.2), CTFFIND4 (4.1.5), RELION (2.1), I-TASSER, SAMUEL (17.05), Coot (0.8.8), Chimera (1.12), Phenix (1.11.1-2575). |

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

### Data

**Policy information about availability of data**

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The atomic structure coordinates are deposited in the Protein Data Bank under the accession number 6MEO and 6MET, and the EM maps in the EMDataBank.
under the accession number EMD-9108 and EMD-9109. All other related data generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Field-specific reporting

Please select the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- Life sciences
- Behavioural & social sciences
- Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/authors/policies/ReportingSummary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Statistical methods were not needed to predetermine sample size for the biochemical and structural studies in this work. Multiple independent EM data sets were collected for structural analysis. All other experiments were repeated multiple times with the similar results. |
| Data exclusions | No data were excluded from analyses. |
| Replication | Multiple EM data sets were collected with very similar quality. All other experiments have been repeated multiple times with excellent reproducibility. |
| Randomization | Experimental groups are not needed for this work, therefore randomization is not relevant. |
| Blinding | The investigators were blinded to group allocation during data collection and/or analysis because groups were not necessary for this work. |

Reporting for specific materials, systems and methods

Materials & experimental systems

| n/a | Involved in the study |
| --- | --- |
| ☒ | Unique biological materials |
| ☒ | Antibodies |
| ☒ | Eukaryotic cell lines |
| ☒ | Palaeontology |
| ☒ | Animals and other organisms |
| ☒ | Human research participants |

Methods

| n/a | Involved in the study |
| --- | --- |
| ☒ | ChIP-seq |
| ☒ | Flow cytometry |
| ☒ | MRI-based neuroimaging |

Antibodies

Hybridoma cells for production of an anti-V3 antibody 447-52D was kindly provided by Dr. Susan Zolla-Pazner, New York University. PE Mouse anti-human CD195 (Clone 2D7/CCR5 (RUO), Catalog # 550632, Lot # 5219800; BD Biosciences, San Jose, CA). Anti-His tagged PE conjugated Mouse IgG (Catalog # IC050P, Lot # LHN0316101; R&D Systems, Minneapolis, MN). Anti-CCR5 antibody (Catalog # AB1889, Lot # 2816560; EMD Millipore Corp, USA).

Validation

Antibody 447-52D was tested for binding to HIV-1 gp120. The BD Biosciences website states PE Mouse Anti-Human CD195 is routinely tested by flow cytometry. For anti-His tagged PE conjugated Mouse IgG, the R&D Systems website lists the following citations: T Carmenate et al., J. Immunol., 2018;0(0); DX Bu et al., Oncotarget, 2018;9(40):25764-25780; Bozza S et al., J Immunol, 2014;193(5):2340-8; Sun Y et al., J. Biol. Chem., 2012;287(19):15837-50. The EMD Millipore website states that anti-CCR5 antibody (Catalog # AB1889) was confirmed by western blot analysis of CCR5 in THP-1 whole cell lysate with anti-CCR5 (NT) at 1:1000 dilution.

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

HEK 293T cells were purchased from ATCC; Expi293F from Thermo Fisher Scientific. 293 T or Expi293F stable cell lines were generated either in Bing Chen’s lab at Boston Children’s Hospital or at Codex Biosolutions.
Authentication
Each cell line was authenticated for protein expression by western blot and/or flow cytometry, and other functional assays, such as cell-cell fusion and chemokine receptor assays.

Mycoplasma contamination
Mycoplasma contamination is routinely tested for our cell culture and no contaminated cells were ever used for our studies.

Commonly misidentified lines
(See ICLAC register)
None.

Flow Cytometry

Plots

Confirm that:
☐ The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
☒ The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a ‘group’ is an analysis of identical markers).
☐ All plots are contour plots with outliers or pseudocolor plots.
☐ A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation
HEK 293T and Expi293F 293T cells stable cell lines was used. For CCR5 expression analysis 1 million of CCR5 expressing cells were washed with PBS and incubated for 30~40 minutes on ice with PE Mouse antihuman CD195 (BD Biosciences, San Jose, CA) in PBS containing 1% BSA. For CCR5-CD4-gp120 complex detection on the cell surface, 1 million of CCR5 expressing cells were washed with PBS and incubated for 30~40 minutes on ice with CD4 and gp120 at concentrations of 4 μg/ml and 10 4 μg/ml respectively in PBS containing 1% BSA. The cells were then washed twice with PBS containing 1% BSA and stained with Anti-His tagged PE conjugated Mouse IgG (R&D Systems, Minneapolis, MN) at 5 μg/ml. All the fluorescently labeled cells were washed twice with PBS containing 1% BSA and analyzed immediately using a BD FACS Canto II instrument and program FACSDIVA (BD Biosciences, San Jose, CA). All data were analyzed by FlowJo (FlowJo, LLC, Ashland, OR).

Instrument
BD FACSCanto II

Software
FlowJo

Cell population abundance
N/A

Gating strategy
Only gating used during analysis was to separate live and single cell populations.

☐ Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.