Adaptation of Subtype A Human Immunodeficiency Virus Type 1 Envelope to Pig-tailed Macaque Cells

Byline: ADAPTATION OF SUBTYPE A HIV-1 ENV TO MACAQUE CELLS

Daryl Humes1,2 and Julie Overbaugh2*

Program in Molecular and Cellular Biology, University of Washington, Seattle, Washington, 1 and Division of Human Biology, Fred Hutchinson Cancer Research Center, Seattle, Washington2

*Corresponding author. Mailing address: Division of Human Biology, Fred Hutchinson Cancer Research Center, 1100 Fairview Ave. N, Mailstop C3-168, Seattle, WA 98109-102. Phone: (206) 667-3524. Fax: (206) 667-1535. E-mail: joverbau@fhcrc.org.

Abstract

The relevance of simian/human immunodeficiency virus (SHIV) infection of macaques to HIV-1 infection in humans depends on how closely SHIVs mimic HIV-1 transmission, pathogenesis, and diversity. Circulating HIV-1 strains are predominantly subtypes C and A, and overwhelmingly require CCR5 for entry, yet most SHIVs incorporate CXCR4-using subtype B envelopes (Envs). While pathogenic subtype C-based SHIVs have been constructed, subtype A-based SHIVs (SHIV-As) constructed to date have been unable to replicate in macaque cells. To understand the barriers to SHIV-A replication in macaque cells, HIVA/Q23/SIVvif was constructed by engineering a CCR5-tropic subtype A provirus to express SIV vif, which counters the macaque APOBEC3G restriction. HIVA/Q23/SIVvif replicated poorly in pig-tailed macaque (Ptm) lymphocytes, but viruses were adapted to Ptm lymphocytes. Two independent mutations in gp120, G312V (V3 loop) and A204E (C2 region), were identified that increased peak virus levels by >100-fold. Introduction of G312V and A204E to multiple subtype A Envs and substitution of G312 and A204 with other residues increased entry into Ptm cells by 10 to 100-fold. G312V and A204E Env variants continued to require CCR5 for entry, but were up to 50 and 200-fold more sensitive to neutralization by IgG1b12 and soluble CD4, and had a 5 to 50-fold increase in their ability to utilize ptmCD4 as compared to their wild-type counterparts. These findings identify the inefficient use of ptmCD4 as an unappreciated restriction to subtype A HIV-1 replication in Ptm cells, and reveal amino acid changes to gp120 that can overcome this barrier.
Introduction

Non-human primate (NHP) models of HIV/AIDS are important tools for studying aspects of HIV-1 transmission, pathogenesis, and immunity. The most commonly used NHPs in these studies are Asian macaques, particularly Rhesus macaques (Rhm; *Macaca mulatta*) and pig-tailed macaques (PtM; *Macaca nemestrina*) (2). HIV-1 is thought to be able to utilize the macaque CD4 and CXCR4/CCR5 receptors for viral entry (13, 28), however HIV-1 replication is restricted by other factors, most notably TRIM5α in Rhm, and the APOBEC3 family of cytidine deaminases in both species (15, 36, 58). Because of these species-specific restrictions, the direct study of HIV-1 infection and pathogenesis in macaques is not possible and instead research has focused on the development of SIV/HIV chimeric viruses (SHIVs). SHIVs have traditionally been constructed by inserting the region of the HIV-1 genome encompassing the *tat, rev, vpu,* and *env* genes into the context of the pathogenic SIV<sub>mac</sub>239 backbone (55). More recently, directed approaches that rely on engineering HIV-1 to encode SIV *gag* and *vif* sequences to specifically counter macaque TRIM5α and APOBEC3 have led to the creation of ‘minimal’ SHIVs that primarily encode HIV-1 proteins (16, 24). The *env* gene is a particularly important component of SHIVs because the envelope (Env) determines viral tropism and is the target of neutralizing antibodies, which are considered an important part of the protective HIV-1 immune response [as reviewed in (37)].

If SHIVs are to be effective as predictors of human disease and vaccine efficacy, they should closely mimic the transmitted strains in human infection. The Envs from most circulating strains of HIV-1 require CCR5 as a co-receptor for entry, whereas many of
the current SHIVs make use of envelopes from CXCR4-tropic or dual-tropic clones, such as NL4-3, HXB2, HIV\textsubscript{SF33} and HIV\textsubscript{89.6} (31, 35, 51, 55). Moreover, the Envs encoded by these SHIVs are highly sensitive to neutralization as compared to circulating HIV-1 variants (4, 5, 38, 60). Among subtype B CCR5-tropic SHIVs (14, 41, 45), SHIV\textsubscript{SF162} passaged isolates are the most commonly used, however the SF162 Env encoded by these SHIVs is also extremely sensitive to neutralization (54, 63). Thus, current SHIVs do not provide a realistic benchmark for neutralizing antibody protection from circulating strains of HIV-1, and many also do not model the dominant CCR5-mediated mode of transmission.

The world-wide epidemic is comprised of very diverse HIV-1 genotypes, termed clades or subtypes. In sub-Saharan Africa, which carries the highest burden of new HIV-1 infections and HIV-1 related deaths, it is subtypes C and A that predominate (17). SHIVs that are infectious to macaques have been generated using subtype C env sequences (9, 56, 57), as well as env sequences from the circulating recombinant CRF\_AE, which is the most common HIV-1 subtype in Southeast Asia, (20, 26). Despite the relative prominence of subtype A strains in the most afflicted regions of the world, attempts to make subtype A-based SHIVs (SHIV-As) have thus far been unsuccessful as the SHIV-As tested to date failed to replicate in macaque cells (19).

In order to gain further insight into barriers to SHIV-A replication in macaque cells, we created HIV\textsubscript{A\textsubscript{Q23}/SIV\textsubscript{vif}}, a minimal SHIV encoding the vif gene from SIV\textsubscript{mac239} in the context of the Q23-17 provirus, which is a CCR5-tropic subtype A HIV-1 molecular
clone obtained soon after seroconversion (48). This minimal SHIV approach takes
advantage of the fact that the APOBEC3-mediated restriction to HIV-1 replication in Ptm
cells can be countered by SIV Vif (15), and, the fact that, in contrast to Rhm TRIM5α,
the Ptm TRIM5 isoforms and TRIMCyp do not antagonize HIV-1 infection (6, 7, 33, 62).
In this study, the replicative properties and adaptation of HIVA\textsubscript{Q23}/SIV\textsubscript{vif} to Ptm cells
were explored. Two adaptive mutations were identified that, when introduced into
different subtype A Envs, permit much more efficient usage of Ptm CD4, resulting in a
dramatic increase in the infectivity of Ptm cells. These findings identify the inefficient
use of Ptm CD4 as a previously uncharacterized barrier to subtype A HIV-1 replication in
Ptm cells, and provide approaches to increase SHIV-A infection in macaque cells.

Materials and Methods

Construction of HIVA\textsubscript{Q23}/SIV\textsubscript{vif}

HIVA\textsubscript{Q23}/SIV\textsubscript{vif}, a full-length replication-competent clone expressing \textit{vif} from SIV\textsubscript{mac}239,
was created from Q23\textsubscript{Δvif} (46). Q23\textsubscript{Δvif} was derived from the Q23-17 full-length
molecular clone (48) and was engineered with unique SalI and MluI restriction sites at
the 5’ and 3’ ends of \textit{vif}, causing a frameshift in the endogenous \textit{vif} gene and allowing
for the insertion and expression of different \textit{vif} variants [as in (53)]. To make
HIVA\textsubscript{Q23}/SIV\textsubscript{vif}, the entire SIV\textsubscript{mac}239 \textit{vif} open reading frame was amplified from
SIV\textsubscript{mac}239\textsubscript{Δenv} (a gift from Dr. David Evans) using forward primer 5’-
GAAGGTCGACATGGAGGAAAAGA-3’ and reverse primer 5’-
AGTGACCGTGTCATGCCAGTATTCCCAA-3’ (restriction sites underlined). The
PCR product was then digested with the SalI and MluI restriction enzymes, ligated into
Q23\textsubscript{Δvif}, and verified by sequencing.
Env clones and mutagenesis

In addition to Q23ENV.17 [(50); referred to throughout as Q23-17], the following plasmids expressing subtype A Envs were used in the study: QF495.23M.ENV.A3 [(4); referred to throughout as QF495.A3], BG505.W6M.ENV.B1 and MG505.W0M.ENV.H3 [(64); referred to throughout as BG505.B1 and MG505.H3, respectively], and Q259.D2.26 and Q259.D2.17 (34). Additionally, the SF162P3 clone, constructed by inserting the predominant V1-V5 sequence from the SHIV SF162P3 isolate into the HIV-1SF162 Env clone, [(21); a gift from Dr. Cecilia Cheng-Mayer] and the SIV Mne CL8 Env clone (47) were used.

Mutations were introduced to the subtype A Env clones by site-directed mutagenesis using primers designed according to the Quik-Change site-directed mutagenesis kit (Stratagene; primer sequences available upon request) to amplify 25 ng of plasmid with Pfu Turbo (Invitrogen) under the following reaction conditions: 95°C for 5 min, followed by 18 cycles of 95°C for 30s, 55°C for 1 min, and 68°C for 16 min. The Env mutants were sequenced through the entirety of the env open reading frame to verify that no undesired nucleotide changes had occurred.

Construction of other full-length molecular clones

Chimeric full-length molecular clones were constructed by digesting the subtype B pNL-DT5R [(24); a gift from Dr. Malcolm Martin] and HIVAQ23/SIVvif with EcoRI (restriction site located in vpr) and XhoI (restriction site located in nef) and ligating the heterologous
fragments. The resulting chimeras were named Q/N_{vpr-nef} and N/Q_{vpr-nef} to indicate the origin of the vpr to nef portion in each clone (Fig. 1b).

The HIV\textsubscript{A\textsuperscript{Q23}}/SIV\textsubscript{vif} provirus was engineered to express different env genes of interest using a previously described method (49). In short, HIV\textsubscript{A\textsuperscript{Q23}}/SIV\textsubscript{vif} was digested with SmaI (restriction site located in vpr) and XhoI to excise an ~3 kb sequence encompassing the Q23-17 env gene. Heterologous env genes were then introduced into HIV\textsubscript{A\textsuperscript{Q23}}/SIV\textsubscript{vif} by digesting the env clones of interest with SmaI and XhoI and ligating the fragment into HIV\textsubscript{A\textsuperscript{Q23}}/SIV\textsubscript{vif}.

Virus production and titration

HEK 293T cells (referred to throughout as 293Ts) were used to produce all virus preparations and were maintained in DMEM medium (Invitrogen) supplemented with 10% heat-inactivated fetal calf serum (FCS) and 2 mM L-glutamine.

Full-length replication-competent viruses were produced by transfecting 293T cells using polyethyleneimine (PEI; Polysciences). Briefly, 2×10\textsuperscript{6} 293Ts were plated 24 hours prior to transfection in a T75 tissue culture flask. The next day 6 μg of DNA was incubated with 60 μg of PEI in 600 μL of serum-free DMEM for 10 minutes before adding to cells. Viral supernatants were collected 72 hours after transfection, filtered through a 0.22 μm filter (Millipore Corporation) and stored at -80°C until use.
The following env-deficient HIV-1 proviruses were used to generate pseudoviruses:

Q23Δenv and pLai3ΔenvLuc2 (a gift from Dr. Michael Emerman), both of which have been described previously (34, 65); and Q23Δenv-GFP, which was constructed by sub-cloning eGFP from pEGFPN1 (Clontech) into Q23Δenv. For this purpose, BamHI and NotI sites that were introduced at nucleotides 31 and 63 in the nef open reading frame of Q23Δenv and the eGFP sequence was introduced using these same restriction sites.

Pseudoviruses were produced by co-transfecting 293T cells with one of the plasmids encoding env-deficient proviruses described above and plasmids encoding env clones at a 2:1 mass ratio. To do this, 2.5×10⁵ 293T cells were plated in each well of a 6-well dish 24 hours prior to transfection. For each well, 1 µg of total DNA was mixed with 10 µg of PEI in 100 µL of serum-free DMEM. In some cases, 3 µL of Fugene 6 (Roche) was used in place of PEI per the manufacturer’s instructions. Pseudoviruses lacking Env (Env(-)) were used to determine the background levels of some assays, and were generated by co-transfecting the empty pCI-neo (Promega) mammalian expression vector in place of an env expressing plasmid. Viral supernatants were harvested 48-72 hours post-transfection and cleared of cellular debris by centrifuging at 1300 rpm for 5 min. In some cases, cleared supernatants were concentrated ~20-50-fold using Amicon Ultra 10K filters (Millipore Corporation). All viral preparations were frozen at -80°C until use.

Viral titers were determined by infecting TZM-bl reporter cells (NIH AIDS Research and Reference Reagent Program) with thawed cell-free virus in the presence of 10 µg/mL of DEAE-dextran. Forty-eight hours later, the cells were fixed and stained for beta-
galactosidase activation, and blue foci were counted to obtain infectious titers. Infectious titers were reported as infectious particles (IP)/mL (5).

Virus replication assays

Full-length proviral clones were assessed for the production of infectious virus in human peripheral blood mononuclear cells (PBMCs), primary Ptm PBMCs, and immortalized Ptm lymphocytes. PBMCs from HIV-negative donors were isolated by the Ficoll gradient method, activated for 72 h with 10 U of phytohemagglutinin M/mL (Roche), and maintained in RPMI 1640 medium (Invitrogen) with 10% heat-inactivated FCS, 2 mM L-glutamine, 100 U of penicillin/ml, 100 µg of streptomycin/ml, and 10 U/mL of interleukin-2 (Roche) for 48 h prior to infection and thereafter. Ptm PBMCs were isolated using a 95% Ficoll gradient, activated for 40 h in with 4 U of phytohemagglutinin M/mL, and maintained in RPMI 1640 medium with 25 mM HEPES, 20% heat-inactivated FCS, 2 mM L-glutamine, 100 U of penicillin/ml, 100 µg of streptomycin/ml, and 100 U/mL of recombinant human interleukin-2 (Roche). The immortalized Ptm lymphocytes used in this study have been described previously [(40); a gift from Drs. Nina Munoz and Hans-Peter Kiem]. Ptm lymphocytes were maintained in IMDM medium (Invitrogen) with 10% heat-inactivated FCS, 2 mM L-glutamine, 100 U of penicillin/ml, 100 µg of streptomycin/mL, and 100 U of interleukin-2/mL (Chiron Corporation).

Viral stocks were mixed with either 3×10^6 stimulated donor human PBMCs, 2×10^6 Ptm PBMCs, or 1×10^6 immortalized Ptm lymphocytes at a multiplicity of infection (MOI) of...
0.02 or 0.2 in a final volume of 250 µL. After spinoculation (43) at room temperature for
2 to 3 hours at 1200×g, cells were washed three times in 1.5 mL of the appropriate media
and resuspended into duplicate 600 µL cultures in a 48-well dish. Cultures were
maintained for 15 days, with approximately two-thirds of the medium being replaced
every 3 days. The p24\text{gag} levels were determined by measuring cleared culture
supernatants with a p24\text{gag} Antigen ELISA kit (ZeptoMetrix, Buffalo, NY).

Luciferase assays
Infections of 8×10^4 immortalized Ptm lymphocytes were performed in triplicate by
spinoculation with luciferase reporter viruses for 2 hours at an MOI of 0.02 in a final
volume of 100 µL. Infections were allowed to proceed for 72 hours before lysis with
Brite-Glo reagent (Invitrogen) according to the manufacturer’s guidelines. Luciferase
activity was immediately read on the Fluoroskan Ascent FL luminometer (Thermo
LabSystems) with a 1000 ms integration period. Infections with Env(-) pseudoviruses
were used to determine background levels for the assay.

Long-term culturing of HIVA\text{Q23/SIV}_{vif} and sequencing of outgrowth variants
Cultures of 1×10^6 immortalized Ptm lymphocytes were infected with HIVA\text{Q23/SIV}_{vif} at
an MOI of 0.2 in an initial volume of 1 mL in 12-well dishes. Cells were maintained at a
concentration of 0.8 to 3×10^6 cells/mL without discarding cells and with at least two-
thirds of the media being replenished every 4-5 days. Supernatant was assessed for the
presence of virus every 4-5 days by infecting TZM-bl cells.
Using the QIAgen Blood DNA Kit (QIAgen), total DNA was extracted from cells in cultures in which viral outgrowth was identified. The number of integrated proviral copies was determined using a previously described real-time PCR assay (3). To sequence viral variants, 1000 copies of the integrated provirus were amplified with 4 nested PCRs, resulting in overlapping amplicons spanning the entirety of the HIV-1 genome. The primers and conditions used for this are available upon request. The nested PCRs were performed in triplicate, and the PCR products were detected as single prominent bands by gel electrophoresis. All reactions were treated with ExoSap (Amersham Biosciences) and the amplicons were sequenced directly without gel purification.

Infection of Ptm lymphocytes in the presence of TAK779

TAK779 treated immortalized Ptm lymphocytes were pre-incubated with 1 µM of TAK779 (NIH AIDS Reference and Reagent Program) for 2 hours at 37°C and maintained thereafter in 1 µM of TAK779. Triplicate infections of 8×10^4 TAK779-treated or untreated cells were then performed by spinoculation for 2 hours with pseudotyped luciferase reporter viruses at an MOI of 0.02 in a final volume of 100uL. Infected cultures were maintained for 72 hours before lysis and subsequent measurement of luciferase activity. Percent inhibition by TAK779 was determined by comparing relative luciferase levels in TAK779-treated cells versus untreated cells.

Neutralization assays.
Neutralization assays were performed using the TZM-bl neutralization assay as described previously (5). Approximately 500 IPs of Q23Δenv-derived pseudoviruses were incubated with 6 serial three-fold dilutions of IgG1b12 [(8); referred to throughout as b12] or soluble CD4 (sCD4) in a 96-well plate. One hour later, 1×10^5 TZM-bl cells were added to each well with DEAE-dextran to a final concentration of 10 µg/mL. The relative levels of infection were determined by assessing β-galactosidase activity in triplicate wells after 48 h. Median inhibitory concentrations (IC_{50}s) were defined as the concentration of b12 or sCD4 that resulted in 50% inhibition of β-galactosidase, and were calculated using the linear fit model.

Construction and transient transfection of CD4 and CCR5 expression plasmids

Human CCR5 and Ptm CCR5 (referred to throughout as huCCR5 and ptmCCR5, respectively) in the pBABE-puro vector were described previously (11, 25). To clone human and Ptm CD4 (referred to throughout as huCD4 and ptmCD4, respectively), total RNA was isolated from human PBMCs and Ptm lymphocytes using the Qiagen RNeasy Mini Kit. Total cDNA was obtained by reverse-transcribing 2 µg of RNA with SuperScript II RT (Invitrogen) using oligo-dT primers. The CD4 open-reading frame was PCR-amplified using forward primer 5'-GATGGATCCATGAACGGGGAGTCCC-3' with reverse primer 5’-

GATGGATCCATGAACGGGGGAATCCC-3' for huCD4 and forward primer 5’-

GATGGATCCATGAACGGGGAATCCC-3’ with the same reverse primer for ptmCD4. The PCR products were digested with BamHI and SalI (underlined in the primers), cloned into the pBABE-puro plasmid (39), and verified by sequencing.
CD4 and CCR5 expression plasmids were co-transfected in equal amounts into 293T cells using Fugene 6 at a ratio of 3 µL of transfection reagent to 1 µg of DNA as per the manufacturer’s protocol. In some cases, transfections were performed with 6 µg of DNA in T75 tissue culture flasks with 18 µL of Fugene in a total volume of 200 µL of serum-free DMEM.

**Analysis of receptor expression levels by flow cytometry**

CD4 and CCR5 expression levels were determined by flow cytometry using APC conjugated mouse anti-human CD4 antibody (cat. No. 551980, BD Biosciences) and PE conjugated mouse anti-human CCR5 (cat. no. 550632, BD Biosciences). Briefly, 1-2×10^5 cells were washed in PBS/2% FBS, and incubated in ~100 µL of PBS/2% FBS with 2 uL of the anti-CD4 antibody and/or 5 µL of the anti-CCR5 antibody at room temperature for 30 mins. The cells were then washed once in 1mL of PBS/2% FBS and resuspended in 200 µL PBS/2% FBS for analysis by flow cytometry.

**Results**

**Replication of HIVA_Q23/SIV_vif in immortalized Ptm lymphocytes**

HIVA_Q23/SIV_vif, a minimal SHIV derived from the subtype A CCR5-tropic Q23-17 provirus, was tested for its ability to replicate in human PBMCs and immortalized Ptm lymphocytes. NL-DT5R, a CXCR4-tropic minimal SHIV that is derived from NL4-3 and encodes the vif gene and the cyclophilin A (CypA)-binding loop from SIVmac239, was used as a positive control for replication in Ptm cells [(24); Fig. 1a]. In human PBMCs, HIVA_Q23/SIV_vif achieved peak levels of ~300 ng p24gag/mL, slightly higher than NL-
DT5R, which reached peak levels of ~100 ng p24\textsuperscript{gag}/mL. In Ptm cells, however, while NL-DT5R reached levels of ~300 ng p24\textsuperscript{gag}/mL, HIVA\textsubscript{Q23}/SIV\textsubscript{vif} reached peak levels that were ~300-fold lower at ~1 ng p24\textsuperscript{gag}/mL, despite the fact that infections with HIVA\textsubscript{Q23}/SIV\textsubscript{vif} were performed at a 10-fold greater MOI (Fig. 1b).

To determine which regions of the HIVA\textsubscript{Q23}/SIV\textsubscript{vif} genome were responsible for its impaired infection of Ptm lymphocytes, reciprocal chimeras between HIVA\textsubscript{Q23}/SIV\textsubscript{vif} and NL-DT5R were constructed and evaluated for their ability to replicate in human and Ptm cells (Fig. 1a). In human PBMCs, both chimeric viruses replicated to levels of p24\textsuperscript{gag} that were within the same range as HIVA\textsubscript{Q23}/SIV\textsubscript{vif} and NL-DT5R (Fig. 1b). In Ptm lymphocytes, Q/N\textsubscript{vpr-nef} replicated to similar levels as NL-DT5R, indicating that the 5'LTR and gag-pol region of HIVA\textsubscript{Q23}/SIV\textsubscript{vif} functioned for virus replication in Ptm cells. Conversely, N/Q\textsubscript{vpr-nef} reached a peak level of p24\textsuperscript{gag} that was >2 logs lower than NL-DT5R, even with infections performed at a 10-fold greater MOI (Fig. 1b). This suggested that the 3’ region of HIVA\textsubscript{Q23}/SIV\textsubscript{vif}, encompassing the entirety of tat, rev, vpu and env, as well as parts of vpr and nef, was responsible for the low levels of replication in Ptm cells.

Entry of viruses bearing subtype A Envs into immortalized Ptm lymphocytes.

Given that the viral determinant for impaired replication of HIVA\textsubscript{Q23}/SIV\textsubscript{vif} in Ptm cells included the env gene, the Q23-17 Env was examined for its ability to mediate entry into Ptm cells. The Q23-17 Env was compared to the SF162P3 Env, which is derived from SHIV\textsubscript{SF162P3}, a CCR5-tropic isolate known to establish persistent spreading infection in
Ptms (21). Pseudotyped luciferase reporter viruses, which were normalized based on MOI, infected Ptms 30-fold less efficiently when carrying the Q23-17 Env as compared to those pseudotyped with the SF162P3 Env (Fig. 2).

To determine whether the Q23-17 Env was typical of subtype A Envs, 5 additional CCR5-tropic subtype A Envs, obtained from recently infected individuals, were tested for their ability to infect immortalized Ptms lymphocytes. Viruses carrying the BG505.B1, Q259.d2.26 and Q259.d2.17 Envs mediated entry into Ptms cells to levels that were comparable to Q23-17, while the infectivity of viruses carrying the QF495.A3 and MG505.H3 Envs was below the background levels of the assay (Fig. 2). Overall, these results suggested that inefficient entry into Ptms cells is a common characteristic of subtype A Envs.

Adaptation of HIVAQ23/SIVvif to immortalized Ptms lymphocytes and identification of adaptive amino acid changes.

To determine if HIVAQ23/SIVvif could be adapted to replicate in immortalized Ptms lymphocytes, the virus was maintained in multiple long-term cultures in two independent experiments. In the first experiment, viral outgrowth was seen in 1 of 9 cultures after approximately 35 days in culture, with viral supernatant reaching titers of >$10^6$ IP/mL in TZM-bl cells (data not shown). Sequencing of the integrated proviral genome in triplicate from cells in this culture revealed only a single G to T mutation that was clearly present in all sequences, which encoded a glycine to valine change at position 312 of the Env surface unit protein, gp120 (G312V, HXB2 numbering). Introduction of the G to T
mutation to HIVA\textsubscript{Q23}/SIV\textsubscript{vif} resulted in virus replication that achieved a $>2$ log increase in p24\textsubscript{gag} levels as compared to wild-type (Fig. 3a). However, the HIVA\textsubscript{Q23}/SIV\textsubscript{vif} G312V molecular clone showed intermediate levels of replication compared to the viral quasispecies, with the peak p24\textsubscript{gag} level of the adapted viral quasispecies being $\sim2$ log higher than the G312V molecular clone. This indicated that the G312V amino acid change may only confer a portion of the full replication potential of the adapted viral quasispecies in Ptm cells. While no other dominant mutation was identified in the genome of the adapted virus that could readily explain these differences, the possibility that other compensatory mutations arose in the viral quasispecies over the 15-day period of the replication assays cannot be ruled out.

In a second experiment, replicating virus was found in 2 out of 30 cultures after approximately 32 days of culturing, with titers of $>10^5$ IP/mL (data not shown). Sequencing of the integrated proviral genome from cells in each of the two independent cultures harboring outgrowth virus revealed an identical adaptive mutation; a single C to A nucleotide change resulting in a predicted alanine to aspartic acid change at position 204 of gp120 (A204E, HXB2 numbering). Introduction of this same C to A nucleotide change into HIVA\textsubscript{Q23}/SIV\textsubscript{vif} resulted in a $>4$ log increase in p24\textsubscript{gag} levels as compared to the wild-type virus (Fig. 3b), which was similar to the levels of replication observed with the adapted viral quasispecies, thus indicating that the A204E amino acid change is responsible for the increased ability of the second adapted virus to infect Ptm cells.
Viruses bearing the envelope protein with the G312V and A204E amino acid changes were also tested for replication in primary Ptm PBMCs from two different donor animals. Peak replication was achieved at day 3 or 6 for the wild-type HIVA\textsubscript{Q23/SIV}\textsubscript{vif} and at day 6 for the G312V and A204E variants. The G312V change increased peak p24\textsubscript{gag} levels by approximately 4-fold compared to wild-type and the A204E p24\textsubscript{gag} levels increased by approximately 10-fold as compared to wild-type (Fig. 3c). The levels of replication of these variants were considerably lower (~10-100-fold) than the replication levels of the CXCR4-tropic minimal SHIV, NL-DT5R (not shown).

**Effect of amino acid changes at G312 and A204 on the ability of the Q23-17 Env to mediate entry into immortalized Ptm lymphocytes.**

To examine how the G312V and A204E changes were affecting entry into Ptm cells, reporter pseudoviruses carrying the Q23-17 Env with the G312V or A204E change were used to infect Ptm lymphocytes. Levels of entry were ~100-fold higher than wild-type for the Q23-17 G312V variant, and ~50-fold higher for the Q23-17 A204E variant. These levels were generally slightly higher than those of the SF162P3 Env, in the case of the G312V substitution; and similar to the SF162P3 Env, in the case of the A204E substitution (Fig. 4). Additionally, a double mutant was generated to determine whether there were any synergistic effects between the G312V and A204E substitutions. However, pseudoviruses bearing the Q23-17 double-mutant Env were not infectious in TZM-bl cells and so were not examined for entry into Ptm cells (data not shown).
The adaptive A204E and G312V changes that were observed in culture are rare among HIV-1 sequences, with only three examples of the G312V substitution and seven examples of the A204E substitution observed in the >10,000 HIV-1/SIVcpz Env sequences surveyed (http://www.hiv.lanl.gov). Furthermore, changes to G312 and A204 have not been reported in any infectious SHIVs constructed to date (data not shown).

The G312 residue, which is the first amino acid of the GPG(R/Q) motif located at the tip of the V3 loop (30), is conserved in >92% of Env sequences from all HIV-1 subtypes and >96% in subtype A Envs. The most common amino acid substitutions at G312 are alanine, arginine, and, specific to subtype A, histidine. The A204 residue, which is located adjacent to the β3 strand of the bridging sheet in the C2 region of gp120 (52), is conserved in >98% of all HIV-1 subtypes and is invariant in subtype A. The most common substitutions for A204 are serine and threonine.

A panel of mutants encoding the most common amino acid substitutions at G312 and A204 was made to examine how these substitutions compared to the adaptive G312V and A204E substitutions for their ability to mediate entry into Ptm cells. The titers of luciferase reporter pseudoviruses carrying the mutant Q23-17 Envs were comparable to those carrying the wild-type Q23-17 in HeLa-derived TZM-bl cells, with the exception of the G312R variant, whose limited infectivity precluded it from further study (data not shown). In Ptm cells, the level of entry of the G312A mutant was comparable whereas the G312H mutant had a >15-fold increased level of entry. Substitutions of serine and threonine at A204 had more modest outcomes; the A204S and A204T mutants
mediated entry into Ptm lymphocytes at levels that were only 3 to 5-fold greater than wild-type Q23-17 (Fig. 4).

To see if the A204E change was eliciting its effect due to the introduction of a negative charge, A204 was substituted with aspartic acid (A204D), which also bears a negative charge. Notably, the viruses carrying the A204D Env variants typically showed an approximately 100-fold decrease in infectivity on TZM-bl cells as compared to the wild-type Q23-17 Env (data not shown). However, when equal MOIs of the A204D virus were used to infect Ptm cells, luciferase levels were comparable to the A204E variant that was adapted in culture (Fig. 4). All together these results indicate that it is not the presence of a specific amino acid at position 204 (E) or G312 (V) that confer increased entry into Ptm lymphocytes; other amino acid changes at these positions can also impact infectivity in Ptm cells.

Effects of G312V and A204E amino acid changes in other subtype A Envs on entry and replication in Ptm cells

To examine whether the effects of the G312V and A204E changes were context-specific, the changes were introduced individually to each of the subtype A Envs tested in Fig. 2 and the mutated variants were assayed for their ability to mediate entry into Ptm lymphocytes. Most of the reporter viruses carrying the G312V and A204E Envs had infectious titers that were comparable to the parental wild-type in TZM-bl cells. This was not true, however for the Q259.d2.26 G312V and A204E variants, which were 10 to 20-fold less infectious than wild-type, and the Q259.d2.17 G312V and A204E variants,
whose low titers precluded them from use in the assay (data not shown). In all cases where the mutants retained infectivity in human cells, introduction of the G312V and A204E changes increased entry into Ptm cells compared to the wild-type Envs. These increases in entry ranged from a 10-fold increase, in the case of the Q259.d2.26 A204E variant, to a >100-fold increase, in the case of the MG505.H3 A204E variant (Fig. 5a). Despite the increase in entry, the G312V and A204E variants did not all mediate entry to the same degree as the SF162P3 Env, with levels of entry that were as much as 10-fold less, in the case of Q259.d2.26 (Fig. 5a).

To determine whether increases in entry predicted increased replication in Ptm lymphocytes, full-length molecular clones of HIVA/Q23/SIV_vif expressing the BG505.B1 or Q259.d2.26 Envs and their associated A204E and G312V variants were tested. Replication-competent viruses harboring the BG505.B1 Env and its variants were able to establish levels of infection reaching ~300 ng p24_gag/mL in human PBMCs, similar to those seen previously with HIVA/Q23/SIV_vif. The results in Ptm cells were also reminiscent of HIVA/Q23/SIV_vif, with viruses harboring the BG505.B1 G312V and A204E variants reaching peak p24_gag levels that were more than 3 logs greater than wild-type (Fig. 5b, left panel). However, the results for Q259.d2.26 and its G312V and A204E variants did not follow this trend. The G312V and A204E amino acid changes resulted in a 1.5 to 2 log reduction in p24_gag levels in human PBMCs as compared to the virus expressing the wild-type Q259.d2.d26 Env (Fig. 5b, right panel). These results were mirrored in Ptm lymphocytes, where the G312V and A204E changes did not appreciably
increase spreading infection compared to the wild-type virus, reaching maximum p24
gag levels of only 1 ng/mL.

Influence of the G312V and A204E amino acid changes on co-receptor usage

To establish if the G312V and A204E amino acid changes were exerting their effects by causing a change in co-receptor usage, Ptm lymphocytes were treated with saturating amounts of the CCR5 antagonist TAK779, and infected with luciferase pseudoviruses carrying the G312V and A204E Env variants. Infection mediated by the CCR5-tropic positive control SF162P3 was 100% inhibited by TAK779, and, as expected for a CXCR4-tropic Env, treatment with TAK779 had relatively little effect on infection mediated by the NL-DT5R Env (Table 1). Much like SF162P3, entry by the G312V and A204E variants was greatly inhibited (98 – 100%) by TAK779 (Table 1), thus indicating that Envs with the G312V and A204E changes continued to require CCR5 as a co-receptor for entry into Ptm lymphocytes.

Neutralization of the G312V and A204E variants by b12 and sCD4

The b12 monoclonal antibody and sCD4 were used to probe differences that the G312V and A204E variants may be causing in Env conformation and interaction with CD4. None of the viruses bearing the wild-type Envs were sensitive to neutralization by b12 (IC_{50} > 50 µg/mL), as has been observed previously (5, 64). The G312V and A204E amino acid changes had variable effects on neutralization by b12, depending on the viral context (Table 2). For example, the QF495.A3, Q259.d2.26, and MG505.H3 G312V and A204E Env variants were susceptible to neutralization, with IC_{50} values ranging from...
~0.6 µg/mL for the QF494.A3 A204E variant to ~8.6 µg/mL for the MG505.H3 G312V variant. Conversely, much like their wild-type counterparts, the G312V and A204E variants of Q23-17 and BG505.B1 were resistant to b12 neutralization.

The effects of the amino acid changes on sensitivity to sCD4 were more dramatic and consistent. The wild-type Envs were all relatively insensitive to sCD4, with IC$_{50}$s > 40 µg/mL. Introduction of the G312V and A204E changes rendered each Env highly susceptible to neutralization by sCD4, with IC$_{50}$ values ranging from ~8 µg/mL to < 0.2 µg/mL, the latter representing a greater than 200-fold increase in susceptibility compared to wild-type (Table 2). There did not appear to be a correlation between the sensitivities of the G312V and A204E variants to b12 and to sCD4. This was exemplified by the Q23-17 and BG505.B1 variants, which were insensitive to b12, but exquisitely sensitive to sCD4.

Use of Ptm CD4 and CCR5 by G312V and A204E variants

The G312V and A204E Env variants were next examined to see if they showed any differences in their ability to mediate entry using Ptm CD4 or CCR5. To do this, 293T cells were transiently transfected with CD4 and CCR5 expression plasmids in all possible combinations: huCD4/huCCR5, huCD4/ptmCCR5, ptmCD4/ptmCCR5, and ptmCD4/huCCR5 (Fig. 6a). In a given experiment, 20-50% of the cells were found to be double positive for CD4 and CCR5 expression, and the levels of expression for the receptors were similar across the four receptor combinations (Fig. 6a). The cells were infected with GFP reporter pseudoviruses carrying different Envs, and GFP-positive cells
were counted in triplicate wells, with the resulting titers being normalized to the huCD4/huCCR5 combination. Viruses bearing the SF162P3 Env, as well as those bearing the Env from SIV Mne CL8, a molecular clone that establishes persistent infection in Ptms (44), were used as positive controls. These positive control viruses were able to infect cells expressing any of the four combinations of CD4 and CCR5 to comparable levels (Fig. 6b).

Viruses carrying each of the 5 wild-type Envs infected cells expressing ptmCCR5 to similar levels as those expressing huCCR5, but showed a 10 to 30-fold decrease in infection in cells expressing ptmCD4 versus cells expressing huCD4 (Fig. 6b). The G312V and A204E Env variants were also unchanged in their ability to mediate infection using ptmCCR5 compared to huCCR5, but mediated increased levels of infection via ptmCD4 by as much as 50-fold, typically reaching the levels of infection that were comparable to those in cells expressing huCD4 (Fig. 6b). An exception to this was the MG505.H3 G312V variant, which increased ptmCD4-mediated entry by as little as 5-fold, still ~10-fold lower than entry mediated by huCD4. Overall, however, these findings suggested that the subtype A Envs used in this study were limited in their capacity to infect cells expressing ptmCD4 and that the G312V and A204E increased the efficiency with which the Envs are able to utilize ptmCD4.

**Expression of CD4 on immortalized Ptm lymphocytes**

To determine whether differences in the level of CD4 expression may contribute to differences in infectivity of the adapted HIVAQ23/SIVsif variants in Ptm versus human
lymphocytes, the expression of CD4 in the immortalized Ptm lymphocytes was compared to the expression of CD4 in primary Ptm PBMCs and in human PBMCs. Expression of CD4 was equivalent between all of the cells types (Fig 7a), suggesting that differences in infectivity cannot be explained by differences in cell surface CD4 levels. There are differences in amino acid sequence, between the human and Ptm CD4 molecules, including in regions that play a role in envelope binding, as shown in Fig. 7b.

Discussion

In this study, the limited ability of subtype A Envs to use ptmCD4 for entry was identified as a major barrier to the replication of SHIV-As in macaque cells. A minimal SHIV-A, HIVA_{Q23}/SIV_{vif}, replicated poorly in Ptm lymphocytes, but variants displaying increased replication in Ptm lymphocytes were selected after long-term culturing. Two independent adaptive amino acid changes in Env, G312V and A204E, conferred more efficient entry into Ptm cells when introduced into the Q23-17 parental Env. These same changes conferred high levels of replication in Ptm lymphocytes when introduced into the parental HIVA_{Q23}/SIV_{vif} proviral clone. Importantly, increased entry was also observed when either of these changes was independently introduced into multiple subtype A Envs, suggesting that these amino acid positions have a general impact on the interactions between subtype A Env and ptmCD4. Env variants encoding G312V and A204E maintained CCR5-tropism, but were much more sensitive to neutralization by sCD4 and mediated more efficient entry by ptmCD4. These findings implicate Env/CD4 interactions in the restriction of SHIV-A replication in macaque cells, and provide insight into specific amino acid positions in gp120 that can enhance these interactions.
Prior studies with subtype B minimal SHIVs suggested that the inclusion of the \textit{vif} gene from SIV\textsubscript{mac}239 in HIV\textsubscript{A\textsubscript{Q23}}/SIV\textsubscript{vif} would be sufficient for the virus to evade APOBEC3-mediated restriction, thus allowing for replication in Ptm cells (15). However, HIV\textsubscript{A\textsubscript{Q23}}/SIV\textsubscript{vif} replicated poorly in immortalized Ptm lymphocytes, in contrast to NL-DT5R, a CXCR4-tropic minimal SHIV that encodes both the \textit{vif} gene and the CypA-binding loop of SIV\textsubscript{mac}239 (24), which achieved high levels of replication. Similar results were observed when these SHIVs were used to infect primary Ptm lymphocytes (data not shown). The SIV CypA binding loop is required to evade Rhm TRIM5\textalpha, but should not be necessary for replication in Ptm lymphocytes because the capsid-directed TRIM5 isoforms and TRIMCyp expressed by Ptm do not restrict HIV-1 (6, 7, 15, 33, 62). Indeed, the lack of restriction to the HIV-1 capsid in Ptm cells was further verified here using chimeras between NL-DT5R and HIV\textsubscript{A\textsubscript{Q23}}/SIV\textsubscript{vif}, which showed that the HIV\textsubscript{A\textsubscript{Q23}}/SIV\textsubscript{vif} capsid supported replication in Ptm lymphocytes. Instead, the chimeras demonstrated that it was the 3' portion of HIV\textsubscript{A\textsubscript{Q23}}/SIV\textsubscript{vif}, including \textit{tat}, \textit{rev}, \textit{vpu}, and \textit{env}, that limited the replication of the SHIV-A in Ptm lymphocytes. These findings were consistent with previous studies using traditional SHIVs encoding the 3' elements of HIV-1, which implicated \textit{env} as the main determinant for infection of macaque cells (19). The identification of the Q23-17 Env as the reason for reduced replication in Ptm cells was definitively shown when mutations encoding the G312V or A204E amino acid changes to gp120, which were identified in viruses selected for increased replication in immortalized Ptm lymphocytes, were introduced into the parental clone and the resulting viruses replicated in immortalized Ptm lymphocytes to levels that were comparable to
Viruses bearing the G312V or A204E mutations also showed increased replication compared to the parental virus in primary Ptm PBMCs, but the differences were much more modest than those observed in immortalized CD4 lymphocytes, perhaps reflecting the fact that CD4/CCR5 positive lymphocytes comprise only a fraction of the PBMCs.

The increases in entry into Ptm lymphocytes by the G312V and A204E variants in luciferase reporter assays were not necessarily directly correlated with increases in virus spread, as exemplified by the G312V change in the context of Q23-17 and BG505.B1. The G312V changes conferred similar increases in entry in the single-cycle reporter assay as the A204E mutants, yet in the viral replication assays, peak replication levels were much lower for a full-length clone encoding the Q23-17 G312V Env as compared to one encoding the BG505.B1 G312V Env. The same immortalized Ptm lymphocytes were used for both the single cycle and replication assays, ruling out the possibility of differences in cell targets. This suggests that differences in infectivity may be due to differences in how the viruses were generated. Depending on the Env being expressed, pseudoviruses have been shown to express artificially high amounts of Env, both in the processed and unprocessed forms (18), and this can result in higher levels of infectivity compared to replication competent virus (49).

The G312 residue, located at the tip of the V3 loop, and the A204 residue, located adjacent to the β3 strand of the bridging sheet, are in regions of gp120 that are known to mediate interaction with the co-receptor (22, 52). However, the increased ability of the
G312V and A204E variants to infect Ptm cells was not due to a change in co-receptor use or an increased ability to mediate infection by ptmCCR5. Surprisingly, subtype A Envs were deficient in their ability to mediate infection of cells expressing ptmCD4, and the G312V and A204E amino acid changes rescued this deficiency. Flow cytometric analyses demonstrated that the differences in infectivity could not be explained by differences in cell-surface CD4 expression. These results argue that the barrier to HIVαQ23/SIV_vif replication in Ptm lymphocytes was due to inefficient use of ptmCD4, and that the G312V and A204E changes arose to circumvent this barrier.

A notable property of the G312V and A204E variants was their increased sensitivity to sCD4, with both mutations conferring increased sensitivity to sCD4 in all Envs tested, in most cases, by more than 100-fold. This finding suggests that sensitivity to sCD4 may predict how well a particular envelope uses ptmCD4 for entry. If this is the case, then most circulating HIV-1 Env variants, which tend to be relatively insensitive to sCD4 [e.g. (5, 32)], may not promote efficient entry via ptmCD4. However, these findings also raise the interesting possibility that sensitivity to sCD4 may provide a means to identify representative transmitted Env variants that would be the best candidates for a successful SHIV. In support of this hypothesis, it is significant that many of the existing CCR5-tropic SHIVs incorporate HIV-1 Envs that are relatively sensitive to soluble CD4, including the SF162 lineage (10, 27), ADA (60), BaL (12) and the subtype C isolate HIV2873i (56).
One model that may explain how changes to G312 and A204 are eliciting their effect is by increasing exposure of the CD4-binding site, which could then allow for better usage of ptmCD4. G312 is found in the V3 loop, which has been shown to be a determinant for increased sensitivity to sCD4 (23, 42, 59), implying that changes to the V3 loop may participate in quaternary interactions in Env that could lead to a more exposed CD4-binding site. A204 is adjacent to the highly conserved C205 residue, whose replacement has been shown to increase the susceptibility of the HIV-1 Env to sCD4, presumably by abrogating a highly conserved disulfide bond, resulting in a more open Env conformation (61). It is possible that changes to A204, particularly the introduction of a negative charge as in the A204E and A204D variants, may also be serving to modulate this disulfide bond or may be opening up the Env structure via other steric interactions. The increased exposure of the CD4-binding site in the G312V and A204E variants is further supported by the sensitivity that some of the variants display to the b12 monoclonal antibody, whose epitope overlaps the CD4-binding site (8, 66). The high degree of conservation of the G312 and A204 residues may indicate that they play some role in maintaining the structural integrity of the envelope trimer. This may explain why the G312V/A204E double mutant, and, in some contexts the single G312V or A204E changes (for example, in the case of the Q259.D2.17 Env) result in an envelope that does not support efficient entry or viral replication.

Little is known regarding how differences in macaque and human CD4 impact the ability of HIV-1 Envs to mediate infection of macaque cells, although one study has concluded that ptmCD4 was not a barrier to HIV-1 infection (13).
of HIV-1 was used in this previous study, and these results are consistent with observations that CXCR4-tropic subtype B SHIVs are infectious in macaque cells. The differences between this study and the data presented here most likely reflect differences in the biology of lab-adapted strains compared to CCR5-tropic variants cloned directly from infected individuals, and serves as a reminder of the difficulty of extrapolating findings with lab-adapted HIV variants to more relevant, circulating strains of HIV-1.

The primary determinants of HIV Env-CD4 interaction have been mapped to the D1 and D2 domains of human CD4, and human and ptmCD4 differ at 17 amino acid positions in these domains. Structural studies indicate that F43 and R58 from CD4 form contacts with gp120 (29). There is an amino acid change at R58 (K in Ptm) that could alter Ptm CD4-Env interactions, but it is a conservative amino acid difference. There are other, non-conservative amino acid differences at S23 (N in Ptm) and N52 (S in Ptm). Although these residues have not been recognized as contact residues in structural analyses, they have been identified in mutagenesis studies as being critical for gp120 binding, (1). Thus, disruption of CD4-gp120 binding, either by modulation of direct amino acid interactions, or indirectly, though effects on binding affinity, may explain the differences in HIV-mediated entry between human and ptmCD4 observed here.

Prior studies have shown that SHIV-As were not infectious in Rhm cells (19), and our findings may provide an explanation for these results. The published sequence of Rhm CD4 is identical to ptmCD4 at each of the four residues noted above (data not shown). Further studies are needed to determine whether Rhm CD4 presents a similar barrier to
infection as ptmCD4 and whether amino acid changes at positions A204 and G312, which are at highly conserved across all subtypes, can overcome this barrier. Such studies may also help define the precise changes that alter HIV Env/macaque CD4 interactions.

The usefulness of SHIV models, particularly as tools to examine the biology of HIV-1 transmission and strategies to prevent infection, will depend on how well they mimic HIV-1 transmission and early infection in humans. There are numerous barriers to HIV-1 infection in macaques, and SHIV proviruses tested to date have required further adaptation in the animal to increase replication. The improved understanding of host restriction factors has permitted more targeted approaches to developing infectious SHIVs although the initial chimeric viruses did not replicate to high levels in infected animals (15, 16, 24). Here, we identify differences in CD4 as a barrier to HIV-1 infection of pig-tailed macaque cells, and show that a single amino acid change in Env is sufficient to surmount this limitation, at least for subtype A Envs. These data indicate it may be possible to develop SHIVs that are derived from more relevant HIV-1 variants with only minor modifications. However, in using these findings to develop more relevant SHIV models, it will be important to consider how changes that permit efficient CD4 interaction impact other key biological properties of the envelope, such as sensitivity to neutralization. Identifying inefficient use of CD4 as a barrier to HIV-1 infection of macaque cells provides a critical first step in the process of designing SHIVs based on biologically relevant HIV-1 variants.
Acknowledgements

We would to thank like Drs. Catherine Blish, Dara Lehman, Erica Lovelace, and Shiu-Lok Hu for many helpful discussions; Dr. Jason Kimata for comments on the manuscript; and Stephanie Rainwater for constructing Q23Δenv-GFP. This study was supported by NIH grants AI38518 and R01 HD058304 to JO.

References

   Sattentau, P. R. Clapham, R. A. Weiss, J. S. McDougal, and C. Pietropaolo.
   1989. Identification of the residues in human CD4 critical for the binding of HIV.
   Cell. 57:469-481.


3. Benki, S., R. S. McClelland, S. Emery, J. M. Baeten, B. A. Richardson, L.
   Lavreys, K. Mandaliya, and J. Overbaugh. 2006. Quantification of genital
   human immunodeficiency virus type 1 (HIV-1) DNA in specimens from women
   with low plasma HIV-1 RNA levels typical of HIV-1 nontransmitters. J. Clin.

   monoclonal antibody resistance among early subtype A, C, and D HIV-1

   2007. HIV-1 subtype A envelope variants from early in infection have variable
   sensitivity to neutralization and to inhibitors of viral entry. AIDS. 21:693-702.
   doi: 10.1097/QAD.0b013e32805e8727.


   neutralization of primary isolates of HIV-1 by a recombinant human monoclonal


54. Seaman, M. S., H. Janes, N. Hawkins, L. E. Grandpre, C. Devoy, A. Giri, R. T. Coffey, L. Harris, B. Wood, M. G. Daniels, T. Bhattacharya, A. Lapedes,


**Tables**

**TABLE 1. Inhibition of G312V and A204E variants by TAK779**

<table>
<thead>
<tr>
<th>Clone</th>
<th>Residue Change</th>
<th>Percent Inhibition*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q23-17</td>
<td>G312V</td>
<td>100 ± 0</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>98 ± 1</td>
</tr>
<tr>
<td>QF495.A3</td>
<td>G312V</td>
<td>99 ± 1</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>MG505.H3</td>
<td>G312V</td>
<td>100 ± 0</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>BG505.B1</td>
<td>G312V</td>
<td>100 ± 0</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>99 ± 0</td>
</tr>
<tr>
<td>Q259.D2.26</td>
<td>G312V</td>
<td>100 ± 0</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>SF162P3</td>
<td>--</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>NL-DT5R</td>
<td>--</td>
<td>15 ± 21</td>
</tr>
</tbody>
</table>

*Percent inhibition of luciferase activity in cells treated with 1uM TAK779 compared to untreated cells (avg ± SD for two independent experiments).
TABLE 2. Neutralization of G312V and A204E variants by b12 and sCD4

<table>
<thead>
<tr>
<th>Envelope Clone</th>
<th>Residue Change</th>
<th>IC50 (µg/mL) b12</th>
<th>IC50 (µg/mL) sCD4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q23-17</td>
<td>--</td>
<td>&gt;50 b</td>
<td>&gt;50 a</td>
</tr>
<tr>
<td></td>
<td>G312V</td>
<td>&gt;50 b</td>
<td>0.8 ± 0.7</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>&gt;50 b</td>
<td>0.3 ± 0.2</td>
</tr>
<tr>
<td>QF495.A3</td>
<td>--</td>
<td>&gt;50 b</td>
<td>48 ± 2.8 d</td>
</tr>
<tr>
<td></td>
<td>G312V</td>
<td>1.4 ± 0.1</td>
<td>&lt;0.2 c</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>0.6 ± 0.0</td>
<td>&lt;0.2 c</td>
</tr>
<tr>
<td>MG505.H3</td>
<td>--</td>
<td>&gt;50 b</td>
<td>&gt;50 a</td>
</tr>
<tr>
<td></td>
<td>G312V</td>
<td>8.6 ± 0.4</td>
<td>6.9 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>1.5 ± 0.4</td>
<td>0.7 ± 0.0</td>
</tr>
<tr>
<td>BG505.B1</td>
<td>--</td>
<td>&gt;50 b</td>
<td>&gt;50 a</td>
</tr>
<tr>
<td></td>
<td>G312V</td>
<td>&gt;50 b</td>
<td>&lt;0.2 c</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>&gt;50 b</td>
<td>0.3 ± 0.2</td>
</tr>
<tr>
<td>Q259.D2.26</td>
<td>--</td>
<td>&gt;50 b</td>
<td>47 ± 4.5 d</td>
</tr>
<tr>
<td></td>
<td>G312V</td>
<td>2.0 ± 0.6</td>
<td>0.2 ± 0.1 e</td>
</tr>
<tr>
<td></td>
<td>A204E</td>
<td>1.8 ± 0.5</td>
<td>&lt;0.2 e</td>
</tr>
</tbody>
</table>

aExpressed as avg ± SD for two experiments unless otherwise noted
bIC50s from duplicate experiments were both >50
cIC50 value of <0.2 from one experiment was set to 0.1 and avg ± SD was reported
dIC50 value of >50 from one experiment was set to 50 and avg ± SD was reported
eIC50s from duplicate experiments were both <0.2.
Figure Legends

FIG 1. Infection of human PBMCs and Ptm lymphocytes with minimal SHIVs. (a) Schematic representation of the contribution of Q23-17, NL4-3 and SIVmac239 sequences to the minimal SHIVs used for replication experiments. E and X represent the EcoRI and XhoI sites used to make N/Q\textsubscript{vpr-nef} and Q/N\textsubscript{vpr-nef}. (b) p24\textsuperscript{gag} levels are shown as a function of time post-infection in human PBMCs and Ptm lymphocytes. The data points represent the average measurement from duplicate infected cultures. The figure key is shown in the top plot, and viruses with an asterisk were used at a 10-fold greater MOI for infection of Ptm cells. The results are representative of at least three independent experiments.

FIG 2. Single-cycle infection of Ptm lymphocytes with luciferase pseudoviruses bearing subtype A Envs. The y-axis shows relative light units (RLUs) in cells infected with the virus indicated on the x-axis. The dashed line represents background RLU levels for the experiment observed in cells infected with Env(-) pseudovirus. Error bars represent the standard deviation obtained from triplicate wells. The results are representative of at least three independent experiments.

FIG 3. Infection of human PBMCs and immortalized Ptm lymphocytes with HIV\textsubscript{Q23}/SIV\textsubscript{vif} carrying the G312V change (a) or the A204E change (b). The p24\textsuperscript{gag} levels are shown as a function of time post-infection in human PBMCs (top graphs) and Ptm lymphocytes (bottom graphs). The data points represent the average measurement from duplicate cultures. The figure key is shown in the top plots, and ‘Adapt. Sup.’ refers to the uncloned adapted viral quasispecies obtained by long-term culturing from which the mutations were isolated. The results are representative of at least three independent experiments. (c) Infection of primary Ptm PBMCs from two different donors with parental HIV\textsubscript{Q23}/SIV\textsubscript{vif} (light grey diamonds), HIV\textsubscript{Q23}/SIV\textsubscript{vif} G312V (dark grey triangles), or HIV\textsubscript{Q23}/SIV\textsubscript{vif} A204E (black diamonds). The p24\textsuperscript{gag} levels are shown as a function of time post-infection and data points represent the average measurement from duplicate cultures.

FIG 4. Single-cycle infection of Ptm lymphocytes with luciferase pseudoviruses bearing Q23-17 Env mutants. The y-axis shows relative light units (RLUs) in cells infected with the virus variant indicated on the x-axis. The dashed line represents background RLU levels for the experiment observed in cells infected with Env(-) pseudovirus. Error bars represent the standard deviation obtained from triplicate wells. ‘wt’ refers to the wild-type Q23-17 envelope, and ‘ND’ is not determined due to insufficient titer. The results are representative of at least two independent experiments.

FIG 5. (a) Single-cycle infection of Ptm lymphocytes with luciferase pseudoviruses bearing G312V and A204E Env variants. The y-axis shows relative light units (RLUs) in cells infected with the virus indicated on the x-axis. The dashed line represents background RLU levels for the experiment observed in cells infected with Env(-) pseudovirus. Error bars represent the standard deviation obtained from
triplicate wells. (b) Infection of human PBMCs and Ptm lymphocytes with HIVA Q23/SIV vif expressing BG505.B1 (left panel) and Q259.D2.26 (right panel) Env variants. The p24 levels are shown as a function of time post-infection in human PBMCs (top graphs) and Ptm lymphocytes (bottom graphs). The data points represent the average measurement from duplicate cultures. The figure key is shown in the top plots. ‘wt’ refers to the wild-type Env to which the G312V or A204E change was introduced. Both (a) and (b) are representative of three independent experiments.

FIG 6. (a) Transient expression of human and Ptm CD4 and CCR5 in 293T cells. The y-axis represent CD4 expression and the x-axis represents CCR5 expression as determined by flow cytometry. The CD4/CCR5 combination is denoted at the top of each plot and ‘pBabe-puro’ denotes cells transfected with the empty pBabe-puro vector. These plots are representative of three replicate experiments. (b) Single-cycle infection of 293T cells expressing human versus Ptm CD4 and CCR5 with GFP reporter pseudoviruses bearing wild-type subtype A Env and G312V and A204E variants. The y-axis represents viral infection relative to the huCD4/huCCR5 combination, which is set to 1 for reference. The name of the parental virus is indicated at the top of each plot, and the amino acid change is indicated on the x-axis (‘wt’ refers to the wild-type Env variant). In general 20-300 GFP-positive cells were enumerated per well, depending on the dilution, the virus and the CD4/CCR5 pair. No GFP-positive cells were observed when cells were infected with Env(-) pseudoviruses (not shown). Three replicates were performed, and each replicate included all of the receptor pairings with the exception of one in which the ptmCD4/huCCR5 pair was not included. The final replicate for the ptmCD4/huCCR5 pair was later performed in a separate experiment. The error bars represent the standard deviation of the mean from all of the data amassed in the replicate experiments.

FIG 7. (a) CD4 expression on human PBMCs, Ptm PBMCs and immortalized Ptm lymphocytes. Live cells were gated on CD4 expression and CD4-positive cells are shown in the histogram. Shaded curves represent unstained cells and, in the PBMC expression profiles, the black and grey curves represent profiles of cells from two different donors. (b) Amino acid alignment of the D1 and D2 domains of human and Ptm CD4. Triangles above the alignment denote the beginning residue of the D1, D2, or D3 (not labelled) immunoglobulin-like domains. Dots in the Ptm sequence indicate conserved residues, and in positions where the Ptm and human sequence differ, the corresponding residue encoded by Ptms is shown. The residues that are highlighted in grey have previously been found to be important for gp120 binding (1).
Fig. 1

(a) Diagram showing the comparison of different strains of HIV-1, with markers for NL-DT5R, HNA_{env}^{SN_{ef}}, NQ_{env}^{SN_{ef}} and QN_{env}^{prof}

(b) Graph showing the p24 levels in Human PBMCs and Ptm Lymphocytes post-infection. Different strains are indicated with markers for NL-DT5R, HNA_{env}^{SN_{ef}}, NQ_{env}^{SN_{ef}} and QN_{env}^{prof}.
Fig. 2
Fig. 3

a) G312V

- Human PBMCs
  - Adapt. Sup. 1
  - H0A33S
  - H0A33S
  - G312V

- Ptm Lymphocytes
  - Adapt. Sup. 1
  - H0A33S
  - H0A33S

b) A204E

- Human PBMCs
  - Adapt. Sup. 2A
  - H0A33S
  - H0A33S

- Ptm Lymphocytes
  - Adapt. Sup. 2B
  - H0A33S
  - H0A33S

C

- Ptm PBMCs Donor #1
  - Adapt. Sup. 1
  - H0A33S

- Ptm PBMCs Donor #2
  - Adapt. Sup. 2B
  - H0A33S
Fig. 4
Fig 5.
Fig 6.
Fig 7.

![Diagram showing CD4-APC fluorescence in different cell types: Human PBMCs, Ptm PBMCs, Ptm lymphocytes.](image)

### b

**D1**

<table>
<thead>
<tr>
<th>Human</th>
<th>Pigtail</th>
</tr>
</thead>
<tbody>
<tr>
<td>KKVVLGKGGDTVELCTASQKSI</td>
<td>N</td>
</tr>
</tbody>
</table>

**D2**

<table>
<thead>
<tr>
<th>Human</th>
<th>Pigtail</th>
</tr>
</thead>
<tbody>
<tr>
<td>LMDQGFLKIKNLDTSYICVEDEQKEKQLV</td>
<td>C’sM</td>
</tr>
</tbody>
</table>

**D3**

<table>
<thead>
<tr>
<th>Human</th>
<th>Pigtail</th>
</tr>
</thead>
<tbody>
<tr>
<td>PGSSPSVTQCRGRPNQGQRSQSLQDSQ</td>
<td>K’G’R’P’R’S’D’T</td>
</tr>
</tbody>
</table>