Natural cystatin C fragments inhibit GPR15-mediated HIV and SIV infection without interfering with GPR15L signaling

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GPR15 is a G protein-coupled receptor (GPCR) proposed to play a role in mucosal immunity that also serves as a major entry cofactor for HIV-2 and simian immunodeficiency virus (SIV). To discover novel endogenous GPR15 ligands, we screened a hemofiltrate (HF)-derived peptide library for inhibitors of GPR15-mediated SIV infection. Our approach identified a C-terminal fragment of cystatin C (CysC95-146) that specifically inhibits GPR15-dependent HIV-1, HIV-2, and SIV infection. In contrast, GPR15L, the chemokine ligand of GPR15, failed to inhibit virus infection. We found that cystatin C fragments preventing GPR15-mediated viral entry do not interfere with GPR15L signaling and are generated by proteases activated at sites of inflammation. The antiretroviral activity of CysC95-146 was confirmed in primary CD4+ T cells and is conserved in simian hosts of SIV infection. Thus, we identified a potent endogenous inhibitor of GPR15-mediated HIV and SIV infection that does not interfere with the physiological function of this GPCR.

G protein-coupled receptors | GPR15 | immunodeficiency viruses | chemokines | cystatin C

G protein-coupled receptors (GPCRs) constitute the largest family of membrane proteins involved in the transduction of signals from the extracellular environment into the cell and play key roles in immune responses, homeostasis, metabolism, and organogenesis (1, 2). Besides their physiological roles, some GPCRs also represent important coreceptors for HIV and/or simian immunodeficiency virus (SIV) entry. HIV-1, the main causative agent of AIDS, utilizes C-C chemokine receptor 5 (CCR5) and CXC chemokine receptor 4 (CXCR4) as major entry cofactors (3–8). The chemokine ligands CCL5 (also known as RANTES) and CXCL12 (also named SDF-1) inhibit CCR5- or CXCR4-mediated HIV-1 infection, respectively. Thus, we have previously taken advantage of HIV-1 entry to examine complex blood-derived peptide libraries for novel naturally occurring ligands of CCR5 and CXCR4 (9, 10). Initially, we identified a truncated form of the chemokine C-C ligand 14 (CCL14) as a CCR5 agonist and potent inhibitor of CCR5-tropic HIV-1 strains (11, 12). More recently, we discovered a small fragment of human serum albumin (EPI-X4) as an effective and specific inhibitor of CXCR4-mediated HIV and SIV infection (13). While HIV-1 coreceptor utilization is mainly restricted to CCR5 and CXCR4, HIV-2 and SIVs are more promiscuous in their entry cofactor usage. For example, many HIV-2 and SIV strains utilize BOB/GPR15 and Bonzo/STRL-33/CXCR6 in addition to CCR5 and/or CXCR4 for viral entry into CD4+ target cells (14–20). GPR15 is a GPCR reported to regulate T cell trafficking to the colon that may play a role in intestinal homeostasis and inflammation (21, 22). Recently, an agonistic C-C chemokine ligand of GPR15, named GPR15L, has been characterized (23, 24). GPR15L is expressed in colon and cervical epithelia and might play a role in mucosal immunity.

To discover novel endogenous GPR15 ligands, we screened a hemofiltrate (HF)-derived peptide library containing essentially all peptides and small proteins circulating in human blood in their final processed and physiologically relevant forms (10) for inhibitors of GPR15-mediated SIV infection. Multiple rounds of peptide separation and antiviral screening identified a C-terminal fragment of cystatin C (named CysC95-146) as a potent and specific inhibitor of GPR15-dependent HIV and SIV infection. Cystatin C is a small (13 kDa) basic protein that is produced by all nucleated cells (25) and represents the most abundant and potent inhibitor of GPR15-dependent HIV and SIV infection. In contrast, GPR15L, the chemokine ligand of GPR15, failed to inhibit virus infection. We found that cystatin C fragments preventing GPR15-mediated viral entry do not interfere with GPR15L signaling and are generated by proteases activated at sites of inflammation. The antiretroviral activity of CysC95-146 was confirmed in primary CD4+ T cells and is conserved in simian hosts of SIV infection. Thus, we identified a potent endogenous inhibitor of GPR15-mediated HIV and SIV infection that does not interfere with the physiological function of this GPCR.

Significance

G protein-coupled receptors (GPCRs) are involved in many physiological processes and important drug targets. However, most therapeutic agents targeting GPCRs, such as the coreceptors of HIV-1 CCR5 and CXCR4, also interfere with their signaling function. Here, we used primate lentiviruses as tools to discover novel endogenous ligands of GPR15, a coreceptor for HIV-2 and SIV proposed to play a role in mucosal immunity. We found that C-terminal fragments of cystatin C generated by immune-activated proteases inhibit GPR15-mediated HIV and SIV infection without interfering with GPR15L chemokine signaling. Thus, we identified an endogenous bioactive peptide that specifically prevents the detrimental activity of a GPCR (i.e., virus infection) without compromising its physiological signaling function.

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extracellular inhibitor of cysteine proteases (26). It is found in virtually all tissues and body fluids and commonly used as a marker of renal function (27). Accumulating evidence suggests a role of cystatin C in inflammation, neutrophil chemotaxis, and resistance to bacterial as well as viral infections (28–31). We show that cystatin C fragments preventing GPR15-dependent HIV and SIV infection are generated by proteases activated during antiviral immune responses. Unexpectedly, the recently discovered chemokine ligand of GPR15, GPR15L (23, 24), had no significant effect on viral entry. In addition, CysC95-146 prevented SIV and HIV-2 infection without interfering with GPR15L-mediated signaling. Our data support that naturally occurring cystatin C fragments are capable of blocking GPR15-mediated primate lentiviral infection without interfering with the physiological signaling function of this GPCR.

Results

Identification of a GPR15-Specific SIV Inhibitor. To identify novel ligands of GPR15, peptide libraries were generated from up to 10,000 L of hemofiltrate derived from individuals with chronic renal failure by cation exchange separation followed by reverse-phase (RP) chromatography (10, 32). Initially, the hemofiltrate was applied to a large cation exchange column and eluted using eight buffers with increasing pH ranging from 2.5 to 9.0. Subsequently, the resulting pH pools (P1 to P8) were separated into ∼40 to 50 peptide-containing fractions (F1 to F50) by RP chromatography. The final library comprised about 360 peptide fractions representing essentially the entire blood peptidome in a highly concentrated, salt-free, and bioactive form. Screening of this peptide library using the infectious SIVmac239 molecular clones (IMCs) of SIVsmm (Fig. 2A) and human osteosarcoma (GHOST) cells stably expressing CD4 and GPR15 (14) identified several neighboring fractions from pH pool 4 that blocked GPR15-mediated cytopathic activity (41, 42); and 7312A, originally isolated from an individual from Côte d’Ivoire with dual HIV-1 and HIV-2 infection, suggesting that it is specific for GPR15 (Fig. 2B) and did not display significant cytotoxic effects (Fig. 2C).

SIVsmm is the precursor of HIV-2 (37), which is endemic in West Africa and has infected about 1 to 2 million people (38). To evaluate a possible role of CysC95-146 in HIV-2 infection, we examined its effect on infection by three HIV-2 strains: ROD10, representing a derivative of the first reported infectious HIV-2 clone (39, 40); ST, an attenuated HIV-2 strain with low in vitro cytopathic activity (41, 42); and 7312A, originally isolated from an individual from Côte d’Ivoire with dual HIV-1 and HIV-2 infection (43). HIV-2 ROD10 and ST belong to the group A of HIV-2 that is most widespread in the human population, while 7312A represents a recombinant form of groups A and B (44). CysC95-146 inhibited all HIV-2 strains in a dose-dependent manner with IC50 values ranging from 1.3 to 7.0 μM (Fig. 2D).

A Natural Cystatin C Fragment Inhibits GPR15-Dependent SIV and HIV Infection. To verify that the identified peptide is responsible for the antiviral activity, we chemically synthesized the S2 C-terminal amino acid residues of cystatin C (indicated in Fig. 1C). The synthetic peptide, referred to as CysC95-146, inhibited GPR15-mediated SIVmac239 infection in a dose-dependent manner with a mean 50% inhibitory concentration (IC50) of ~0.5 μM (Fig. 2A). Potent inhibition was confirmed for three divergent infectious molecular clones (IMCs) of SIVsmm (Fig. 2B and SI Appendix, Fig. S1), which naturally infects sooty mangabeys (36) and frequently utilizes GPR15 as an entry cofactor (20). CysC95-146 had little if any effect on CCR5- or CXCR6-mediated virus infection, suggesting that it is specific for GPR15 (Fig. 2B) and did not display significant cytotoxic effects (Fig. 2C).

Fig. 1. Identification of a C-Terminal cystatin C fragment inhibiting GPR15-mediated SIVmac infection. (A) The gray bars indicate the efficiency of SIVmac239 infection of GHOST-GPR15 cells in the presence of the hemofiltrate peptide library fractions compared to the absence of peptide (100%), and the black line indicates the peptide/protein elution profile. Fractions used for further purification are indicated in red and highlighted by an arrow. + indicates the absence in the peptide fraction. – shows uninfected cells. (B) MALDI-TOF spectrum of the active fraction obtained after the fifth round of purification. (C) Amino acid sequence of human cystatin C. The signal peptide (green), the isolated peptide (red), and putative C-C bridges are indicated. The cleavage site to generate CysC95-146 is indicated by a red arrow.
CysC95-146 specifically inhibits GPR15-mediated SIV and HIV infection. (A) GHOST-GPR15 cells were infected with a SIVmac239 luciferase reporter construct in the presence of CysC95-146. Experiments shown in all panels were performed at least in triplicates and curves show mean values ± SEM. (B) GHOST cells engineered to express GPR15, CXCR6, or CCR5 were infected with different SIV strains. Values show the percentage of virally infected (GFP+) cells in the presence of increasing concentrations of CysC95-146 compared to the percentage of infected cells obtained in the absence of peptide (100%). The dotted line indicates the percentage of eGFP+ cells obtained after infection of the parental GHOST cell line in the absence of peptide. (C) CysC95-146 is not cytotoxic. GHOST-GPR15 seeded in 96-well F-bottom plates were incubated with increasing amounts of peptide for 3 d at 37 °C. Metabolic activity was analyzed by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) and CellTiter-Glo assay. (D and E) Inhibition of (D) the indicated HIV-2 molecular clones or (E) HIV-1 ZP6248 by CysC95-146. In E, SIVmac239 is shown as positive control for comparison. Experiments were performed as described in B.

However, maximal inhibition of HIV-2 entry did not exceed ~80%. One reason for this might be GPR15-independent infection of GHOST cells. Indeed, HIV-2 ROD10 infected the parental GHOST cell line that expresses low levels of CXCR4 but not GPR15 or other known entry cofactors with significant efficiency (Fig. 2D and SI Appendix, Fig. S1).

HIV-1 is less promiscuous in coreceptor usage than HIV-2. It usually utilizes only CCR5 during chronic infection and frequently involves the ability to use CXCR4 during or after AIDS progression (45, 46). Some HIV-1 isolates, however, may also utilize GPR15 for productive infection and replication especially at high expression levels (19, 47, 48). Interestingly, it has been reported that the transmitted/founder HIV-1 IMC ZP6248 is severely impaired in CCR5 and CXCR4 coreceptor usage but capable of infecting cell lines expressing alternative coreceptors including GPR15 (48). CysC95-146 inhibited infection by the HIV-1 ZP6248 molecular clone in GPR15-GHOST cells with an IC50 of 1.1 μM (Fig. 2E). Altogether, our results showed that CysC95-146 specifically inhibits GPR15-dependent SIV and HIV infection.

The Inhibitory CysC95-146 Fragment Binds to GPR15-Expressing Cells. To examine whether CysC95-146 specifically targets GPR15-expressing cells, we generated N- and C-terminal fusions of this GPCR with the enhanced green fluorescent protein (eGFP). N-terminal tagging with eGFP resulted in mislocalization of the parental GHOST cell line that expresses low levels of CXC4 but not GPR15 or other known entry cofactors with significant efficiency (Fig. 2D and SI Appendix, Fig. S1). In contrast, N-terminal truncations of CysC95-146 by up to 12-aa residues (CysC107-146) (Fig. 4A) did not disrupt its antiviral activity. However, a peptide containing an additional 10 residues at its N terminus (CysC85-146) (Fig. 4B) displayed little inhibitory effect on SIVmac239 entry (Fig. 4A). Further truncations at the N or C terminus resulted in reduced activity or fully disrupted inhibition of GPR15-mediated SIVmac239 infection. Notably, some of the cystatin C fragments analyzed, such as CysC107-146, were even more potent than CysC95-146 in inhibiting GPR15-mediated SIVmac239 infection (Fig. 4C). Altogether, the results showed that a large variety of C-terminal cystatin C fragments prevent GPR15 at the cell surface.

A Variety of C-Terminal CysC Fragments Prevent GPR15-Mediated Lentiviral Infection. To further examine the specificity of the antiviral activity of CysC95-146, we determined whether full-length cystatin C and other C-terminal fragments also affect primate lentiviral infection. Our results showed that full-length cystatin C displayed little inhibitory effect on SIVmac239 entry (Fig. 4A). In contrast, N-terminal truncations of CysC95-146 by up to 12-aa residues (CysC107-146) (Fig. 4A), as well as expansion by up to 6 residues (89 to 146) (Fig. 4B) did not disrupt its antiviral activity. However, a peptide containing an additional 10 residues at its N terminus (CysC85-146) did not inhibit SIVmac entry into GPR15-GHOST cells (Fig. 4B). Comprehensive analyses of a variety of C-terminal cystatin C fragments revealed that residues 107 to 140 are sufficient for antiviral activity (Fig. 4C). Further truncations at the N or C termini resulted in reduced activity or fully disrupted inhibition of GPR15-mediated SIVmac239 infection. Notably, some of the cystatin C fragments analyzed, such as CysC107-146, were even more potent than CysC95-146 in inhibiting GPR15-mediated SIVmac239 infection (Fig. 4C). Altogether, the results showed that a large variety of C-terminal cystatin C fragments prevent GPR15-mediated SIVmac infection.

Proteolytic Generation of Antiviral CysC Fragments. CysC95-146 was isolated from a hemofiltrate-derived peptide library, suggesting that it naturally circulates in the blood stream. To further
As well as trypsin, chymase, and napsin A (Fig. 5 C, we treated the full-length protein with cathepsins C, D, and G, fragments that can be achieved in vivo.

To examine this, we developed an MS-based method for the quantification of CysC95-146 in hemofiltrate (SI Appendix, Fig. S3A). These analyses showed that CysC95-146 was present at concentrations of \( \sim 10.7 \text{ ng/mL} \) (236 pmol) in the original hemofiltrate fraction (SI Appendix, Fig. S3 B and C). This concentration is lower than the IC\(_{50}\). However, hemofiltrate is significantly diluted compared to blood. In addition, material may have been lost during sample preparation and other C-terminal fragments are also antivirally active (Fig. 4). In addition, cystatin C levels are elevated in HIV-infected patients compared to healthy individuals (49, 50). Thus, our measurements most likely underestimate the concentrations of antiviral C-terminal cystatin C fragments that can be achieved in vivo.

To examine the generation of antiviral peptides from cystatin C, we treated the full-length protein with cathepsins C, D, and G, as well as trypsin, chymase, and napsin A (Fig. 5A). These proteases represent major components of the endo- and lysosomal protein degradation machinery (26, 51) and some of them are efficiently released from immune cells during infectious and inflammatory processes (52, 53). Treatment of cystatin C with cathepsin D, trypsin, pepsin, chymase, and napsin A resulted in the generation of peptides with sizes similar to CysC95-146 (Fig. 5A). Products obtained after digestion with cathepsin D, chymase, and napsin A significantly inhibited GPR15-mediated SIVmac infection (Fig. 5B). In addition, mass spectrometry of the digestion products confirmed the presence of several antivirally active cystatin C fragments in cathepsin D, chymase, and napsin A-digested samples, although only napsin A generated CysC95-146 originally isolated from human serum (Fig. 5C).

Cathepsin D is an important component of the lysosomal protein degradation pathway in virtually all cells (54) and released from immune cells during inflammatory processes (52). Chymase is a serine protease that is mainly produced by activated mast cells and elevated in some viral infections (55). Napsin A is an aspartic proteinase that is abundantly expressed in normal lung and kidney tissue and a marker for some neoplasia (56). Altogether, these results show that cystatin C fragments which inhibit GPR15-mediated HIV and SIV infection are detectable in blood-derived human hemofiltrate and can be generated by proteases that are present and activated at sites of infection and inflammation.

**GPR15L Does Not Prevent SIV Entry.** For a long time, GPR15 had remained an orphan receptor but recently an agonistic chemokine ligand (named GPR15L) that modulates lymphocyte recruitment to epithelia has been identified (23, 24). It is well established that the chemokine ligands of the main entry cofactors of HIV-1, CCL5/RANTES and CXCL-12/SDF-1, inhibit CCR5- or CXCR4-mediated HIV-1 infection, respectively (reviewed in ref. 57). Unexpectedly, GPR15L did not display an inhibitory effect on GPR15-mediated SIVmac infection (Fig. 6B), although it induced down-regulation of GPR15 from the surface of GHOST-GPR15 and CEM-M7 cells (Fig. 6B and SI Appendix, Fig. S4).

Receptor internalization was strongly reduced when the cells were kept on ice, indicating that it mainly resulted from endocytosis and not from competition with the antibody used for staining. In

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contrast to GPR15L, CysC95-146 did not affect cell surface expression of GPR15 (Fig. 6C). In addition, CysC95-146 did not induce calcium release in Chinese hamster ovary (CHO) cells engineered to express human GPR15 together with the promiscuous G protein Gα16 (23), while GPR15 signaling was confirmed for GPR15L (Fig. 6D). Altogether, the results showed that CysC95-146 prevents SIV infection without altering GPR15 cell surface expression, while down-modulation of GPR15 by GPR15L was insufficient to cause significant inhibitory effects on SIV entry.

**CysC95-146 Does Not Interfere with GPR15L Signaling.** The results outlined above suggested that CysC95-146 might inhibit GPR15-mediated viral entry without interfering with the physiological signaling activity of this GPCR. Indeed, calcium flux assays performed in the presence of constant quantities of GPR15L and increasing doses of various CysC fragments revealed that C-terminal CysC fragments did not reduce the signaling activity of GPR15L (Fig. 6F). Conversely, GPR15L did not enhance the antiviral activity of CysC95-146 (Fig. 6F). To obtain insights into the region(s) in GPR15 targeted by CysC95-146, we examined the ability of this antiviral peptide to compete with GPR15 antibodies. CysC95-146 competed with Ab367902 targeting the extracellular N terminus as well as (less efficiently) with Ab188938 and Ab188939. CysC95-146 competed with Ab8104 targeting the extracellular C terminus as well as (less efficiently) with Ab188938 and Ab188939. CysC95-146 prevented SIV infection without altering GPR15 cell surface expression, while down-modulation of GPR15 by GPR15L was insufficient to cause significant inhibitory effects on SIV entry.

**The Antiviral Activity of CysC95-146 Is Conserved in Simian Hosts of SIV.** HIV-1 and HIV-2 are the result of at least 13 independent zoonotic transmissions of SIVs from great apes or sooty mangabeys to humans that occurred in the last century (37, 59). In contrast, SIVs have infected nonhuman primate species for many thousands or even millions of years, possibly since primate speciation (60). GPR15 is a major SIV entry cofactor in the two best-studied simian species naturally infected with SIV, i.e., sooty mangabeys and African green monkeys (16, 20). GPR15 also represents a major entry cofactor of SIVmac in macaques, the best
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established nonhuman primate model for AIDS in humans. To examine whether C-terminal cystatin C fragments may play a role in SIV infection, we examined whether their ability to inhibit GPR15-mediated viral entry is conserved in monkeys. Cystatin C is the most ancestral member of the cystatin family of inhibitors of cysteine peptidases and found in many vertebrates (61). Sequence alignments revealed that the C terminus of human cystatin C is fully conserved in great apes and that the macaque ortholog differs only in two amino acids from its human counterpart (Fig. 7A). We chemically synthesized the macaque CysC95-146 variant and found that it inhibits SIVmac239 infection as efficiently as the human version (Fig. 7B). Thus, the antiviral activity of CysC95-146 is conserved in simian hosts of SIV infection.

Effect of CysC95-142 and GPR15L on SIV and HIV Infection of Primary T Cells. To determine whether inhibition of GPR15-mediated HIV or SIV infection by CysC95-146 is relevant in primary viral target cells, we infected phytohaemagglutinin (PHA)-stimulated peripheral blood mononuclear cells (PBMCs) from three human donors in the presence and absence of this peptide. Since PBMCs express various SIV and HIV coreceptors, we performed the experiment in the presence and absence of AMD3100 and Maraviroc, preventing CXCR4- and CCR5-mediated viral entry, respectively. Fluorescence-activated cell sorting (FACS) analyses allowed us to determine the percentages of infected cells by intracellular p24 or p27 antigen staining and showed that PHA stimulation induced CCR5, GPR15, and CXCR6 cell surface expression (SI Appendix, Fig. S7 A and B). CysC95-146 had no inhibitory effect on a CCR5-tropic derivative of HIV-1 NL-4-3 and the CXCR4-tropic HIV-2 ROD9 molecular clone (SI Appendix, Fig. S7C). However, the peptide significantly reduced SIVmac239 infection of human PBMCs in both the presence and absence of additional inhibitors (SI Appendix, Fig. S7C). To further examine this, we analyzed the effects on three additional virus strains, i.e., the GPR15-tropic HIV-1 ZP6248 IMC, HIV-2 7312, and SIVsmm L1, which are capable of using CCR5, GPR15, and CXCR6 as entry cofactors (similarly to SIV-mac239). CysC95-146 clearly reduced PBMC infection by all three HIV-1, HIV-2, and SIVsmm strains (SI Appendix, Fig. S7D), suggesting that GPR15-mediated entry contributes to primate lentiviral infection of primary CD4+ T cells.

To analyze possible effects on spreading infection, we pre-treated PHA-stimulated human PBMCs with the various inhibitors prior to virus exposure. Infectious virus production was determined by infection of TZM-bl indicator cells with PBMC culture supernatants obtained on different days postinfection (dpi). Predictably, AMD3100 blocked CXCR4-tropic HIV-1, while Maraviroc fully prevented CCR5-tropic HIV-1 replication (Fig. 7C). In comparison, GPR15L, CysC95-142, and the CXCR6 chemokine ligand CXCL16 had no significant effect on CXCR4- or CCR5-tropic HIV-1 replication. Exposure of PBMCs to the GPR15-tropic HIV-1 ZP6248 strain did not result in significant replication, precluding meaningful analysis of inhibitors. However, GPR15L, CysC95-146, and CXCL16 all reduced replication of SIVmac239 in human PBMC cultures, albeit less efficiently than Maraviroc (Fig. 7C). CysC95-146 was more effective than GPR15L and CXCL16 and suppressed infectious virus yield on average by ∼60% (Fig. 7D). FACS analyses revealed that CysC95-146 did not affect GPR15 cell surface expression while GPR15L reduced it by ∼40% (SI Appendix, Fig. S8 A and B). Unexpectedly, GPR15L induced significant down-modulation of CXCR4 and competed with the 12G antibody that

Fig. 5. Treatment with various proteases generates antiviral cystatin C fragments. (A) Human cystatin C protein was digested with the indicated proteases. Digestion products were separated by sodium dodecyl sulphate–polyacrylamide gel electrophoresis (SDS-PAGE) and visualized by Coomassie Brilliant Blue staining. As controls, nondigested cystatin C as well as the synthesized CysC95-146 were included. (B) Proteases and nondigested protein were removed by ultrafiltration with different kilodalton cutoffs for purification. GHOST-GPR15 cells were then treated with the digestion products or CysC95-146 or full-length cystatin C and subsequently infected with a SIVmac239 luciferase reporter virus. Infection was measured at 3 dpi as described before. Results are displayed as means ± SD of one experiment in triplicate. ∗P < 0.05; ∗∗P < 0.01 (Mann–Whitney U test, unpaired t test, nonparametric). (C) Heat map visualization of identified cystatin C fragments in samples digested with the indicated proteases by mass spectrometry.
targets ECL-2 of CXCR4 for binding to this GPCR (SI Appendix, Fig. S8B and C). However, in contrast to the small molecule inhibitor AMD3100, neither CysC95-146 nor GPR15L inhibited CXCR4-mediated HIV-1 infection (SI Appendix, Fig. S8D). Altogether, these results show that CysC95-146 suppresses GPR15-mediated primate lentiviral infection in primary human target cells and revealed an unexpected effect of GPR15L on CXCR4 cell surface expression.

We hypothesized that CysC95-146 may only reduce SIVmac239 replication by ~60% because this virus also utilizes CCR5 for entry into primary T cells. To examine this, we screened healthy uninfected individuals for the presence of the Δ32/Δ32 deletion in the CCR5 gene that is present in about ~1% of Caucasians and disrupts functional CCR5 expression (62). We identified one donor containing homozygous deletions (Fig. 7E) and performed infection experiments in Δ32/Δ32 PBMCs in the presence of various antiviral agents. Predictably, R5-tropic HIV-1 did not replicate in PBMCs lacking CCR5. Replication of SIVmac239 was almost entirely prevented by CysC95-146 but hardly affected by GPR15L, Maraviroc, AMD3100, or CXCL16 (Fig. 7F). On average, CysC95-146 reduced production of infectious SIVmac239 by ~95% but had no inhibitory effect on X4-tropic HIV-1 (Fig. 7G). Vice versa, AMD3100 reduced infectious yield of X4 HIV-1 by 94% but had no significant effect on SIVmac239. Thus, in the absence of CCR5, CysC95-146 prevents SIV replication in primary T cells almost entirely, suggesting potent inhibition of GPR15-dependent virus entry.

Discussion

In the present study, we identified CysC95-146 and related C-terminal fragments of cystatin C as effective and specific endogenous inhibitors of GPR15-dependent HIV and SIV infection. In contrast to CysC95-146, GPR15L, the recently discovered...
Chemokine ligand of GPR15 (23, 24), displayed little if any inhibitory effect on HIV and SIV entry. This came as a surprise because the chemokine ligands of CCR5 and CXCR4 inhibit R5- or X4-tropic HIV-1 infection, respectively (63, 64). Notably, C-terminal fragments of cystatin C prevent SIV and HIV-2 infection without interfering with GPR15L-mediated signaling activity of GPR15. In contrast, small molecule inhibitors of CCR5- and CXCR4-dependent HIV-1 entry, such as Maraviroc or AMD3100, antagonize chemokine signaling via these GPCRs (65, 66). To our knowledge, CysC95-146 and related peptides are the first agents preventing GPCR-mediated infection by lentiviral pathogens without interfering with the signaling function of the corresponding chemokine receptor. Specific targeting of the detrimental function but not the signaling activity is of significant interest since GPCRs are involved in many physiological and pathological processes and the target of about 30% of all current drugs.

Cystatin C is produced by all nucleated cells, found in all tissues and body fluids, and represents the most abundant cysteine protease inhibitor (27). It is best known as a marker for renal failure. However, accumulating data also support an important role of cystatin C in the immune response against various exogenous or endogenous pathogens (28, 31, 49). The plasma levels in healthy individuals are about 0.1 \( \mu M \). However, cystatin C is induced in HIV-infected individuals and reaches blood plasma levels up to 0.5 \( \mu M \) under conditions of renal failure, infection, and inflammation (35, 67, 68). This concentration approximates the IC\(_{50}\) of antiviral cystatin C fragments and it is

![Fig. 7.](https://doi.org/10.1073/pnas.2023776118)

**Fig. 7.** The antiviral activity of CysC95-146 is conserved in monkeys and affects HIV-2 and SIV infection in human T cells. (A) Alignment of cystatin C amino acid sequences from the indicated species. Dots indicate identity to the human sequence and dashes identify gaps introduced to optimize the alignment. The region corresponding to the CysC95-146 is shaded. (B) Antiviral activity of human and monkey-derived CysC95-146 peptides. GHOST-GPR15 cells were incubated with increasing amounts of human and monkey CysC95-146 for 2 h at 37 °C prior to infection with SIVmac239 Firefly luciferase (F-Luc). At 3 dpi, infection was analyzed via F-Luc reporter assay. The experiment was performed in triplicates. (C) CysC95-146 shows antiviral activity against GPR15-mediated SIVmac239 replication in human primary cells. To examine possible effects on spreading infection, we isolated and stimulated human PBMCs and treated them with the various compounds (CysC95-146, GPR15L, AMD3100, MVC, and CXCL16) prior to virus exposure. Infectious virus production was determined by infection of TZM-bl indicator cells with PBMC culture supernatants obtained at different days post-infection. (D) Calculated area under the curve (AUC) for the virus replication data obtained in C. ***P < 0.001, **P < 0.01 (Welch's t test, unpaired). (E) Verification of homozygous deletions in the CCR5 gene of a \( \Delta32/\Delta32 \) PBMC donor. (F) Replication kinetics of SIVmac239 in \( \Delta32/\Delta32 \) PBMCs in the presence of various antiviral agents. Experimental details and symbols are provided in C. (G) Calculated AUC for the virus replication of SIVmac239 (see F) and X4 HIV-1 in \( \Delta32/\Delta32 \) PBMCs.

Chemokine ligand of GPR15 (23, 24), displayed little if any inhibitory effect on HIV and SIV entry. This came as a surprise because the chemokine ligands of CCR5 and CXCR4 inhibit R5- or X4-tropic HIV-1 infection, respectively (63, 64). Notably, C-terminal fragments of cystatin C prevent SIV and HIV-2 infection without interfering with GPR15L-mediated signaling activity of GPR15. In contrast, small molecule inhibitors of CCR5- and CXCR4-dependent HIV-1 entry, such as Maraviroc or AMD3100, antagonize chemokine signaling via these GPCRs (65, 66). To our knowledge, CysC95-146 and related peptides are the first agents preventing GPCR-mediated infection by lentiviral pathogens without interfering with the signaling function of the corresponding chemokine receptor. Specific targeting of the detrimental function but not the signaling activity is of significant interest since GPCRs are involved in many physiological and pathological processes and the target of about 30% of all current drugs.

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8 of 11 | PNAS Hayn et al.

https://doi.org/10.1073/pnas.2023776118

Natural cystatin C fragments inhibit GPR15-mediated HIV and SIV infection without interfering with GPR15L signaling
conceivable that the local levels at sites of infection and inflammation might exceed the systemic plasma levels. In fact, we found that peptides blocking GPR15-mediated SIV entry are generated from cystatin C by treatment with cathespin D, chymase, and napsin A (Fig. 5). These proteases are secreted by lysosomal exocytosis (69) or via specialized secretory granules (53) during immune responses and activated under acidic conditions. Acidification is a hallmark of inflammatory tissues (70) and thought to play a key role in innate immunity (71).

The generation of the active CysC fragments shows notable parallels to the generation of the CXCR4 antagonist and X4 HIV-1 inhibitor EPI-X4, which is generated from albumin by cathespin D and E under acidic conditions (13). Albumin is more abundant than cystatin C but the IC_{50} of EPI-X4 is also 10-fold higher than that of CysC95-146. Similarly, it has been reported that proteolytic processing of chemokine (C-C motif) ligand 14 (CCL14), commonly also known as hemophilic CC chemokine 1 (HCC-1), by trypsin-like serine proteases generates a potent CCR5 agonist (CCL14[9-74]) that efficiently inhibits R5-tropic HIV-1 strains (11, 12). Similar to cystatin C and albumin, the nonfunctional full-length CCL14 precursor is present at high concentrations in normal plasma. These results suggest that EPI-X4, CCL14[9-74], and active cystatin C fragments are all preferentially generated at sites of infection and inflammation, where they might act locally to cooperatively inhibit CXCR4, CCR5, and GPR15-mediated HIV or SIV infection, respectively. It is tempting to speculate that the combination of such endogenous inhibitors may have driven promiscuous coreceptor usage of SIVs that have most likely been infecting primary species for millions of years (72, 73).

We have previously shown that structure–activity relationship (SAR) studies enhance the activity of endogenous peptides by several orders of magnitude and offer perspectives for therapeutic applications (13, 74). For example, an optimized derivative of a natural 20-residue fragment of α1-antitrypsin that targets the gp41 fusion peptide of HIV-1 was safe and effective in human individuals (75). In addition, optimized derivatives of the endogenous CXCR4 antagonist Epi-X4 prevent atomic detergent and airway inflammation in preclinical mouse models (76). CysC95-146 is relatively large but tolerates truncations without loss of activity (Fig. 4). We will perform structural and molecular modeling studies and SAR analyses to determine the minimal active size of C-terminal cystatin C fragments and to increase their activity by rational design of derivatives predicted to interact more strongly with GPR15.

Our results show that CysC95-146 prevents GPR15-mediated HIV-2 and SIV infection, while GPR15L displayed little if any inhibitory activity although it induced down-modulation of GPR15 from the cell surface. It is known that both receptor removal from the cell surface as well as competitive inhibition by occupation of the interaction site(s) of the HIV envelope glycoprotein by chemokines might contribute to inhibition of CCR5- or CXCR4-dependent HIV-1 infection (77, 78). Our data support that competition by C-terminal cystatin C fragments is more effective than GPR15L-induced down-regulation of GPR15 in inhibiting lentiviral infection in both GHOST indicator and primary CD4+ T cells, suggesting that only a certain threshold is required for viral entry. Further structure–function analyses are required to fully elucidate the antiviral mechanism and the interaction(s) of CysC95-146 and GPR15L with GPR15. Our preliminary results from antibody competition assays and molecular modeling analyses suggest that CysC95-146 interacts most strongly with the N-terminal region of GPR15 (Fig. 6f). The observed antiviral effect agrees with previous results showing that the N-terminal domains of CCR5 and CXCR4 are targeted by the HIV-1 envelope glycoproteins and play a key role in membrane fusion (79, 80). Consistent with published data on CCR5 and HIV-1, we found that antibodies targeting the N-terminus or ECL-1 of GPR15 prevented SIVmac239 infection (Fig. 6g). GPR15L differs in GPCR interaction from prototype chemokines (23) and may interact with more C-terminal domains of GPR15. Differential GPR15 interaction sites also explain why CysC95-142 did not affect GPR15L-mediated signaling (Fig. 6d) and, vice versa, the chemokine ligand did not enhance the inhibitory effect of the CysC95-146 fragment (Fig. 6f).

Antiviral cystatin C fragments may have some relevance in humans since GPR15 is a common coreceptor of HIV-2 that infects about 1 to 2 million people mainly in Sub-Saharan Africa and is also used by some HIV-1 strains. We show that CysC95-146 inhibits HIV-2 and the highly unusual HIV-1 ZP6248 strain not only in indicator cell lines but also in primary human cells. Our results also demonstrate that the antiviral activity of CysC95-146 is conserved in monkeys and support that the peptide inhibits SIV infection of primary human cells. HIV entered the human population only about a century ago and was hence clearly not a driving force in the evolution of endogenous peptide blocks GPR15-mediated entry. In comparison, SIVs have infected nonhuman primate species for many thousands if not millions of years (37). GPR15 coreceptor usage is found in diverse groups of primate lentiviruses (81) and most likely represents an ancient function. Thus, it is tempting to speculate that the evolution of inhibitors of GPR15-mediated viral entry by inflammation and infection-associated proteases might have been driven by efficient primate lentiviruses. Finally, our results suggest that GPR15 allows SIV to replicate in Δ32/Δ32 PBMCs in the absence of CCR5 (Fig. 7 E–G). In humans, the Δ32/Δ32 genotype is associated with a reduced risk of the acquisition of HIV-1 infection via the sexual route (62, 82). Notably, sooty mangabeys, the original host of SIVsmm/HIV-2, also frequently lack functional CCR5 expression (20). However, the prevalence of natural SIVsmm infection was not significantly reduced in animals lacking functional CCR5 most likely due to efficient coreceptor usage of GPR15 and CXCR6.

Our identification of CysC95-146 provides proof of concept that some ligands of GPCRs can block pathogens without interfering with their physiological signaling function. Notably, this is not the case for inhibitors of CCR5- and CXCR4-mediated HIV infection, and this precludes e.g., usage of AMD3100 for the treatment of chronic diseases since proper CXCR4 signaling is critical for many physiological processes. After the CXCR4 antagonist EPI-X4 (13), CysC95-146 is another example of the proteolytic generation of a peptide virus inhibitor from an abundant precursor protein by proteases that are activated under acidic conditions. It is conceivable that generation of antimicrobial effects by proteolytic generation of abundant precursors might provide a more effective and rapid means to generate innate immune effectors than de novo synthesis. Further studies to clarify whether generation of antimicrobial molecules by proteolysis of abundant precursor proteins by proteases activated during infection and inflammation represents a common concept of innate immune defense seem warranted.
Fluorescence-Based Calcium Release Assays. GPR15 signaling efficiency was determined as described previously (23).

Statistical Methods. The mean activities were compared using Student’s t test. Similar results were obtained with the Mann–Whitney U test. The software package Statview version 4.0 (Abacus Concepts) was used for all calculations.

Data Availability. All data generated in this study are included in the paper and SI Appendix.

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1. A. J. Venkatakrishnan et al., Molecular signatures of G-protein-coupled receptors. Nature 494, 185–194 (2013).
2. B. C. Heng, J. Aubel, M. Fussenegger, An overview of the diverse roles of G-protein coupled receptors (GPCRs) in the pathophysiology of various human diseases. Bio-technol. Adv. 31, 1676–1694 (2013).
3. H. Deng et al., Identification of a major co-receptor for primary isolates of HIV-1. Nature 381, 661–666 (1996).
4. Y. Feng, C. C. Broder, P. E. Kennedy, HIV-1 entry cofactor: Functional cdNA cloning of a seven-transmembrane, G protein-coupled receptor. Science 272, 872–877 (1996).
5. T. Dragić et al., HIV-1 entry into CD4+ cells is mediated by the chemokine receptor CCCKR5. Nature 381, 667–673 (1996).
6. G. Alkhatib et al., CC CKR5: A RANTES, MIP-1, MIP-1 receptor as a fusion cofactor for macrophage-tropic HIV-1. Science 272, 1955–1958 (1996).
7. H. Choe et al., The beta-chemokine receptors CC3 and CC5 facilitate infection by macrophage-tropic HIV-1. Nat. Med. 11, 1135–1146 (2005).
8. B. J. Doranz et al., A dual-tropic primary HIV-1 isolate that uses fusin and the beta-chemokine receptors CKR-5, CKR-3, and CKR-2B as fusogenic cofactors. Cell 85, 1149–1158 (1996).
9. M. Bosso, L. Ständker, F. Kirchhoff, J. Münch, Exploiting the human peptidome for novel antimicrobial and anticancer agents. Bioorg. Med. Chem. 26, 2719–2726 (2018).
10. J. Münch, L. Ständker, W.-G. Forssmann, F. Kirchhoff, Discovery of modulators of HIV-1 infection from the human peptidome. Nat. Rev. Microbiol. 12, 715–722 (2014).
11. M. Metheux et al., The natural proteolytic processing of hemofiltrate CC chemokine 1 generates a potent CC chemokine receptor (CCR5) and CC5 agonists with anti-HIV properties. J. Exp. Med. 192, 1501–1508 (2000).
12. J. Münch et al., Hemofiltrate CC chemokine [9-74] causes effective internalization of CC5 and is a potent inhibitor of R5-tropic human immunodeficiency virus type 1 strains in primary T cells and macrophages. Antimicrob. Agents Chemother. 46, 982–990 (2002).
13. O. Zirafi et al., Discovery and characterization of an endogenous CXCXR4 antagonist. Cell Rep. 11, 737–747 (2015).
14. A. Mörner et al., Primary human immunodeficiency virus type 2 use multiple coreceptors for viral entry. J. Virol. 72, 5425–5432 (1998).
15. H. K. Deng, D. Unutmaz, V. N. KewalRamalan, D. R. Littman, Expression cloning of new receptors used by simian and human immunodeficiency viruses. Nature 388, 296–300 (1997).
16. N. E. Riddick et al., Simian immunodeficiency virus utilizes human and sooty mangabeys but not rhesus macaque STRL33 for efficient entry. J. Virol. 74, 5075–5082 (2000).
17. M. Farzan et al., Mammalian natural killer cell receptors are major targets for R5 primary isolates of human immunodeficiency virus type 1. J. Exp. Med. 196, 405–411 (1997).
18. S. M. Owen et al., Genetically divergent strains of human immunodeficiency virus type 2 use multiple coreceptors for viral entry. J. Virol. 72, 5425–5432 (1998).
19. C. C. Broder, P. E. Kennedy, HIV-1 entry cofactor: Functional cdNA cloning of a seven-transmembrane, G protein-coupled receptor. Science 272, 872–877 (1996).

Yan et al.
Natural cystatin C fragments inhibit GPR15-mediated HIV and SIV infection without interfering with GPR15 signaling

Hayn et al.
61. P. de Sousa-Pereira et al., Evolution of C, D and S-type cystatins in mammals: An extensive gene duplication in primates. PLoS One 9, e103850 (2014).
62. M. Samson et al., Resistance to HIV-1 infection in caucasian individuals bearing mutant alleles of the CCR-5 chemokine receptor gene. Nature 382, 722–725 (1996).
63. C. C. Bleul et al., The lymphocyte chemottractant SDF-1 is a ligand for LESTR/fusin and blocks HIV-1 entry. Nature 382, 829–833 (1996).
64. E. Oberlin et al., The CXC chemokine SDF-1 is the ligand for LESTR/fusin and prevents infection by T-cell-line-adapted HIV-1. Nature 382, 833–835 (1996).
65. G. A. Donzella et al., AMD3100, a small molecule inhibitor of HIV-1 entry via the CXCR4 co-receptor. Nat. Med. 4, 72–77 (1998).
66. P. Dorr et al., Maraviroc (UK-427,857), a potent, orally bioavailable, and selective small-molecule inhibitor of chemokine receptor CCR5 with broad-spectrum anti-human immunodeficiency virus type 1 activity. Antimicrob. Agents Chemother. 49, 4721–4732 (2005).
67. B. Bhasin et al., HIV viremia and T-cell activation differentially affect the performance of glomerular filtration rate equations based on creatinine and cystatin C. PLoS One 8, e22028 (2013).
68. C. T. Longenecker et al.; AIDS Clinical Trials Group Study A5224s Team, Reductions in plasma cystatin C after initiation of antiretroviral therapy are associated with reductions in inflammation: ACTG A5224s. J. Acquir. Immune Defic. Syndr. 69, 168–177 (2015).
69. A. Rodríguez, P. Webster, J. Ortego, N. W. Andrews, Lysosomes behave as Ca²⁺-regulated exocytic vesicles in fibroblasts and epithelial cells. J. Cell Biol. 137, 93–104 (1997).
70. F. Okajima, Regulation of inflammation by extracellular acidification and proton-sensing GPCRs. Cell. Signal. 25, 2263–2271 (2013).
71. K. Rajamäki et al., Extracellular acidosis is a novel danger signal alerting innate immunity via the NLRP3 inflammasome. J. Biol. Chem. 288, 13410–13419 (2013).
72. A. A. Compton, H. S. Malik, M. Emerman, Host gene evolution traces the evolutionary history of ancient primate lentiviruses. Philos. Trans. R. Soc. Lond. B Biol. Sci. 368, 20120496 (2013).
73. R. J. Gifford et al., A transitional endogenous lentivirus from the genome of a basal primate and implications for lentivirus evolution. Proc. Natl. Acad. Sci. U.S.A. 105, 20362–20367 (2008).
74. J. Münch et al., Discovery and optimization of a natural HIV-1 entry inhibitor targeting the gp41 fusion peptide. Cell 129, 263–275 (2007).
75. W.-G. Forsmann et al., Short-term monotherapy in HIV-infected patients with a virus entry inhibitor against the gp41 fusion peptide. Sci. Transl. Med. 2, 63re2 (2010).
76. M. Harms et al, An optimized derivative of an endogenous CXCR4 antagonist prevents atopic dermatitis and airway inflammation. bioRxiv:10.1101/2020.08.28.272781 (29 August 2020).
77. M. A. Lobritz et al., Multifaceted mechanisms of HIV inhibition and resistance to CCR5 inhibitors PSC-RANTES and Maraviroc. Antimicrob. Agents Chemother. 57, 2640–2650 (2013).
78. A. Steen, T. W. Schwartz, M. M. Rosenkilde, Targeting CXCR4 in HIV cell-entry inhibition. Mini Rev. Med. Chem. 9, 1605–1621 (2009).
79. H. Golding et al., CCR5 N-terminal region plays a critical role in HIV-1 inhibition by Toxoplasma gondii-derived cyclophilin-18. J. Biol. Chem. 280, 29570–29577 (2005).
80. N. Zhou et al., Structural and functional characterization of human CXCR4 as a chemokine receptor and HIV-1 co-receptor by mutagenesis and molecular modeling studies. J. Biol. Chem. 276, 42826–42833 (2001).
81. D. Unutmaz, V. N. KewalRamani, D. R. Littman, G protein-coupled receptors in HIV and SIV entry: New perspectives on lentivirus-host interactions and on the utility of animal models. Semin. Immunol. 10, 225–236 (1998).
82. R. Liu et al., Homozygous defect in HIV-1 coreceptor accounts for resistance of some multiply-exposed individuals to HIV-1 infection. Cell 86, 367–377 (1996).