Inhibition of HIV transmission in human cervicovaginal explants and humanized mice using CD4 aptamer-siRNA chimeras

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The continued spread of the HIV epidemic underscores the need to interrupt transmission. One attractive strategy is a topical vaginal microbicide. Sexual transmission of herpes simplex virus type 2 (HSV-2) in mice can be inhibited by intravaginal siRNA application. To overcome the challenges of knocking down gene expression in immune cells susceptible to HIV infection, we used chimeric RNAs composed of an aptamer fused to an siRNA for targeted gene knockdown in cells bearing an aptamer-binding receptor. Here, we showed that CD4 aptamer-siRNA chimeras (CD4-AsiCs) specifically suppress gene expression in CD4+ T cells and macrophages in vitro, in polarized cervicovaginal tissue explants, and in the female genital tract of humanized mice. CD4-AsiCs do not activate lymphocytes or stimulate innate immunity. CD4-AsiCs that knock down HIV genes and/or CCR5 inhibited HIV infection in vitro and in tissue explants. When applied intravaginally to humanized mice, CD4-AsiCs protected against HIV vaginal transmission. Thus, CD4-AsiCs could be used as the active ingredient of a microbicide to prevent HIV sexual transmission.

Introduction
The CAPRISA004 study, which demonstrated partial protection from sexual transmission of HIV-1 by vaginally applied tenofovir gel (1), has galvanized interest in developing an HIV microbicide. One of the major obstacles confronting this and other strategies for interrupting transmission is the transience of protection, requiring topical application just before sexual exposure and raising associated problems with compliance (2). Soon after RNAi was found to operate in mammalian cells, multiple groups showed that RNAi could be harnessed to inhibit HIV infection in vitro (3–7). Moreover, siRNAs directed against conserved viral gene sequences or the HIV receptor or coreceptor inhibit diverse viruses from multiple clades (3, 8, 9). Although knocking down the HIV receptor CD4 inhibits HIV transmission (3), targeting the CD4 gene would likely interfere with mounting effective immune responses and is therefore not desirable. CCR5, the HIV coreceptor responsible for virtually all sexual transmission of HIV (10, 11), is a more attractive RNAi target. CCR5 antagonists (12) have already proven useful at preventing HIV transmission in nonhuman primates (13) and humans (14–17). Humans bearing homozygous CCR5 mutations that abrogate CCR5 function are resistant to HIV infection and do not lead to any significant immune dysfunction (18–22). siRNAs directed against CCR5 efficiently silence gene expression for several weeks in vitro and in nondividing macrophages, which suggests that gene knockdown might be used to induce durable resistance to HIV infection, circumventing the need to apply a microbicide just before each sexual encounter (8). In fact, sexual transmission of another virus, herpes simplex virus type 2 (HSV-2), can be blocked in mice for at least a week by intravaginal (IVAG) application of siRNAs targeting HSV-2 genes and the HSV-2 receptor, nectin-1 (23, 24).

Translation of these promising results for blocking HSV-2 transmission to HIV prevention, however, must first overcome the hurdle of in vivo siRNA delivery to the immune cells that HIV infects, principally CD4+ T cells and macrophages, which are resistant to most transfection techniques. Although cholesterol-conjugated siRNAs are efficiently taken up by epithelial cells throughout the genital tract (including deep in the lamina propria, resulting in protection against lethal HSV-2 infection in mice; ref. 24), these reagents do not knock down gene expression in T lymphocytes or macrophages when applied IVAG to mice (E. Basar, unpublished observations). We previously developed a method for cell-specific siRNA transfection of immune cells that uses a fusion protein composed of a cell-targeting antibody fragment joined to a protamine peptide that binds nucleic acids (25, 26). siRNAs mixed with the fusion protein are taken up by and knock down gene expression in cells bearing the cognate surface receptor, both in vitro and in tissues after intravenous injection. Modifications of this approach effectively inhibit HIV infection in humanized mice (27). However, antibody-based fusion proteins are expensive to manufacture, are potentially immunogenic, and may require refrigerated storage, making them ill-suited for use in a microbicide for resource-poor settings.

Chimeric RNAs, composed of an siRNA fused to an aptamer (a structured RNA selected to bind a cell surface ligand with high affinity), provide an attractive alternative for in vivo gene knockdown (28–31). Aptamer-siRNA chimeras (AsiCs) efficiently transfect and knock down gene expression in cells bearing the surface receptor recognized by the aptamer. Intravenous injection of AsiCs incorporating aptamers targeting prostate surface membrane Ag...
Transcripts were eluted from denaturing SDS-PAGE gels and analyzed by column chromatography and native SDS-PAGE gels (Supplemental Figure 3), before annealing the antisense (active) siRNA strand. A chimera using an aptamer targeting PSMA (28) was synthesized as a binding control, while scrambled siRNA sequences controlled for gene silencing specificity. CD4-AsiCs engineered with one of the aptamers was consistently more effective than the other (clone 9 vs. clone 12; Supplemental Figure 1, B and C; data not shown; and ref. 33); therefore, unless otherwise indicated, clone 9 was used for subsequent experiments.

CD4-AsiCs are taken up by primary CD4+ cells in vitro. To test for siRNA uptake into CD4+ cells, primary monocyte-derived macrophages (MDMs) and CD4+ T cells, freshly isolated from the blood of healthy donors, were incubated with CD4-AsiCs labeled with Cy3 at the 3′ terminus of the antisense strand. CD4-AsiCs were efficiently and uniformly taken up by both MDMs (Supplemental Figure 4, A and B) and CD4+ T cells (Figure 1, B and C). PSMA-AsiCs were not internalized without transfection (complexation with cationic lipids for MDMs, electroporation for T cells). When added to resting PBMCs, Cy3-labeled CD4-AsiCs were selectively taken up by CD4+ monocytes and T cells, but only to a limited extent by CD8+ T cells (Figure 1D). The small subpopulation of circulating CD8+ T cells that took up the Cy3-labeled CD4-AsiCs may represent recently activated CD8+ T cells that express low levels of CD4 (data not shown). We also cannot rule out that staining might be due to cell surface binding without internalization.

CD4-AsiCs knock down target gene expression in primary CD4+ cells in vitro in a dose-dependent manner. Our next goal was to evaluate CD4-AsiC–mediated gene silencing. Primary CD4+ cells were treated with CD4-AsiCs bearing a CCR5 siRNA, and CCR5 expression was quantified by flow cytometry. CCR5 was knocked down in both MDMs (Supplemental Figure 4, C and D) and CD4+ T cells (Figure 1, E and F). Gene silencing was specific: CCR5 expression was unchanged when chimeras containing a scrambled siRNA sequence or the PSMA aptamer were tested, or when cells were incubated with the CD4 aptamer alone. As expected, transfection of CCR5 PSMA-Asi-Cs knocked down CCR5 expression in both cell types. Although silencing in CD4+ T cells was uniform, we observed 2 populations of MDMs: knockdown was virtually complete in one, whereas CCR5 was only partially downmodulated in the other. These 2 populations were also seen with lipid transfection. The reason for this is unclear, but is unlikely to be secondary to differential uptake, since uptake of Cy3-labeled CD4-AsiCs was uniform in macrophages (Supplemental Figure 4B).

Dose-dependent uptake of Cy3-labeled CD4-AsiCs and CCR5 knockdown occurred in both HeLa cells expressing CD4 and CCR5 (data not shown) and primary MDMs (Supplemental Figure 5). Uptake and gene silencing in MDMs after 72 hours were confirmed by fluorescence microscopy, in which Cy3 uptake was found to coincide with CCR5 knockdown, and neither uptake nor gene silencing occurred in MDMs incubated with Cy3-labeled siRNAs on their own (Supplemental Figure 4E). CD4-AsiCs could be readily designed to silence other genes, including the nuclear envelope gene lamin A (Supplemental Figure 6), the pan-leukocyte marker CD45 (Supplemental Figure 7), and the mitotic spindle gene EG5 (data not shown). Dose-dependent silencing was restricted to CD4+ cells, as assessed by immunoblot to measure protein and quantitative RT-PCR.
Figure 2
CD4-AsiCs inhibit HIV replication in vitro. MDMs (A–C) and CD4+ T cells (D and E) were infected for 48 hours with HIV-1env and HIV-1luc, respectively, and then treated with CD4-AsiCs or PSMA-AsiCs bearing gag and vif (g/v) siRNAs. Scrambled siRNA chimeras and CD4 aptamers were controls. Transfection controls used OF (A and B) or nucleofection (D and E). (A and D) Intracellular p24 expression, relative to infectant mock-treated cultures, was measured by flow cytometry 48 hours later. CD4-AsiCs inhibited HIV replication (mean ± SEM normalized to mock; n = 3; *P < 0.05, **P < 0.005, 2-tailed t test). Insets show representative histograms of MDMs and CD4+ T lymphocytes treated with gag and vif/CD4-AsiCs (gray, uninfected; blue, infected mock-treated; red, infected CD4-AsiC–treated). (C) HIV infection was also evaluated by fluorescence in situ hybridization (original magnification, ×60) using FITC-labeled probes complementary to HIV genomic RNA. HIV RNA was virtually undetectable in MDMs treated with 4 μM total final concentration of CD4-AsiCs. (B and E) To evaluate gene silencing independently of the effect of the CD4 aptamer on blocking viral entry, HIV replication was assessed by infection with VSV(G)-pseudotyped virus containing a luciferase reporter gene. Primary MDMs and CD4+ T cells were pretreated for 48 hours before infection with 4 μM mixtures of CD4- or PSMA-AsiCs targeting gag and vif or containing scrambled siRNAs. Luciferase activity, measured 48 hours later, was significantly inhibited in cells treated with CD4-AsiCs directed against either viral or luciferase genes (mean ± SEM normalized to mock; n = 3; *P < 0.05. **P < 0.005).

(qRT-PCR) for mRNA. Based on these results, we expect that CD4-AsiCs could be used to manipulate expression of virtually any gene in human CD4+ immune cells.

CD4-AsiCs are Dicer substrates and are processed intracellularly into functional siRNAs that use the RNAi pathway to direct mRNA cleavage. To understand the mechanism of CD4-AsiC–mediated silencing, we first tested whether these chimeras are substrates for the endoribonuclease Dicer, which processes longer endogenous RNA precursors to short 20- to 25-nt RNAs as part of the RNAi pathway in the nuclease Dicer, which processes longer endogenous RNA precursors to short 20- to 25-nt RNAs as part of the RNAi pathway in the nuclease Dicer, they were virtually completely digested to an fragment of the expected size was detected 72 hours after adding 32P-end-labeled CD4-AsiCs also resulted in their final concentration of CD4-AsiCs. (HIV RNA was virtually undetectable in MDMs treated with 4 μM total final concentration of CD4-AsiCs) and CD4-AsiCs are Dicer substrates and are processed intracellularly into functional siRNAs that use the RNAi pathway to direct mRNA cleavage (Supplemental Figure 8C and ref. 35). To confirm that CD4-AsiCs are processed by Dicer to release functional siRNA, we used recombinant Dicer, they were virtually completely digested to an expected approximately 21- to 23-nt cleavage product that migrated like a CCR5 siRNA (Supplemental Figure 8A). Treatment of primary CD4+ T cells with 32P-end-labeled CD4-AsiCs also resulted in their processing to an approximately 21- to 23-nt duplex RNA (Supplemental Figure 8B), which suggests that similar Dicer cleavage also occurs within cells. Gene silencing by CD4-AsiCs also depended on intracellular Dicer expression in HCT-116 cells, since knockdown of lamin A by CD4-AsiCs only occurred in WT cells, but not in Dicer−/− cells (Supplemental Figure 8C and ref. 35). To confirm that Asic-mediated silencing was caused by siRNA-directed cleavage of target gene mRNA, modified S′-rapid amplification of cDNA ends (S′-RACE; ref. 36) was used to analyze RNA isolated from primary MDMs treated with CCR5 CD4-AsiCs for CCR5 mRNA cleavage fragments (Supplemental Figure 9). An amplified CCR5 RNA fragment of the expected size was detected 72 hours after adding the CCR5 CD4-AsiCs. Sequencing of the amplified fragments confirmed that cleavage occurred 10 nt from the S′ end where the CCR5 antisense strand bound, at the expected site (37). Thus, we conclude that CD4-AsiCs are processed by Dicer to release functional siRNA duplexes that direct target mRNA cleavage via the RNAi pathway.

CD4-AsiCs inhibit HIV replication in primary CD4+ cells in vitro. Inhibiting HIV infection would be a stringent test of effective gene knockdown in CD4+ cells. We first tested whether treatment with a mixture of CD4-AsiCs against HIV gag and vif (Figure 2) or against CCR5 (Supplemental Figure 10) could suppress viral replication in an established HIV infection. Primary cells were infected with HIV-1 48 hours before CD4-AsiC treatment. Viral production was assessed by flow cytometry analysis of intracellular p24 capsid Ag (p24-Ag) or by p24 Ag ELISA of the culture supernatant. CD4-AsiCs against viral genes inhibited HIV replication in MDMs and CD4+ T cells (Figure 2, A and D). siRNAs against CCR5 also inhibited HIV infection in primary MDMs (Supplemental Figure 10), and siRNAs against viral genes inhibited infection in HeLa-CD4 and Jurkat cells in a dose-dependent manner (Supplemental Figure 11). There was no antiviral effect using chimeras containing PSMA aptamers or scrambled siRNAs. Viral inhibition and RNA internalization were confirmed by FISH for HIV RNA and Cy3 fluorescence, respectively, in MDMs treated with a cocktail of gag, vif, and Cy3-labeled CCR5 CD4-AsiCs, but not with a cocktail of PSMA-AsiCs (Figure 2C).

These CD4 aptamers bind CD4 principally via its V4 domain, but also partially block the receptor’s V1 gp120-binding domain (33, 38). Therefore, CD4-AsiCs could inhibit HIV infection by 2 mechanisms: by blocking HIV entry via inhibition of HIV gp120 binding to CD4, or by knocking down either viral genes or host genes required for viral entry or replication. In fact, prefection treatment of cells with CD4-AsiCs bearing scrambled siRNAs or with CD4 aptamers lacking an siRNA tail partially inhibited infection in HeLa-CD4+ cells (data not shown), which suggests that blocking HIV binding to CD4 by the aptamer contributes to overall inhibition of HIV infection. To evaluate the gene silencing component of chimera-mediated HIV suppression on its own, we pretreated cells with Asis before challenging with a CD4-independent single-round VSV(G)-pseudotyped HIV-1 virus encoding a luciferase reporter gene. In both MDMs and CD4+ T cells, CD4-AsiCs bearing gag and vif or lucifase siRNAs significantly decreased luciferase activity compared with all control conditions (Figure 2, B and E). Therefore, the CD4-AsiCs encoding either viral or CCR5 siRNAs likely inhibit HIV infection by both blocking entry and gene knockdown.

CD4-AsiCs do not alter expression of lymphocyte activation markers. CD4-AsiCs might be a useful tool for genetic manipulation of hard-to-transfect CD4+ cells in order to study the effect of knocking down one gene at a time. Ideally, treatment with CD4-AsiCs should not alter CD4 surface expression or cause lymphocyte activation, which would confound interpretation of gene silencing experiments. Activation of CD4+ T cells by CD4-AsiCs would also make them undesirable for preventing or treating HIV infection, since activated T cells are more susceptible to infection. Since CD4-AsiCs contain a single receptor binding site and T cell activation requires receptor cross-linking, we would not expect CD4-AsiC treatment to activate the cells. To evaluate this, uninfected CD4+ T cells treated with CD4-AsiCs directed against exogenous viral genes were assayed by flow cytometry over 2 days for changes in surface expression of CD4 and other markers of lymphocyte activation. Neither CD4 nor other cell surface receptors (CD3, CD45, CD25, and CD69) were significantly changed compared with mock-treated controls (Supplemental Figure 12).

CD4-AsiCs inhibit HIV replication in polarized cervicovaginal explants. To assess whether CD4-AsiCs can penetrate the vaginal epithelium and specifically knock down gene expression in CD4+ cells in intact human tissue, we treated the apical surface of polarized agarose-embedded human cervicovaginal explants, obtained from normal hysterectomy specimens (39, 40), with Cy3-labeled CCR5 CD4-AsiCs twice in a 24-hour interval. Cy3 fluorescence and CCR5 expression were measured in isolated CD4+ and CD8+ T cells and in CD4+...
Figure 3
CD4-AsiCs inhibit HIV replication in human cervicovaginal explants. (A–D) Cy3-labeled CD4-AsiCs or PBS was applied to the epithelium of agarose-embedded polarized explants 72 and 48 hours before flow cytometry of single-cell suspensions analyzed for Cy3 uptake (B) and CCR5 knockdown (C). Uptake was highest in CD3+CD4+ T cells and CD4+CD14+ macrophages, and CCR5 knockdown occurred only in CD4+ populations. (D) MFI (mean ± SEM normalized to mock; *P < 0.05, **P < 0.01, 2-tailed t test). Data from quadruplicate samples from 1 donor are representative of results from 3 donors. (E-H) HIV challenge experiment. Polarized vaginal explants from 3 donors were pretreated with indicated total concentrations of CD4-AsiCs targeting CCR5 (F), gag and vif (G), or all 3 genes (H) for 48 hours before HIV-1BaL challenge. Viral replication, measured by p24 Ag ELISA of the lower transwell culture medium, showed specific dose-dependent inhibition by CD4-AsiCs against CCR5 (F) or gag and vif (G) compared with mock-treated controls. At the same total concentration, AsiCs against CCR5 and gag and vif (H) was more effective than either CCR5 or viral CD4-AsiCs alone. PSMA-AsiCs did not inhibit viral replication. Shown are p24 Ag levels (mean ± SEM normalized to mock; *P < 0.05, **P < 0.01, 2-tailed t test). (I and J) The CD4 aptamer was compared with the gag and vif chimera by treating vaginal explants with serial 2-fold dilutions (1–8 μM) of each. p24 Ag ELISA was measured and normalized as above. Data are representative of independent experiments from 2 donors, performed in quadruplicate (*P < 0.05, **P < 0.0001, 1-way ANOVA with Dunnett multiple-comparison test).

To determine whether antiviral CD4-AsiCs can block HIV transmission in intact vaginal tissue, the epithelial surface of polarized explants was treated twice with CD4-AsiCs containing siRNAs targeting CCR5, gag and vif, or all 3 genes before challenge with HIVBaL 48 hours later. Mucosal viral replication, as assessed by p24 Ag ELISA of the explant culture medium, was inhibited by CD4-AsiC treatment in a dose-dependent manner (Figure 3, F-H). The triple cocktail administered at the same total concentration was more effective than the CCR5 or the antiviral CD4-AsiCs alone. A triple cocktail of PSMA-AsiCs did not inhibit HIV transmission to the tissue. To assess the antiviral contribution of the CD4 aptamer, we compared viral inhibition by the CD4 aptamer and the CD4-AsiCs encoding gag and vif siRNAs in a dose response study. The CD4 aptamer on its own inhibited HIV transmission in situ, but was about 2- to 4-fold less effective than the CD4-AsiCs (Figure 3, I and J), which suggests that although blocking CD4 binding contributes to HIV inhibition, the major antiviral activity of the CD4-AsiC is due to gene knockdown.

CD4-AsiCs do not trigger an innate immune response in human cervicovaginal tissue. Depending on their sequence, concentration, chemical modifications, and delivery vehicle, siRNAs can trigger innate immune sensors that recognize foreign nucleic acids or cause cytotoxicity and tissue damage (41). Moreover, the demonstrated antiviral effect could be a side effect of IFN induction by TLR or RIG-I activation. To examine whether CD4-AsiCs have any adverse or unanticipated off-target effects, we treated polarized human macrophages by flow cytometry 2 days after the second treatment (Figure 3, A–D). Cy3-labeled CD4-AsiCs were specifically taken up by CD4+ T cells and macrophages in situ and uniformly knocked down CCR5 expression. Despite nominal uptake by CD8+ T cells, no gene silencing was observed in this population. Neither uptake nor CCR5 silencing was seen in tissues treated with PSMA-AsiCs (Figure 3D). Topical application of CD4-AsiCs encoding a CD45 siRNA to explants also specifically knocked down CD45 expression only in CD4+ T cells and macrophages (data not shown).

Moreover, CD4-AsiCs did not induce IFNβ or IFNγ, inflammatory cytokines (IL6, IL8, and IL12), or IFN-responsive genes (IP10, OAS1, and STAT1), as measured by sensitive qRT-PCR of RNA extracted from whole tissue harvested at their expected peaks, 6 and 24 hours after treatment. Contrary to a previously published report (28), PSMA-AsiCs stimulated IFNB expression at 6 hours and IP10 at 24 hours, likely reflecting the higher sensitivity of qRT-PCR relative to ELISA for measuring cytokine induction.

CD4-AsiCs suppress target gene expression in CD4+ cells in vivo. To validate the antiviral properties of CD4-AsiCs in vivo, we used the BLT humanized mouse model (42). A mixture of CD4-AsiCs bearing CD45 siRNAs and Cy3-labeled CCR5 siRNAs were applied IVAG to NOD/SCID-BLT mice, 72 hours and 48 hours prior to sacrifice, each at doses ranging from 5 to 80 pmol (Figure 4A). To assess cell-specific siRNA delivery and gene knockdown in vivo, Cy3 fluorescence and CD45 and CCR5 expression in subpopulations of CD4+ and CD8+ cells were evaluated by flow cytometry of single-cell suspensions of vaginal tissue and compared with mock-treated control mice (Figure 4, B and C). Cy3 fluorescence uptake was uniform in CD3+CD4+ T cells and CD14+CD4+ tissue macrophages, but was absent in tissue CD3+CD8+ T cells. Dose-dependent CCR5 and CD45 knockdown was observed in CD4+ T cells, but not in CD8+ T cells. At the highest CD4-AsiC dose, cell surface CD45 and CCR5 MFI in CD4+ T cells was reduced 12- and 5-fold, respectively, compared with mock-treated controls.

Topically applied CD4-AsiCs inhibit vaginal transmission of HIV to humanized mice. To test whether topically applied CD4-AsiCs protect humanized mice from vaginal HIV challenge, 4 NSG-BLT mice were treated with a combination of CD4-AsiCs targeting CCR5, gag, and vif prior to IVAG challenge with 10^5TCID50 HIVBaL (Figure 4D), a viral dose that uniformly infects control mice (data not shown and refs. 43, 44). The CCR5 AsiCs were administered 48 and 24 hours prior to challenge, and the viral gene-targeting AsiCs were administered 24 hours before and 4 hours after IVAG challenge. We treated 4 mice with equivalent doses of CD4 aptamers lacking siRNA conjugates and 4 mice with PBS using the same dosing regimen. In the 12 weeks of observation after challenge, all the aptamer- and AsiC-treated mice survived, while 2 of the mock-treated mice died (Figure 4E). Whereas all the control and aptamer-treated mice became infected and had detectable HIV p24 antigenemia 2–4 weeks after infection, none of the CD4-AsiC-treated mice developed detectable p24 antigenemia (Figure 4F). Plasma viral burden, assessed by sensitive qRT-PCR assay for HIV gag mRNA, was detected at very low levels and only episodically several months later in the 2 of 4 CD4-AsiC–treated mice that became infected, a significant reduction compared with the mock-treated or aptamer-treated animals (Figure 4G). All the control mice and 2 of the 4 aptamer-treated mice showed dramatic depletion of circulating CD4+ T cells, whereas all mice treated with CD4-AsiCs and the other 2 aptamer-treated mice maintained relatively normal CD4 counts and stable CD4/CD8 ratios (Figure 4H and Supplemental Figure 14).

Although we did not directly determine the cause of death of the mock-treated mice, the profound depletion of their circulating...
CD4+ T cells in the 2 weeks prior to death may have been responsible. Thus, topical application of a mixture of CD4-AsiCs targeting CCR5 and HIV genes provided protection against vaginal challenge. Although the detection of plasma viremia and development of CD4+ T cell depletion were delayed in the aptamer-treated mice, their viral load was not significantly less than in mock-treated mice by the end of the observation period. Therefore, the CD4 aptamer, which inhibits HIV from binding to CD4, was substantially less effective than the chimera, which can both block binding and suppress entry and viral replication by gene knockdown.

**Discussion**

Delivery remains a significant obstacle to the clinical development of siRNA-based drugs. Although cholesterol-conjugated siRNAs silence gene expression without apparent toxicity in mucosal epithelial cells and can be used to prevent HSV-2 transmission (24), this approach would not inhibit HIV transmission, since cholesterol-conjugated siRNAs do not transfect the cells that HIV infects. siRNAs can be delivered into immune cells by receptor-mediated endocytosis, by either complexing siRNAs to antibody fusion proteins or encapsulating siRNAs into liposomes or other nanoparticles bearing targeting antibodies or ligands to cell surface receptors (25–27, 45). Here we demonstrated that an alternate approach—chimeric RNAs composed of an aptamer linked to an siRNA sequence, which mimics endogenous miRNA structures. Some of these changes to prevent sexual HIV transmission in humans, which is very inefficient, requiring hundreds of exposures for each transmission event, and where usually only a single virion is able to establish a foothold in the host (46). Protection was achieved here with a highest dose of approximately 0.2 mg/kg (120 pmol), which is a feasible dose for small RNA drugs, but which might be reduced even further by drug optimization. Future studies in humanized mice will evaluate how long gene silencing in the genital tissue and protection lasts in order to determine whether RNAi-based microbicides could be intermittently dosed with acceptable compliance.

Targeted delivery has the potential dual advantage of reduced toxicity to bystander cells and reduced effective dose. What delivery approach is preferable for clinical or research purposes is difficult to determine ab initio and may depend on the application. Chimeric RNAs have the advantage of being a single molecule rather than a complex mixture, are less likely to be immunogenic than proteins, and are more straightforward to purify and less costly to produce than RNAs that need to be formulated with proteins, nanoparticles, or liposomes. CD4-AsiCs were shown to knock down 2 viral genes, 1 transgene (lacZ) and 4 host genes (CCR5, CD45, lamin A, and EG5) and can likely be designed to inhibit the expression of any gene. The kinetics of target gene suppression may differ between targets, depending on target gene mRNA and protein stability. For example, CCR5 and CD45 surface protein expression was not appreciably reduced until 72 hours after CD4-AsiC treatment. However, mRNA levels of these genes, when measured by qRT-PCR, declined within a day.

This study builds on previous studies that used intravenous injection of PSMA-AsiCs to transfect human prostate cancer cells in an orthotopic mouse tumor model (28, 29) or that used HIV gp120–AsiCs to transfect HIV-infected CD4+ cells in vitro and in vivo (30–32). Based on these studies, the CD4-AsiCs developed here likely could be used for systemic gene silencing in circulating immune cells. If CD4-AsiCs perform as well as we expect, they may be a potent tool for genetic manipulation (which has been so powerful for understanding mouse immunology by use of knockout mice) to study the role of individual molecules in complex human immune responses in humanized mice in vivo. Because CD4-AsiCs do not appear to perturb CD4 cell surface expression or alter other immune receptors that are sensitive indicators of immune activation, it should be possible to test the effect of knocking down one gene product at a time. The lack of cellular toxicity of CD4-AsiCs might also make them an attractive alternative to electroporation for in vitro transfection of CD4+ T cells. The clone 9 aptamer also recognizes rhesus macaque CD4, and the corresponding CD4-AsiC knocks down gene expression in rhesus PBMCs in vitro (L.A. Wheeler, unpublished observations), which suggests that CD4-AsiCs could also be used to study SIV or SHIV infection and immune responses in nonhuman primates.

The micromolar concentrations of CD4-AsiCs used here for in vitro gene knockdown and HIV infection inhibition studies are somewhat higher (about 2–10 times more) than those published in some previous reports (3, 24, 28, 31). The higher concentration could be due to differences in target cell type. The CD4-AsiCs have not been optimized, and improvements in CD4-AsiC design or synthesis could increase silencing efficiency. Modifications might include optimizing the aptamer or siRNA sequence, altering the linker joining the aptamer to the siRNA, or modifying the ratio of siRNA strands, or replacing the double-stranded siRNA with a stem-loop that mimics endogenous miRNA structures. Some of these changes include optimizing the aptamer or siRNA sequence, altering the linker joining the aptamer to the siRNA, or modifying the ratio of siRNA strands, or replacing the double-stranded siRNA with a stem-loop that mimics endogenous miRNA structures. Some of these changes...
have been used in earlier studies, where the optimal AsiC construct depended on the particular siRNA sequence (29, 31). Studies are in progress to understand how CD4-AsiCs are taken up and delivered to the cytosol, where they interact with the endogenous RNAi machinery. To optimize CD4-AsiC design, it will be helpful to compare not only the efficiency of gene silencing, but also the efficiency of intracellular processing to the mature siRNA and its incorporation into the RNA-induced silencing complex. It may also be helpful to shorten the construct to make chemical synthesis more practical. We plan to test whether any part of the aptamer sequence can be deleted without losing binding affinity. Another approach would be to synthesize the aptamer and siRNA separately, using complementary adapter sequences to join them (30, 31).

Despite the high concentrations required in vitro, CD4-AsiC-mediated in vivo silencing and protection from HIV infection required significantly lower doses (about 7–25 times less) than were used to inhibit HSV-2 transmission with lipopolysaccharide or cholesterol-conjugated siRNAs (23, 24). The stability of the CD4-AsiCs over 36 hours in human vaginal fluid suggests that further stabilization for topical use may not be required. However, the in vivo half-life in the blood and other body fluids and within cells, as well as the efficiency and durability of gene knockdown, might be improved by further chemical modifications, such as introduction of 2′-OCH3 to purines on the active strand. For systemic use, chemical conjugation to cholesterol or polyethylene glycol might also improve circulating half-life (29, 36).

The pathways used to deliver RNAs into cells, whether by aptamers or by other delivery methods, remain poorly understood. Our preliminary studies suggest that Cy3-labeled CD4-AsiCs are initially taken up into early endosomes and then escape to the cytosol. Endocytosis might be triggered by activation of the CD4 receptor by aptamer binding or occur via the continuous basal internalization of cell surface receptors. The latter pathway may be more likely, since CD4 cell surface expression is not appreciably altered by CD4-AsiC treatment and since CD4-AsiCs are monomeric and are not expected to crosslink the receptor. Although lack of perturbation of CD4 surface expression would be ideal for using CD4-AsiCs as a research tool, a divalent or polyvalent reagent that activates temporary CD4 internalization might also have advantages for HIV prevention or therapy. This might add a third mechanism for inhibiting HIV cellular transmission (removal of the viral receptor from the cell surface) to the other 2 mechanisms demonstrated in the present study (gene silencing and partially blocking the virion binding site on CD4). Once in the endosomes, how the RNA is released into the cytosol is unknown. Understanding the uptake and trafficking within cells of CD4-AsiCs will undoubtedly facilitate design improvements.

**Methods**

**Chimera synthesis.** Aptamer-siRNA chimera synthesis was modified from previously described methods (28, 33). Primers and template DNA (Supplemental Table 1) were commercially synthesized (IDT). RNA intermediates, transcribed in vitro using Epicentre’s DuraScribe kit, were resolved on 15% PAGE gels [Invitrogen] and eluted into either buffer A for in vitro studies (1% LiClO4, 7 mM triethylamine; Sigma-Aldrich) or buffer B for in vivo studies (10 mM Tris, pH 7.5–8.0, 50 mM NaCl, 1 mM EDTA) prior to ethanol precipitation and desalting using a G25 column (GE). RNAs were mixed in a 1:1 molar ratio with commercially synthesized active siRNA strands (Dharmacon), heated to 90°C, and allowed to cool slowly to room temperature. In some cases, the active strand was synthesized with a Cy3 label at its 3′ end. Vaginal stability assay. 2 nmol CCR5 CD4+AsiC synthesized using 2′-fluoro-pyrimidines, chemically stabilized cholesterol-conjugated CCR5-siRNA, and unmodified 21-mer CCR5 siRNA in 100 μl PBS were added to 100 μl of vaginal fluid obtained from a healthy preovulatory donor. At regular intervals, 20 μl was removed, resuspended in TRIzol (Invitrogen) for RNA extraction, and frozen at –80°C prior to resolution by PAGE analysis 24 hours after the final collection time point. RNA content was analyzed by densitometry, and amounts were calculated relative to RNA content at time 0.

**Cell lines.** HeLa, Jurkat, and K562 cells (ATCC) were cultured as previously described (8, 25). HeLa-CD4 and HeLa-MAGI CCR5 cell lines, obtained from the NIH AIDS Reagent Program, were maintained in DMEM supplemented with 0.5 mg/ml G418. HCT-116 cells (ATCC) were cultured according to the supplier’s instructions. Dicer− − HCT-116 cells were a gift of B. Vogelstein (Johns Hopkins Medical Institutions, Baltimore, Maryland, USA; ref. 35).

**Primary cells.** Primary cells from the blood of healthy donors were isolated by Ficoll (GE) density centrifugation and cultured in H10 medium (RPMI 1640 [Cellgro] containing 10% Human AB Serum [GemCell], 100 U/ml penicillin, and 100 μg/ml streptomycin sulfate). In some cases, CD4+ cells were separated using immunomagnetic beads (Miltenyi). CD4+ T cells and MDMs were prepared as previously described (8). CD4+ T cells were cultured in H10 containing 60 IU/ml IL-2 and were activated using 4 μg/ml PHA (Difco). Resting PBMCs were cultured in H10 containing 4 μg/ml IL-15.

**Viruses.** HIVint and HIVint virus were obtained from the NIH AIDS Reagent Program. HIVint was generated by infecting pooled PBMCs that had been stimulated with PHA (4 μg/ml) in H10 plus 60 IU/ml IL-2 (Chiron). HIVint was propagated as previously described (47). p24 Ag levels in culture supernatants were measured by HIV-1 p24 Antigen ELISA kit (Perkin Elmer). VSV(G)-pseudotyped HIV-Luc (provided by A. Engelman; Dana-Farber Cancer Institute, Boston, Massachusetts, USA) was generated in 293T cells as previously described (48). Viral stocks of the HIV-1R5–CSF molecular clone were produced through transfection of HEK293 cells as previously described (49). Supernatant virus was concentrated 1:50 using the PEG-it Virus Precipitation Solution (System Biosciences) per the manufacturer’s protocol.

**HIV-1 infection in vitro and in situ.** Cells were infected with the indicated HIV-1 isolates using an MOI of approximately 1. Infection of Jurkat and CD4+ T cells was by spinoculation at 1,200 g for 2 hours in the presence of 2 mg/ml Polybrene. HeLa-CD4 and MDM cells were infected by incubating the cells with virus at 37°C for 48 hours. Cells were infected with the single-round VSV(G)-pseudotyped virus for 6–8 hours at 37°C, washed, and incubated again at 37°C in fresh media for approximately 48 hours prior to analysis as previously described (48). For HIV infection of polarized explants, HIVint (~100 ng p24) was applied to the apical surface of agarose-embedded explants in 200 μl, and the explants were then incubated at 37°C. Viral replication in the tissue was assessed by measuring p24 Ag in the lower transwell chamber using the HIV-1 p24 Antigen ELISA kit (Perkin Elmer). siRNA transfection. HeLa-CD4 and MDM cells were transfected with Oligofectamine per the manufacturer’s protocol. CD4+ T cells, Jurkat cells, and K562 cells were transfected by AMAXA according to the manufacturer’s protocol (Lonza). See Supplemental Table 2 for all siRNA sequences.

**Flow cytometry.** Direct immunostaining of CD3, CD4, CD8, CD14, CD45, and CCR5 was performed using 1:20 dilutions of murine mAb for 30–60 minutes at 4°C (BioLegend). Cells were stained in PBS containing 0.5% FCS, 1 mM EDTA, and 25 mM HEPES. Samples were washed twice in the same buffer. Data for 1- and 2-color staining experiments were acquired using FACSCalibur (BD Biosciences); for multicolor experiments, data were acquired using FACS-Canto II (BD Biosciences). All data analysis was performed using FlowJo (Treestar Inc.).
Fluorescence microscopy. Fluorescence microscopy was performed as previously described (24) using primary antibodies (BioLegend) and secondary donkey anti-mouse antibodies (Invitrogen). All images shown were acquired using a ×60 oil objective.

Intracellular p24 staining. Intracellular staining, performed as previously described (8), was analyzed on a FACSCalibur with Cell Quest software (Becton Dickinson) and/or FlowJo software. The p24 MFI was normalized relative to the mock-treated control.

Immunoblot. Total cell extracts were prepared in RIPA buffer (50 mM Tris-HCl pH 7.4, 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 1% sodium deoxycholate, 0.1% SDS). Protein concentration was measured using the BCA Protein Assay Reagent (Pierce), and 10 μg of total protein was resolved on 10% SDS-PAGE, transferred to cellulose membranes (Immobilon-P, Millipore), and probed with mouse mAb against lamin A (BioLegend). The membranes were developed using SuperSignal West Femto (Pierce). For loading control, 10 μg of total protein was resolved on 10% SDS-PAGE and probed using anti-tubulin mouse mAb (Sigma-Aldrich).

In vitro Dicer cleavage assay. Recombinant Dicer cleavage assay was performed using the Turbo Dicer siRNA Generation Kit (Genlantis) according to the manufacturer’s protocol.

S′-RACE and sequence analysis. S′-RACE was performed using the First Choice RLM-RACE Kit (Ambion) according to the manufacturer’s protocol, modified to use 45 cycles of amplification for the nested PCR reaction. The nested PCR product was cloned using standard M13 and T7 primers into the PGEM-T Easy vector (Promega) for sequencing at the Dana-Farber Harvard Cancer Center Sequencing Core (Boston, Massachusetts, USA).

Luciferase assay. The luciferase activity of VSV-G(pseudotyped) reporter virus–infected samples was measured 48 hours after transfection using the Luciferase Assay System (Promega) and a Top count NXT microplate (Becton Dickinson) and/or FlowJo software. The p24 MFI was normalized relative to the mock-treated control.

Cytokine and cytokine induction. An equimolar mixture of CD4-AsiC or PMA-AsiC targeting gag and vif (total concentration 4 μM) was applied to the apical surface of the explant, and RNA was extracted using TRIzol reagent (Invitrogen) 6 and 24 hours later. Treatment with poly(I:C) (50 μg/ml; Invitrogen) served as a positive control. qRT-PCR was performed as above (see Supplemental Table 3 for primers). mRNA expression was normalized to GAPDH, then normalized relative to mock-treated controls.

In vivo treatment and HIV challenge. All in vivo experiments were performed using immunodeficient mice bearing human bone marrow (either NOD/SCID or NSG) following reconstitution with CD34+ cells from human fetal liver and surgical human thymic graft (i.e., BLT mice) as previously described (42). Uptake and silencing was assessed in NOD/SCID-BLT mice following 2 IVAG treatments of Cy3-labeled AsiCs, at the indicated doses, 72 hours and 48 hours prior to sacrifice. Vaginal tissue was extracted, and a single-cell suspension was isolated by collagenase digestion and stained as described above. HIV protection was assessed in NSG-BLT mice treated IVAG (a) 48 hours before challenge with 10 μl PBS containing 80 pmol CCR5 CD4-AsiCs; (b) 24 hours before challenge with 10 μl PBS containing 40 pmol each of CCR5 gag and vif CD4-AsiCs; and (c) 4 hours after challenge with 10 μl PBS containing 40 pmol each of gag and vif CD4-AsiCs. Female mice were challenged with atrumatic IVAG instillation of 10⁵ TCID₅₀ HIVJR-CSF in 10 μl PBS as previously described (44). Animal work was approved by the Animal Care and Use Committees of Massachusetts General Hospital and Harvard Medical School.

Analysis of HIV infection. Blood was obtained by venipuncture at weekly intervals for 12 weeks after HIV challenge. Cells were pelleted by centrifugation, and plasma was stored at –80°C until analysis. Cell pellets were twice treated with rbc lysis buffer (Sigma-Aldrich), washed with flow cytometry buffer described above, and stained for CD3, CD4, and CD8. Viral RNA was extracted from 75 μl plasma using the QiaAmp Viral RNA kit (QIAGEN) according to the manufacturer’s instructions. cDNA was reverse transcribed using SuperscriptIII (Invitrogen) and HIV-gag–specific primers (Supplemental Figure 2) according to the manufacturer’s protocol. qRT-PCR was performed as described above. The remaining serum was aliquoted for p24 Ag ELISA (Perkin Elmer), performed according to the manufacturer’s instructions.

Statistics. Data for most experiments were analyzed by Student’s t test. All P values are for 2-tailed significance tests. For analysis of data based on independent experiments using samples from multiple donors, 1-way ANOVA with Dunnett multiple-comparison test was performed using GraphPad Prism (GraphPad Software). Assessment of HIV infection was by 2-way ANOVA with Dunnett multiple-comparison test. P values less than 0.05 were considered significant. The limit of detection was calculated using a previously described method (50) and is shown as the average of the calculated limit of detection for each individual assay.
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