Genome editing of CCR5 by AsCpf1 renders CD4\(^+\) T cells resistance to HIV-1 infection

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**Abstract**

**Background:** The chemokine receptor CCR5 is one of the co-receptor of HIV-1 infection. People with homozygous CCR5\(\Delta32\) deletion resist HIV-1 infection, which makes the CCR5 an important target for HIV-1 gene therapy. Although the CRISPR/Cas9 has ever been used for HIV-1 study, the newly developed CRISPR/AsCpf1 has never been utilized in HIV-1 co-receptor disruption. The CRISPR/Cpf1 system shows many advantages over CRISPR/Cas9, such as lower off-target, small size of nuclease, easy sgRNA design for multiplex gene editing, etc. Therefore, the CRISPR/Cpf1 mediated gene editing will confer a more specific and safe strategy in HIV-1 co-receptor disruption.

**Results:** Here, we demonstrated that CRISPR/AsCpf1 could ablate the main co-receptor of HIV-1 infection-CCR5 efficiently with two screened sgRNAs via different delivery strategies (lentivirus, adenovirus). The edited cells resisted R5-tropic HIV-1 infection but not X4-tropic HIV-1 infection compared with the control group in different cell types of HIV-1 study (TZM.bl, SupT1-R5, Primary CD4\(^+\) T cells). Meanwhile, the edited cells exhibited selective advantage over unedited cells while under the pressure of R5-tropic HIV-1. Furthermore, we clarified that the predicted off-target sites of selected sgRNAs were very limited, which is much less than regular using sgRNAs for CRISPR/Cas9, and no evident off-target was observed. We also showed that the disruption of CCR5 by CRISPR/AsCpf1 took no effects on cell proliferation and apoptosis.

**Conclusions:** Our study provides a basis for a possible application of CCR5-targeting gene editing by CRISPR/AsCpf1 with high specific sgRNAs against HIV-1 infection.

**Keywords:** CCR5, HIV-1, CRISPR/AsCpf1, HIV-1 resistance, Selective advantage

**Background**

Acquired immunodeficiency syndrome (AIDS), caused by HIV-1 (human immune deficiency virus type 1) infection has threatened the health of people all over the world. According to the statistic data from World Health Organization in 2018, 37.9 million people are living with HIV globally, and 23.3 million people (62%) were receiving highly active antiretroviral therapy (HAART) (https://www.who.int/hiv/data/en/).

However, HAART can not eradicate the HIV-1 virus, for the integration of HIV-1 into host genome. Moreover, the HAART has many limitations, such as long term drug taking, high cost and side effects (hepatic lesion, cardiovascular diseases, etc.) [1–3]. Therefore, it is necessary...
and inevitable to look for alternative and more effective therapies to cure HIV-1 infection.

HIV-1 infection of host cells is a multi-step process, the virus entry into cells begin with viral envelop protein gp120 binding to CD4 and one of the co-receptors CCR5 or CXCR4 on cell surface, the co-receptor CCR5 or CXCR4 is required for different HIV-1 strains, with CCR5 for CXCR4(R5)-tropic strain and CXCR4 for CXCR4(X4)-tropic strains [4, 5]. The genomic RNA of the virus will be converted to double-strand DNA after virus entering cells by membrane infusion. The viral DNA will integrate into target cell genome to be the provirus, the provirus can be transcribed as viral RNAs, which will be packaged to be a new virus particle. Previous reports have clarified that individuals with homozgyous deletion of co-receptor CCR5 (CCR5Δ32) are resistant to HIV-1 infection [6–8]. Meanwhile, clinical reports of two patients (Berlin Patient, London Patient) with HIV-1 infection and leukaemia received HLA-matched and homozgyous CCR5Δ32 bone marrow transplant have been proved that they got clinic-defined cure with undetectable HIV-1 [9–11]. Therefore, the co-receptor CCR5 has been a reasonable target for gene editing against HIV-1 infection.

Over last decades, several genome editing tools have been developed and utilized for diseases study and cure, such as zinc finger nuclease (ZFN), transcription activator like effector nucleases (TALEN), which have been proven to be efficient CCR5 gene editing tools [12–16]. In the year of 2013, Feng Zhang and George Church et al. developed Clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR associated nuclease 9 (CRISPR-Cas9) gene modification technique, which has resulted revolution in gene editing [17, 18]. CRISPR/Cas9 technology takes several advantages over ZFN and TALEN, such as easy design, high efficiency, etc. Hence, the technology has also been applied to mediate HIV-1 co-receptor editing [19–22]. Li et al. has reported that they have disrupted CCR5 in different CD4+ T cells, which has protected the edited cells from HIV-1 (R5-strains) infection. Meanwhile, analyzing the most effective 3 sgRNAs and their corresponding 15 potential off-target sites revealed that no significant editing efficacy in these sites [23]. For the co-receptor CXCR4, Hou et al. has proven that the disruption of CXCR4 by CRISPR/SpCas9 in genome level confers the edited cells resistant to HIV-1(X4-strains) infection and no obvious effects on off-target and proliferation, Wang et al. has verified the phenomenon with CXCR4 modification by CRISPR/SaCas9 [20, 24]. Some works about simultaneous editing of HIV-1 co-receptor CCR5 and CXCR4 by CRISPR/Cas9 have also been reported, Yu et al. and our previous work have confirmed that the two genes could be disrupted simultaneously in genome level and the edited cells could resist R5-tropic strain and X4-tropic strain concurrently with survival advantage over unedited cells under mixed HIV-1 infection pressure [25, 26]. Recently, Xu et al. have reported they have utilized CRISPR/Cas9 to edit CCR5 gene in HSPCS and transplant the cells to a acute lymphoblastic leukemia (ALL) and HIV-1 bearing 27 years old male in China. The ALL was complete remission, and donor cells carrying the disrupted CCR5 persisted for more than 19 months without related adverse events. The percentage of edited cells with CCR5 ablation increased by a small degree during a short of anti retroviral therapy pause [27].

The CRISPR/Cpf1 technology was first reported in 2016 and the nuclease Cpf1 has three different origins, namely Acidaminococcus (AsCpf1), Lachnospiraceae bacterium (LbCpf1), Francisella novicida (FnCpf1) [28, 29]. Compared with the well-known CRISPR/spCas9, CRISPR/Cpf1 displays many difference, such as the protospacer-adjacent motif (PAM) sequence of Cpf1 is 5′-TTTN-3′ but not 5′-NGG-3′ of spCas9, which makes the sgRNAs more specific. The size of nuclease Cpf1 is much smaller than SpCas9, indicating the feasibility of the expressing the system in adenovirus (Adv) or adenovirus associated virus (AAV). The sgRNA using by Cpf1 to bind target sequence consists of CRISPR RNA (crRNA) only without trans-activating CRISPR RNA (tracrRNA) in spCas9, for the nuclease can process the maturity of crRNA by itself, which means the multiple-target design will be much more easier than spCas9 [28, 30]. The CRISPR/Cpf1 technology, like CRISPR/spCas9, has also been utilized in biological and medical study. For example, Dai et al. has edited TRAC and PDCD1 gene in the generation of modular CAR-T cells [31]. Wang et al. has successfully adopted the technology to modify plants phenotype associated genes [32]. However, no report has been published about HIV-1 co-receptor CCR5 or CXCR4 editing with CRISPR/Cpf1.

In this study, we screened and identified two CRISPR/AsCpf1 using sgRNAs with high specificity to target CCR5. With different CD4+T cells (Tzm.bl, SupT1-R5, primary CD4+T cells), we demonstrated that CRISPR/AsCpf1 could ablate CCR5 gene efficiently in all cell types and confer the edited cells resistant to R5-tropic strain but not X4-tropic strain infection. Meanwhile, the edited cells show survival advantage over unedited cells under R5-tropic strain infection pressure.

**Results**

**Lenti-AsCpf1 mediated CCR5 disruption in adherent CD4+ TZM.bl cell line**

To silence CCR5 expression, we designed 5 sgRNAs targeting different sites within CCR5 exon (Fig. 1a, b), and cloned the sgRNAs to lenti-AsCpf1 backbone after U6
promoter. The constructed plasmids and control plasmid were transfected into TZM.bl cell lines, an adherent cell line with high expression of CD4 and HIV-1 co-receptor, CCR5 and CXCR4 on cell surface. The transfected cells were added and incubated with 1 μg/ml puromycin for 24 h after the first day, the treated cells were collected and analyzed with T7E1 assay which cleaves DNA at distorted duplexes caused by mismatches. The related primers are listed in Additional file 1: Table S1. The result showed that the designed #4, #5 could cleave the 828 bp CCR5 PCR amplicon, while the blank and control showed no obvious cleaved bands (Fig. 2a), which indicated the lenti-AsCpf1 could mediate CCR5 ablation in TZM.bl cell line with sgRNA #4 and #5 efficiently. DNA sequencing has also been performed to analyze the indels (insert/deletion) efficacy of each sgRNA, the results showed the #4 leaded 15 out of 20 editing (75.0%) and #5 induced editing ratio was 12/20 (60.0%) (Fig. 2b), which implied that the #4 might be more efficiently in CCR5 disruption than #5. Furthermore, to identify if the CCR5 genome level ablation mediated CCR5 protein expression regulation on cell surface, flow cytometry was performed to measure the CCR5 protein on cell surface (Fig. 2c). The results displayed that #4, #5, compared with positive control, exhibited 44.23% and 31.78% CCR5 protein down-regulation respectively. To investigate if the effect of CCR5 editing with AsCpf1 or spCas9 was different, two CRISPR/spCas9-sgRNA (#1, #2) were constructed and compared with CRISPR/AsCpf1-#4, among which the sgRNAs overlap each other (Additional file 2: Fig S1). According to the results, the CCR5 editing ratio of Cas9-#1, #2 ranged from 60.0% to 70.0%, which showed no difference comparing with AsCpf1-#4 (60.0%). Besides, the top 3 ranking candidates of all predicted off-target sites for Cas9-#1, #2 have been analyzed with DNA sequencing and T7E1, and the results revealed that the selected off-target sites for analysis exhibited no disruption. The results indicated that both AsCpf1 and spCas9, with target sequence overlap each other in this study, could ablate CCR5 in genome level efficiently.

CCR5 ablation could be induced by AsCpf1-sgRNA packaged lentivirus in SupT1-R5 cells

The major target cell infected by HIV-1 in human body is primary CD4+ T cells, a suspension cell type. Thus, we further tested if the CRISPR-AsCpf1 could work in SupT1-R5 cells, which is a common model of suspension cells in HIV-1 study. The AsCpf1-sgRNA -#4/#5
Lenti-CRISPR/AsCpf1 mediated the disruption of CCR5 in TZM.bl cells. 

a 5 sgRNAs for CCR5 editing by CRISPR/AsCpf1 were screened and identified by T7 endonuclease 1 (T7E1).

b The amplicons of #4 and #5 were subjected to DNA sequencing by ligating with PGEM-T easy vector.

c The protein level of CCR5 on cell surface were analyzed by flow cytometry at the third day post-transfection.

**Fig. 2** Lenti-CRISPR/AsCpf1 mediated the disruption of CCR5 in TZM.bl cells. 

- a 5 sgRNAs for CCR5 editing by CRISPR/AsCpf1 were screened and identified by T7 endonuclease 1 (T7E1).
- The amplicons of #4 and #5 were subjected to DNA sequencing by ligating with PGEM-T easy vector.
- The protein level of CCR5 on cell surface were analyzed by flow cytometry at the third day post-transfection.
and control were packaged into lentivirus respectively, and then the virus were used to transduce the SupT1-R5 cells. The treated cells were collected and performed with T7E1 assay after 3 days post transduction. Obvious CCR5 cleaved fragments could be observed of #4 or #5, but the control or blank showed no disruption (Fig. 3a). Consistently, DNA sequencing was performed for the PCR product of CCR5 (Fig. 3b), the result showed 11/20 of #4 (55%) and 8/20 of #5 (40%) were processed with indels after the corresponding lentivirus transduction, which indicated the high efficacy of lenti-CRISPR/AsCpf1-#4/#5 mediated CCR5 disruption. In a parallel assay, 7 days after transduction, the whole protein was collected and analyzed with immunoblotting to detect the CCR5 protein change in each treated cells (Fig. 3c), the result showed that the CCR5 protein levels were markedly down-regulated with transduction of #4 or #5 lentivirus compared to control. In addition, the flow cytometry was also performed to analyze the CCR5 expression on cell surface. The results showed that the CCR5 protein expression level were down-regulated by 34.55% in #4 and 30.91% in #5 compared with control (Fig. 3d). The same results could be observed with another suspension CD4+ T cell-Jurkat T cell line (data not shown).

The edited CD4+ T cells inhibits HIV-1 infection

To test whether the CCR5 modification would render the cells resistant to HIV-1 infection, two HIV-1 variants (HIV-1YU-2, HIV-1NL4-3) were used to infect the edited cell types above and the corresponding control cells. As shown in (Fig. 4a) at day 5 post-infection, the modified TZM.bl cells showed significant R5-tropic variant (HIV-1YU-2) resistance at MOI 0.1 or 0.5 compared with control. However, the edited TZM.bl cells exhibited no difference while X4-tropic variant (HIV-1NL4-3) was used at MOI 0.5. The same experimental phenomenon could be observed with the suspension cell type-SupT1-R5 from day 1 to day 5 post-transduction (Fig. 4b). The results indicated that the disruption of CCR5 mediated by lenti-CRISPR-#4/#5 in TZM.bl and SupT1-R5 cells, rendered the cells resistant to R5-tropic HIV-1 infection but not X4-tropic infection. In addition, as the CXCR4 was another co-receptor of HIV-1 infection, we have also screened and edited CXCR4 in genome successfully with CRISPR/AsCpf1 in TZM.bl cells, and the co-receptors CCR5 and CXCR4 have been ablated simultaneously by co-transfection the corresponding plasmids (AsCpf1-CCR5-#4 and AsCpf1-CXCR4-#2) into TZM.bl cells. Interestingly, the CCR5 and CXCR4 edited cells could resist HIV-1YU-2 and HIV-1NL4-3 infection concurrently, while the single co-receptor edited groups showed no difference when compared with control (Additional file 3: Fig S2).

The edited SupT1-R5 cells gain a selective advantage over unedited cells

In order to rule out the effect of multiple rounds of infection in unedited cells, we performed a selective advantage assay by exposing the treated SupT1-R5 cells to R5-tropic HIV-1YU-2 and prolonging the culture time for 14 days. The culture supernatant (day 1, day 7, day 14) of each group were collected and analyzed with HIV-1 gag p24 measurement (Fig. 5a). From the result, the unmodified control showed a continuous increase of the HIV-1 titer, but the modified groups exhibited a mild rise. Meanwhile, T7E1 assay was performed to evaluate the CCR5 ablation of edited cells and control cells at day 0, day 7 and day 14 (Fig. 5b), the result demonstrated that the edited groups (#4 and #5) displayed an obvious increase of lower migrating bands corresponding to cleavage products, but the control group showed no editing persistently. The results suggested that the #4, #5 modified SupT1-R5 cells were enriched during HIV-1YU-2 infection, and the cells were conferred with increased HIV-1 resistance ability as the infection processing. Thus, the conclusion could be drawn that the #4 or #5 edited SupT1-R5 was resistant to HIV-1YU-2 infection and exhibited a survival advantage over unedited control cells.

CCR5 disruption protects primary CD4+ T cells from HIV-1 infection by Adv-CRISPR/AsCpf1

After successfully editing CCR5 in suspension CD4+ SupT1-R5 cells with AsCpf1- #4/#5 contained lentivirus, we attempted to disrupt CCR5 in primary CD4+ T cells with the packaged lentivirus. Unfortunately, we were not able to disrupt the CCR5 obviously even with different MOI. Thus, we tried to package the AsCpf1-sgRNAs into adenovirus, the transduction efficacy of the adenovirus was more than 30% when compared with control at MOI 100 (Fig. 6a). To evaluate the disruption efficacy of CCR5 by the Adv-AsCpf1-#4/#5 in primary CD4+ T cells, the T7E1 assay was performed after the cells were transduced with adenovirus (MOI 100) at day 5 (Fig. 6b). The lower migrating fragments in #4 and #5 revealed that Adv-AsCpf1-#4/#5 could successfully edit the primary CD4+T cells. DNA sequencing for the CCR5 amplicon was used further to analyze the indels (Fig. 6c). From the statistic, 8/28 of #4 (28.6%) and 6/25 of #5 (24.0%) were edited with different indels respectively, which indicated the Adv-AsCpf1-#4/#5 could disrupt CCR5 in primary CD4+T cells efficiently. To test if the CCR5 ablation in genome level resulted in protein level
Fig. 3 The CRISPR/AsCpf1-#4/#5 expressing lentivirus induced genome editing of CCR5 in SupT1-R5 cells. 

a 300 ng PCR products of #4 and #5 or control were analyzed by T7E1 assay after lenti-CRISPR/AsCpf1 mediated CCR5 disruption at day 3.

b The amplicons of #4 and #5 were treated with poly A adding kit, and the products were further ligated with PGEM-T easy vector and analyzed with DNA sequencing.

c CCR5 protein level of the edited SupT1-R5 cells were detected at day 7 with 10 µg load after packaged lentivirus transduction.

The CCR5 expression on cell surface were assessed with FACS.
regulation, the immunoblotting showed the obvious down-regulation of CCR5 when compared with control (Fig. 6d). Since the CCR5 gene of primary CD4⁺ T cells could be edited by Adv-AsCpf1-#4/#5, like TZM.bl and SupT1-R5 cell line, we next examined whether the edited primary CD4⁺ T cells could defense HIV-1 infection. R5-tropic HIV-1 \textsubscript{YU-2} and X4-tropic HIV-1 \textsubscript{NL4-3} were utilized to infect the cells (Fig. 6e), the supernatants of each group were collected and measured from day 1 to day 7 with p24 ELISA. The results suggested that the CCR5 editing by Adv-AsCpf1-#4/#5 in primary CD4⁺ T cells protected the cells from R5-tropic HIV-1 \textsubscript{YU-2} but not X4-tropic HIV-1 \textsubscript{NL4-3} infection compared with that of cells transduced with empty vector. Moreover, we further investigated whether the editing of CCR5 inhibited proliferation or induced apoptosis of the primary CD4⁺ T cells. The cell counts were monitored post-transduction from day 1 to day 7 and the apoptosis rates were analyzed for each group at day 5. The results displayed that no obvious proliferation and apoptosis change were detected in CCR5 editing groups (#4, #5) when compared with control cells, which indicated that the CCR5 disruption took no effect on general biological process of cells (Fig. 6f, g).

Specific disruption of CCR5 by CRISPR/AsCpf1 does not induce detectable off-targets effects

While the CRISPR/AsCpf1 could mediate efficiently CCR5 disruption in different cells, as the sgRNAs sequence of #4 and #5 we screened and identified may have some similarity with other locus in genome, it may result in inaccurate recognition by sgRNAs of #4 and #5 and cleavage by nuclease AsCpf1. Thus, the off-target analysis of #4 and #5 was performed and 2 off-target sites for #4 and 6 for #5 have been predicted with online tool CCTop (http://crispr.cos.uni-heidelberg.de/). All of the predicted sites were amplified by PCR with genome template from primary CD4⁺ T cells. The amplicons were analyzed with T7E1 assay. As shown in (Fig. 7b), all of the sites exhibited no significant difference of fragments compared to control and blank. Meanwhile, DNA sequencing of all the sites were performed, while no cleavage of these sites were detected (Fig. 7a). The results indicated that the selected sgRNAs were high specific and safe, and no obvious off-targets
Discussion
In this study, we screened and identified two sgRNAs (#4, #5) targeting CCR5 gene by CRISPR/AsCpf1. The CRISPR/AsCpf1-#4/#5 have been clarified that they could work efficiently to disrupt CCR5 gene in different cell types of HIV-1 study (TZM.bl, SupT1-R5, primary CD4+ T cells). The edited cells, compared with unedited control group showed that they could resist R5-tropic HIV-1 but not X4-tropic HIV-1 infection in all cell types. Meanwhile, we have verified that the CCR5 edited SupT1-R5 cells took the selective advantage over unmodified control cells. Furthermore, the safety of the Adv-CRISPR/AsCpf1-#4/#5 mediated CCR5 ablation...
in primary CD4+ T cells also have been investigated and the disruption has no effect on cell proliferation and apoptosis, the off-target analysis of the two sgRNAs demonstrated that all the predicted sites showed no cleavage, which indicated that the disruption was relatively safe for gene therapy in the future.
From previous study, CCR5 is the major co-receptor of HIV-1 entering target cells at the early stage, many studies involving CCR5 disruption have been reported by different gene editing tools, such as ZFN, TALEN and the newly developed CRISPR/Cas9. However, the ZFN and TALEN show many limitations, such as complex design, high off-target rate and not available to most of labs. Several studies associated with CCR5 ablation by CRISPR/Cas9 have been reported recently, for example, Xiao et al. have reported that the CRISPR/SaCas9, a Cas9 version from Staphylococcus aureus, could efficiently edit the CCR5 in different cell types including human.

| Gene    | Target Sequence       | PAM       | Coordinate          | Off-target analysis |
|---------|-----------------------|-----------|---------------------|---------------------|
| CCR5    | TCAGGAGGACGACGACGACG  | TTTA      | chr3:46373046-46373072 | 0/120               |
| CCR2    | TTAAGGAGGACGACGACGACG| TTTA      | chr3:46537707-46357733 | 0/120               |
| MACROD2 | TTAGGAGGACGACGACGACG | TTTA      | chr20:15448182-15448208 | 0/120               |
| CCR5    | GAAATAACATGTCAACCTTGG | TTTA      | chr3:46373188-46373214 | 0/120               |
| DMRT1   | GTAATAAATGTCAACATTTT | TTTT      | chr9:879878-879904  | 0/120               |
| BRINP1  | GAAATAAATGTCAACCTTCA | TTTT      | chr9:119239030-119239056 | 0/120               |
| PAICSP7 | GAAAAACATGTCAACATTTT | TTTA      | chrX:94656240-94656266 | 0/120               |
| RPS4XP12| GAAAAAAATGTCAACATTTT | TTTA      | chr11:121267972-121267998 | 0/120               |
| LRRG66  | GAAAAATATGTCAACATTTT | TTTT      | chr4:52002969-52002995 | 0/120               |
| FAM47B  | CAAGACATTGTGTTAACTTTT | TTTT      | chrX:34971306-34971332 | 0/120               |

**Fig. 7** Specific disruption of CCR5 by CRISPR/AsCpf1 has no effect on detectable off-target. 

- **a** The predicted off-target sites by online tool-CCTop were listed, and the possible indels of each amplicons were analyzed with DNA sequencing.
- **b** The T7 endonuclease 1 was further utilized to cut the products.
CD34+-hematopoietic stem/progenitor cells, and the results in humanized mice have clarified that CCR5 disruption via lenti-CRISPR/SaCas9 renders CD4+ T cells survival from HIV-1 infection [33]. A recent clinical study by Xu et al. via transplanting the CCR5-edited stem cells by CRISPR/SpCas9 to a patient with AIDS and leukemia, has been verified that the CCR5 edited cells could resist HIV-1 infection, for a short time suspension of HAART promoted the editing efficacy. Meanwhile, the disruption efficacy in cells could be detected for 19 months without unspecific targeting and other side effects. Nevertheless, attempts should be made to study some other gene tools with much more easy design and higher safety. CRISPR/Cpf1 is a member of CRISPR/Cas family. Compared to the well known CRISPR/Cas9, the technology displays some differences, such as the recognition of target can be processed without the tracrRNA, for the Cpf1 displays not only endonuclease activity but also exonuclease activity, which assists the mature of crRNA. The PAM sequence of CRISPR/Cpf1 is 5′-TTN-3′ but not 5′-NGG-3′ in CRISPR/SpCas9 or 5′-NGGRT-3′ in CRISPR/SaCas9, which makes the CRISPR/Cpf1 may take advantages over CRISPR/Cas9 in design and off-target concern. According to the studies of Benjamin et al. and Daesik et al., the endonuclease Cpf1 exhibits much more sensitivity to mismatches than spCas9, with no more than two single mismatches resulting in almost complete loss of Cpf1 activity at positions 1–18 (numbered 1–23 in the 5′ to 3′ direction). Furthermore, the two studies have also verified that the potential off-target sites of AsCpf1 were less than spCas9 [34, 35]. Interestingly, in predication of the off-target sites about the two sgRNA(#4, #5) using by CRISPR-AsCpf1 online, only few of the sites were presented (< 10), other than our previous work about CRISPR/Cas9 in off-target sites predication with tens to hundreds candidates. Besides, although the results of the top 3 off-target candidates analysis about spCas9-#1, #2 in our study exhibited no disruption, it should be considered that the sites we selected were less than 2% of all the predicted sites. So our results further indicated the superiority of CRISPR/AsCpf1 in gene editing, and proved that the sgRNAs (#4, #5) we selected are quite specific. Also, our study about CCR5 disruption by CRISPR/Cpf1 was the first report of CRISPR/Cpf1 editing of HIV-1 co-receptor, our results demonstrated the efficacy and specificity of the CRISPR-AsCpf1 sgRNAs (#4, #5). Moreover, the vector we used while investigating the efficacy of the system in primary CD4+ T cells was adenovirus, the vector, not like lentiviral system integrating into genome that may induce cytotoxicity, was relatively safe and efficient.

It should also be noted in this study that CXCR4, as another co-receptor of HIV-1, can function as the co-receptor for R5-tropic HIV-1 infection at late stage or some HIV-1 variants can enter cells via CXCR4 directly but not CCR5, just as the results of our study with additional experiments about CCR5 and CXCR4 editing simultaneously in TZM.bl resisting R5-tropic and X4-tropic HIV-1 infection, which makes the specific CCR5 disruption only may not be sufficient. In addition, the CRISPR/AsCpf1-#4/#5 have not been investigated in vivo, such as in humanized mouse.

Conclusions
In this study, we screened and identified two sgRNAs (#4, #5) for CRISPR/AsCpf1 editing CCR5 in genome level with high specificity. We demonstrated that the CRISPR/AsCpf1-#4/#5 could ablate CCR5 efficiently in different cell types (TZM.bl, SupT1-R5, primary CD4+ T cells), and CCR5 disruption protects the cells from R5-tropic HIV-1 but not X4-tropic HIV-1 without unspecific targeting and effects on cell proliferation and apoptosis. In addition, we clarified that the CCR5 disrupted cells took selective advantages over unedited cells during R5-tropic HIV-1 infection. Our study suggests that CCR5 disruption mediated by CRISPR/AsCpf1 can confer an effective and safe approach in HIV-1 gene therapy and may have potential of clinical application.

Methods
sgRNA design and lenti-CRISPR/AsCpf-sgRNA construction
The sgRNAs design for CCR5 editing was based on online tool-CCTop (http://crispr.cos.uni-heidelberg.de/). 5 top ranking score of sgRNAs targeting CCR5 exon were selected and synthesized with 5′-AGAT and 5′-AAAA overhangs. For the construction of lenti-CRISPR/AsCpf-sgRNA, the plasmid lenti-AsCpf1 (Addgene #PY108) was digested with endonuclease BsmB1 (Fermentas), the overhangs were filled in with T4 ligase.

Cell lines culture and primary CD4+ T cell isolation
The studied TZM.bl and SupT1-R5 cells were cultured as previously described [20, 26, 33]. The primary CD4+ T cells were isolated from healthy donors with Negative CD4+ T Cells Isolation Kits (STEMCELL) according to the manufacturer's instructions. The isolated primary CD4+ T cells were further cultured in CD3/CD28 coated 6-well plate with RPMI containing 10% FBS and 20 IU/ml recombinant human interleukin-2 (IL-2) was added.

Virus (lentivirus, adenovirus) production and transduction
Lentivirus were produced with co-transfection of HEK293T cells with lenti-CRISPR/AsCpf1-#4/#5 or control, psPAX2, pMD2.G. After 60 h, the supernatants were collected and incubated with Lenti-X Concentrator...
and the concentrated virus were further cryopreserved at −80 °C. The target cells were transduced with concentrated lentivirus (MOI of 30) in the presence of 8 μg/ml polybrene (Sigma). The production of Adv-CRISPR/AsCpf1-#4/#5 or control were performed by HANBIO company (Shanghai, China). The Adv-AsCpf1 mediated transduction of primary CD4+T cells was according to instruction of manufacturer. Namely, 5 × 10^5 cells/well in 12-well plate were infected with the adenovirus (MOI = 100) by spinning at 200g for 2 h, and the culture medium was replaced with fresh RMPI1640 after 12 h virus transduction. The transduced primary CD4+T cells were further cultured until downstream analysis.

T7 endonuclease 1 (T7E1) cleavage assay and DNA sequencing

Genomic DNA of the treated cells were extracted with TIANamp Genomic DNA kit (TianGen) as the manufacturer’s instruction. The specific targets were amplified with PCR, and the amplicons were gel extracted. For the T7E1 assay, 300 ng of the amplicon was annealed and then incubated with T7 endonuclease 1 (NEB) for 30 min at 37 °C. The cleavage was analyzed by 1.5% agarose gel electrophoresis. To further analyze the amplicon, the purified fragment was cloned into pGEM-T Easy vector (Promega), the detail of indels (insert/delete) were identified and compared with control.

Luciferase activity assay and p24 gag detection by ELISA

The modified cells were infected with R5-tropic HIV-1 YU-2 or X4-tropic HIV-1NL4-3 at MOI = 0.1 or 0.5, and the cells were further cultured after virus wash. For the detection of luciferase activity, the infected cells were washed and then lysed with 100 μl lysis buffer, and further measure-ment was performed with BrightGlo Luciferase assay according to the instruction of manufacturer (Promega). The supernatants of treated cells at indicated days post-infection were collected and the titer of virus were deter-mined by p24 ELISA kit (ZeptoMetrix) as its instruction.

Flow cytometry analysis

To assess the disruption of CCR5 on the cell surface of TZM.bl, 5 μl FITC-conjugated mouse anti-human CCR5 antibody (Biolegend) were used and co-incubated with 5 × 10^5 cells at the third day post-treatment for 15 min. The incubated cells were further washed for 3 times with 1 × PBS, and then evaluated with CytoFLEX S (Beckman). The data were analyzed by CytExpert software. To evaluate the effects of adv-CRISPR/AsCpf1 mediated CCR5 disruption on the apoptosis of primary CD4+T cells, the apoptosis kit (US EverBright) was utilized. The primary CD4+T cells were incubated with 5 μl Annexin V-PE and 10 μl 7-AAD for 30 min, and the cells were analyzed with FCS as the instruction of manufacturer.

Selective advantage analysis of CCR5-disrupted SupT1-R5 cells after R5-tropic HIV-1 challenge

To assess whether the CCR5 edited cells were conferred with survival advantage over unedited control cells, the SupT1-R5 cells were transduced with associated lentivirus at MOI = 30. After 4 days post-transduction, the treated cells were infected with R5-tropic HIV-1_YU-2 for 8 h. The cells were further cultured for 14 days after the infected cells were washed and replenished with fresh medium. The genomic DNA at day 0, 7, 14 post-infection were extracted and assessed with T7E1 assay. Consistently, the virus titer of HIV-1 were detected by p24 ELISA at day 1, 7 and 14.

Off-target analysis

To evaluate the possible off-target of the selected sgRNAs (#4, #5), the sgRNAs were analyzed by online tool CCTop (http://crispr.cos.uni-heidelberg.de/). According to the predicted results, all of the off-target sites for sgRNAs (#4, #5) were amplified with PCR and assessed with T7E1 assay. For further investigating the possible in/dels, the amplicons of treated cells were performed with DNA sequencing.

Statistical analysis

Statistical analyses were performed using SPSS software, version 16.0. Differences between two groups were analyzed using the Unpaired Student’s t test. *p < 0.05; **p < 0.01, and ***p < 0.001 represents significant differ-ences. NS represents not significant.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s13578-020-00444-w.
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