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<td>Author(s)</td>
<td>Chen, Jia; Huang, Dane; Chen, Wei; Guo, Chaowan; Wei, Bo; Wu, Chongchao; Peng, Zhou; Fan, Jun; Hou, Zhibo; Fang, Yongsheng; Wang, Yifei; Kitazato, Kaio; Yu, Guoying; Zou, Chunbin; Qian, Chuiwen; Xiong, Sheng</td>
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<td>© 2014 Chen et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.</td>
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Linker-Extended Native Cyanovirin-N Facilitates PEGylation and Potently Inhibits HIV-1 by Targeting the Glycan Ligand

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Abstract
Cyanovirin-N (CVN) potently inhibits human immunodeficiency virus type 1 (HIV-1) infection, but both cytotoxicity and immunogenicity have hindered the translation of this protein into a viable therapeutic. A molecular docking analysis suggested that up to 12 residues were involved in the interaction of the reverse parallel CVN dimer with the oligosaccharide targets, among which Leu-1 was the most prominent hot spot residue. This finding provided a possible explanation for the lack of anti-HIV-1 activity observed with N-terminal PEGylated CVN. Therefore, linker-CVN (LCVN) was designed as a CVN derivative with a flexible and hydrophilic linker (Gly-Ser), at the N-terminus. The N-terminal α-amine of LCVN was PEGylated to create 10 K PEG-aldehyde (ALD)-LCVN. LCVN and 10 K PEG-ALD-LCVN retained the specificity and affinity of CVN for high mannose N-glycans. Moreover, LCVN exhibited significant anti-HIV-1 activity with attenuated cytotoxicity in the HaCaT keratinocyte cell line and MT-4 T lymphocyte cell lines. 10 K PEG-ALD-LCVN also efficiently inactivated HIV-1 with remarkably decreased cytotoxicity and pronounced cell-to-cell fusion inhibitory activity in vitro. The linker-extended CVN and the mono-PEGylated derivative were determined to be promising candidates for the development of an anti-HIV-1 agent. This derivatization approach provided a model for the PEGylation of biologic candidates without introducing point mutations.

Introduction
Currently, over 30 million people worldwide are infected with human immunodeficiency virus type 1 (HIV-1) and 1.5 million-1.9 million people died from AIDS-related causes at the end of 2011; approximately 2.2 million-2.8 million people become infected each year, of whom 95% live in low- and middle-income countries [1,2]. Microbicides are promising alternative agents for the prevention of HIV-1 transmission [3]. Cyanovirin-N (CVN), a protein originally isolated from the freshwater cyanobacterium Nostoc ellipsosporum, exhibits specific and potent anti-HIV activity by binding with high affinity to the glycans present on gp120 and gp41 [4,5]. CVN irreversibly inactivates both laboratory-adapted and wild type HIV-1 strains during the viral entry stage. The antiviral effects of CVN also include the inhibition of cell-to-cell fusion, virus-to-cell fusion and cell-to-cell transmission [6]. CVN has generated interest as a promising new generation of microbicides characterized by specific and potent activity, a novel mechanism of action and unusual physicochemical stability.

CVN may be useful in two different clinical applications, either as a targeting agent or as a topical microbicide, to prevent the sexual transmission of HIV-1 by providing a method for female control over the HIV/AIDS epidemic [7]. Because of its cyanobacterial origins, CVN exhibits the limitations that are typical of such proteins in pharmaceutical applications, including a short plasma half-life, proteolysis and immunogenicity. Polyethylene glycol (PEG) is a well-studied polymer that is utilized as a biological compatibles by attenuating both immunogenicity and cytotoxicity, to increase the half-life and to alter the biodistribution [8].
homogeneity and facilitating the preservation of bioactivity, site-specific PEGylation at the N-terminus or on random amines on the side chains of CVN has resulted in inactive molecules [7,9]. The only PEGylated version of CVN that is bioactive is the mutant Q62C, in which glutamine 62 was replaced with a cysteine, and the extra free sulfhydryl was site-specifically PEGylated with maleimide-activated PEG [7]. The in vitro anti-HIV-1 activity of the Q62C mutant was approximately 50% that of wild type (WT) CVN. The 20 kDa PEG-CVN Q62C conjugate demonstrated approximately 80% of the activity observed with CVN WT. The 30 kDa conjugate had nearly no activity. From these reported data, we hypothesized that N-terminal residues and certain lysine residues might exist in or near the glycan binding sites of CVN.

To confirm this hypothesis, molecular docking and experimental approaches were utilized to investigate the binding selectivity of CVN to oligosaccharides with various structures. The protein-ligand complexes of CVN 3GXY with high mannose N-glycans were also docked and analyzed to further characterize the hot spot residues in CVN. This structure-function relationship study suggested that Leu-1 in the N terminus was the most important hot spot residue for binding to Man$_7$GlcNAc$_2$ glycans. Therefore, a rational PEGylation process was designed to avoid blocking the N-terminal hot spot residues.

The well documented (Gly4Ser)$_3$ molecule is a flexible hydrophilic linker peptide that has been utilized to fuse 2 independent polypeptides into a protein with multiple domains and functions [10]. Based on the merits of the (Gly4Ser)$_3$ linker peptide and the conundrum of CVN PEGylation, we extended the addition to the 2 intramolecular disulfide bonds in CVN makes the other known proteins. The extremely low sequence homology in E. coli efficiently expressed in the cytoplasm of E. coli was constructed to efficiently produce soluble LCVN in E. coli. The N-terminal residue of CVN with (Gly4Ser)$_3$ to create linker-CVN (LCVN) and performed site-specific PEGylation of LCVN at the N-terminal amine group using mPEG-aldehyde (ALD). We hypothesized that this PEG-linker-CVN might preserve the bioactivity of CVN by separating the large PEG group from the CVN active site as well as facilitate the preparation of highly homogenous PEGylated products. This strategy avoided introducing a point mutation into the primary sequence of CVN that could alter its bioactivity.

There is no sequence homology greater than 8 contiguous amino acids or 20% of the total sequence between CVN and any other known proteins. The extremely low sequence homology in addition to the 2 intramolecular disulfide bonds in CVN makes the artificial production of this protein in E. coli difficult [6]. In a previous study, biologically functional CVN was efficiently expressed in the cytoplasm of E. coli after fusion to small ubiquitin-related modifier (SUMO) coupled with a hexahis-}

dinidine tag [11]. Utilizing this strategy, the fusion gene his$_6$-sumo-linker-cvn was constructed to efficiently produce soluble LCVN in E. coli. The N-terminal PEGylation of LCVN was performed to create 10 K PEG-ALD-LCVN, a site-specific PEG conjugate of LCVN. Subsequently, the gp120, gp41 and oligosaccharide binding characteristics of LCVN and 10 K PEG-ALD-LCVN were evaluated. The anti-HIV-1 activity and cytotoxicity of these 2 CVN derivatives were determined by MTT and syncytiumformation assays to elucidate the effects of the linker peptide on oligosaccharide binding and the anti-HIV-1 activity of CVN and to explore the feasibility of site-specific PEGylation of pharmaceutical proteins via the (Gly4Ser)$_3$ extension.

### Results

**CVN Targeting to 24 Potential Oligosaccharides that were Selected from a Pool of 6 Types of N-glycans by Molecular Docking**

Molecular docking was performed to determine the binding selectivity of CVN for oligosaccharides with various structures and to clarify the binding modes. The crystalization data for CVN (Figure S1) [http://www.rcsb.org/pdb] was utilized to dock 53 oligosaccharide targets that were selected to represent 6 types of carbohydrate structures. These oligosaccharides included 13 complex N-glycans, 13 high mannose N-glycans, 13 branched and linear oligomannoses, 3 hybrid N-glycans, 3 N-glycans with a core pentasaccharide or related moiety and 8 oligosaccharides originating from glycolipids (Figure 1). The consensus scores (CS) for the 53 oligosaccharides are listed in Table 1. High scores indicated improved biological activity. Several complex and hybrid N-glycans exhibited a high CS. Oligosaccharides No. 1, 2, 4, 10 and 42 were characterized with a CS >0.5. Most high mannose N-glycans (No. 14–26) had a CS between 0.2 and 0.5. Several oligomannose moieties had a CS of zero.

To characterize the CVN binding potential, dozens of high-scoring and commercially available oligosaccharides were selected from each oligosaccharide category for further investigation. According to this priority principle, 7 oligosaccharides (No. 1–2, 4–6, 9 and 10) that belonged to the maximum CS group were selected to represent the complex N-glycans. Nine oligosaccharides (No. 14–17, 19–22 and 24) that belonged to the medium CS group were selected to represent the high mannose N-glycans. From all the branched and linear oligomannoses in the minimum CS group, glycans No. 27–30 and 43 were selected to represent N-glycans with a pentasaccharide core, and No. 46–47 and 52 represented oligosaccharides originating from glycolipids. In total, 24 oligosaccharides (asterisks, Figure 1) were selected to represent the diverse carbohydrate structures in the centrifugal ultrafiltration-HPLC assays.

**Specific Recognition of CVN by Non-reduced Terminal Man$_{\sim 9}$GlcNAc$_2$ Glycans**

To verify the MOE docking data and evaluate the minimum oligosaccharide structure required for high-affinity binding to CVN, centrifugal ultrafiltration-HPLC was performed using fluorescence-labeled oligosaccharides (PA-oligosaccharides). The structures of the 24 oligosaccharides utilized in this study are indicated with asterisks in Figure 1. Before testing LCVN binding to the selected oligosaccharides, the optimum pH for the binding assay was determined using PA-heptasaccharide (No. 19, Figure 1) in 50 mM 2-(N-morpholino)ethanesulfonic acid (MES, pH 5.0), 50 mM sodium phosphate (pH 6.0) and 50 mM Tris-HCl (pH 7.0, 8.0 or 9.0). Similar assays were performed to optimize the reaction time from 20 to 100 min in 50 mM Tris-HCl (pH 7.0). Maximal CVN-oligosaccharide binding was achieved at pH 7.0–9.0 (Figure 2A) after a 60 min incubation period. The binding of CVN to its targets was stable after incubating for 60 min or longer (Figure 2B). Therefore, the binding of CVN to the selected PA-oligosaccharides was assayed at room temperature for 60 min at pH 7.0.

CVN exclusively bound to high mannose N-glycans (No. 15–22) without recognizing other oligosaccharides, including oligomannose (No. 27–30), complex N-glycans (No. 1–10) and oligosaccharides with a pentasaccharide core (No. 43) (Figure 2C). Glycolipid oligosaccharides (No. 46, 47 and 52) did not interact with CVN. The high mannose N-glycans (No. 16–17 and 19–22) bound to CVN with a binding ratio greater than 85% (Figure 2C).
CVN bound with high affinity to Man\textsubscript{7}GlcNAc\textsubscript{2} with 1 non-reduced terminal Man\textsubscript{a}1–2Man moiety in the D1 or D3 arm (No. 17 and 19). Man\textsubscript{8}GlcNAc\textsubscript{2} with 2 exposed Man\textsubscript{a}1–2Man moieties in the D1, D2 or D3 arms (No. 20–22) or Man\textsubscript{9}GlcNAc\textsubscript{2} with 3 Man\textsubscript{a}1–2Man moieties (No. 16) exhibited a strong interaction with CVN. The binding ratio decreased to 37% for oligosaccharide No. 15, which only has 1 non-reducing terminal Man\textsubscript{a}1–2Man moiety in the D1 arm provided by Man\textsubscript{6}GlcNAc\textsubscript{2}. These data clearly suggested that CVN exhibited strict specificity for high mannose N-glycans by recognizing the extended carbohydrate structure of the non-reducing terminal Man\textsubscript{a}1–2Man moieties in the D1, D2 or D3 arms provided by Man\textsubscript{7–9}GlcNAc\textsubscript{2} (No. 16–17 and 19–22).

Oligosaccharide Man\textsubscript{a}1–2Man Binding to the Hot Spot Residues was Critical for the Oligosaccharide-CVN Interaction

To characterize the structural interactions between CVN and its ligands, oligosaccharides No. 22 and 28 were selected to represent an active and a less active group, respectively, for further analysis. The active pocket of CVN 3GXY is located in the gap between the \(\beta\) sheet of chains A and B. The extended structure of the non-reducing terminal mannose moieties in the oligosaccharide bound to the active pocket of CVN, whereas the other part of the oligosaccharide was exposed outside of the pocket (Figure 3). An overview of the protein-ligand interactions for oligosaccharides No. 22 and 28 is presented in Figure 3. For oligosaccharide No. 22, hydrogen bonds formed between the ligand and Leu-1, Lys-3, Thr-7, Glu-23, Thr-25, Tyr-29 and Glu-101 in the active site.
Table 1. MOE docking values for 53 oligosaccharides in the training set.

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x, the docking energy for 3GXY; y, the docking energy for 2PYS. doi:10.1371/journal.pone.0086455.t001

(Figure 3A); for oligosaccharide 28, hydrogen bonds were present between the ligand and Leu-1, Gly-2, Lys-3, Thr-7, Thr-25 and Asn-93 (Figure 3B). Although both the active and the less active oligosaccharides interacted with 6 residues in CVN, the binding energy for oligosaccharide No. 22 was −71.32 kcal/mol, which was lower than that for oligosaccharide No. 28 (−44.16 kcal/mol). The multiple hydrogen bond interactions and the lower binding energy for the oligosaccharide No. 22-CVN complex corresponded with the greater activity of this particular oligosaccharide.

Even with the differences in binding energy, oligosaccharides No. 22 and 28 interacted with the same 4 amino acids, Leu-1, Lys-3, Thr-7 and Thr-25 (Figure 3). For oligosaccharide No. 22, 5 amino acids (Leu-1, Lys-3, Thr-7, Thr-25 and Glu-101) interacted with the extended non-reducing terminal mannose moieties. In contrast, only 1 residue (Thr-25) interacted with the mannose of oligosaccharide No. 28, indicating that the multiple mannoses interacting with multiple residues contributed to the robust activity of oligosaccharide No. 22.

To further characterize the hot spot residues in CVN and its derivatives, protein-ligand complexes of CVN 3GXY with all 6 high mannose N-glycans (No. 16–17 and 19–22) were docked and analyzed by MOE. The frequencies at which the hot spot residues were directly involved in the interactions are presented in Figure 4A. In total, 12 amino acids (Leu-1, Gly-2, Lys-3, Glu-6, Thr-7, Tyr-9, Gln-23, Thr-25, Gly-27, Asn-93, Asp-95 and Glu-101) in CVN were involved in binding to oligosaccharide ligands. Most of these residues could form hydrogen bonds with the ligands. All 6 oligosaccharides bound to Leu-1, and over half of the ligands bound to Gly-2, Lys-3, Glu-6, Glu-23, Asn-93 and Glu-101. The 3D structure formed by the 12 hot spot residues (the binding residues) was defined as a new binding pocket in CVN for oligosaccharides. This binding pocket could be utilized as a reference for further molecular docking studies to select novel ligands.

To illustrate the type of mannose structure that was specifically targeted in the oligosaccharide-CVN (3GXY) binding model and to evaluate the consistency with the centrifugal ultrafiltration-HPLC assay, all the binding moieties in the 6 oligosaccharides were analyzed and summarized as the number of total targeting residues in CVN and the number of Manα1–2Man-targeting
residues involved in binding to each oligosaccharide (Figure 4B). For CVN, 4–10 residues bound to each oligosaccharide with 2–4 Man\textsubscript{1,2}Man-targeting residues. In general, 63% of the binding occurred between hot spot residues and Man\textsubscript{1,2}Man moieties. Oligosaccharide No. 19 was targeted by 10 amino acids, with 3 of these hot spot residues targeting the extended non-reducing terminal Man\textsubscript{1,2}Man moieties. These data were consistent with the centrifugal ultrafiltration-HPLC study, suggesting that CVN specifically recognized the extended non-reducing terminal Man\textsubscript{1,2}Man moieties provided by Man\textsubscript{7,9}GlcNAc\textsubscript{2}.

Because oligosaccharide No. 19 was targeted by most hot spot residues, the 3D model of CVN 3GXY binding to this oligosaccharide is illustrated in Figure 4C. The 10 binding residues were Leu-1, Gly-2, Glu-6, Tyr-9, Gln-23, Thr-25, Gly-27, Asn-93, Asp-95 and Glu-101 (highlighted in light pink). The hydrogen bonds are indicated by dashed lines. Three of these hot

Figure 2. Binding specificities of CVN to PA-oligosaccharides. The optimum (A) pH and (B) reaction time for CVN binding to PA-heptasaccharide (No. 19, Figure 1) were determined via centrifugal ultrafiltration-HPLC, and subsequently the (C) binding activity (specificity) of CVN and the 24 oligosaccharides was measured. The right panel in (C) depicts the non-reducing terminal Man\textsubscript{1,2}Man moieties in the D1, D2 or D3 arms of the Man\textsubscript{7,9}GlcNAc\textsubscript{2} glycans that participated in the binding. Two independent experiments were performed for each PA-oligosaccharide, and the binding activity is presented as the average of the duplicate assays.
doi:10.1371/journal.pone.0086455.g002
spot residues, Gln-6, Tyr-9 and Glu-23, targeted the extended non-reducing terminal Man$_{2-2}$Man moieties. This model provided insight into the CVN-oligosaccharide interaction.

Both LCVN and the PEGylated Product Retained the Specificity and Affinity of CVN for Man$_{7-9}$GlcNAc$_{2}$ Glycans

The structure-function relationship study suggested that Leu-1 in the N terminus of CVN was the most important hot spot residue for binding to Man$_{7-9}$GlcNAc$_{2}$ glycans and that most of the N1 to

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**Figure 3. The mode of CVN binding to oligosaccharides.** An overview of the mode of CVN binding to oligosaccharides (A) No. 22 and (B) No. 28 is presented. The fuzzy blue blob indicates ligand exposure to the solvent. For oligosaccharide 22, which exhibited strong binding, Leu-1, Lys-3, Thr-7, Glu-23, Thr-25, Tyr-29 and Glu-101 were involved in the protein-ligand interaction. For oligosaccharide No. 28, which was characterized by weak binding, the docking simulation indicated that hydrogen bonds formed between the ligand and Leu-1, Gly-2, Lys-3, Thr-7, Thr-25 and Asn-93. doi:10.1371/journal.pone.0086455.g003
with binding ratios of 100%. The binding ratio for oligosaccharide 21, it bound to high mannose N-glycans No. 16, 17, 20 and 22 slightly decreased affinity to high mannose N-glycans No. 19 and (No. 15–22) over other types of oligosaccharides (No. 1–10, 27–30, specificity. The centrifugal ultrafiltration-HPLC assays indicated carbohydrate structures was determined to characterize the glycan CVN to both gp120 and gp41.

ALD-LCVN maintained the glycan-specific binding of native CVN. These data suggested that both LCVN and 10 K PEG-ALD-LCVN were determined to characterize their glycan binding ability. As a positive control, CVN bound to glycosylated gp41 (Figure 6A) and gp120 (Figure 6B) in a dose-dependent manner but did not exhibit any affinity for non-glycosylated gp41 (Figure 6C) or gp120 (Figure 6D). CVN bound more tightly to gp41 than to gp120. LCVN and 10 K PEG-ALD-LCVN had the same binding specificity to the glycosylated substrates (Figure 6A–6D), but their affinities were slightly decreased compared with CVN. These data suggested that both LCVN and 10 K PEG-ALD-LCVN maintained the glycanspecific binding of native CVN to both gp120 and gp41.

The binding of LCVN and 10 K PEG-ALD-LCVN to the 24 oligosaccharides (asterisks, Figure 1) selected to represent diverse carbohydrate structures was determined to characterize the glycan specificity. The centrifugal ultrafiltration-HPLC assays indicated that the 2 proteins exclusively recognized high mannose N-glycans (No. 15–22) over other types of oligosaccharides (No. 1–10, 27–30, 43, 46, 47 and 52) (Figure 6E). Although LCVN exhibited a slightly decreased affinity to high mannose N-glycans No. 19 and 21, it bound to high mannose N-glycans No. 16, 17, 20 and 22 with binding ratios of 100%. The binding ratio for oligosaccharide No. 15, which has 1 non-reducing terminal Man2,3Man moiety in Man3GlcNAc2, and LCVN decreased to <40%, which corresponds with the data for CVN (Figures 6E and 2C). These data clearly suggested that both LCVN and 10 K PEG-ALD-LCVN retained the affinity of CVN for specific oligosaccharides. All 3 versions of CVN had identical target specificity and consistent binding potency to the extended carbohydrate structure of the non-reducing terminal Man2,3Man moieties in the D1, D2 or D3 arms provided by Man2,3GlcNAc2 glycans.

LCVN Cytotoxicity was Significantly Lower and was Further Attenuated by PEGylation

As promising microbicide candidates, CVN and its derivatives would be applied topically on human skin and/or mucosa. The HaCaT keratinocyte cell line and the MT-4 T lymphocyte cell line were utilized to evaluate the in vitro cytotoxicity of LCVN and its derivatives. Both LCVN and 10 K PEG-ALD-LCVN exhibited significantly less cytotoxicity than native CVN (Table 2). For HaCaT cells treated for 24 h, the CC50 values for LCVN and 10 K PEG-ALD-LCVN were 8.59±1.31 μM and >12.00 μM, respectively. The value for native CVN was 1.74±0.22 μM, suggesting that the cytotoxicity of LCVN was approximately 1/6 that of CVN. For 10 K PEG-ALD-LCVN, the cytotoxicity was <1/10 that of CVN. After treating the cells for 48 h, LCVN exhibited less cytotoxicity than CVN, and 10 K PEG-ALD-LCVN was approximately 1/6 as cytotoxic as native CVN. The MT-4 T lymphocyte cell line was sensitive to the various versions of CVN, with both LCVN and the PEGylated product exhibiting significantly reduced toxicity (Table 3). The cytotoxicity of LCVN was approximately 1/4 that for CVN. For 10 K PEG-ALD-LCVN, the ratio was 1/42. These data suggested that LCVN was remarkably less cytotoxic than native CVN and that the in vitro toxicity was further reduced by PEGylation. The addition of the 15-aa extension (linker) at the N-terminus of CVN decreased the cytotoxicity. After PEGylation, this toxicity decreased by approximately 40-fold. These results suggested that further examination of these modified proteins for potential anti-HIV activity would be beneficial.

LCVN Exhibited More Potent Anti-HIV-1 Activity in the Nanomolar Range

The anti-HIV activities of both LCVN and 10 K PEG-ALD-LCVN were determined by the WST-1 method. As presented in Table 3, LCVN and the PEGylated product protected MT-4 cells from infection with HIV-1/IIIB. The IC50 of LCVN was 14.36±1.35 nM and that for native CVN was 21.83±2.79 nM, suggesting that LCVN possessed more anti-HIV activity than native CVN. Although 10 K PEG-ALD-LCVN exhibited significantly less anti-HIV activity than both LCVN and CVN, the cytotoxicity to MT-4 cells was also significantly decreased. Considering activity and cytotoxicity, both LCVN and 10 K PEG-ALD-LCVN exhibited improved safety profiles. The SI values for LCVN and 10 K PEG-ALD-LCVN were approximately 5-fold higher than that for CVN. Among the 3 CVN derivatives, LCVN exhibited the most potent anti-HIV activity, the highest SI value and the lowest cytotoxicity. The derivative 10 K PEG-ALD-LCVN retained the potent anti-HIV activity of CVN in the nanomolar range and possessed the lowest cytotoxicity, which was similar to that of AZT.

LCVN that were involved in the binding are colored in light pink, and their ligands are depicted in green. The hydrogen bonds are illustrated as dashed light pink lines.

Figure 5. Structural schematics of LCVN and the PEGylated product. The N-terminal glycine of LCVN and the serine-leucine joint between the linker and the CVN sequence are indicated. The blank rectangle represents the residual polypeptide of CVN. 10 KD mPEG-ALD was selectively reacted with the N-terminal ω-amine of LCVN at different pKa values to create 10 K PEG-ALD-LCVN.

do:10.1371/journal.pone.0086455.g005
Figure 6. The binding of LCVNs to HIV-1 envelope proteins and oligosaccharides. The binding of LCVNs to glycosylated (A) gp41 and (B) gp120 and non-glycosylated (C) gp41 and (D) gp120 was determined by ELISA. CVN served as the positive control, and BSA was utilized as the negative control. The data points represent the mean ± SD of independent triplicate experiments. (E) The LCVN binding activity (specificity) with the
LCVN Exhibited Significantly Greater Fusion Inhibitory Activity than CVN

HIV-1 spreads efficiently, primarily via cell-to-cell fusion. To determine the fusion inhibitory activity of LCVN and its PEGylated conjugate, a cell-to-cell fusion assay was performed according to the method of Tochikura et al. [12]. In this syncytium formation assay, MOLT-4 cells were co-cultured with HIV-producing MOLT-4/IIIB cells for 24 h in the presence of LCVN or its derivatives. The results demonstrated that the fusion inhibitory activity of LCVN was significantly greater than that of CVN; PEGylation further enhanced this activity in the high and medium dose groups (Figure 7). In the lower dose group (28 nM), 10 K PEG-ALD-LCVN exhibited less inhibitory activity than LCVN, but its activity remained higher than that of CVN. These data confirmed the merits of enhancing bioactivity and attenuating toxicity by adding an N-terminal linker (LCVN) and suggested that the PEG groups in 10 K PEG-ALD-LCVN might interfere with the fusion of HIV-1-positive cells to normal ones by steric hindrance. This hypothesis and these data provided insight into the mechanism of HIV-1 transmission and will aid the discovery and development of novel fusion inhibitory compounds.

Discussion

A microbicide must not damage the mucosa because such damage may increase the risk of HIV-1 infection. Significant efforts had been made to reduce the toxicity and increase the anti-HIV activity of microbicide candidates, such as CVN. In this study, the structure-function relationship for CVN was investigated to identify a more rational structure for a CVN derivative for further optimization. After the first docking of CVN 3GXY and 2PYS to the 53 oligosaccharides, most complex and hybrid N-glycans exhibited a high CS with low or no binding in the experimental assays. By analyzing the protein-ligand binding modes, the nitrogen in the oligosaccharide was determined to interact with the amino acids of CVN, accounting for the majority of the binding free energy. CVN did not interact with oligomannose (No. 27) and had a low affinity for Man5GlcNAc2 glycans with 2 or 3 reducing Man and Man7GlcNAc2 glycans with 1 reducing end. In contrast with the docking data, Glu-41 was the most important hot spot residue, with a frequency of 83%, suggesting that Glu-41 might be one of the critical binding residues in the parallel CVN dimer; PEGylation further enhanced this activity in the high and medium dose groups (Figure 7). In the lower dose group (28 nM), 10 K PEG-ALD-LCVN might interfere with the fusion of HIV-1-positive cells to normal ones by steric hindrance. This hypothesis and these data provided insight into the mechanism of HIV-1 transmission and will aid the discovery and development of novel fusion inhibitory compounds.

Table 2. The cytotoxicity of LCVN and its PEGylated derivatives in HaCaT cells (mean±SD, n = 3).

<table>
<thead>
<tr>
<th>Cytotoxicity (CC50, μM)</th>
<th>24 h</th>
<th>48 h</th>
</tr>
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<tbody>
<tr>
<td>LCVN</td>
<td>8.59±1.31</td>
<td>1.75±0.14</td>
</tr>
<tr>
<td>10 K PEG-ALD-LCVN</td>
<td>&gt;12.00</td>
<td>6.11±0.29</td>
</tr>
<tr>
<td>CVN</td>
<td>1.74±0.22</td>
<td>1.18±0.09</td>
</tr>
</tbody>
</table>

CC50, 50% cell inhibition concentration.
doi:10.1371/journal.pone.0086455.g002

Table 3. The anti-HIV-1 activity and MT-4 cytotoxicity of LCVNs (mean±SD, n = 3).

<table>
<thead>
<tr>
<th></th>
<th>IC50 (nM)</th>
<th>CC50 (nM)</th>
<th>SI</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCVN</td>
<td>14.36±1.35</td>
<td>647.15±93.42</td>
<td>45.06</td>
</tr>
<tr>
<td>10 K PEG-ALD-LCVN</td>
<td>176.40±25.03</td>
<td>7181.07±861.72</td>
<td>40.71</td>
</tr>
<tr>
<td>CVN</td>
<td>21.83±2.79</td>
<td>169.56±13.63</td>
<td>7.77</td>
</tr>
<tr>
<td>AZT</td>
<td>36.55±5.64</td>
<td>&gt;4482</td>
<td>&gt;122.63</td>
</tr>
</tbody>
</table>

IC50, 50% viral inhibitory concentration; CC50, 50% cell inhibitory concentration; SI, the ratio of CC50 to IC50.
doi:10.1371/journal.pone.0086455.t003

24 oligosaccharides was assayed by centrifugal ultrafiltration-HPLC. Two independent experiments were performed for each PA-oligosaccharide, and the binding activity is presented as the average of the duplicate assays.
doi:10.1371/journal.pone.0086455.gp006
**Figure 7. LCVN fusion inhibitory activity.** (A) Phase-contrast micrographs were obtained 24 h after co-culturing normal MOLT-4 and HIV-1-positive cells in the presence of LCVN. The HIV-1-induced multinucleated giant cells are indicated by black arrows. (B) The relative fusion inhibition rates (%) for LCVN and its derivatives were calculated (**P < 0.01 vs. CVN, * P < 0.05 vs. CVN). doi:10.1371/journal.pone.0086455.g007
A comparison of the docking and the crystallography data for the 2 types of CVN dimers with the experimental glycan/oligosaccharide binding assay data suggested that (i) the simulations of the CVN-oligosaccharide complexes were highly consistent with the interaction mode suggested by crystallography and (ii) the simulation had high fidelity with the experimental assay. An analysis of the docking of CVN to the oligosaccharides suggested that (i) Leu-1, Gly-2, Lys-3, Thr-2, Glu-23, Asn-93 and Asp-95 were particularly important for the binding of reverse parallel CVN to its targets, with Leu-1 being the most predominant hot spot residue; and (ii) in parallel CVN dimers, Glu-41, Asn-42, Asn-53, Thr-57 and Lys-74 were the most important hot spot residues. This binding model provided a possible explanation for the reduction in bioactivity for N-terminal PEGylated CVN and also supported our strategy of utilizing PEGylation in conjunction with a linker to separate the large PEG group from the oligosaccharide binding site in CVN. Furthermore, it was deduced that CVN might be more inclined to form a reverse parallel structure in solution because N-terminal PEGylated CVN was reported to be fully inactive, but all the hot spot residues are located in the center of parallel CVN [7].

Gp120 is responsible for target cell tropism and viral attachment via an interaction with the cell surface receptor CD4 and the co-receptors CCR5 or CXCR4 [21, 22]. The binding of gp120 to its receptor and co-receptor induces a cascade of refolding events in gp41 that bring the viral and cell membranes together [23]. CVN binds with high affinity to glycosylated gp120 and gp41. The stronger binding to gp41 than gp120 suggested that CVN interferes with the process of HIV receptor recognition and membrane fusion. In addition, CVN may act at the stage of membrane fusion by binding to gp41, thereby inhibiting the gp41 refolding events.

Based on the knowledge above, LCVN was designed and further modified at the N-terminus using a site-specific method and 10 KD mPEG-ALD to maintain the integrity of the binding sites in CVN. LCVN exhibited greater anti-HIV-1/IIIB activity in the MTT and fusion inhibitory assays and lower cytotoxicity than native CVN. The enhanced bioactivity of LCVNs may have resulted from (i) the (Gly2Ser)3 linker contributing to correct folding and the proper biological function of the linker-tagged protein [24]; (ii) the increased molecular weight and the enhanced thermostability that amplified the steric hindrance of CVN, increasing the fusion inhibitory activity [25]; and (iii) the hydrophilicity of the flexible linker, which could interfere with the structural integrity of the viral envelope. It would be interesting to fully elucidate the mechanism by which LCVN exhibited enhanced anti-HIV-1 activity.

The anti-HIV-1/IIIB activity of 10 KD mPEG-ALD was significantly decreased in the WST-1 assay, but this LCVN derivative exhibited more potent fusion inhibitory activity than native CVN. WST-1 is a substrate that measures the metabolic activity of viable cells, so the WST-1 assay indirectly evaluates the anti-viral activity of a tested compound. The fusion inhibitory assay directly measures the antiviral activity of CVN because this assay simulates the actual process of HIV-1 transmission between normal and HIV-1-infected cells. Therefore, the fusion inhibitory assay is more pertinent for studying the antiviral mechanism of action of CVN. The fusion inhibitory activity of 10 K PEG-ALD-LCVN was greater than that of CVN. These data strengthened the therapeutic potential for 10 K PEG-ALD-LCVN and suggested that the molecular weight of CVN and its derivatives might be crucial for antiviral activity. The steric hindrance effect might be involved in the antiviral mechanism of action of 10 K PEG-ALD-LCVN. Other groups have reported that derivatives with a higher molecular weight, such as the engineered CVN dimer, have greater antiviral activity than monomeric forms of CVN [26]. An interesting recent report demonstrated that fusions of CVN and the gp41 membrane-proximal external region (MPER) peptide joined by a (Gly2Ser)3 linker (where x is 4 or 6) induce specific, irreversible lysis of pseudotyped HIV-1 virions and fully infectious HIV-1 virions [27]. Both fusion components, CVN and MPER, are required for the cell-free virolysis of HIV-1. Considering the merits of the PEG-linker-CVN that we demonstrated here, it would be interesting to create a chimERIC CVN derivative with N-terminal PEGylation and a C-terminal MPER fusion and to explore the potential of this novel agent, PEG-linker-CVN-linker-MPER, as a tri-acting virucidal entry inhibitor of HIV-1.

It would be prudent to test the anti-HIV-1 activity of LCVN and its PEGylated conjugate on additional HIV-1 strains. Here, we only utilized HIV-1/IIIB as a model strain to evaluate the potential of LCVN and the PEG-LCVN conjugate. It has been well validated that CVN irreversibly inactivates a broad range of laboratory-adapted HIV strains and clinical isolates with different tropisms at the nanomolar level [28, 29]. For example, the EC50 values for CVN against the HIV-1 R5 strains HIV-1(Ba-L), HIV-1(Ada-M) and HIV-1(89.6) are 17 nM, 1.7 nM and 36.8 nM, respectively [28, 29]. Our data demonstrated that LCVN and 10 K PEG-ALD-LCVN retained the specificity and potency of the anti-HIV-1 activity of CVN and suggested the therapeutic potential of these CVN derivatives against R5 and other HIV-1 strains.

**Conclusions**

A linker-extended CVN derivative, LCVN, and its PEGylated product, 10 K PEG-ALD-LCVN, were rationally designed and constructed after molecular docking and experimental approaches. Twelve residues were determined to be involved in the targeting of the reverse parallel CVN dimer to oligosaccharide ligands, among which Leu-1 (the N-terminal leucine in the B chain of the CVN dimer) was the most important hot spot residue. Eight residues were suggested to interact with the oligosaccharides in the parallel CVN dimer, with Glu-41 being one of the most important hot spot residues. Both LCVN and 10 K PEG-ALD-LCVN retained the oligosaccharide specificity of CVN binding to high mannose N-glycans with >1 terminal Manα1→Man moieties in gp120 and gp41. It was exciting that the CVN derivatives exhibited potent anti-HIV activity with remarkably decreased cytotoxicity. The improved biological compatibility of these 2 CVN derivatives suggested that these modifications could produce promising microbicidal candidates and provide a template for a universal strategy for the PEGylation of biologic candidates without introducing point mutations. The CVN-oligosaccharide interaction analysis provided a possible explanation for the loss of anti-HIV-1 activity with N-terminal PEGylated CVN and suggested the dominant conformation of the CVN dimer in solution.

**Materials and Methods**

**Chemicals, Reagents and Media**

All the chemicals and reagents were obtained from Sigma (St. Louis, MO, USA) unless otherwise stated. All the media and supplements, including fetal bovine serum (FBS), were purchased from Invitrogen (New York, NY, USA) unless otherwise stated. Recombinant LC3N and the PEGylated product 10 K PEG-ALD-LCVN (Figure 5) were prepared in-house by a process modified from Gao et al. [11].
Cell Culture
The immortal human HaCaT keratinocyte cell line, purchased from the China Center for Type Culture Collection (Wuhan University, Wuhan, China), was propagated in Eagle’s minimal essential medium (MEM) supplemented with 10% FBS, 1.0 mM sodium pyruvate, 0.1 mM non-essential amino acids and 1.5 g/L sodium bicarbonate. The MT-4 T lymphocyte cell line (NIH AIDS Reagent Program, Germantown, MD, USA) was cultured in RPMI 1640 medium containing 10% FBS and 0.22% sodium bicarbonate. The cells were cultured at 37 °C in a humidified atmosphere with 5% CO2.

Enzyme-linked Immunosorbent Assay (ELISA)
The target binding activity of LCVN to HIV-1 gp120 and gp41 was determined using an ELISA-like assay as described previously [30]. Briefly, 100 ng of gp120 or gp41 derived from HIV-1 PM1H2 (BPB, Beijing, China) was coated onto a 96-well plate, which was subsequently incubated with serially diluted LCVN or 10 K PEG-ALD-LCVN. The bound proteins were detected using a rabbit-anti-CVN polyclonal antibody (1:10000, in-house preparation) and goat-anti-rabbit IgG-HRP (1:3000) followed by chromogenic development with 3, 3', 5, 5'-tetramethylbenzidine (TMB) at λ450.

Molecular Docking
CVN exists predominantly as a monomer in solution and as a domain-swapped dimer in crystals, producing both parallel (head-to-head) and reverse parallel dimer conformations. The crystal structures of parallel CVN (2PYS) and reverse parallel CVN (3GXY) (Figure S1) were downloaded from the RCSB Protein Data Bank (http://www.rcsb.org/pdb) [31]. To determine the optimum scoring function, docking experiments were performed for all the protein-ligand complexes using 3 molecular docking platforms (Flex_X [32], CDocker (DS 2.1, Accelrys) [33] and MOE [32]). The active sites were identified using the crystallographic ligand for all the datasets. All the docking experiments reported here were performed with the default parameters. Based on the ligand-protein binding energy, the 30 top-ranked docking poses were retained for further study.

The optimum docking program for CVN was selected using the root-mean-square deviation (RMSD) and the scores of the re-docking of the ligands to the known CVN crystal structures. After re-docking the ligands to 2PYS and 3GXY, the RMSD values with Flex_X ranged from 0.1 to 9.2, and the re-docking scores with CDocker were >0 kcal/mol. For MOE, the RMSD values were <1, and the docking scores were <-35.6 kcal/mol, indicating that MOE was the most appropriate program for CVN docking.

MOE has 2 docking placement methods, Alpha Triangle matcher and Proxy Triangle [34]. The active site was minimized using the AMBER 99 force field in MOE with the default parameters. All the oligosaccharides were docked, employing Triangle Matcher as the placement method and London dG as the first scoring function. The refinement was set to force field (AMBER 99), and the docked poses were energy-minimized in the receptor pocket. Affinity scoring was utilized to assess and rank the receptor-ligand complexes. A low docking score correlated with increased binding affinity.

To screen novel and bioactive targets of CVN, the 2D structures of 53 oligosaccharides were converted into 3D structures. With the energies minimized, the moieties were docked into the binding sites of 2PYS and 3GXY by MOE. A consensus score (CSi) was calculated from the normalized docking score of 3GXY (X_i) and 2PYS (Y_i) to objectively rank the 53 oligosaccharides with a high degree of confidence.

\[
X_i = \frac{-x_i - \min(-x)}{\max(-x) - \min(-x)} \\
Y_i = \frac{-y_i - \min(-y)}{\max(-y) - \min(-y)} \\
CS_i = \sqrt{X_i^2 + Y_i^2}
\]

Function X_i is the normalized docking score of an oligosaccharide to 3GXY, and x_i is the (un-normalized) docking score of an oligosaccharide to 3GXY. Function Y_i is the normalized docking score of an oligosaccharide to 2PYS, and y_i is the (un-normalized) docking score of an oligosaccharide to 2PYS. Min(-x) and Min(-y) are the minimum scores among the values determined for the 53 oligosaccharides. Max(-x) and Max(-y) are the maximum scores among the values determined for the 53 oligosaccharides.

Centrifugal Ultrafiltration-HPLC
A centrifugal ultrafiltration-HPLC assay was utilized to determine the oligosaccharide binding properties of the LCVNs as described by Katoh et al. [35]. Briefly, the LCVNs were incubated with pyridylaminated (PA)-oligosaccharides (Takara Bio Inc., Otsu, Shiga, Japan) in 50 mM Tris-HCl (pH 7.0) at room temperature for 60 min. The reaction mixture was centrifuged (14,000 x g for 15 min) in a centrifugal ultrafiltration tube (Sartorius Stedim, Boston, MA, USA) with a molecular weight cut-off value of 5,000 Daltons. The unbound PA-oligosaccharides (O_unbound) and the total amount of added PA-oligosaccharides (O_added) were quantified from the peak area detected on a TSKgel ODS 80 TM column (4.6 x 150 mm) (Tosoh Corporation, Tokyo, Japan) at λ250/400 for the coupled fluorescence group. The bound PA oligosaccharide (O_bound) was defined as the volume of O_added minus that of O_unbound. The binding activity was expressed as the ratio of O_bound to O_added and presented as % binding.

MTT Assay
The in vitro cytotoxicity of the LCVNs was determined via a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay in HaCaT cells. LCVNs were serially diluted from 12 μM to 0.375 μM, added to a monolayer of HaCaT cells in 96-well plates and incubated for either 24 or 48 h. The MTT solution was added for color development. The absorbance was measured at λ570/630, and the data were plotted to obtain the 50% cell inhibitory concentration (IC50).

WST-1 Assay
The in vitro anti-HIV-1 activity of the LCVNs was determined using the water-soluble tetrazolium salt 2-(4-iodophenyl)-3-(4-nitrophenoxy)-5-(2,4- disulphophenoxy)-2H-tetrazolium (WST-1) in a P3 laboratory. For this assay, 2-fold dilutions of LCVNs were mixed with HIV-1/IIIB (100 TCID50/50 μL), added to MT-4 cells (104 cells/100 μL/well) in 96-well microplates and incubated for 96 h. WST-1 was added to quantitate the number of viable cells at λ570/630. The cytotoxicities of the LCVNs in MT-4 cells were simultaneously determined in virus-free wells. Azidothymidine (AZT) was utilized as a positive control. The 50% inhibitory concentration (IC50), 50% cytotoxicity concentration (CC50) and selectivity index (SI, the ratio of CC50 to IC50) were determined.

Cell-to-cell Fusion Assay
Cell-to-cell fusion assays and cell-to-cell virus transmission assays, also known as syncytium formation assays, were performed with a co-culture system comprised of MOLT-4 (ATCC® CRL-1582™, Manassas, VA, USA) and MOLT-4/IIIB cells as...
previously described [36]. MOLT-4/IIIB cells are chronically HIV-1-infected MOLT-4 cells that allow HIV-1 replication and induce syncytium formation between uninfected MOLT-4 cells. MOLT-4 cells (2.5 × 10^5/250 μL) and MOLT-4/IIIB cells (2.5 × 10^5/250 μL) were mixed and seeded in 24-well plates (Falcon). LCVN and PEGylated LCVN were diluted 4-fold with RPMI-1640 containing 10% fetal calf serum and antibiotics. Individually cultured MOLT-4 and MOLT-4/IIIB cells and co-cultures of MOLT-4 and MOLT-4/IIIB cells without LCVN or PEGylated LCVN were utilized as controls, and co-cultured cells treated with LCVN or PEGylated LCVN at three different concentrations (452, 113 and 28 nM) were examined as the test conditions. Cytopathic effects indicated by the presence of syncytia were observed and recorded using a microscope after 24 h at 37°C with 5% CO₂. The number of viable cells was determined by trypan blue dye exclusion, and the fusion index (FI) was calculated as the following: \( \text{FI} = \left[ \frac{1}{\text{cell number in a test well}} - \frac{1}{\text{cell number in control well}} \right] \times 100 \). The fusion inhibition rate (FIR) (MOLT-4 only) was calculated as the following: \( \text{FIR} = \frac{\text{cell number in control well} - \text{cell number in a test well}}{\text{cell number in control well}} \) × 100. Where FI was the fusion index of the test sample and FI(0) was that of the co-cultured control [12].

References

Supporting Information

Figure S1 The three-dimensional (3D) structures of CVN utilized in this study. 2PYS, 1IIY and 2RDK are parallel domain-swapped dimers of CVN, and 3GXY and 3GZ8 are reverse-paralleldomain-swapped dimers of CVN. The structure coordinates of the protein-ligand complexes were retrieved from the Protein Data Bank (PDB) for the comparative molecular docking studies.

Table S1 Hot spot residues in CVN that target different ligands as determined by structural resolution approaches.

Author Contributions

Conceived and designed the experiments: SX JC. Performed the experiments: JC DH WC CG BW CW ZP. Analyzed the data: SX JC. Contributed reagents/materials/analysis tools: KK JF ZH YW GY CZ YF. Wrote the paper: SX JC DH WC.