Interaction between Cell-Penetrating Peptides and Acid-Sensitive Anionic Oligopeptides as a Model for the Design of Targeted Drug Carriers

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ABSTRACT: Overcoming the nonspecific cellular uptake of cell-penetrating peptides (CPPs) is a major hurdle in their clinical application. Using pH as the activation switch, histidine-glutamic acid (HE) dipeptide repeats were fused to CPPs to trigger the membrane-penetrating activity at mildly acidic pH environments (i.e., pH 6.5 or below) while masking the internalization at neutral pH (i.e., pH 7.0 or above). In this study, a series of recombinant GST-fusion proteins containing an HE oligopeptide sequence (i.e., (HE)$_n$ with $n = 8$, 10, or 12) and a cationic CPP (i.e., YG(RG)$_6$, YGR$_6$G$_6$, or Tat) were engineered for a pH-sensitive study comparing their cellular uptake and surface binding in cultured HeLa cells. Circular dichroism (CD) spectroscopy was performed to correlate differences between CPPs in secondary structure with the pH sensitivity. YGR$_6$G$_6$ with clustered arginine residues exhibited greater pH sensitivity in cellular uptake than YG(RG)$_6$ with separated arginine residues. Increasing the stretch of HE repeats decreased cellular uptake and surface binding for both YG(RG)$_6$ and YGR$_6$G$_6$. The ratio of cellular internalization at pH 7.5 vs 6.0 was not changed by the presence of serum. CD spectral data revealed that both (HE)$_{10}$-Tat and (HE)$_{10}$-YGR$_6$G$_6$ exhibited an unordered secondary structure, whereas (HE)$_{10}$-YG(RG)$_6$ adopted an antiparallel β-sheet conformation. This β-sheet conformation presumably stabilized the association of (HE)$_{10}$ with YG(RG)$_6$, leading to weakened pH sensitivity of (HE)$_{10}$-YG(RG)$_6$. On the other hand, the random-coiled structures, that is, (HE)$_{10}$-YGR$_6$G$_6$ and (HE)$_{10}$-Tat, both showed higher pH sensitivity as determined in cell experiments. The data presented in this study provide a basis for the future design of pH-sensitive HE-CPP carrier for targeted drug delivery.

KEYWORDS: cell-penetrating peptide, arginine-rich peptide, pH sensitive delivery, circular dichroism

INTRODUCTION

Rational utilization of the power of cell-penetrating peptides (CPPs) has been pursued for decades since HIV’s trans-activator of transcription (Tat) protein joined the fascinating family as the first unveiled CPP model.¹ CPPs, also known as protein transduction domains (PTDs) or membrane transduction peptides (MTPs), generally consist of 30 or less amino acids and are categorized as amphipathic or cationic depending on their sequences. Receptor-independent internalization and noninvasive entry into cells are the two major advantages that draw increasing attention to the field of CPP-based intracellular delivery systems. A variety of cargos, including bioactive molecules,² DNA/siRNA,⁴,⁵ quantum dots,⁶ and nanoparticles,⁷,⁸ have been delivered using different types of CPPs in vitro and in vivo. Additionally, some studies have shown that CPPs could also promote gastrointestinal absorption of protein drugs⁹,¹⁰ as well as direct brain delivery via intranasal administration.¹¹

However, lack of specificity is the major challenge for CPPs to be employed clinically.¹²⁻¹⁴ Many approaches have been investigated to confer CPPs with the ability of targeting delivery, but no break-through technology has yet been achieved. The nonspecific in vivo biodistribution of CPPs is

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mostly caused by the cationic characteristics of the oligopeptide.\textsuperscript{15} Therefore, one of the most promising approaches is to reversibly mask the positive charges in a CPP with a polyanionic counterpart. The selective activation of oligoanion-masked CPP can be achieved by specific proteolysis,\textsuperscript{16} light activation\textsuperscript{17} or differences in the microenvironment\textsuperscript{18} at target site. Recently, we have designed a recombinant co-oligopeptide containing Model Amphipathic Peptide (MAP, KLALKLALKAKALKLA) as the CPP sequence, and 10-mer of histidine-glutamic acid repeats ((HE)\textsubscript{10}) as a pH-sensitive blocking sequence used on an amphipathic CPP could also be applied for the design of activatable CPPs with either clustered or mixed positive charges in the oligopeptides. MAP is an amphipathic peptide that shows high binding and cellular internalization at pH 6.8 or below.\textsuperscript{19} In addition to solid tumor tissues, endosomal/lysosomal compartments\textsuperscript{21} as well as the acidic bioenvironments. By conjugating with a ligand, HE-CPP under an inactive form at mildly acidic pH could be activated under mildly acidic pH conditions. In cultured HeLa cells, it exhibited a low surface binding and cellular internalization at pH 7.4 but high surface binding and cellular internalization at pH 6.8 or below.\textsuperscript{19} Furthermore, the construct showed high accumulation and retention for up to 24 h near the tumor site in a xenograft breast cancer mouse model.\textsuperscript{20} Therefore, we wanted to determine if the same pH-sensitive masking of HE repeats, leading to restoration of the membrane-permeability of the CPP, could weaken the masking effect of HE repeats, leading to restoration of the membrane-permeability of the CPP.

In this study, the systematic design of anionic oligopeptides for neutralizing the cationic charges in oligoarginine CPPs is investigated. Oligoarginine exhibits many differences from amphipathic CPPs like MAP, such as different intracellular localization,\textsuperscript{23} and lacks a secondary structure.\textsuperscript{24} Similar to extracellular pH conditions in tumor tissues, the acidic microenvironments at the sites of infection or inflammation could weaken the masking effect of HE repeats, leading to activation of the CPP and the masking sequences. In addition, the cationic charge distribution of the CPP, either as clustered or evenly mixed sequence, may also affect the neutralizing efficiency.\textsuperscript{25} Using HE oligopeptide with various lengths, the masking effect on oligoarginine at a pH range between 6.0 and 7.5 was evaluated for the design of activatable CPPs with either clustered or mixed positive charges in the oligopeptides.

## EXPERIMENTAL SECTION

### Plasmid Construction and Production of Protein

The pGEX-4T-1 vectors (GE Healthcare Life Sciences, Piscataway, NJ) were utilized in this study to clone all plasmids. Similar to our previous design,\textsuperscript{19} the fusion protein contains glutathione S-transferase (GST) as a protein cargo fused to an HE oligopeptide sequence (HE\textsubscript{n} with n = 8, 10, or 12), a short pentaglycine linker (G\textsubscript{5}), and a arginine-rich CPP (YG(RG)\textsubscript{6}, YGR\textsubscript{5}G\textsubscript{6}, or Tat peptide (YGKRRRRQQRRR)). In order to allow for further characterization of the HE-CPP peptide sequences, a tyrosine residue was incorporated in the CPP sequence for quantitation, while a thrombin cleavage recognition sequence (Leu-Val-Pro-Arg-Gly-Ser, LVPR\textsubscript{GS}) was reserved between GST and HE oligopeptide. The ssDNAs of (HE)\textsubscript{n} (n = 8, 10, and 12), G\textsubscript{5}-YG(RG)\textsubscript{6}, and G\textsubscript{5}-YGR\textsubscript{5}G\textsubscript{6} were synthesized by ValueGene Inc. (San Diego, CA). After annealing ssDNAs into dsDNA, the dsDNAs coding for (HE)\textsubscript{n} (n = 8, 10, or 12) and G\textsubscript{5}-YG(RG)\textsubscript{6} or G\textsubscript{5}-YGRG\textsubscript{6} were successively introduced into the pGEX-4T-1 vector downstream from the amino-terminal of GST sequence through BamH I, EcoR I, and Not I restriction enzyme cleavage sites. Due to the two-step clones and the EcoR I restriction enzyme cleavage site, an extra amino acid residue, phenylalanine (Phe, F), was present between the HE sequence and G\textsubscript{5} linker. After ligation and transformation, colony PCR was performed, and then plasmids were duplicated overnight in Escherichia coli (E. coli) DH5\textalpha competent cells in lysogeny broth (LB) media containing 75 μg/mL ampicillin and 20 mM L-Glucose at 37 °C. After sequencing (GeneWiz, San Diego, CA), the plasmids were transformed into E. coli expression strain BL21. The recombinant protein was expressed and purified using glutathione (GSH) agaroose beads as the procedure described in our published report.\textsuperscript{19} Samples collected during purification were analyzed by SDS-PAGE followed by Coomassie blue staining. The recombinant proteins produced for this study are listed in Table 1.

| Table 1. Yield and Purity of Recombinant GST-Fusion Proteins |
|---------------------------------------------------------------|
| **oligopeptide sequence** | **M.W. (kDa)** | **Yield** (mg) | **pI** |
|---------------------------|----------------|---------------|-------|
| GST-(HE)\textsubscript{8}EFG\textsubscript{5}YG(RG)\textsubscript{6} | 30.15 | 49.4 | 6.77 |
| GST-(HE)\textsubscript{10}EFG\textsubscript{5}YG(RG)\textsubscript{6} | 30.68 | 43.5 | 6.62 |
| GST-(HE)\textsubscript{12}EFG\textsubscript{5}YG(RG)\textsubscript{6} | 31.22 | 25.5 | 6.53 |
| GST-(HE)\textsubscript{8}EFG\textsubscript{5}YGR\textsubscript{5}G\textsubscript{6} | 30.15 | 40.9 | 6.77 |
| GST-(HE)\textsubscript{10}EFG\textsubscript{5}YGR\textsubscript{5}G\textsubscript{6} | 30.68 | 39.1 | 6.62 |
| GST-(HE)\textsubscript{12}EFG\textsubscript{5}YGR\textsubscript{5}G\textsubscript{6} | 31.22 | 21.1 | 6.53 |
| GST-(HE)\textsubscript{8}EFG\textsubscript{5}YGRRKRRQRRR | 31.48 | 15.8 | 6.71 |
| GST-(HE)\textsubscript{10} | 30.34 | 39.7 | 6.20 |

*Amount per 500 mL LB media. The purity was >90% for all preparations, as determined by band density comparison in SDS-PAGE with Coomassie blue staining. *Isoelectric point (pI), calculated using an online software program.\textsuperscript{26}*

### Iodination of Protein

The purified recombinant GST-fusion proteins were radiolabeled with Na\textsuperscript{125}I (PerkinElmer Health Sciences, Chicago, IL) using the chloramine T method as previously described.\textsuperscript{26} \textsuperscript{125}I-Protein was purified by size exclusion chromatography using a gravity-flow column packed with Sephadex G50 beads (GE Healthcare Life Sciences). Reaction solution was loaded on the column prebalanced with PBS, and liquid flow-through was collected in fractions and monitored by measuring the radioactivity. Finally, fraction(s) containing \textsuperscript{125}I-proteins were stored at −20 °C until use.

### Cellular Uptake and Surface Binding Assay

Human cervical carcinoma (HeLa) cells were originally maintained at 37 °C in a 95% humidified atmosphere of 5% CO\textsubscript{2} in RPMI 1640 medium (Mediatech, Manassas, VA) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine, 50 μ/mL penicillin and 50 μg/mL streptomycin. Two or three days before cellular uptake and surface binding assay, appropriate number of HeLa cells was seeded into 6-well plates. After being grown to confluence, cell monolayers were treated with serum-free or serum-containing (1–10% FBS) medium for 10 min followed by incubation in serum-free or serum-containing (1–10% FBS) medium at pH 6.0, 6.5, 7.0, or 7.5 containing 5 μg/
ml $^{125}$I-labeled protein and protease inhibitor cocktail. After 1 h of treatment at 37 °C, the cell monolayers were washed three times with 1 mL of ice-cold PBS and subsequently treated with trypsin-EDTA at 37 °C for 3 min to detach the cells as well as remove the surface-bound (noninternalized) protein from the cell membrane. Cells were collected and centrifuged (PR-J centrifuge, International Equipment Company, Nashville, TN) at 1500 rpm for 3 min to separate intracellular (cell pellet) from surface bound (supernatant) fractions. The cell pellet was washed twice with ice-cold PBS and then dissolved in 1 mL of 1 N NaOH. The $^{125}$I-proteins were detected using a γ-counter (Packard, Downers Grove, IL), and total cell protein was determined by performing the micro-BCA assay with a Pierce micro-BCA assay kit (Thermo Fisher Scientific, Rockford, IL).

**Peptides Preparation and Circular Dichroism Spectroscopy.** In order to study the structure changes of the HE-fused CPP peptides under various pH conditions without the interference by GST protein domain, (HE)$_{10}$-YG(RG)$_n$ and (HE)$_{10}$-Tat were isolated from the respective GST-fusion proteins by column thrombin cleavage during GST agarose chromatography. After cleavage, the GST-free peptides were purified using HisPur Ni-NTA resins and dialyzed against 10 mM phosphate buffer followed by Tricine-SDS-PAGE analysis and sterilization by filtration (0.22 μm micro pore membrane). The peptide concentrations were determined by their absorption at 280 nm (UV 160U spectrophotometer, Shimadzu, Japan). For CD analysis, peptides were diluted to 25 mM in PBS at different pH ranging from 6.0 to 7.5. A J-815 CD spectrometer equipped with a PTC-423S/15 Peltier thermostatic cell holder (Jasco, Easton, MD) was utilized to measure the spectra. All spectra were recorded from 250 to 160 nm in a 10 nm quartz cuvette at 20 °C using a 0.5 nm data interval and a scanning speed of 100 nm/min. The CD spectra were plotted as an average ellipticity (mdeg) versus wavelength (nm).

**Statistical Analysis.** All data are given as mean ± standard deviation (SD). Mean and SD values were calculated from three independent measurements per treatment group. The Student’s $t$ test was exploited to perform comparison between data sets, where statistically significant differences were assigned for $P$ values of <0.05.

■ RESULTS

**Expression and Purification of Recombinant GST-Fusion Protein.** Plasmids encoding GST-(HE)$_n$-CPP were successfully constructed in pGEX-4T-1 vector (Figure 1A) according to the positive sequencing results. Recombinant GST-fusion proteins were expressed in *E. coli* BL-21 and purified by GSH affinity chromatography. Expression and purification was confirmed by Coomassie blue-stained SDS-PAGE, where gel of purified GST-(HE)$_n$-YG(RG)$_n$ (Figure 1B) and GST-(HE)$_n$-YG(RG)$_6$ (Figure 1C), $n = 8, 10$, and 12, showed gradient positions with varied HE repeats and approximately 90% purity based on the band densities. The yield of each protein obtained from 500 mL LB culture (Table 1) indicated that prolonging HE-repeats in the oligopeptide sequence decreased expression yield.

**Cell Uptake and Surface Binding Study.** To investigate the impact of the length of HE repeats on the pH sensitivity of CPP-incorporated recombinant GST-fusion proteins, cellular uptake and surface binding assay were performed. Confluent HeLa cell monolayers were grown in 6-well plates and treated with $^{125}$I-radiolabeled proteins in serum-free RPMI 1640 media, at various pH (6.0, 6.5, 7.0, and 7.5) for 1 h at 37 °C. The data (Figure 2) demonstrated that incorporation of HE oligopeptide significantly affected cell uptake and surface binding in a pH-dependent manner.

As seen in the cell uptake figures (Figures 2A, C, and E), the internalization of the fusion proteins increased with decreasing pH. The fusion protein with mixed arginine configuration, GST-(HE)$_{10}$-YG(RG)$_6$ showed a 5-, 4-, and 3-fold increase for $n = 8, 10, 12$, respectively, at pH 7.5 vs 6. With a clustered arginine configuration, GST-(HE)$_{10}$-YG(RG)$_6$ was more sensitive to pH changes, where a 5-, 6-, and 4 fold increase for $n = 8, 10$, and 12, respectively, was observed at pH 7.5 vs 6. The different lengths of (HE) repeats also affected the pH-dependent uptake patterns. The fusion proteins with 8 and 10 (HE) repeats showed higher uptake than the 12-(HE) repeat peptide across the entire pH range for both GST-(HE)$_{10}$-YG(RG)$_6$ and GST-(HE)$_{10}$-YG(RG)$_6$. Furthermore, the 12-(HE) repeat peptides showed less of a difference in the clustered vs mixed configuration than the 8- and 10-(HE) repeats. The surface binding of the fusion proteins showed a weaker pH-dependency than the cell uptake, where a 2-fold difference was noted at pH 7.5 vs 6 for all fusion proteins tested (Figures 2B, D, and F). As a control, the cell uptake and surface binding of GST-(HE)$_{10}$ was also measured. The results shown in Figure...
3 indicate that both binding and uptake of GST-(HE)10 was not dependent on pH, and was low across the pH range tested.

Comparison with Tat[47−57]. Cell-penetrating Tat[47−57] peptide is one of the most extensively studied CPPs. The cell internalization and binding of GST-(HE)12-Tat was compared to the blocked oligoarginine construct at a ∼1:1.5 ratio of cationic amino acids in the CPP sequence (R, K) to number of (HE) repeats (i.e., GST-(HE)12-Tat vs GST-(HE)10-YGR6G6). In the cell experiments (Figure 4), GST-(HE)12-Tat exhibited a stronger pH sensitivity in both cell uptake (4.4-fold increase at pH 6.0 versus pH 7.5) and surface binding (2.0-fold increase at pH 6.0 versus pH 7.5). Therefore, the pH sensitivity was comparable to the arginine in a clustered configuration, which is consistent with the Tat peptide sequence.

Influence of Serum on Cell Uptake and Surface Binding. To mimic physiological conditions, FBS was added in the media at different concentrations from 1−10%, and the cell uptake and surface binding were assayed at pH 6.0 and 7.5. As expected, the surface binding and uptake of both GST-(HE)10-YGR6G6 and GST-(HE)10-YG(RG)6 decreased with increasing concentrations of FBS (Figures 5 and 6). However, the ratio of either the surface binding or uptake at pH 6.0 versus 7.5 was similar in the presence of 1−10% FBS (Table 2). Therefore, serum did not affect the pH sensitivity of the constructs.

Circular Dichroism Spectroscopy. The secondary structure of the HE-peptides was analyzed by CD spectroscopy.28−31 As shown in Figures 7B and 8, (HE)10-YGR6G6 and (HE)10-Tat exhibited a random-coiled structure with a minimum at 198 nm. The (HE)10-YG(RG)6 peptide, on the other hand, exhibited a β-sheet conformation with a minimum at 218 nm and maximum at 195 nm (Figure 7A). The content of β-sheet conformation (HE)10-YG(RG)6 was evaluated at various pH, and the β-sheet content increased with increasing pH. The CD spectra showed an isosbestic point at 205 nm, consistent with a two-state transition from random coil to β-sheet. For both for (HE)10-YGR6G6 and (HE)10-Tat, pH had limited influence on the secondary structure (Figure 8).

## DISCUSSION

It is well established that positively charged amino acid residues, that is, arginine and lysine, are essential for both the binding and uptake of CPPs in mammalian cells. However, the polycationic charge is also responsible for the nonspecific cellular interaction and, consequently, the cytotoxicity of CPPs that limits the in vivo application in drug delivery. Therefore, the reversible masking of the positive charge has been a promising approach to avoid the nonspecific binding of CPPs to cell surface. In this report, we used oligoarginine peptides, with arginyl residues in a clustered (i.e., YGR6G6) or a mixed (i.e., YG(RG)6) configuration, as the CPP models to investigate the influence of the negative charges generated from a pH-dependent anionic HE oligopeptide on the binding and uptake of CPP in cultured HeLa cells.

Due to the number of glutamic acid residues, a longer HE oligopeptide should provide a higher masking efficiency, which
should yield a lower cellular binding of the CPP. In our preliminary studies, we found that oligopeptides with a ∼1:1 ratio of HE-to-cationic amino acid (R/K) showed a lack of masking effect at pH 7.5, with lower pH sensitivity. Therefore, in this study, an excess of negative charges in HE repeats was designed to ensure a complete neutralization of the six positive charges in YG(RG)6 or YGR6G6 at pH 7.5. As shown in Figure 2E, both YG(RG)6- and YGR6G6-fused proteins showed a decrease in cell uptake and surface binding with an increase of the length of HE oligopeptide. When linked to a 12-mer HE oligopeptide, the cell absorption capacity of YG(RG)6 and YGR6G6 was immensely restrained, leading to a minimal cell uptake and a poor pH sensitivity in the pH range from 7.5 to 6.5 (Figure 2E). Although GST-(HE)12-YGR6G6 and GST-(HE)10-YGR6G6 showed a high uptake, GST-(HE)10-YGR6G6 actually had the greatest pH sensitivity consistent with the estimated pI of (HE)n-YGR6G6 fused oligopeptides (Table 1). Although (HE)n-YGR6G6 and (HE)n-YG(RG)6 had a similar pI with n = 8, 10, or 12, they exhibited a totally different pH-dependent profile in cellular internalization. The different profile is possibly due to the different interaction patterns between HE and CPP with either clustered or mixed positive charges. However, regardless of the length of HE oligopeptide moiety, all three fusion peptides with YGR6G6 showed a significantly higher cell uptake at pH 6 than that of their YG(RG)6 counterpart. At pH 6.5 (i.e., pH = pKₐ for histidine), half of the anionic charges in HE peptide will have been neutralized by the protonated histidyl residues, which, according to our data, is sufficient to begin to recover the binding and uptake of the CPP sequence. The significantly higher cellular uptake of clustered Arg-peptide than that of mixed Arg-peptide at pH 6.5 or lower is possibly due to the fact that the insertion of glycine residues between the arginine residues will result in a decreased cell uptake of the oligoarginine.25 However, the difference in cell uptake between these two HE fusion peptides, especially at pH 6.5, appeared much greater than that what has been observed between YGR6G6 and YG(RG)6. It is likely that the difference is also due to the presence of (HE)n sequence in the fusion oligopeptides.
which may influence the overall conformation of the fusion peptides. Therefore, the difference in conformation between (HE)\textsubscript{10}-YGR\textsubscript{6}G\textsubscript{6} and (HE)\textsubscript{10}-YG(RG)\textsubscript{6} was further investigated by using CD spectroscopy.

In order to further evaluate the oligopeptides for further use in in vivo studies, the pH sensitivity was evaluated in vitro in the presence of FBS. As shown in Figures 5 and 6, increasing amounts of FBS led to decreased cell uptake and binding. This result was expected because the addition of FBS has been shown to decrease the nonspecific binding and the electrostatic adsorption of CPPs.\textsuperscript{32,33} However, the presence of up to 10% FBS did not affect the pH sensitivity of the constructs because the ratio of cell uptake and of surface binding at pH 6.0 versus pH 7.5 was similar across the 0−10% FBS range (Table 2).

CD spectra have been studied on some arginine-rich peptides with cluster of arginine residues, that is, HIV-1 Tat,\textsuperscript{34,35} R\textsubscript{7},\textsuperscript{35} and R\textsubscript{9},\textsuperscript{34,36} to assess their secondary structures. However, no evidence showed that they have an ordered structure regardless of solvent/buffer systems used in those studies. As shown in Figure 7B, (HE)\textsubscript{10}-YGR\textsubscript{6}G\textsubscript{6} tends to have an unstructured conformation in the range from pH 6.0 to 7.5, with a minimum absorption around 198 nm and no positive peak.\textsuperscript{36,37} However, the spectra of (HE)\textsubscript{10}-YG(RG)\textsubscript{6} as shown in Figure 7A, indicated an antiparallel β-sheet conformation,\textsuperscript{28} whereas its

Table 2. Effect of Serum on pH Sensitivity of GST-Fusion Proteins

| sequence            | % FBS | ratio (pH 6.0/pH 7.5)\textsuperscript{a} | cell uptake | surface binding |
|---------------------|-------|------------------------------------------|-------------|-----------------|
| GST-(HE)\textsubscript{10}R\textsubscript{6}G\textsubscript{6} | 0%    | 7.2                                      | 2.2         |
|                     | 1%    | 5.4\textsuperscript{b}                   | 2.6         |
|                     | 5%    | 6.5                                      | 2.6         |
|                     | 10%   | 4.9\textsuperscript{b}                   | 2.2         |
| GST-(HE)\textsubscript{10}(RG)\textsubscript{6} | 0%    | 5.8                                      | 2.7         |
|                     | 1%    | 5.1                                      | 2.8         |
|                     | 5%    | 5.8                                      | 3.4         |
|                     | 10%   | 5.1                                      | 2.9         |

\textsuperscript{a}Ratios were calculated based on the results shown in Figures 5 and 6.

\textsuperscript{b}Indicates a statistically significant difference (p = 0.03) compared to the ratio in the respective 0% FBS control. All other differences compared to the respective 0% FBS controls were nonsignificant (p > 0.05).
YG(RG)₆ fusion peptides. Figure 7A shows that the majority of (HE)₁₀-YG(RG)₆ still adopts the same conformation even at uptake and binding assay (Figures 4A and 4B). Figure 7B shows a high pH sensitivity between pH 7.0 to pH 6.0 in cell of (HE)

Consequently, the maintenance of an antiparallel β-sheet conformation of (HE)₁₀-YG(RG)₆ is only slightly anionic (net charge = −pH 6.0, that is, when the net charge of the (HE)₁₀ moiety in the (mdeg) versus wavelength (nm) after deduction of background. (HE)₁₀-YG(RG)₆ and (HE)₁₀-YGR₆G₆ were isolated from their corresponding recombinant GST-fusion proteins, GST-(HE)₁₀-YG(RG)₆ and GST-(HE)₁₀-YGR₆G₆ after overnight thrombin treatment. After buffer exchange and desalting, isolated peptides were diluted to 25 mM in 10 mM phosphate buffer saline (PBS). CD spectra were determined as described in the Experimental Section. Data were plotted as ellipticity (mdeg) versus wavelength (nm) after deduction of background.

In conclusion, our results indicate that HE oligopeptide can be conjugated to CPPs with both clustered and mixed positive charges to form CPP derivatives with a reactivation in mildly acidic environment. However, a higher pH sensitivity can be expected in clustered than in mixed positive charge CPPs. In general, the increase of the length of the HE-repeat can decrease the nonspecific absorption of CPP at neutral pH, but it will also decrease the reactivation at acidic pH. Therefore, when designing a reversible masking polyanion to decrease the nonspecific binding of CPPs in drug delivery across biological barriers, both the total charges of the anions and the molecular conformation of the ionic complex should be considered.

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**Notes**

The authors declare no competing financial interest.

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In our study, the insertion of glycine as a spacer between Arg to form a β-hairpin conformation by other means. Because the CD measurements were obtained from isolated HE-CPP peptides without GST, the drastic difference in the spectra between (HE)₁₀-YGR₆G₆ and (HE)₁₀-YG(RG)₆ as shown in Figure 7A clearly suggests a difference in the structure of these 2 peptides. Glycine is the most conformationally unrestrained amino acid, and it has been utilized as shown in Figure 7A clearly suggests a difference in the structure of these 2 peptides. Glycine is the most conformationally unrestrained amino acid, and it has been utilized between Arg to form a β-hairpin conformation by other means. In addition, the stability of the CPP-sequence and the carboxylic group of glutamic acid interaction between the guanidine group of arginine residues in the HE-sequence. In conclusion, our results indicate that HE oligopeptide can be conjugated to CPPs with both clustered and mixed positive charges to form CPP derivatives with a reactivation in mildly acidic environment. However, a higher pH sensitivity can be expected in clustered than in mixed positive charge CPPs. In general, the increase of the length of the HE-repeat can decrease the nonspecific absorption of CPP at neutral pH, but it will also decrease the reactivation at acidic pH. Therefore, when designing a reversible masking polyanion to decrease the nonspecific binding of CPPs in drug delivery across biological barriers, both the total charges of the anions and the molecular conformation of the ionic complex should be considered.

**Figure 7.** Circular dichroism (CD) spectra of (HE)₁₀-YG(RG)₆ (A) and (HE)₁₀-YGR₆G₆ (B) at pH 6.0 (gray line), 6.5 (long dashed black line), 7.0 (regular black line), and 7.5 (bold black line). (HE)₁₀-YG(RG)₆ and (HE)₁₀-YGR₆G₆ were isolated from their corresponding recombinant GST-fusion proteins, GST-(HE)₁₀-YG(RG)₆ and GST-(HE)₁₀-YGR₆G₆ after overnight thrombin treatment. After buffer exchange and desalting, isolated peptides were diluted to 25 mM in 10 mM phosphate buffer saline (PBS). CD spectra were determined as described in the Experimental Section. Data were plotted as ellipticity (mdeg) versus wavelength (nm) after deduction of background.

**Figure 8.** Circular dichroism (CD) spectra of (HE)₁₀-YG(RG)₆ (bold line), (HE)₁₀-YGR₆G₆ (regular line) and (HE)₁₀-Tat (long dashed line) at pH 6.0 (A) and 7.5 (B). GST-(HE)₁₀-YG(RG)₆, GST-(HE)₁₀-YGR₆G₆ and GST-(HE)₁₀-Tat could be eluted form corresponding GSH columns with PBS with or without 250 mM acetate. Buffer exchange and desalting were then performed. Afterward, isolated peptides were diluted in 10 mM PBS at a final concentration of 25 mM. CD spectra were determined as described in the Experimental Section. Data were plotted as ellipticity (mdeg) versus wavelength (nm) after deduction of background.

The authors declare no competing financial interest.

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