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The ultimate goal of our research is the regulation of cellular functions by designed peptides and proteins. Current research subjects include (1) development of novel intracellular delivery systems aiming at elucidation and control of cellular functions using designed membrane permeable peptide vectors, (2) elucidation of the DNA binding and recognition modes of C2H2-type zinc finger proteins and design of artificial transcription factors with various DNA binding specificities, and (3) design of stimulation-responsive artificial peptides and proteins.

**Research Activities (Year 2009)**

**Publications**


**Presentations**

“Chemical and Biological Factors that Affect the Internalization of Arginine-Rich Cell-Penetrating Peptides”, Futaki S, PepVec2009 Meeting on “Intracellular Delivery of Therapeutic Molecules: From Bench to Bedside” Montpellier, France, 1 November 2009.


“Creation of Zinc Finger-Based Artificial Transcription Factors”, Imanishi M, Department Seminar, School of Pharmacy, University of Maryland, Baltimore, USA, 21 November 2009.


**Grants**


Imanishi M, Creation of Artificial Transcription Factors towards Construction of Artificial Genetic Circuit, Grant-
Cytosolic Targeting of Macromolecules Using a pH-Dependent Fusogenic Peptide in Combination with Cationic Liposomes

pH-Sensitive peptides and polymers have been employed as additives to enhance the cytosolic delivery of drugs and genes by facilitating their endosomal escape. However, little attention has been paid to the intracellular fate of these peptides and polymers. In this study, we explored the possibility of utilizing GALA, a pH-sensitive fusogenic peptide, as a cytosol-targeting vehicle. In combination with cationic liposomes, Lipofectamine 2000 (LF2000), the feasibility of this approach for the cytosolic targeting of proteins and nanoparticles was exemplified through the delivery of avidin (68 kDa) and streptavidin-coated quantum dots (15-20 nm) in serum-containing medium. The use of cationic liposomes is critical to enhance the cell-surface adhesion of the GALA conjugates and eventual endosomal uptake. Circular dichroism studies suggest that the GALA can be liberated from cationic liposomes at a reducing pH to form a helical structure and this may eventually lead to disruption of the endosomal membrane to achieve an efficient leakage of the GALA conjugates into the cytosol.

Figure 1. Concept of cytosolic targeting using GALA as an addressing vehicle in combination with cationic liposomes.

Cobalt(II)-Responsive DNA Binding of a GCN4-bZIP Protein Containing Cysteine Residues Functionalized with Iminodiacetic Acid

Endowment of novel functions inducing that of metal switch can be attainable through structural design of peptides and proteins. We previously reported that helical peptides having a pair of iminodiacetic acid (Ida) derivatives of lysine at positions $i$ and $i+2$ induce critical helix destabilization in the presence of metals to lead functional switch of peptides. However, due to the lack of the methodology to effectively introduce the Ida moieties at specific positions in proteins, the application of this concept has been limited to synthetic peptides.

We present a new method for introducing the Ida moieties into proteins. This employs specific modification of cysteines by treatment with a new functionalization agent, $N$-(2-tosylthioethyl) iminodiacetic acid (Ts-S-IDA). The practicability of this approach was exemplified through the metal-responding switching of the DNA binding of the yeast transcription factor GCN4-derived proteins bearing Ida moieties. Two pairs of Ida moieties were incorporated in the leucine zipper segment of the GCN4-bZIP protein in such a way that the Ida moieties of each pair were in $i$ and $i+2$ positions. Complex formation of the Ida groups in the leucine zipper of the GCN4-bZIP protein induced metal-assisted DNA binding switch of the protein to the target DNA.

Figure 2. Preparation of Ida-modified cysteine in GCN4-bZIP protein mutant and conceptual scheme of metal-assisted DNA binding switch of GCN4-bZIP protein modified with Ida.

in-Aid for Young Scientists (B), 1 April 2008–31 March 2010.

Nakase I, Receptor Target and Efficient Internalization of Therapeutic Molecules into Cells Using Membrane Permeable Peptides, Grant-in-Aid for Young Scientists (B), 1 April 2009–31 March 2011.

Awards

Azuma Y, Best Poster Award, “Metal-Induced DNA-Binding Switch of bZIP Proteins Modified with Iminodiacetic Acid (Ida)” The 19th Symposium on Role of Metals in Biological Reactions, Biology and Medicine (SRM2009), Suita, 12 June 2009.