Liver cell specific targeting by the preS1 domain of hepatitis B virus surface antigen displayed on protein nanocages

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Abstract: Protein nanocages are self-organized complexes of oligomers whose three-dimensional architecture can be determined in detail. These structures possess nanoscale inner cavities into which a variety of molecules, including therapeutic or diagnostic agents, can be encapsulated. These properties yield these particles suitable for a new class of drug delivery carrier, or as a bio-imaging reagent that might respond to biochemical signals in many different cellular processes. We report here the design, synthesis, and biological characterization of a hepatocyte-specific nanocage carrying small heat-shock protein. These nanoscale protein cages, with a targeting peptide composed of a preS1 derivative from the hepatitis B virus on their surfaces, were prepared by genetic engineering techniques. PreS1-carrying nanocages showed lower cytotoxicity and significantly higher specificity for human hepatocyte cell lines than other cell lines in vitro. These results suggested that small heat-shock protein-based nanocages present great potential for the development of effective targeted delivery of various agents to specific cells.

Keywords: protein nanocages, drug delivery system, hepatocyte cell lines specific, hepatitis B virus

Introduction

As most drugs have both beneficial and unfavorable effects pertaining to chemotherapy, which might also be tissue dependent, it is necessary to deliver therapeutic agents selectively to their target sites.1,4 Conventional chemotherapeutic agents diffuse nonspecifically throughout the body where they affect both malignant and normal cells; unfortunately, 95% of all new potential therapeutics have poor pharmacokinetic and biopharmaceutical properties.3 To overcome these problems, there is a serious need to develop effective drug delivery systems (DDS) that distribute therapeutically active drug molecules only to the desired site of action without affecting healthy organs and tissues.

Various strategies for site-specific drug delivery have led to the development of drug carriers. In particular, DDS based on liposomes6,7 and synthetic polymers8–11 have been extensively studied as novel drug-packaging strategies for cancer chemotherapy. In the last decade, nanotechnological innovations have played an important role in size control and surface modification of nanomaterials, and the resulting properties play a critical role in target specificity for tumor tissues via improved pharmacokinetics and pharmacodynamics and in allowing active intracellular delivery characteristics.12 Traditional DDS materials, including liposomes and synthetic polymers, can be manufactured in bulk at low cost and with a wide diversity of backbones, surface
 chemicals, and molecular weights; however, their aggregate structure and compositional sequence cannot be accurately controlled and therefore such materials offer disadvantages in their poor responsiveness to cellular behavior that require sequence specificity and structural stringency to be sensitive to signals exhibited by certain cell biochemical processes or states.

On the other hand, natural or recombinant protein-based nanocapsules or nanocages are also being actively investigated as new types of nanomaterial. These nanocapsules are self-organized protein oligomer complexes with nanoscale inner cavities and whose three-dimensional architecture can be specified in great detail. Thus, they have the potential to offer special advantages in developing new functions that are responsive to biochemical signals exhibited by various cellular processes. Such specific interactions derive from the incorporated amino acid sequences that can be designed to address multiple functional requirements for biomaterial applications. Recently, we constructed stimulus-responsive nanomaterial by incorporating a genetically engineered small heat shock protein (sHSP) that can change its conformation in response to dual stimuli, a protease signal and temperature, in contrast to the wild type.

Here, a novel targeted drug delivery method is presented involving a protein-based nanocage composed of HSP16.5, a sHSP from a hyperthermophilic archaeon. This protein forms a 400 kDa homogeneous complex comprised of 24 subunits that self-organizes under physiological conditions to form a nanoscale, hollow, spherical capsule with small pores, contain very high thermal stability, and offer the potential for a variety of molecules, including drugs and magnetic resonance imaging contrast agents, to be encapsulated in the cavity. These unique properties yield HSP16.5 suitable for a drug carrier, but this protein exhibits no significant organ or tissue specificity in body distribution following intravenous administration in mice. To address this problem, a recombinant fusion protein was constructed from HSP16.5 and a target-specific peptide using genetic engineering. Examinations of the crystal structure of the HSP16.5 complex have revealed that the C-terminal region is on the exterior surface of the capsule-like structure. Thus, a recombinant HSP16.5 was designed and produced by an in-frame fusion, with the sequence for a targeting peptide added in this region. Specificity for liver cells-targeted delivery of the nanocapsule was achieved using a preS1 peptide for the targeting peptide (Figure 1). This peptide is derived from the envelope proteins of the hepatitis B virus (HBV), with a known affinity for primary hepatocytes and hepatocyte cell lines. The mechanism of hepatocyte infection induced by HBV is not absolutely clear; however, specific attachment of viruses to cells is the essential first step in virus entry into cells. Several studies suggested that amino acids 21-47 of preS1 play a major role in the cell attachment. For example, Dash et al reported that this 27-mer preS1 peptide (21-47) binds to HepG2 cells in a dose-dependent manner with high affinity. Furthermore, this binding is possibly tissue-specific, since only weak to insignificant levels of binding were obtained with Vero cells, HeLa cells, and human peripheral lymphocytes. This paper describes the characterization of 27-mer preS1 peptide-carrying nanocapsules and their potential for biomedical applications, including targeted drug delivery and bioimaging.

![Figure 1](https://www.dovepress.com/)

**Figure 1** Schematic illustration of genetically engineered nanocages for hepatocyte selective targeting.
Materials and methods
DNA cloning and expression
Plasmid pET-HSPG41C-preS1, a bacterial expression vector containing the preS1 gene fused in frame with the HSP16.5 variant (HSPG41C), was cloned by polymerase chain reaction (PCR). The HSPG41C mutant presents unique reactive cysteine residues on the interior surface of the assembled cage for the attachment of cargo molecules. This mutant was constructed according to the guidelines noted in a previous paper, and is used as a PCR template. The primers for HSPG41C-preS1 were 5'-taatacgactcactataggg-3' (forward primer), 5'-caggctctCTAGGGTTGAAGTCCAGTC-CGGGTTGTAGAGTTACCGAAAGCGGGGTTCGACGTGGTTGTCGGGGAAGAAACCCACCGGGCTAGCtcatgttgattcctttcttaattgagga-3' (reverse primer), and the latter primer contained a preS1 gene coding region (capital letters) and BamHI restriction site (underlined). The PCR-generated 0.63-kb fragment was digested with NdeI and BamHI and the resulting DNA fragment ligated into NdeI and BamHI-digested pET21a vector (Novagen; Merck Japan, Tokyo, Japan). Correct insertion of the fragment was verified by DNA sequencing and plasmids then amplified and purified by standard techniques.

The resulting vector was allowed to express HSPG41C-preS1 in Escherichia coli BL21-Gold(DE3) cells (Novagen) using a T7 expression system. Bacterial cells of BL21 gold (DE3) containing pET-HSPG41C-preS1 were grown in 2 × YT medium (Sigma-Aldrich, St Louis, MO) with 100 mg/mL ampicillin at 37°C. Recombinant protein production was initiated by addition of isopropyl thiogalactopyranoside (IPTG; Wako Pure Chemical Industries, Ltd, Tokyo, Japan) and is used as a PCR template. The primers for HSPG41C-preS1 were 5'-taatacgactcactataggg-3' (forward primer), 5'-caggctctCTAGGGTTGAAGTCCAGTC-CGGGTTGTAGAGTTACCGAAAGCGGGGTTCGACGTGGTTGTCGGGGAAGAAACCCACCGGGCTAGCtcatgttgattcctttcttaattgagga-3' (reverse primer), and the latter primer contained a preS1 gene coding region (capital letters) and BamHI restriction site (underlined). The PCR-generated 0.63-kb fragment was digested with NdeI and BamHI and the resulting DNA fragment ligated into NdeI and BamHI-digested pET21a vector (Novagen; Merck Japan, Tokyo, Japan). Correct insertion of the fragment was verified by DNA sequencing and plasmids then amplified and purified by standard techniques.

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Protein purification and characterization
Cells were pelleted by centrifugation at 4°C, the pellet resuspended in 8 mL of sample buffer (20 mM KH2PO4-KOH, pH 7.0, 2 mM DTT, and 1 mM ethylenediaminetetraacetic acid), and the resulting suspension sonicated on ice (200 watts, 45 seconds). DNase I and RNase A were then added to final concentrations of 5 and 1 mg/mL, respectively. The cell lysate mixture was incubated at 4°C for 30 minutes, and the insoluble material removed by centrifugation (20,000 g, 20 minutes) at 4°C. Purification of the recombinant protein was performed by ion-exchange chromatography, in which the supernatant was loaded on a HiLoad 26/10 Q Sepharose HPTM anion-exchange column (Amersham Pharmacia Biotech Inc; GE Healthcare, Little Chalfont, UK) and HSPG41C-preS1 eluted with sample buffer containing 1M NaCl. Fractions containing HSPG41C-preS1, confirmed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), were loaded onto a silica-based GPC column TSKgel G3000SW (Tohosch Corporation, Tokyo, Japan) and HSPG41C-preS1 eluted with storage buffer (25 mM NaH2PO4/Na2HPO4, pH 7.0, and 0.1 M NaCl). The resulting purified HSPG41C-preS1 fractions were analyzed by SDS-PAGE using 12% gel according to standard protocol. The recombinant HSPG41C-preS1 protein cages were characterized by matrix-assisted laser desorption/ ionization time-of-flight mass spectrometry (Autoflex Speed; Bruker Daltonics Japan, Yokohama, Japan) and dynamic light scattering (Malvern Nanosizer ZS; Malvern Instruments).

Cytotoxicity assays
HepG2, Huh-7, and HeLa cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM; Wako Pure Chemical Industries, Ltd), supplemented with 1% (v/v) antibiotic-antimycotic mix (Gibco 15240-062; Life Technologies Japan Ltd., Tokyo, Japan) and 10% (v/v) fetal bovine serum (FBS; GE Healthcare, Little Chalfont, UK). MCF-7 cells were cultured in RPMI-1640 (Sigma-Aldrich) with 1% (v/v) antibiotics and 10% FBS and at 37°C in a humidified 5% CO2 atmosphere.

The cellular toxicity of the nanocages in vitro was assessed using the CellTiter-Glo® luminescent cell viability assay (Promega Corporation, Madison, WI) according to the manufacturer’s instructions. This assay is a homogeneous method designed to determine the viable cell numbers in culture based on quantitation of available adenosine triphosphate, signaling the presence of metabolically active cells at the assay end point. Cells (1 × 104 cells/well) were cultured in 96-well microtiter plates in complete growth medium in the presence of increasing nanocapsule concentrations. For chronic toxicity assessment, nanocapsules were applied to cells at various concentrations and incubated for 16 hours prior to assay for viable cells.

Fluorescence assay
Prior to a fluorescence assay, HSPG41C-preS1 and HSPG41C (used as a control) were labeled with fluorescent dye. Briefly, nanocapsules were incubated with a 5-fold molar excess of Alexa488-maleimide (Invitrogen; Life Technologies, Carlsbad, CA) in pH 8.0 phosphate buffer at 50°C overnight and then excess dye removed by ultrafiltration. Uptake or transfection was assessed by
For investigating nanocapsule cellular localization, HepG2, Huh-7, HeLa, and MCF-7 cells were seeded at $1 \times 10^4$ cells/well in μ-Slides (μ-Slide 6 well; ibidi GmbH, Munich, Germany), incubated with colorless DMEM containing 1% antibiotic-antimycotic mix and 10% FBS (PAA; GE Healthcare) overnight. Then, fluorescent-labeled nanocapsules of HSPG41C-preS1 or HSPG41C control were added to wells at a final 1 μM concentration and, after additional 6 hours of incubation, non-incorporated nanocapsules removed by washing cells twice with phosphate buffered saline and adding fresh medium. Fluorescence confocal laser scanning microscopy was performed with a Bio-Rad Radiance 2100 confocal laser scanning microscopy system (Bio-Rad Laboratories Japan, Tokyo, Japan) attached to a Nikon TE2000-U microscope (Nikon Corporation, Tokyo, Japan). All fluorescence images were acquired under the same conditions of objective lens, laser excitation intensity, photomultiplier gain, and imaging size. Statistical significances of the differences were determined by Student’s t-test.

![Figure 2](image.png)

**Figure 2** Dynamic light scattering analysis of protein nanocages. (A) HSPG41C nanocage controls; (B) HSPG41C-preS1 nanocages.
Results and discussion
Expression and characterization of preS1-carrying protein nanocages

Expression vectors encoding the HSPG41C-preS1 or HSPG41C were constructed and recombinant sHSP16.5 proteins produced in *E. coli* BL21-Gold(DE3) and purified by sequential anion exchange chromatography followed by size exclusion chromatography under native conditions (Figure S1). Purified proteins, separated by SDS-PAGE, appeared as a single band by Coomassie blue staining. The observed molecular weight of purified HSPG41C-preS1 (m/z = 19676.2 Da), analyzed by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry with a sinapic acid matrix, was largely in agreement with calculations (m/z = 19668.2 Da) (Figure S2). The protein nanocage size range and distribution were measured by means of dynamic light scattering. These results indicated that the average nanocage diameter of the HSPG41C-preS1 and HSPG41C were 14.4 and 12.7 nm, respectively, with a narrow size distribution (Figure 2). These results demonstrated that HSPG41C-preS1 cages were slightly larger than HSPG41C parent cages lacking the targeting peptide.

**In vitro cellular cytotoxicity of nanocages**

As cell cytotoxicity is an important factor in selecting materials for drug carriers, the nanocages produced here were characterized regarding their effect on cell viability under the same conditions as used for fluorescence assay of cellular uptake experiments. Neither HSPG41C nor HSPG41C-preS1 had any appreciable cytotoxic effect under these conditions (Figure 3); similar results were obtained from wild type HSP16.5 protein under the same conditions (data not shown). These results showed that the single amino acid substitution mutation, contained in HSPG41C, and the cage surface modification with the

![Cell viability measurements after treatment with protein nanocages against various cell lines. (A) HSPG41C nanocage controls; (B) HSPG41C-preS1 nanocages.](https://www.dovepress.com/)

*Figure 3 Cell viability measurements after treatment with protein nanocages against various cell lines. (A) HSPG41C nanocage controls; (B) HSPG41C-preS1 nanocages.*
preS1 peptide of wild type HSP16.5, contained on HSPG41C-preS1, did not significantly affect their cytotoxicity.

**Hepatocyte specific delivery of protein nanocages in vitro**

The mechanism for the entry of HBV particles into target cells, in particular into hepatocytes, is not yet understood. However, the preS1 surface antigen of HBV is known to play an important role in initial HBV attachment to hepatic cell lines, with the preS1 domain N-terminal segment believed to be essential. To demonstrate that preS1 fusion into the HSP16.5 protein nanocage produced a new nanocage binding affinity towards hepatocyte cell lines, the specific target cells of HBV preS1 molecule, transfection assays were performed.

Fluorescent-labeled HSPG41C-preS1 nanocages were observed to efficiently internalize to human hepatoma cell lines HepG2 and Huh-7, which bear specific HBV receptors compared to internalization to the other cell lines (Figure 4). Incubation with these nanocages resulted in specific

**Figure 4** Fluorescence confocal laser scanning microscopy of various cells with fluorescently-labeled HSPG41C-preS1 nanocages. Alexa488-labeled HSPG41C-preS1 nanocages added to culture media with $1 \times 10^4$ cells of Huh-7 (A), HepG2 (B), HeLa (C), and MCF7 (D), and extensively washed with PBS before CLSM analysis.

**Abbreviations:** PBS, phosphate buffered saline; CLSM, confocal laser scanning microscopy.

**Figure 5** Fluorescence confocal laser scanning microscopy of various cells with fluorescently-labeled HSPG41C nanocages. Alexa488-labeled HSPG41C nanocages added to culture media with $1 \times 10^4$ cells of Huh-7 (A), HepG2 (B), HeLa (C), and MCF7 (D), and extensively washed with PBS before CLSM analysis.

**Abbreviations:** PBS, phosphate buffered saline; CLSM, confocal laser scanning microscopy.
accumulation of these proteins in hepatic cell cytoplasm but not in the nucleus. On the other hand, the control fluorescent-labeled HSPG41C nanocages were slightly taken up by Huh-7, HepG2, and MCF-7 under the same conditions (Figure 5). Nanoparticles are mostly internalized into cells by endocytosis. Endocytosis can be classified into four groups: clathrin-mediated endocytosis (CME), caveolae-mediated endocytosis, macropinocytosis, and clathrin- and caveolae-independent endocytosis. We investigated the endocytic pathway of HSPG41C-preS1 nanocages using three types of chemical compound that inhibited specific endocytic pathways (Figure S3). Chlorpromazine inhibits CME, amiloride inhibits macropinocytosis, and filipin III inhibits caveolae-mediated endocytosis. When HepG2 cells were treated with amiloride and filipin, 5% and 6% decreases in cellular uptake of HSPG41C-preS1 nanocages were detected compared with untreated cells. Chlorpromazine treatment showed a marked decrease (23%) in the uptake of HSPG41C-preS1 nanocages. These results suggest that HSPG41C-preS1 nanocage was taken up by HepG2 cells by various types of endocytosis, but the main uptake pathway was CME.

To determine the HSPG41C-preS1 nanocage internalization process into hepatoma cells, competition assays using synthetic preS1 peptide, encompassing amino acid residues 21-47 of preS1, were performed. Fluorescent-labeled HSPG41C-preS1 nanocage incorporation in the presence of various concentrations of synthetic preS1 peptides was intensity normalized to 100 for untreated Huh-7 (Figure 6). In the case of human hepatoma-derived cell lines HepG2 and Huh-7, synthetic preS1 peptides inhibited incorporation of labeled HSPG41C-preS1 nanocages in a dose-dependent manner. On the other hand, the presence of synthetic preS1 peptides did not significantly influence interaction between these nanocages and HeLa cells. These results suggested that the cell binding observed for these nanocages was due specifically to the presence of the preS1 moiety on their surfaces and the internalization of the nanocages was achieved by interaction with preS1 receptors on the hepatocyte cell lines.

**Conclusion**

In summary, genetic incorporation of hepatocyte binding preS1 peptide onto the exterior surface of nanocages conferred cell-specific targeting capabilities to this protein cage architecture. These protein nanocages were efficiently internalized into hepatocyte cell lines, mainly through clathrin-mediated endocytosis, without cytotoxicity, and were localized at cytoplasm. Furthermore, the binding of HSPG41C-preS1 nanocages to HepG2 cells was prevented by synthetic preS1-(21-47) peptide in a dose-dependent manner, suggesting that this sequence could be directly responsible for nanocage attachment to the cellular surface. In fact, it is known that the preS1 domain of HBV is modified with myristic acid at the N-terminal region and that this modification is important for efficient infectivity, and thus it appears that it would be necessary to conjugate the HSPG41C-preS1 nanocages with a similar lipid to enhance specificity in hepatocyte targeting in vivo. This strategy was effective for producing suitable nanocages that were directed to and taken up by specific cell types, target organs, or cancer
cells, which is an important development in such applications, as ligand-mediated active binding to sites and cellular uptake are particularly valuable to therapeutic agents that are not easily taken up by cells. Furthermore, these nanocages are self-organized complexes of protein oligomers and possess nanoscale inner cavities, whose three-dimensional architecture can be specified in detail and into which a variety of molecules, including therapeutic or diagnostic agents, could be encapsulated.

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Disclosure

The authors report no conflict of interest in this work.

References

Supplementary figures

**Figure S1** Gel permeation chromatography purification of HSPG41C nanocages and HSPG41C-preS1 nanocages. GHSPG41C nanocages (A); HSPG41C-preS1 nanocages (B). The peaks observed at 13.31 minutes in (A) and 13.26 minutes in (B) were collected individually. The collected fractions were then analyzed by SDS-PAGE using 12% gel according to the standard protocol (C).

Note: Lane 1, molecular weight standards; lane 2, HSPG41C nanocages; lane 3, HSPG41C-preS1 nanocages.

Abbreviation: SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis.

**Figure S2** MALDI-TOF mass spectrum of HSPG41C nanocages and HSPG41C-preS1 nanocages. GHSPG41C nanocages (A); HSPG41C-preS1 nanocages (B).

Abbreviation: MALDI-TOF, matrix-assisted laser desorption/ionization time of flight.
Figure S3 Inhibition of cellular uptake of HSPG41C-preS1 nanocages by endocytosis inhibitors.

Notes: HepG2 cells were harvested on poly-L-lysine-coated 48-well plates at an initial density of 50,000 cells/well and grown overnight. Cells were treated with DMEM containing chlorpromazine (10 µg/mL), amiloride (500 µM), or filipin III (10 µg/mL) for 1 hour (all from Sigma-Aldrich, St Louis, MO). Then 84 nM of fluorescent-labeled HSPG41C-preS1 nanocages solution containing the above inhibitors was added to cells. Two hours after transfection, cells were rinsed three times with PBS and then lysed in lysis buffer (pH 7.5, 20 mM Tris-HCl, 2 mM EDTA, and 0.05% Triton-X 100). Lysate solutions were replaced on black-bottomed 96-well plates and the fluorescence intensity of each sample measured using a Microplate Reader (ARVO MX 1420; Perkin Elmer Inc, Waltham, MA). Data are means ± SEM of three independent experiments.

Abbreviations: CPZ, chlorpromazine; Amil, amiloride; Filip, filipin; DMEM, Dulbecco’s Modified Eagle’s Medium; PBS, phosphate buffered saline; HCl, hydrochloride; EDTA, ethylenediaminetetraacetic acid; SEM, standard error of the mean.