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Engineering Human Epidermal Growth Receptor 2-Targeting Hepatitis B Virus Core Nanoparticles for siRNA Delivery in Vitro and in Vivo

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Supporting Information

ABSTRACT: Hepatitis B virus core (HBC) particles acquire the capacity to disassemble and reassemble in a controlled manner, allowing entrapment and delivery of drugs and macromolecules to cells. HBC particles are made of 180–240 copies of 21 kDa protein monomers, assembled into 30–34 nm diameter icosahedral particles. In this study, we aimed at formulating HBC particles for the delivery of siRNA for gene silencing in vitro and in vivo. We have previously reported recombinant HBC particles expressing ZHER2 affibodies, specifically targeting human epidermal growth receptor 2 (HER2)-expressing cancer cells (ZHER2ΔHBC). siRNA was encapsulated within the ZHER2ΔHBC particles following disassembly and reassembly. The ZHER2ΔHBC–siRNA hybrids were able to secure the encapsulated siRNA from serum and nucleases in vitro. Enhanced siRNA uptake in HER2-expressing cancer cells treated with ZHER2ΔHBC–siRNA hybrids was observed compared to the nontargeted HBC–siRNA hybrids in a time- and dose-dependent manner. A successful in vitro polo-like kinase 1 (PLK1) gene knockdown was demonstrated in cancer cells treated with ZHER2ΔHBC–siPLK1 hybrids, to levels comparable to commercial transfecting reagents. Interestingly, ZHER2ΔHBC particles exhibit intrinsic capability of reducing the solid tumor mass, independent of siPLK1 therapy, in an intraperitoneal tumor model following intraperitoneal injection.

KEYWORDS: virus-like particles, hepatitis B virus core particles, human epidermal growth factor receptor 2, affibody, gene silencing

1. INTRODUCTION

RNA interference (RNAi) is a biological mechanism whereby small noncoding double-stranded RNAs (dsRNA) interfere with the expression of a particular gene that shares homologous sequence with dsRNA. Once small interference RNA (siRNA), 20–30 nucleotide dsRNA, is present in the cytoplasm of the cell, it is incorporated into a protein complex known as RNA induced silencing complex (RISC). The whole construct can induce a selective endonucleolytic cleavage of the complementary target mRNA, which propagates gene silencing. Gene silencing has been of great interest as a powerful therapeutic strategy to regulate diseases of disordered gene origin, including cancer. However, the therapeutic applications of siRNA have been limited, due to its high molecular weight and anionic charge, which causes difficulty for siRNA to penetrate the cellular membranes by passive diffusion mechanisms. In addition, siRNA activity is greatly limited by rapid enzymatic degradation due to the presence of ribonucleases (RNases) in serum.

In order to achieve efficient cytoplasmic delivery of siRNA in vitro and in vivo, delivery systems using nanoparticles have been employed by researchers, including nonviral delivery systems such as cationic liposomes, polymers, and carbon nanotubes, and viral delivery systems such as viral-like particles (VLPs). VLPs are hollow protein capsids derived from viruses but lacking the viral genomes. They are ideal bio-nanocarriers due to their biocompatibility, regulated structure, and homogeneous size. Many medical applications, including vaccine development and delivery of peptides and nucleic acids have been approached using VLPs. Hepatitis B virus core (HBC) particles are one of the VLPs that have been investigated by researchers as nanocarriers for drug delivery. HBC particles are hollow nanoparticles (30–34 nm in diameter), consisting of 180–240 copies of core monomers. Like other VLPs, HBC particles are noninfectious but still inherit the essential features of the hepatitis B virus, including the cell-penetrating ability. They also have the capability to assemble and disassemble in a controlled manner allowing

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encapsulation of biomolecules including oligonucleotides. In addition, HBC particles can be genetically modified with targeting molecules on the surface, which is beneficial for site-specific delivery. These features make HBC particles suitable candidates for delivery of therapeutics.

In this study, we aimed at formulating the HBC particles that specifically target human epidermal growth receptor 2 (HER2)-expressing cancer cells, for the delivery of siRNA and gene silencing in vitro and in vivo (Scheme 1). Polo-like kinase (PLK1), a serine/threonine kinase that has key roles in cell division and checkpoint regulation of mitosis, was chosen as the targeting gene. PLK1 is known to overexpress in several types of cancer and siRNA-mediated silencing of PLK1 in cancer cells has been shown to result in reduced cell viability with induction of apoptosis after depletion of PLK1 protein. We hypothesized that the HER2-targeting HBC particles would be able to encapsulate and deliver the functional siRNA (siPLK1) in vitro and induce gene silencing in cancer cells in vitro causing cell death. We also hypothesized that the encapsulation strategy is likely to improve the longevity of siRNA in vivo following local administration. HBC particles were prepared and characterized using E. coli expression system. siRNA was encapsulated into HBC particles using three different methods: disassembly and reassembly in the presence of urea, osmotic shock, or disassembly and reassembly by dithiothreitol (DTT)/CaCl2 treatment. siRNA encapsulation efficiency was determined by gel retardation and aldehyde sulfa latex beads assay. Cancer cells were treated with HBC-siRNA hybrid particles, and the cell uptake was assessed by flow cytometry. Cell cytotoxicity and gene silencing of cancer cells treated with HBC-siPLK1 hybrid particles were assessed using MTT assay and immunostaining followed by gel electrophoresis at 100 V for 45 min for gel retardation assay. Samples were loaded onto 1% agarose gel and underwent gel electrophoresis at 100 V for 45 min for gel retardation assay. Samples were loaded onto 1% agarose gel, underwent gel retardation assay, and then imaged using the ChemiDoc MP system as described previously.

2. EXPERIMENTAL SECTION

2.1. Materials. Lithium chloride and ethylene glycol-bis(β-aminoethy1 ether)-N,N,N',N'-tetracetic acid (EGTA) were obtained from Amresco (USA). Ethylenediaminetetraacetic acid (EDTA) disodium and dithiothreitol (DTT) were obtained from forMax (Med) (UK). Calcium chloride (anhydrous) and urea (carbamide) were obtained from Melford (UK). Ribonuclease A (from bovine pancreas), MTTR (3,4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide), and dimethylsulfoxide (DMSO) were obtained from Sigma-Aldrich (UK). Bovine serum albumin, lyophilized powder, ≥96% (agarose gel electrophoresis), was obtained from Sigma-Aldrich (Germany). BD OptEIA Mouse TNFα (Mono/Mono) ELISA kit was obtained from BD Biosciences, USA. Xenolight d- luciferin potassium salt was obtained from PerkinElmer (EU). GAPDH (14C10) rabbit mAb, anti-rabbit IgG HRP-linked antibody, and anti-mouse IgG HRP-linked antibody were obtained from Cell Signaling Technology (UK). Anti-PLK1 antibody (35–206) was obtained from Abcam (UK). Heat inactivated newborn calf serum (FBS) was obtained from First Link Limited (UK). Lipofectamine2000, aldehyde/sulfate latex (4% w/v, 4 μm), Alexa Fluor 488 NHS ester (succinimidyl ester), MEM Glutamax Supplement, DMEM high glucose pyruvate, advanced RPMI 1640 medium, PBS (10X) pH 7.4, trypsin-EDTA (0.05%) penicillin–streptomycin (10,000 U/mL), l-glutamine (200 mM), GlutaMax Supplement, SnakeSkin dialysis tubing (10000 MWCO, 35 mm) and Pierce 16% formaldehyde (w/v, methanol-free), Pierce BCA Protein Assay Kit were purchased from Thermo-Fisher Scientific (UK). Noncoding siRNA (siNEG) (anti-sense sequence, 5′-CAUCGUGAAGUGAAGCGCAA-3′), Atto-655-labeled siNEG, and siPLK1 (sequence 5′-CUUUGUAGAAGAAGACUCGdTdT-3′) were obtained from Eurogentec (Belgium). Precision Protein StreptTactin-HRP conjugate was purchased from Bio-Rad Laboratories, USA. Heparin (5000 IU/mL) (Sigma, UK) was obtained from St. Thomas’s Hospital, UK. Plasmids pET-22b(+) -dc149-His6 and pET-22b(+) -dc149-Z342-His6 were obtained from Associate Professor Chiaki Ogino, Chemical and Science Engineering Department, Kobe University, Japan.

2.2. Expression, Purification, and Assembly of HBC Particles. Expression, purification, and assembly of ΔHBC and ZHER2-ΔHBC particles were prepared as previously published by Suffian et al. and detailed in SI. Protein concentrations of assembled ΔHBC and ZHER2-ΔHBC particles were measured using NanoDrop ND-1000 UV–vis spectrophotometer (Thermo Fisher Scientific, USA).

2.3. siRNA Encapsulation in HBC Particles. Encapsulation of siRNA into ZHER2-ΔHBC particles was performed using three methods mentioned below. Fluorescently labeled nonfunctional siRNA, siNEG (0.05 μg) with emission at 650 nm, and ZHER2-ΔHBC particles with emission at 525 nm were used to assess encapsulation efficiency.

2.3.1. Method I: Disassembly/Reassembly Using Urea/NaCl. Fluorescently labeled ZHER2-ΔHBC particles (30, 60, 120: 1 (1 HBc/siNEG molar ratio), 240, 480 μg) were dialyzed against 1 L of PBS and incubated for 1 h at 20 °C, and then dialyzed against 1 L of ultrapure water containing 50 mM Tris, 1 mM NaCl overnight at 4 °C to reassemble the particles. Samples were concentrated to 10 μL final volume using Vivaspin tubing (30, 60, 120 μL) and then imaged using ChemiDoc MP system as described previously. Samples were loaded onto 1% agarose gel, underwent gel retardation assay, and then imaged using ChemiDoc MP system as described previously.

2.3.2. Method II: Osmotic Shock. Fluorescently labeled ZHER2-ΔHBCAF480 particles (30, 60, 120, 240, 480 μg) were mixed with 0.25 μg of siNEG (0.25 μg) was mixed with ZHER2-ΔHBCAF480 in its disassembled form, incubated for 1 h at 20 °C, and then dialyzed against 1 L of ultrapure water containing 50 mM Tris, 1 mM NaCl overnight at 4 °C to reassemble the particles. Samples were concentrated to 100 μL final volume using Vivaspin tubes (MWCO 10 kDa). To assess the degree of complexation, samples were loaded onto 1% agarose gel, followed by gel electrophoresis at 100 V for 45 min for gel retardation assay. Gel was then visualized under an Alexa Fluor 647 and Alexa Fluor 488 channels using ChemiDoc MP system (Bio-Rad Laboratories, USA).

2.3.3. Method III: Disassembly/Reassembly Using E. coli. Fluorescently labeled ZHER2-ΔHBCAF480 particles (30, 60, 120, 240, and 480 μg) were mixed with 0.25 μg of siNEGAF480 in 1% PBS and incubated for 10 min at 37 °C. Osmotic shock was achieved by 5X dilution with distilled water. Samples were incubated for 20 min at 37 °C then concentrated to 100 μL volume using Vivaspin tubes (MWCO 10 kDa). Samples were loaded onto 1% agarose gel, underwent gel retardation assay, and then imaged using ChemiDoc MP system as described previously.
retardation assay then imaged using ChemiDocMP system as described previously.

In all methods, samples were buffer-exchanged to 1X PBS prior to addition to cells to maintain isotonicity.

2.4. Flow Cytometry Analysis of HBc–siNEG Hybrid Particles. To form siNEG–Atto-655 encapsulation within \( \Delta HBcAF488 \) particles, flow cytometry analysis of HBc–siNEG particles was performed. HBc particles were fluorescently labeled with Alexa Fluor 488 as previously described.\(^{19}\) HBc–siNEG hybrid particles were encapsulated at 1:1 HBc/siNEG molar ratio by the three different encapsulation methods. Prepared HBc–siNEG hybrid particles were reacted with aldehyde/sulfate latex (4% w/v, 4 μm) microbeads (Invitrogen, USA) as previously described for other proteins, with some modification.\(^{27}\) HBc–siNEG hybrid particles (40 μL) were incubated with 10 μL of aldehyde/sulfate latex (4% w/v, 4 μm) microbeads in a final volume of 500 μL for 15 min at room temperature. Then, final concentration of 10 μM BSA from 100 μM stock solution was added to each sample and incubated for another 15 min at room temperature. Then, an incubation in 1 mL of PBS was conducted for another 75 min with gentle shaking; reaction was stopped by incubation with 1 mL of 100 mM glycine made in PBS for 5 min, and the cell pellet was resuspended in 250 μL of 3% serum/PBS. HBcAF488 particles at HBc/siNEG molar ratio 1:1 were prepared using method III (found to be the most efficient encapsulation method) as described above. After complexes were formed, FBS was then added to the complexes at a final concentration of 50% v/v, and samples were incubated at 37 °C for 0, 24, or 48 h. After incubation, the reaction was stopped by adding EDTA to a final concentration of 10 mM. Nonincubated and co- incubated naked siNEG with FBS and EDTA were used as positive and negative controls, respectively. Samples were subjected to a gel retardation assay (2% agarose gel, 100 V, 30 min, GelRed 0.005% v/v). Gel was then imaged under UV light using ChemiDocMP system (Bio-Rad Laboratories, USA).

2.5. In Vitro siRNA Stability in Serum. The degree of enzymatic degradation induced by serum nucleases was studied by an incubation of noncoding siRNA (siNEG) encapsulated within HBc particles with fetal bovine serum (FBS). The HBc–siNEG hybrid particles at HBc/siNEG molar ratio 1:1 (ratio at which complete siRNA encapsulation was achieved) were prepared using method III (found to be the most efficient encapsulation method) as described above. After complexes were formed, FBS was then added to the complexes at a final concentration of 50% v/v, and samples were incubated at 37 °C for 0, 24, or 48 h. After incubation, the reaction was stopped by adding EDTA to a final concentration of 10 mM. Nonincubated and co- incubated naked siNEG with FBS and EDTA were used as positive and negative controls, respectively. Samples were subjected to a gel retardation assay (2% agarose gel, 100 V, 30 min, GelRed 0.005% v/v). Gel was then imaged under UV light using ChemiDocMP system (Bio-Rad Laboratories, USA).

2.6. In Vitro siRNA Protection against RNase. The hybrid particles at HBc/siNEG molar ratio 1:1 were prepared using method III as described above. After forming the complexes, RNase A was added to the complexes at a final concentration of 6.6 μg/mL. Samples were then incubated at 37 °C for 0, 24, or 48 h. After incubation, the reaction was stopped by adding EDTA to a final concentration of 50 mM. Naked siNEG (0.25 μg) nonincubated and co- incubated with FBS and EDTA were used as positive and negative controls, respectively. Samples were subjected to a gel retardation assay (2% agarose gel, 100 V 30 min, GelRed 0.005% v/v). Gel was then imaged under UV light using ChemiDocMP system (Bio-Rad Laboratories, USA).

2.7. Cell Culture. HER2-positive cells (HER2\(^{+}\)), MDA-MB-435-MLE and SKBR-3 cells, and HER2-negative (HER2\(^{-}\)) MDA-MB-468 cells were selected in the present study. Their HER2 expression level was confirmed previously.\(^{20}\) MDA-MB-435-MLE cells (human melanoma in origin) were obtained by transfecting MDA-MB-435 cells (ATCC, HTB-12) with MUC1, luciferase, and ErbB2 genes as previously reported.\(^{21}\) MDA-MB-435-MLE cells were cultured in DMEM media; MDA-MB-468 human breast adenocarcinoma (ATCC, HTB-132) and SKBR-3 human breast carcinoma (ATCC, HTB-30) were cultured in advanced RPMI media. All culture media were supplemented with 10% FBS, 50 U/mL penicillin, 50 μg/mL streptomycin, and 1% l-glutamine at 37 °C in 5% CO\(_2\). All cells were routinely grown in 75 cm\(^2\) canted-neck tissue culture flasks and passaged twice a week using Trypsin/EDTA at 80% confluency.

2.8. In Vitro Cell Uptake Studies of HBc–siRNA Hybrid Particles. siNEG\(_{Atto655}\), initial stock solution was prepared at 20 μM. Prior to in vitro cell uptake, siRNA complex preparations were carried out as follows: 1.0 μg (250 μL of siNEG\(_{Atto655}\)) in 5% dextrose was mixed with an equal volume of Lipofectamine2000 (4.0 μg) in 5% dextrose at 4:1 nitrogen (N)/ phosphate (P) charge ratio. ΔHBcAF488 or Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\) hybrid particles were prepared using method III (found to be the most efficient encapsulation method) at 1:1 HBc/siNEG molar ratio (containing 480 μg of HBc and 1.0 μg of siNEG), buffer-exchanged to 1X PBS, to maintain isotonicity, and concentrated to 500 μL final volume using Vivaspin MWC0 10 kDa. MDA-MB-468 (HER2\(^{+}\)) and SKBR-3 (HER2\(^{-}\)) cells were seeded at a density of 1 x 10\(^5\) cells/well in 24-well plates and allowed to attach overnight. After washing with PBS buffer, cells were then treated with naked siNEG\(_{Atto655}\), Lipofectamine-siNEG\(_{Atto655}\) complexes, or ΔHBcAF488−siNEG\(_{Atto655}\) or Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\) hybrid particles in serum-free media for 1,4, or 24 h. For 24 h treatment, after 4 h incubation with the hybrids, FBS was added to each well to reach final FBS concentration of 10% v/v, and incubated for another 20 h. At the end of incubation period, cells were washed twice with PBS buffer, trypsinized, and centrifuged at 50 g for 5 min, and the cell pellet was resuspended in 250 μL of PBS buffer. The internalization of HBc–siNEG hybrid particles\(_{Atto655}\) and Lipofectamine-siNEG\(_{Atto655}\) treatments were used as a negative and positive control, respectively. The data were analyzed using BD CellQuest software (BD Biosciences, USA). The effect of the presence of serum on the HBc–siRNA hybrid particle uptake was examined by performing the same treatments described above in complete serum-containing media for 1 or 24 h.

2.9. In Vitro Cytotoxicity Studies of HBc–siRNA Hybrid Particles. MDA-MB-468 (HER2\(^{+}\)) or SKBR-3 (HER2\(^{-}\)) cells were seeded at a density of 5 x 10\(^3\) cells/well in 96-well plates and allowed to attach overnight at 5% CO\(_2\) and 95% air at 37 °C. After washing with PBS buffer, cells were treated with lipoplexes (siNEG or siPLK1 complexed with Lipofectamine2000 (4:1 N/P ratio)), ΔHBcAF488−siNEG, ΔHBc−siPLK1, Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\), or Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\)−siPLK1, all at 1:1 HBc/siRNA complex ratio, containing increasing concentrations of siRNA at 5, 10, or 20 nM in serum-free media for 4 h. After 4 h, the formulation was discarded, cells were washed with PBS buffer, complete medium was added, and cells were incubated for another 68 h. Cell viability was examined by MTT (3-(4, 5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay. Briefly, the medium from each well was removed after 72 h incubation and replaced with 120 μL of MTT solution (at a concentration of 5 mg/mL in PBS with a further 1:6 dilutions in media prior to use). Cells were incubated for 3 h in 5% CO\(_2\) and 95% air at 37 °C. Solution was discarded, and formazan was then solubilized in 200 μL of DMSO. The absorbance was read at 570 nm (A\(_{570nm}\)) using a FLUOSstar OPTIMA plate reader (BMG Labtech, Germany).

The effects of the serum on the HBc–siRNA hybrid particle cytotoxicity were examined by treating cells with the same complexes and conditions mentioned above but in complete media instead. Cell viability was examined by MTT assay as described previously.

2.10. In Vitro PLK1 Gene Silencing in HER2+ Cells. SKBR-3 and MDA-MB-435-MLE cells were seeded at a density of 5 x 10\(^6\) cells/well in 6-well plates and allowed to attach overnight at 5% CO\(_2\) and 95% air at 37 °C. After washing with PBS buffer, cells were then treated with lipoplexes (siNEG or siPLK1 complexed with Lipofectamine2000 (4:1 N/P ratio)) or Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\) or Z\(_{HER2}ΔHBcAF488−siNEG\(_{Atto655}\)−siPLK1 at 1:1 HBc/siRNA molar ratio containing 20 nM siRNA in serum-free media for 4 h, and then serum was added to the treatment at a final concentration of 10% v/v, and cells were incubated for another 20 or 44 h. After 24 or 48 post-transfection, cells were washed twice with ice-cold PBS buffer before lysis. Lysates were examined by Western Blotting analysis to determine the level of PLK1 expression (as described in section 2.15).
2.11. Animal Studies and Tumor Inoculation. All animal experiments were performed in compliance with the UK Home Office (1989) Code of Practice for the housing and care of Animals used in Scientific Procedures. MDA-MB-435-MLE cells (HER2+) were used as the in vivo tumor model. Male NSG mice aged 4–6 weeks (Charles River, UK) were injected intraperitoneally with MDA-MB-435-MLE cells (5.0 × 10⁶ cells in 100 μL PBS), and tumor progression was observed by whole body bioluminescence imaging using IVIS Lumina series III In Vivo Imaging Device (IVIS) (Caliper Life Sciences, PerkinElmer, USA). In each imaging session, mice were administered d-luciferin at 150 mg/kg subcutaneously. Animals were visualized at 12 min postluciferin injection to ensure consistent photon flux and analyzed using Living Image software (Caliper Life Sciences, PerkinElmer, USA). Photon flux from the tumor is proportional to the number of live cells expressing luciferase.

2.12. Therapy Study. On day 4 post-tumor inoculation, NSG mice were divided into 4 treatment groups (n = 7): PBS, ZHER2-ΔHBc−siNEG, ΔHBc−siPLK1, and ZHER2-ΔHBc−siPLK1 treatment. Doses used in therapy experiments were 4 μg of siNEG or siPLK1 per injection per mouse; all treatments were injected intraperitoneally. A total of seven doses of each treatment were given at 3 day intervals on days 4, 7, 10, 13, 16, 19, and 22 post-tumor inoculation. Tumor growth was observed by bioluminescence imaging twice weekly (days 4, 7, 11, 14, 18, 21, and 25), as described above. Mice were sacrificed on day 28. Only solid adherent tumor nodules were excised and weighed, while tumor cells suspended in ascites were discarded.
2.13. Determination of TNF-α Concentration with ELISA. Animals from the therapy study were sacrificed on day 28 (6 days after the last injection), and sera was analyzed for mouse tumor necrosis factor α (TNF-α). A separate group of tumor-bearing mice was injected with lipopolysaccharide (LPS, 3 mg/kg) intraperitoneally, and their sera were collected 1 h postinjection as positive controls. Sera were diluted 1:2 using 1× PBS and analyzed using a BD OptEIA Mouse TNF (Mono/Mono) ELISA set as per the manufacturer’s protocol.

2.14. Organ Histology. Major organs (heart, lung, liver, spleen, and kidney) and tumors were harvested and immediately fixed in 10% neutral buffered formalin at 5 mm² pieces. Tissue pieces were paraffin-embedded and sectioned for hematoxylin and eosin (H&E) staining according to the standard histological protocols at the Royal Veterinary College, U.K. The stained sections were analyzed with a Leica DM 1000 LED Microscope (Leica Microsystems, U.K.) coupled with CDD digital camera (QImaging, U.K.).

2.15. Gene Silencing Evaluation by Western Blot. For in vitro studies, cells were collected directly into a lysis buffer (50 mM Tris-HCl, pH 8.0, 1.0% Triton X100, 150 mM NaCl, 5 mM EDTA) and incubated on ice for 10 min. For in vivo studies, at sacrifice on day 28, collected tumors were crushed directly into the lysis buffer and incubated on ice for 30 min. Samples were then separated by centrifugation at 20 800 × g for 15 min at 4 °C. After the collection of supernatant, total protein concentration was measured using BCA protein assay kit. A total of 200 μg of protein was then taken from each supernatant and separated on 15% SDS-PAGE gels for electrophoretic transfer onto a nitrocellulose membrane. After blocking in BSA, membranes were immunoblotted with mouse anti-GAPDH antibody at 1:10 000 dilution for secondary antibody. For GAPDH, membranes were normalized to GAPDH in each sample and correlated to the reference) detection separately. For PLK1, membranes were immunoblotted with rabbit monoclonal PLK1 antibody at 1:1000 dilution as primary antibody, followed by HRP-linked anti-rabbit at 1:10000 dilution and Precision Protein StreptTactic-HRP conjugate at 1:10 000 dilution for secondary antibody. For GAPDH, membranes were immunoblotted with mouse anti-GAPDH antibody at 1:1000 dilution (primary antibody), followed by mix of HRP-linked antimouse at 1:10000 dilution (secondary antibody) and Precision Protein StreptTactic-HRP conjugate at 1:100000 dilution. The specific bands were identified using enhanced chemiluminescence (ECL) detection system. The membrane was visualized using the ChemiDocMP (Bio-Rad Laboratories, USA) and analyzed with Image Lab (Bio-Rad Laboratories, USA) software. The quantity of PLK1 transcript was normalized to GAPDH in each sample and correlated to the normalized expression in naive cells to calculate the fold changes.

2.16. Statistics. In all experiments, data were presented as mean ± SD, where n denotes the number of repeats. Significant differences were determined using one-way ANOVA. The t-value, degrees of freedom, and two-tailed significance (p-value) were resolved as *p < 0.05, **p < 0.01, and ***p < 0.001.

3. RESULTS

3.1. Disassembly/Reassembly of Hbc Particles by DTT/CaCl2 (Method III) Is the Most Efficient Method for siRNA Encapsulation with Complete Complexation at 1:1 siRNA to Particle Molar Ratio. The encapsulation of siRNA into Hbc particles was first optimized. The recombinant ZHER2−ΔHbc particles were expressed, purified, and assembled. The assembled ZHER2−ΔHbc particles were then fluorescently labeled with Alexa Fluor 488 succinimidyl esters (ZHER2−ΔHbcAF480) as described previously.20 Fluorescently labeled noncoding siRNA (siNEGs) was encapsulated into ZHER2−ΔHbcAF480 particles using three different methods, disassembly/reassembly by urea (method I), osmotic shock (method II), or disassembly/reassembly by DTT/CaCl2 (method III), to form Hbc−siRNA hybrid particles; all methods are schematically described as shown in Figure S1. In method I, urea was used as a denaturing agent to weaken the protein−protein interactions between Hbc monomers, thus loosening the particle formation, which is expected to allow the siRNA to penetrate into the Hbc particles.29,30 Additionally, the high salt concentration of NaCl added in the assembly stage is expected to increase the electrostatic repulsion between the protein monomers, promoting the ZHER2−ΔHbcAF480 assembly in an organized manner forming of Hbc−siRNA hybrid particles.31 In method II, an osmotic shock was induced to encapsulate siRNA following reported protocols with some modifications. An osmotic shock or osmotic stress introduced by a sudden change in the solute concentration around the particles is expected to cause the particles’ pores to swell up, allowing the entry of the siRNA into the particles.33 This method is not expected to cause disassembly of the particles. In method III, Hbc particles were disassembled in the presence of a chelating agent, EGTA, and a reducing agent, DTT.34 An EGTA−DTT combination is therefore expected to completely disrupt the Hbc particles; EGTA has high affinity for Ca2+. As the assembly of Hbc particles is highly Ca2+-dependent, Hbc was then reassembled by adding Ca2+ to increase the electrostatic repulsions between the protein monomers.35,36 The size of empty Hbc particles was confirmed to be ~30 nm.20

Figure 2. Stability studies of Hbc−siNEG hybrid particles against serum and RNase. Gel retardation assay for Hbc−siNEG hybrid particles subjected to (A) serum; (B) RNase at Hbc/siNEG 1:1 molar ratio. The hybrid particles were prepared at a fixed siNEG amount of 0.25 μg. Hbc−siNEG hybrid particles were incubated with 50% v/v FBS (0, 24, or 48 h) or RNase (0, 24, or 48 h) at 37 °C. EDTA was used to inactivate serum proteins.
The siRNA encapsulation profiles using the three different methods were assessed and compared using gel retardation assay as shown in Figure 1A. The green color is assigned to HBc while blue color indicates either free or complexed siRNA. Decreasing signals of free siRNA bands (blue color) observed at the bottom of the gel is a sign of complexation. In method I, siRNA encapsulation efficiency was improved with increasing complexation molar ratios of HBc/siRNA from 2:1 to 4:1. Some free siRNA was still observed even at 4:1 HBc/siRNA molar ratio. In methods II and III, complete encapsulation was achieved at HBc/siRNA molar ratio 4:1 and 1:1, respectively. No free siRNA was detected at the bottom of the gel in such conditions with the concomitant observation of increased fluorescence in the wells.

Another technique, the bead coupling assay, schematically presented in Figure S2, was employed to confirm siRNA incorporation into ZHER2ΔHBc−siNEG particles by flow cytometry using FL-1 (Z-ΔHBcAF488) and FL-4 (siNEGAtto655) channels. Using this method, siRNA encapsulated using method III showed the highest amount of siRNA encapsulation, compared to method I and method II (Figure 1B). This result is in agreement with the results obtained by gel retardation assay. It was concluded that approximately 1 mol of siRNA can be encapsulated in 1 mol of HBc particles using method III. Method III was used in the subsequent studies.

3.2. HBc−siRNA Hybrids Are Stable against RNase and in Serum.

The ability of HBc−siRNA hybrids to protect siNEG from degradation by nuclease was evaluated by challenging the hybrid particles against serum. Naked siNEG and HBc−siNEG hybrid particles were incubated at 37 °C in 50% v/v FBS for 0, 24, or 48 h. As shown in Figure 2A, FBS alone exhibited weak signals shown in the middle of the gel.
(first lane) that are located differently from the signals of either free or encapsulated siNEG occurring at the bottom of the gel or at the wells, respectively, shown in other lanes. After 24 h incubation with serum, naked siNEG was almost degraded, with complete degradation of siNEG observed at 48 h incubation.

In contrast, HBc–siNEG particles exhibited strong signals for encapsulated siNEG in the well at all time points. Interestingly, free siNEG signals were also present at the bottom of the gel, even at the 0 h time point, suggesting a partial release of siNEG from the hybrids in the presence of serum. These results suggest that complexation with HBc particles improves siRNA stability in the presence of serum.

The ability of HBc–siRNA hybrids to protect siRNA against a ribonuclease (RNase) was evaluated by incubating the hybrids with RNase for 0, 24, or 48 h at 37 °C. As shown in Figure 2B, naked siRNA was completely degraded after 24 h incubation with RNase. In contrast, strong signals were observed in the wells of HBc–siNEG particles even after 48 h of incubation with the RNase, indicating that the HBc particles can protect siNEG from enzymatic degradation.

3.3. HER2-Targeting HBc–siRNA Hybrids Are Selectively Taken Up in HER2-Positive Cancer Cells in Vitro.

The intracellular uptake of HBc–siRNA hybrids in MDA-MB-468 (HER2⁻) and SKBR-3 (HER2⁺) cells was evaluated by flow cytometry via FL-1 (ZHER2ΔHBcAF488) and FL-4 (siNEGAtto655) channels and expressed as fold increase in median fluorescent intensity (MFI) (Figure 3). All cells were treated with naked siNEGAtto655, Lipofectamine–siNEGAtto655 complexes, ΔHBcAF488–siNEGAtto655 or ZHER2ΔHBcAF488–siNEGAtto655 hybrids for 1, 4, or 24 h. As shown in Figure 3, treatment of SKBR-3 cells with ZHER2ΔHBcAF488–siNEGAtto655 hybrids promoted an apparent fold increase at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars) in contrast to lower uptake measured at both FL-1 and FL-4 channels in a time-dependent manner (last green bars)

Figure 4. Cell viability of cancer cell lines treated with siRNA–HBc particles in vitro. (A) MDA-MB-468 and SKBR-3 cell lines were treated with siNEG and siPLK1 at increasing concentration (5, 10, and 20 nM) in serum free media for 4 h followed by incubation in complete media for another 68 h. Treatment groups include 5% dextrose, 1×PBS, lipoplexes (siNEG or siPLK1 in complex with Lipofectamine2000 (4:1 N/P)), ΔHBc–siNEG, ΔHBc–siPLK1, ZHER2ΔHBc–siNEG, or ZHER2ΔHBc–siPLK1, all at 1:1 HBc/siRNA molar ratio. Cell viability was analyzed and expressed as a percentage of naive cells, determined by MTT assay. (B) Percentage cell viability (SKBR-3 cells) after 72 h transfection ±10% serum at 20 nM final concentration. Cell viability was analyzed and expressed as a percentage of naive cells, determined by MTT assay. Values are expressed as mean ± SD (n = 6). *p < 0.05, **p < 0.01, ***p < 0.001, relative to naïve (one-way ANOVA test).
hybrids in HER2+ SKBR-3 cells. The intracellular uptake of HBc−siRNA hybrid particles in vitro was also studied in the presence of serum. Results suggested that the presence of serum slightly but significantly reduced the uptake of the hybrids overall (FL-1, FL-4) in both cell types (Figure S3).

3.4. HER2-Targeting HBc−siPLK1 Hybrids Are Functionally Active in Vitro in the Presence or Absence of Serum. The cytotoxic effect of HBc−siRNA hybrids was assessed in MDA-MB-468 (HER2−) and SKBR-3 (HER2+) cells in vitro. Nonfunctional siNEG was replaced with functional siRNA specific to PLK1 gene (siPLK1), in order to assess the gene-silencing efficacy. Cells were treated with 5, 10, or 20 nM siNEG or siPLK1 encapsulated in nontargeted ΔHBc or targeted ZHER2−ΔHBc particles for 72 h, and cell viability was determined by MTT assay. Lipofectamine2000−siPLK1 complexes were used as a positive control (Figure 4A). As expected, cells treated with ΔHBc−siPLK1 showed no significant toxicity in both cell lines. These results are in agreement with those obtained for intracellular uptake studies for ΔHBc particles alone as these particles lose their non-specific binding due to lack of the arginine-rich domain.20 A significant dose-dependent reduction in cell viability was observed for Lipofectamine2000−siPLK1 in both cell lines. Treatments with ΔHBc−siPLK1 did not cause significant cytotoxicity in both cell lines. In contrast, treatments with ZHER2−ΔHBc−siPLK1 induced 30–40% cell death in SKBR-3 cells but not MDA-MB-468 cells. Interestingly, when SKBR-3 cells were treated with ZHER2−ΔHBc−siNEG, a significant amount, approximately 10%, cell death was observed. This could be due to some intrinsic toxicity exhibited by ZHER2−ΔHBc, due to the presence of the affibody, that is, HER2 receptor blockade activity, since such toxicity was not observed when ΔHBc hybrids were tested under same conditions.

The cytotoxic effect of HBc−siRNA hybrids in HER2+ cells was further studied with subject to serum competition. Cell viability of SKBR-3 cells treated with 20 nM of siNEG or siPLK1 encapsulated in ZHER2−ΔHBc particles was assessed by MTT assay at 72 h postincubation. Lipofectamine2000−siPLK1 complexes were used as a positive control (Figure 4B). Similar reduction in cell viability was seen for cells treated with Lipofectamine2000−siPLK1 regardless of the presence of serum. A slight reduction in cytotoxicity was observed for ZHER2−ΔHBc−siPLK1 in the presence of serum (***p < 0.001), but this was significantly lower than equivalent siNEG group.

3.5. Efficient PLK1 Knock-down Is Confirmed in Vitro by Western Blotting. To confirm if the reduction in cell viability in the case of ZHER2−ΔHBc−siPLK1 hybrid treatment was associated with PLK1 gene silencing, the levels of PLK1 protein expression in vitro were examined using Western blotting. HER2+ cells, SKBR-3 and MDA-MB-435-MLE cells, were incubated with ZHER2−ΔHBc−siPLK1 hybrids for 24 or 48 h. GAPDH, a housekeeping gene, was used as an internal control. Lipofectamine2000−siPLK1 complexes were included...
as a control transfecting agent. There was a clear difference in gene silencing efficacy between the two vectors in SKBR-3 cells, where \( Z_{HER2} \cdot \Delta HBc \cdot siPLK1 \) hybrids showed more efficient PLK1 gene knockdown than Lipofectamine2000–siPLK1 complexes as measured by PLK1 protein level (Figure 5A,B). In the case of \( Z_{HER2} \cdot \Delta HBc \cdot siPLK1 \) hybrids transfection, PLK1 expression was reduced to 33.4% ± 1.5% and 20.4% ± 0.6% at 24 and 48 h post incubation, respectively, compared to naive cells. In the case of Lipofectamine2000–siPLK1 complexes, the protein levels were 53.6% ± 0.5% and 33.0% ± 0.6% after the same incubation times (*\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \), relative to group of PBS (one-way ANOVA test).

3.6. \( Z_{HER2} \cdot \Delta HBc \) and Not siPLK1 Treatment Results in Reduction in Intraperitoneal Solid Tumor Mass in Mice. Representative images for in vivo bioluminescent imaging are shown. (A) Representative images for in vivo bioluminescent imaging are shown. (B) Growth curves of MDA-MB-435-MLE tumors. (C) Weight of tumors excised from mice after sacrifice. Data expressed as mean ± SD (\( n = 7 \). *\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \), relative to group of PBS (one-way ANOVA test).

Figure 6. Assessment of tumor bioluminescence and solid weight tumors in HER2-expressing intraperitoneal tumor mouse models. NSG mice (\( n = 7 \)) were intraperitoneally injected with 5.0 × 10^6 MDA-MB-435-MLE cells. Mice were treated with PBS, \( Z_{HER2} \cdot \Delta HBc \cdot siNEG \) (4 \( \mu g \) of siNEG per injection per mouse), \( \Delta HBc \cdot siPLK1 \) (4 \( \mu g \) of siPLK1 per injection per mouse) and \( Z_{HER2} \cdot \Delta HBc \cdot siPLK1 \) (4 \( \mu g \) of siPLK1 per mouse), intraperitoneally. Seven treatments were given intraperitoneally at 3 day intervals, commencing on day 4 post-tumor inoculation. Tumor growth was observed by whole body imaging. (A) Representative images for in vivo bioluminescent imaging are shown. (B) Growth curves of MDA-MB-435-MLE tumors. (C) Weight of tumors excised from mice after sacrifice. Data expressed as mean ± SD (\( n = 7 \). *\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \), relative to group of PBS (one-way ANOVA test).
ΔHBc−siPLK1 (***p < 0.001). No significant difference between these two treatments was observed. Bioluminescence signals are expected to be directly proportional to the tumor metabolic activity. It is however worth mentioning that the signals here are obtained from both semiadherent in ascites and adherent cells forming tumor nodules in the peritoneal cavity, due to the nature of this tumor type. The metabolic activity of singly suspended cells is likely to be higher than those obtained from tumor nodules because the latter are likely to develop a necrotic core. To measure the solid tumor load, the weight of total tumors modules collected post-mortem was measured. Interestingly, contrary to the trends obtained from bioluminescence imaging, weights of solid tumors treated with ZHER2−ΔHBc−siNEG (0.72 ± 0.14 g, **p < 0.01) were significantly lower than those of the PBS group (1.01 ± 0.14 g) (Figure 6C). The level of PLK1 protein expression in HER2-expressing tumors was examined from the protein lysate of MDA-MB-435-MLE tumors. PLK1 proteins were extracted from the solid tumor nodules, collected on day 28 at sacrifice, and the PLK1 expression was evaluated by Western blotting (Supporting Information Figure S4A). ΔHBc−siPLK1 and ZHER2−ΔHBc−siNEG were included to account for any nonspecific toxicity of the siRNA or the ZHER2−ΔHBc particles, respectively. The PLK1 expression level of PBS group was normalized to 100% ± 1.49% relative to expression of GAPDH. Intraperitoneal treatment of ZHER2−ΔHBc−siPLK1 hybrid particles showed slight but significant reduction in the PLK1 protein levels.

Figure 7. Microscopic examination of major organs and tumor sections of MDA-MB-435-MLE tumor-bearing NSG mice. Mice were treated with PBS, ZHER2−ΔHBc−siNEG (4 μg of siNEG/mouse), ΔHBc−siPLK1 (4 μg of siPLK1/mouse), and ZHER2−ΔHBc−siPLK1 (4 μg of siPLK1/mouse), intraperitoneally. Seven treatments were given intraperitoneally at 3 day intervals, commencing on day 4 post-tumor inoculation. At day 28, the mice were sacrificed, and the organs were immediately fixed in 10% neutral buffered formalin as 5 mm3 pieces. These pieces were then paraffin-embedded for hematoxylin and eosin stains (H&E) and analyzed by microscopy. No signs of toxicity were observed in heart, lungs, liver, spleen, and kidneys. Red arrows point to the necrotic area in tumor section (pinkish area). Scale bars are 50 μm.
mice receiving 7 doses of Z HER2-

Significantly high levels of TNF-\(\alpha\) (1048 ± 320 pg/mL, ***\(p < 0.001\)) were observed in the serum from mice treated with LPS (Supporting Information, Figure S6). On the other hand, no significant differences in serum TNF-\(\alpha\) levels were found in mice receiving 7 doses of \(Z_{\text{HER2}}\Delta\text{HBc}–\text{siNEG}, \Delta\text{HBc}–\text{siPLK1},\) or \(Z_{\text{HER2}}\Delta\text{HBc}–\text{siPLK1} (12–21 pg/mL)\) intra-peritoneally within 28 days compared to the PBS group (13 ± 7 pg/mL).

Histological examination of the major organs (heart, lung, liver, spleen, and kidney) with H&E staining exhibited no obvious histological changes compared to control animals (Figure 7). In mice treated with \(Z_{\text{HER2}}\Delta\text{HBc}–\text{siPLK1},\) tumors showed regions of necrosis (Figure 7, Supporting Information, Figure S7, red arrows), which was not observed in tumors from control and \(Z_{\text{HER2}}\Delta\text{HBc}–\text{siNEG}\) treated mice. Both early stage (fragmented and small nucleus) and late stage (ghost shells without nucleus) necrosis were found in necrotic areas, indicating an accumulative therapeutic effect of \(Z_{\text{HER2}}\Delta\text{HBc}–\text{siPLK1}\) throughout the therapy time course.

4. DISCUSSION

In this study, we validated the utilization of recombinant HER2-targeting HBc particles as an efficient delivery carrier for siRNA. To the best of our knowledge, studies on delivering siRNA using HBc particles simultaneously with a specific HER2 targeting to cancer cells have not yet been reported. However, many studies have used HBc particles to deliver small molecules including oligonucleotides38 and drugs39 to cancer cells. Lee et al. reported the use of the HBc particles encapsulated with a green fluorescent protein by dissociating the HBc particles in the presence of urea and subsequently reassembling by dialysis to remove the denaturant.39 In another study, Beterams et al. successfully packaged up to 240 subunits of 17 kDa nuclease into the interior of recombinant HBc particles. Interestingly, the packaged nuclease was enzymatically active and the structure of the particles was not affected.40

HBc particles like other virus-like particles have the capacity to load RNA. Encapsulating and loading nucleic acids within HBc particles is simpler than any other small molecules. This is due to the properties of nucleic acids, which are usually bigger, and the protein capsids have emerged to carry and encapsulate a similar type of cargo, in this case, their own viral genomes. In addition, dissimilar to the lipid-based, metal-based, and polymer-based nanocarriers, HBc particles are highly uniform, can be reproduced with an actual number of monomer proteins, and can be organized in a consistent arrangement. HBc particles are expected to possess a significantly reduced lot-to-lot variability and exact cargo release properties. Furthermore, once the HBc particles managed to internalize in the targeted cells, they are able to disassemble and unleash all of the therapeutic molecules at once, compared with polymeric nanoparticles, which degrade and slowly release the loaded molecules over time. Even though slow cargo release might occasionally be propitious, in many cases, especially in cancer treatment, an immediate cargo release is preferred. HBc particles also have the capability to be functionalized chemically or genetically to exhibit fragments like antibody or peptide, for specific cellular targeting.41 Such characteristics make them distinguishable from other types of nanocarriers for nucleic acid delivery. These make HBc particles have more advantages than other nanoparticles. Taking advantage of the ability of HBc particles to encapsulate small biomolecules into their interior, we aimed at formulating recombinant HBc particles for delivery of siRNA capable of binding and possibly blocking HER2 receptors, allowing a combinatorial approach to kill HER2 cancer cells.

We demonstrated that siRNA encapsulation using method III showed the highest amount of siRNA encapsulated into HBc particles at the 1:1 HBc/siRNA molar ratio, compared to method I and method II. Method I was not able to encapsulate a higher amount of siRNA compared to method III even though urea is a known denaturing agent for its compatibility with protein folding. This could be due to the incomplete reassembly caused by NaCl. The ionic strength from the monovalent sodium ions might be able to trigger the HBc protein assembly,42 but not in an amount sufficient to increase the electrostatic charges between HBc protein monomers to cause complete reassembly of the proteins. Our result showed an enhancement of siRNA encapsulation using the osmotic shock in method II compared to method I. However, complete siRNA encapsulation was only achieved at 4:1 HBc/siRNA molar ratio. Studies reported that encapsulation of nucleic acids using osmotic shock is pH-dependent. Braun et al. reported that the loading of oligonucleotide and plasmid DNA into polyoma VP1 virus-like particles achieved the highest encapsulation efficiency only at acidic pH of 5.38 In our study, pH 7.4 was maintained throughout the experiment, which could be the reason for the lower siRNA encapsulation than expected. Nevertheless, we successfully encapsulated siRNA into HBc particles using method III, a new combination method involving the use of EGTA and DTT. EGTA and DTT were required for the particle disassembly as EGTA has high affinity to Ca\(^{2+}\). HBc particle assembly is highly Ca\(^{2+}\)-dependent,35,36 and DTT has the ability to break the disulfide bringing between HBc dimers.35

The blue intensity reflects quantitatively the amount of free migrating siRNA. Blue signals in the well are only semi-quantitative (larger aggregates may float out of the wells during electrophoresis) and are representative of siRNA in the complexed form. In method I, it was not surprising to see a reduction in the amount of free migrating siRNA as the amount of HBc increased indicating an increased degree of siRNA complexation. In case of methods II and III, the complexation was efficient and occurred at all ratios used so that no free siRNA could be seen at all the tested ratios.

Instability poses a major problem in siRNA applications due to its degradation by the activity of serum or enzymes. We tested the stability of HBc–siRNA hybrid particles by challenging the hybrid particles with serum or RNase. Agarose gel results indicated that the HBc–siNEG hybrid particles confer great siRNA protection ability under several test
conditions. In this study, HBc particles were expected to be able to protect siNEG from enzymatic degradation, but substantial signals can be seen at 0 h either due to partial instability or the presence of some surface-absorbed siRNA, which may have been detached in the presence of serum. The results explained the reduced uptake of siRNA in the presence of serum (Figure S3).

siRNA is an anionic macromolecule that is unable to enter cells by passive diffusion mechanisms, and thus an appropriate delivery system is needed to enhance its cellular uptake efficiency. Many studies have reported successful siRNA delivery in vitro using nanoparticles. For example, Lipofectamine2000, a cationic lipid formulation, is widely used for in vitro siRNA transfection, which succeeded in transfecting siRNA in a mouse’s vascular endothelial cells.44 Recently, Pereira et al. reported the use of carbon nanotube–liposome hybrids to deliver siRNA in human epithelial lung carcinoma A549 cells.45 In our study, a successful delivery of siRNA in vitro was achieved using ZHER2ΔHBcAF488−siNEGΔA0655 hybrids in two different HER2-expressing cancer cell lines. SKBR-3 cells, overexpressing HER2 receptors on cell membrane surface, showed an enhanced uptake of ZHER2ΔHBcAF488−siNEGΔA0655 hybrids compared to ΔHBcAF488−siNEGΔA0655 hybrids. This could be attributed to the presence of ZHER2Δ affibody expressed on the particle surface that specifically binds the HER2 receptor.28,45 In competition experiments using serum, intracellular uptake of HBc−siRNA hybrids was still taking place but with a slight reduction, which was also observed with the positive control, Lipofectamine2000−siNEGΔA0655 complexes. The enhanced cellular uptake of the ZHER2ΔHBcAF488−siNEGΔA0655 hybrids revealed ZHER2ΔHBc as a promising system for the delivery of genetic material. Comparison and quantification of uptake was done by flow cytometry. Confocal laser scanning microscopy, to confirm site of localization inside the cells, could not be performed on MDA-MB-435 cells due to the semiadherent nature of the cells.

In the cytotoxicity assessments, a significant 10% cell death was observed for ZHER2ΔHBc−siNEG hybrids, whereas ΔHBc−siNEG hybrids did not show any significant cytotoxicity. When the nonfunctional siNEG was substituted with a siRNA sequence specific for PLK1 gene, a further significant reduction in cell viability was achieved. The consequent quantitatively assessments by Western blotting of PLK1 protein expression confirmed the PLK1 knockdown after ZHER2ΔHBc−siPLK1 hybrid treatments using the two HER2 positive cell lines, SKBR-3 or MDA-MB-MB435-MLE. The gene-silencing efficacy was even more efficient compared to Lipofectamine2000 transfection, which is encouraging for further in vivo investigation.

The choice of the location of the tumor model and route of administration was based on our previous study where tumor uptake of ∼10% of the total injected HBc dose per gram of tissue was achieved.28 Furthermore, even though the orthotopic inoculation of cancer cells in the mammary fat pad contributes a better model to resemble the early stages of breast cancer advancement, the IP tumor mouse model provides a much more precise representation of clinical disease progression of tumors, which includes metastatic dissemination. While combination of HER2-targeted HBc particles and siPLK1 (or siNEG equivalent formulation) resulted in apparent increase in cancer cell population based on whole body bioluminescence imaging, tumor weight measurement of solid tumor nodules showed contradicting results where ZHER2ΔHBc-based treatments (siNEG or siPLK1) displayed significant reduction in tumor weights. This discrepancy in findings using both techniques was unexpected. A possible explanation could be the fact that the cell line used in this study is a semiadherent cell line.28 Due to this property, tumor cells can be present in ascites fluid, also known as malignant ascites, or as adherent tumor nodules in vivo.46 It is expected that the bioluminescence signals measured are the sum of suspended cells and larger adherent tumor nodules. The discrepancy seen between the bioluminescence and tumor weight measurements may be due to the shift in tumor status from adherent nodules (low bioluminescence) to suspension form (higher bioluminescence). This seems to be primarily a function of the ZHER2Δ affibody and not the siPLK1 treatment. Further studies are needed to understand how ZHER2Δ affibody–receptor binding may influence cell to cell communication and adhesion.

Immune response is always the main concern when it comes to gene therapy. Most of the toxicity associated with VLP nanocarriers arises from the innate and adaptive immune response to their capsids. Systemic administration of VLPs will cause the release of pro-inflammatory cytokines and chemokines, including TNF-α.47,48 Here, to abolish the immune response that might be induced from our HBc particles, we modified the major immunodominant region located between amino acids 75 and 83 with our ZHER2Δ affibody. This particular sequence has the capacity to exhibit a strong immune response.49,50 An establishment of targeting moieties at this sequence should be able to abolish a strong immune response by the host.51−53 The ELISA assay results showed no significant TNF-α level was observed in mice treated with recombinant HBc particles.

Western blotting data hinted that PLK1 silencing worked very slightly (∼10%), but this is obviously not sufficient or therapeutically acceptable. It is acknowledged that the siRNA dose injected into mice was low compared what has been used in other studies. The siPLK1 dose was relatively low (0.2 mg of siRNA/kg per injection) compared to what have been reported in other studies with systemic siPLK1 administration.54,55 The limited dose was due to the low siRNA encapsulation efficiency obtained with the studied particles. Future work should focus on methods to improve siRNA loading so that higher siRNA doses, at lower nanocarrier dose, can be achieved. Taking together the results of therapy (weight measurements on solid tumor nodules) and Western blotting studies, one may think that both the intrinsic bioactivity of ZHER2ΔHBc and siPLK1 silencing took place. The overall potency however is minimal and needs further improvement. The histology analysis suggested that the administration of HER2-targeted HBc particles might induce apoptosis or necrosis to the solid tumors itself.

Another possible factor of the low efficacy might be due to challenges encountered with intraperitoneal delivery route of nucleic acids. It has been reported that local intraperitoneal delivery results in rapid clearance of the nanocarriers from the peritoneal cavity. This rapid clearing from the cavity to the systemic circulation most probably will limit the amount of nanocarriers that reach the actual target cells or tissues.40 In addition, IP fluid possesses a complex biological environment, so stability of our hybrids should also assessed under conditions mimicking those of intraperitoneal fluid for a better representation of in vitro to in vivo correlation.57
5. CONCLUSION

We developed a new siRNA delivery carrier composed of recombinant HBc particles and Z\textsubscript{HER2} affibody. Z\textsubscript{HER2} affibody on the particle surface allowed the recombinant HBc particles to be recognized by the HER2 receptors overexpressed on the cancer cell surface and be subsequently internalized. Furthermore, gene silencing studies with siPLK1 revealed that HBc particles were able to adequately silence PLK1 genes to commensurate levels to cationic lipid nanocarriers in vitro. Following intraperitoneal administration in intraperitoneal xenografts, combination of HER2-targeted HBc particles and siPLK1 did not produce any delay of tumor growth in vivo; however, targeted HBc particles induced a reduction in the weight of solid tumors. Mice, after treatments did not show elevated TNF-\alpha levels in serum, and no obvious histological changes in major organs were observed post-mortem indicating that our HBc particles are biocompatible in vivo.

\section*{ASSOCIATED CONTENT}

\begin{itemize}
  \item Supporting Information
    The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsanm.8b00480.

    Additional details of materials, siRNA encapsulation efficiencies by aldehyde/sulfate latex beads assay, in vitro cellular uptake of HBc-siRNA hybrids subjected to serum competition, confirmation of siNEG\textsubscript{Ado655} encapsulation with beads coupling method, in vitro uptake studies in MDA-MB-468 and SKBR-3 cancer cells in presence or absence of serum, PLK1 gene silencing analysis in MDA-MB-435-MLE tumors, assessment of total body weight of HER2-expressing intraperitoneal tumors, microscopic examination of tumor sections of MDA-MB-435-MLEx tumors, TNF-\alpha cytokine levels in serum after treatment, microscopic examination of tumor sections of MDA-MB-435-MLEx tumors, NSG mice (PDF)
\end{itemize}

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\end{itemize}

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