Immobilization of a carbon nanomaterial-based localized drug-release system using a bispecific material-binding peptide

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Introduction: Inorganic materials are widely used in medical devices, such as artificial hearts, vessels, and joints, in stents, and as nanocarriers for drug-delivery systems. Carbon nanomaterials are of particular interest due to their biological inertness and their capability to accommodate molecules. Several attempts have been proposed, in which carbon nanomaterials are used as nanocarriers for the systemic delivery of drugs.

Materials and methods: We developed a drug-delivery system in which oxidized single-walled carbon nanohorns (oxSWNHs) were immobilized on a titanium (Ti) surface using material-binding peptides to enable localized drug delivery. For this purpose, we utilized a bispecific peptidic aptamer comprising a core sequence of a Ti-binding peptide and a SWNH-binding peptide to immobilize oxSWNHs on Ti.

Results: Scanning electron microscopy was used to confirm the presence of oxSWNHs adsorbed onto the Ti surface, and a quartz crystal microbalance was used to evaluate the binding process during oxSWNH adsorption. The oxSWNHs-ornamented Ti substrate was nontoxic to cells and released biologically active dexamethasone over a sustained period.

Conclusion: This oxSWNHs-immobilized system can be used to modify the surface of Ti in implants and be loaded with drugs that stimulate osteogenesis and bone regeneration.

Keywords: drug carrier, drug delivery, carbon nanomaterial, carbon nanohorn, peptide aptamer

Introduction

Controlled surface modification of substrates is critical for the development of biomaterials that can interact with cells.1–4 Surface modifications can be classified as physical, chemical, or biological.5 In the chemical approach, various types of drug compounds are used as components of modifiers, which induce highly specific reactions in the biological system. To develop a functionalized surface that can release drugs into the microenvironment adjacent to the materials, we must first establish a system to load the drug and slowly release it from the carrier.

Titanium (Ti) is widely used as a biomaterial for dental implants, artificial joints, and stents.6 For developing next-generation titanium implants, several modification methods have been proposed to reduce the time required for osseointegration. These methods include the use of sintered metal beads, surface blasting, acid or alkaline treatment, anodization, and ion implantation.7–10 Surface chemistry is a pivotal factor in the regulation of interactions between the biomaterial and cells/tissue.11–14 In addition to these physical and chemical methods, modification using natural proteins, such as collagen, cell-adhesive molecules, or growth factors, has also been attempted.5 For example, Kashiwagi et al.15 ornamented Ti surfaces with modified bone-morphogenetic protein.
(BMP) that had an affinity for Ti. Different methodologies have been designed to control drug release from Ti.\textsuperscript{15,16} One strategy relies on surface modification. Usually, the rough Ti surface contributes to enhanced drug loading as compared with plane Ti. For example, a Ti plate containing nanopores shows enhanced drug-loading capability.\textsuperscript{17} Another approach involves attachment of therapeutic molecules directly to the Ti surface. Therapeutic molecules are chemically attached to a pure Ti surface\textsuperscript{18} or to self-assembled monolayers, which act as a glue.\textsuperscript{19} Additionally, drug coatings comprising drugs and hydroxyapatite\textsuperscript{20} or polymer\textsuperscript{21} show controlled drug-release profiles in vitro and in vivo.

Carbon nanomaterials have potential applications as carriers in drug-delivery systems\textsuperscript{22–25} because of their favorable characteristics, including stability, inertness, and a large surface area for drug loading.\textsuperscript{30–33} A single-walled carbon nanohorn (SWNH) is a spherical aggregate with a diameter of ~50 nm to ~100 nm and comprising ~2,000 single-walled graphene tubules with a nonuniform diameter of 2 nm to 5 nm and a length of 40 nm to 50 nm.\textsuperscript{34} SWNHs have an extensive surface area and intertubule interstices that can accommodate large amounts of adsorbed molecules.\textsuperscript{35} Moreover, the surface area can be enlarged by oxidization, which causes the formation of nanowindows in the SWNH walls,\textsuperscript{36} through which a variety of small molecules can infiltrate into the inner space.\textsuperscript{37} Oxidized SWNHs (oxSWNHs) can open two to four holes that allow the passage of various materials, including drugs, which can then be stored inside the aggregate.\textsuperscript{37,38} The ability of oxSWNHs to load and release drugs, such as dexamethasone (DEX),\textsuperscript{28} doxorubicin,\textsuperscript{40} and cisplatin,\textsuperscript{41} has also been shown, and no acute toxicity has been reported for SWNHs in animals,\textsuperscript{33,42,43} highlighting their utility for biological applications.

In this study, we used a Ti-binding peptide (TBP-1)\textsuperscript{44} and SWNH-binding peptide (NHBP-1)\textsuperscript{45} to immobilize a drug-delivery system on a Ti surface. TBP-1 is a material-binding peptide isolated as an aptamer against Ti. Although originally isolated as a 12-mer peptide (RKLPDAPGMHTW), further mutational analyses revealed that a hexapeptide sequence (RKLPDPA; named minTBP-1) is sufficient to mediate binding. The arginine, proline, and aspartate in this sequence (R1, P4, and D5, respectively) are critical for Ti recognition, suggesting that electrostatic interactions constitute the predominant binding force.\textsuperscript{46} NHBP-1 is a 12-amino acid aptamer (DYFSSPYEQLF) isolated as a specific SWNH binder using a peptide phage system. Non-covalent modification with a conjugate of polyethylene glycol and NHBP-1 enhances the dispersibility of hydrophobic oxSWNHs under aqueous conditions.\textsuperscript{41} Recent studies of peptide aptamers have highlighted the importance of valency on the part of the displayed peptide for target affinity with the conjugate. Multivalent display of peptide aptamers increases the affinity of the conjugate for the target,\textsuperscript{47,48} and in one instance, multivalency increased conjugate adsorption.\textsuperscript{2} We speculated that TBP–NHBP generated by fusing minTBP-1 and NHBP-1 would be adsorbed onto oxSWNHs, enabling a multivalent display of minTBP-1 on oxSWNHs and allowing the latter to be adsorbed onto the Ti surface. To test this possibility, we synthesized TBP–NHBP and developed a method for immobilizing oxSWNHs on the Ti surface using this peptide. We also tested the ability of DEX-loaded oxSWNHs immobilized on a Ti surface to release the drug in a sustained fashion and examined how this affects the activation of preosteoblasts grown on the functionalized Ti surface.

**Materials and methods**

**Adsorption of oxSWNHs onto Ti plates**

oxSWNHs were prepared by oxidation of as-grown SWNHs in dry air, with the temperature increased from room temperature to 550°C at a rate of 1°C/min.\textsuperscript{49} oxSWNHs were dispersed in dimethylformamide (DMF) at 10 mg/mL, and after sonication at 130 W (Bioruptor® UCD-200; Cosmo Bio Co. Ltd, Tokyo, Japan), 5 µL of the oxSWNH suspension was diluted by adding 175 µL of 10% DMF in water (purified with a Biocel A10 filter; Merck Millipore, Billerica, MA, USA), followed by brief sonication. A 20-µL volume of TBP–NHBP (RKLPDAGGDYFSSPYEQLF; 1 mM in DMF) was added to the solution, which was vortexed for 10 min at room temperature. TBP–NHBP comprised the peptide aptamers minTBP-1 (RKLPDPA) and NHBP-1 (DYFSSPYEQLF) connected via a GG linker. The custom peptide was constructed via solid-phase peptide synthesis by Anygen Co. Ltd (Gwangju, Korea). The final concentrations of oxSWNHs and TBP–NHBP in the mixture were 0.25 mg/mL and 100 µM, respectively. Ti plates (Japan Industrial Specification H4600; 99.9 mass % Ti, 10×10×1 mm; Shinkinzoku Co. Ltd., Osaka, Japan) were incubated for 1 h at room temperature in 500 µL of the oxSWNHs/TBP–NHBP mixture in 24-well culture plates (BD Biosciences, Sparks, MD, USA), washed three times with water filtered through a 0.22-µm membrane (Millipore), and then dried under vacuum for 8 h. The ratio of TBP–NHBP to oxSWNHs was evaluated in pilot experiments under various conditions. oxSWNHs adsorbed onto Ti plates were quantified using an electronic balance (UMX2; Mettler-Toledo, Billerica, MA, USA) and observed onto Ti plates were quantified using an electronic balance (UMX2; Mettler-Toledo, Billerica, MA, USA) and observed using a Biocell A10 filter; Merck Millipore, Billerica, MA, USA), followed by brief sonication. A 20-µL volume of TBP–NHBP (RKLPDAGGDYFSSPYEQLF; 1 mM in DMF) was added to the solution, which was vortexed for 10 min at room temperature. TBP–NHBP comprised the peptide aptamers minTBP-1 (RKLPDPA) and NHBP-1 (DYFSSPYEQLF) connected via a GG linker. The custom peptide was constructed via solid-phase peptide synthesis by Anygen Co. Ltd (Gwangju, Korea). The final concentrations of oxSWNHs and TBP–NHBP in the mixture were 0.25 mg/mL and 100 µM, respectively. Ti plates (Japan Industrial Specification H4600; 99.9 mass % Ti, 10×10×1 mm; Shinkinzoku Co. Ltd., Osaka, Japan) were incubated for 1 h at room temperature in 500 µL of the oxSWNHs/TBP–NHBP mixture in 24-well culture plates (BD Biosciences, Sparks, MD, USA), washed three times with water filtered through a 0.22-µm membrane (Millipore), and then dried under vacuum for 8 h. The ratio of TBP–NHBP to oxSWNHs was evaluated in pilot experiments under various conditions. oxSWNHs adsorbed onto Ti plates were quantified using an electronic balance (UMX2; Mettler-Toledo, Billerica, MA, USA) and observed using a Biocell A10 filter; Merck Millipore, Billerica, MA, USA).
Quartz crystal microbalance (QCM) measurements
The binding of oxSWNHs to the Ti surface was evaluated with a QCM (QCM-D300; Q-Sense AB, Göteborg, Sweden) and a Ti sputter-coated QCM sensor that was first cleaned for 10 min using an ultraviolet/ozone cleaner surface-treatment system (Pro Cleaner™, BioForce Nanosciences, Inc., Salt Lake City, UT, USA). Measurements were performed at 25°C, and data were collected at 14.8 MHz. The sensor was first equilibrated with the mixture containing oxSWNHs and TBP–NHBP1 peptide (100 µM) in 10% DMF, followed by washing with 0.5 mL of 10% DMF. The second and subsequent layers were formed in the same manner. Data were analyzed using Q-tools software (https://qtools.co).

Cell viability
Cell viability was qualitatively analyzed by fluorescence microscopy. MC3T3-E1 cells (ATCC) were seeded on Ti plates (10×10×1 mm) in 24-well plates (BD Biosciences) at a density of 1×10⁴ cells/well and cultured for 48 h. MC3T3E1 cells were maintained in α-minimal essential medium (α-MEM; Gibco®, Thermo Fisher Scientific, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (FBS; JRH Biosciences, Lenexa, KS, USA), 100 µg/mL penicillin (Banyu Pharmaceutical Co., Tokyo, Japan), and 100 U/mL streptomycin (Meiji Seika Kaisha, Tokyo, Japan) in continuous culture at 37°C under a humidified atmosphere with 5% CO₂ and 95% air. After three washes with phosphate-buffered saline (PBS; Gibco-BRL, Carlsbad, CA, USA), adherent cells were stained with 2 µM calcein-AM and 4 µM propidium iodide (Dojindo Laboratories, Kumamoto, Japan) in PBS and washed twice with PBS. The stained cells were photographed under a DMLP reflection microscope (Leica Microsystems, Bensheim, Germany) equipped with a DC480 or DFC350 FX digital camera.

Adsorption of DEX-oxSWNHs onto Ti plates
oxSWNHs (500 µg/mL) were dispersed by sonication in 10% DMF, and cell-culture-grade DEX (1,000 µg/mL; Sigma-Aldrich, St Louis, MO, USA) in 10% DMF was added at a volumetric ratio of 1:1. The mixture was vortexed for 10 min and incubated overnight at room temperature. DEX-oxSWNHs were mixed with TBP–NHBP and vortexed for 10 min. Ti plates (10×10×1 mm) in 24-well culture plates were incubated for 1 h at room temperature in 500 µL of the above mixture, washed three times with water filtered through a 0.22-µm membrane (Milllex-GS; Millipore), and dried under vacuum for 8 h. The amount of DEX was quantified with [³H]-DEX (1.48 TBq/mmol; Amersham Bioscience, Piscataway, NJ, USA) using an LS6500 scintillation counter (Beckman Coulter, Fullerton, CA, USA).

Analysis of in vitro release of DEX from DEX-oxSWNHs on a Ti surface
DEX-oxSWNHs were prepared as described using [³H]-DEX and incubated at 37°C in PBS, Roswell Park Memorial Institute (RPMI) 1640 medium (Gibco), or α-MEM supplemented with 5% FBS, 100 µg/mL penicillin, and 100 U/mL streptomycin. Samples were collected at predetermined time points, and the amount of [³H]-DEX released into the supernatant was quantified using a scintillation counter. For cumulative-release experiments, the supernatant was replaced with fresh medium at each time point.

DEX-responsive promoter assay
An expression vector containing the luciferase-reporter gene under the control of a DEX-responsive element (pBV2-MMTV-LUC) was gifted by Drs H Noguchi and I Abe of the University of Shizuoka (Shizuoka, Japan).39 ST2 cells (RIKEN BioResource Center) were seeded at a density of 2.5×10⁴ cells/well on Ti plates (10×10×1 mm) in 24-well plates. ST2 cells were maintained in RPMI 1640 supplemented with 10% FBS, 100 µg/mL penicillin, and 100 U/mL streptomycin in continuous culture at 37°C under a humidified atmosphere with 5% CO₂ and 95% air. After incubating for 1 day, cells were co-transfected with pBV2-MMTV-LUC and thymidine kinase promoter-driven Renilla luciferase DNA (Promega Corporation, Fitchburg, WI, USA) using FuGENE6 (Promega). After an additional 24 h, luciferase activity was measured using the dual-luciferase reporter assay system (Promega) and an AutoLumat LB953 lumimeter (Berthold Technologies, Oak Ridge, TN, USA) according to the manufacturer’s instructions.

Alkaline phosphatase (ALP) assay
MC3T3-E1 cells were seeded at a density of 1.0×10⁴ cells/well on Ti plates (10×10×1 mm) in 24-well plates and grown until they reached 70%–80% confluence. The culture medium was changed to a differentiation-inducing medium consisting of α-MEM supplemented with 5% FBS, 100 U/mL penicillin, 100 µg/mL streptomycin, 50 µM ascorbic acid, 10 mM β-glycerophosphate, and 20 ng/mL recombinant human bone morphogenetic protein-4 (Genzyme/TECHNE, Cambridge, MA, USA). The cells were incubated for 5, 7, 10, or 14 days, during which time the medium was refreshed every 3 days. Medium changes were performed carefully to ensure that oxSWNHs or DEX-oxSWNHs adhering to the Ti plates were not disturbed and removed along with the medium.
To evaluate ALP activity, the medium was removed, and cells were washed three times with Tris-buffered saline (TBS) comprising 20 mM Tris (pH 7.4) and 150 mM NaCl. The cells were harvested by scraping them into 200 μL TBS containing 1% Triton X-100 on ice and sonicated on ice for 5 min. The resulting cell lysates were centrifuged at 16,100 g for 20 min at 4°C, and the supernatant (50 μL) was mixed with p-nitrophenosphate. The amount of p-nitrophenol released after 45 min was measured by spectrophotometry (405 nm) using a Model 550 microplate reader (Bio-Rad Laboratories Inc., Hercules, CA, USA). The protein content of the supernatant was assayed by the Lowry method using a bicinchoninic acid protein assay kit (Pierce, Rockford, IL, USA). ALP activity was expressed as the amount of released p-nitrophenol normalized to total protein concentration.

Statistical analysis
Differences between groups were evaluated by one-way analysis of variance. Any P-values <0.05 and <0.01 were considered significant.

Results
Immobilization of oxSWNHs/TBP–NHBP on a Ti surface
We generated TBP–NHBP (RKLPDAGDYGFSYPYEQFL), a fusion of minTBP-1 and NHBP-1 connected via a GG linker, to mediate the binding of oxSWNHs to the Ti surface (Figure 1). TBP–NHBP was mixed with oxSWNHs in 10% DMF to obtain the dispersed oxSWNHs/TBP–NHBP complex, which was adsorbed onto the Ti surface by incubating Ti plates in dispersed solutions of the complex. Three samples [oxSWNHs without peptide (oxSWNHs), a mixture of minTBP-1 and oxSWNHs (oxSWNHs/TBP), and a mixture of NHBP and oxSWNHs (oxSWNHs/NHBP)] were adsorbed onto the Ti plates and served as the controls. Figure 2A shows the SEM image of the oxSWNHs/TBP-NHBP-adsorbed particles on the Ti surface. When oxSWNHs were adsorbed onto Ti plates with TBP–NHBP, the weight of the plate increased relative to the controls (Figure 2F). By contrast, the adsorption of oxSWNHs without TBP–NHBP onto the Ti surface was negligible (Figure 2B and F). When oxSWNHs alone were incubated with the Ti sensor (Figure 3D), these results demonstrated the necessity of pre-mixing TBP–NHBP and oxSWNHs for adsorption of oxSWNHs onto the Ti surface.
When TBP–NHBP alone was used, adsorption onto Ti plates was observed (Figure 3E); however, use of oxSWNHs/TBP–NHBP resulted in an increase in $D$ (Figure 3F). Even if cleaning was performed prior to each flow of oxSWNHs/TBP-NHBP, an increase in $D$ was observed, although when TBP-NHBP only was used, a minimal increase in $D$ was observed. It is possible that this resulted due to the unstable adsorption of some of the oxSWNHs to Ti in the chamber. Therefore, the minTBP-1 portion of TBP–NHBP also contributed to the immobilization of oxSWNHs on the Ti surface.

We then investigated whether oxSWNHs/TBP–NHBP on Ti was toxic to cells by culturing MC3T3-E1 cells on oxSWNHs/TBP–NHBP on the Ti plates. The live/dead assay using calcein/propidium-iodide staining revealed good cell viability after 2 days of culture (Figure 4A and B). These results demonstrated that uncoated Ti and oxSWNHs/TBP–NHBP on Ti exhibited comparable effects on cell viability. SWNHs do not contain contaminating metals, as they are prepared by laser ablation from pure graphite without a metallic catalyst, which reduces the risk of toxicity.

**In vitro release of DEX from DEX-oxSWNHs/TBP–NHBP on Ti**

Controlled release from a drug-carrier complex is an important property of drug-delivery systems. In vitro binding and release of the anti-inflammatory glucocorticoid DEX by oxSWNHs was previously reported. Here, we evaluated the in vitro drug-release potential of oxSWNHs on Ti according to the cumulative doses of [$^3$H]-DEX released from DEX-oxSWNHs on Ti plates in PBS, $\alpha$-MEM, and RPMI medium at 37°C (Figure 5). The initial release of DEX from Ti was slower in $\alpha$-MEM or RPMI medium than in PBS. Additionally, DEX from Ti in $\alpha$-MEM or RPMI medium was released in a sustained fashion for up to 2 weeks.

**Biological activity of DEX released from DEX-oxSWNHs/TBP–NHBP on Ti**

DEX is a synthetic agonist of glucocorticoid receptor (GR), which is a nuclear receptor that binds to glucocorticoid response elements (GREs) to activate target-gene transcription. We examined the responsiveness of GREs to DEX released from DEX-oxSWNHs/TBP–NHBP on Ti using the reporter plasmid pBV2-MMTV-LUC containing a luciferase gene under the control of GREs. To this end, we transfected ST2 cells with pBV2-MMTV-LUC, followed by incubation with DEX-oxSWNHs/TBP–NHBP, DEX, or oxSWNHs/TBP–NHBP on Ti. DEX and DEX-oxSWNHs/TBP–NHBP activated luciferase expression, with the latter having a greater effect, whereas the cells alone and oxSWNHs/TBP–NHBP had no effect (Figure 6). Additionally, DEX-oxSWNHs/TBP-NHBP showed an even greater activation than DEX alone on Ti.
DEX promotes the differentiation of preosteoblasts into mature osteoblasts, a process characterized by ALP expression. Additionally, preosteoblastic MC3T3-E1 cells treated with DEX show upregulated expression of ALP, as well as BMP-4, which is a potent inducer of osteoblast differentiation and bone formation. Here, we found that treatment of MC3T3-E1 cells with DEX-oxSWNHs/TBP–NHBP adsorbed on Ti in the presence of BMP-4 increased ALP expression from day 5 to day 14, whereas oxSWNHs/TBP–NHBP had no effect (Figure 7). Moreover, on day 14, the activity was higher for DEX-oxSWNHs/TBP–NHBP as compared with that observed in control samples ($P<0.05$). Our results indicated that DEX released from DEX-oxSWNHs on Ti plates was biologically active.

**Figure 3** Time-dependent changes in $f$ and $D$ determined using a QCM. (A–F) oxSWNHs/TBP–NHBP (A), oxSWNHs (B), TBP–NHBP (C), oxSWNHs/TBP–NHBP (D), TBP–NHBP (E), and TBP–NHBP and oxSWNHs/TBP–NHBP (F) deposited on a Ti-coated sensor. Values on the longitudinal axis represent negative $\Delta f$ and positive $\Delta D$ values. Changes associated with oxSWNHs/TBP–NHBP, oxSWNHs, and TBP–NHBP adsorption are indicated by green, gray, and purple lines, respectively. Green, gray, and purple arrowheads indicate the time points at which oxSWNHs/TBP–NHBP, oxSWNHs, and TBP–NHBP, respectively, were infused into the measurement chamber. Vs represent the time points at which the wash solution (10% DMF) was infused.

**Abbreviations:** oxSWNHs, oxidized single-walled carbon nanohorns; TBP, Ti-binding peptide; NhBP-1, SWNH-binding peptide; DMF, dimethylformamide.
Discussion

Ti and its alloys are frequently used for bone and dental implants, and their junction with adjacent bone tissue is paramount. Ti-based materials are also used as scaffolds for biomolecules that control cellular activities, such as adhesion, differentiation, and proliferation. Ti implants should ideally have osteoinduction capacity, representing the ability to stimulate the formation of new bone from progenitor cells. Various nanomaterials, including carbon nanotubes (CNTs), multi-walled CNTs, and SWNHs, promote bone formation by interacting with osteoblasts. In the present work, we combined the TBP–NHBP peptide with oxSWNHs to obtain dispersed oxSWNHs/TBP–NHBP complexes, which were adsorbed onto the Ti surface. Incorporation of the TBP–NHBP peptide conferred the binding characteristics of minTBP-1 to oxSWNHs and mediated their uniform bidimensional adsorption onto the Ti surface. NHBP-1 binds to oxSWNHs, but has low solubility, whereas fusion

Figure 4 Viability of MC3T3-E1 cells cultured for 2 days on oxSWNHs/TBP–NHBP on Ti. Viable cell counts were determined relative to cells grown on uncoated Ti (none). (A) Live (green, left panel) and dead (red, right panel) cells were stained with calcein-AM and propidium iodide, respectively. White scale: 100 µm. (B) Counts of live cells on uncoated Ti and oxSWNHs/TBP–NHBP-conjugated Ti. Error bars indicate standard deviation (n=4).

Abbreviations: oxSWNHs, oxidized single-walled carbon nanohorns; TBP, Ti-binding peptide; NHBP-1, SWNH-binding peptide.

Figure 5 In vitro release of DEX from DEX-oxSWNHs/TBP–NHBP on Ti. Time course of cumulative release of [3H]-DEX from DEX-oxSWNHs/TBP–NHBP on Ti plates in PBS (blue line), RPMI medium/5% FBS (green line), and α-MEM/5% FBS (red line) at 37°C over 14 days. The inset shows cumulative release of DEX from DEX-oxSWNHs/TBP–NHBP on Ti plates in PBS (blue line), RPMI medium/5% FBS (green line), and α-MEM/5% FBS (red line) at 37°C for 24 hours. Error bars indicate standard deviation (n=3).

Abbreviations: DEX, dexamethasone; PBS, phosphate-buffered saline; RPMI, Roswell Park Memorial Institute; α-MEM, α-minimal essential medium; oxSWNHs, oxidized single-walled carbon nanohorns; TBP, Ti-binding peptide; NHBP-1, SWNH-binding peptide.

Figure 6 Effects of DEX-oxSWNHs on GR transcriptional activity. ST2 cells were transfected with pBv2-MMTV-Luc and incubated with oxSWNHs/TBP–NHBP, DEX, or DEX-oxSWNHs/TBP–NHBP on Ti plates. Error bars indicate standard deviation (n=5). **P < 0.01.

Abbreviations: oxSWNHs, oxidized single-walled carbon nanohorns; TBP, Ti-binding peptide; NHBP-1, SWNH-binding peptide; DEX, dexamethasone.
with highly soluble minTBP-1 enhances the dispersibility of oxSWNHs. This is the likely reason for the uniform adsorption of oxSWNHs onto the Ti surface without aggregation. We also showed that oxSWNHs/TBP–NHBP were adsorbed onto the Ti surface by monitoring the changes in $f$ and $D$ in a Ti plate, observing that pre-mixing TBP–NHBP and oxSWNHs was important for oxSWNHs adsorption. The decrease in $f$ and increase in $D$ were not saturated within the observation range, suggesting that surface treatment can be readily controlled. Furthermore, oxSWNHs/TBP–NHBP immobilized on Ti were nontoxic to cultured cells, even after 2 days of culture and in contrast with CNTs. SWNHs contain no metal impurities, because SWNHs are synthesized through laser ablation or arc discharge using pure graphite rods without any metal catalysts. Therefore, SWNHs have negligible toxicity both in vitro and in vivo, which facilitates fabrication of the functionalized nanoplatforms for biomedical applications with low toxicity.

To assess the potential of oxSWNHs on Ti for drug delivery, we compared the release of the anti-inflammatory glucocorticoid DEX in PBS, α-MEM, and RPMI medium at 37°C. DEX was released from Ti at a slower rate in α-MEM and RPMI than in PBS; moreover, in cell-culture media, the release was sustained for up to 14 days. Variable solubility of DEX in PBS and cell-culture media could account for the differences in the release profiles. Additionally, GREs were activated by DEX and especially DEX-oxSWNHs/TBP–NHBP, with adsorption of the latter onto Ti increasing ALP activity in MC3T3-E1 cells relative to control samples on day 14. Therefore, DEX was not delivered in a single burst at the implantation site, but was steadily released for a sustained effect.

The results of the in vitro assays indicated that DEX released from DEX-oxSWNHs on Ti plates was biologically active. To date, there have been no studies on the effectiveness of DEX delivered by oxSWNHs/TBP–NHBP on Ti for inducing osteogenesis. The majority of studies on Ti-implant-surface treatments have focused on physicochemical surface modifications to improve osseointegration and interactions between surfaces and tissues. In the Ti-surface-modification method that has been reported, a special apparatus, such as a plasma spray apparatus, was required. However, the method described in this study constitutes a simple surface-reforming method that mixes oxSWNHs, TBP–NHBP, and a drug together, followed by incubation of Ti with the mixed solution, which enables it to transport the function of the drug. Because oxSWNHs are capable of transporting various drugs, we believe that it will be possible to perform surface modifications of Ti by enabling the drug to support oxSWNHs/TBP–NHBP according to the clinical purpose. Our findings have provided a basis for the development of new biomaterials for use in dental implants and as osteoblast scaffolds for bone regeneration.

**Conclusion**

We have successfully developed a solid-phase drug-delivery system where oxSWNHs were immobilized on a base material using TBP–NHBP, a bispecific material-binding peptide combining functions of minTBP-1 and NHBP-1. Through the use of TBP–NHBP, oxSWNHs, which otherwise could not be attached to Ti, were successfully immobilized on the Ti surface. SEM was used to confirm the presence of oxSWNHs adsorbed onto the Ti surface, and a QCM was used to evaluate the binding process during oxSWNH adsorption. Moreover, we were able to impart the function of sustained drug release to a substrate by using oxSWNHs. The oxSWNHs/TBP–NHBP-immobilized Ti surface was nontoxic to cells and could be used to deliver DEX, an anti-inflammatory agent, over a sustained period. The released DEX was confirmed as biologically active, suggesting the potential use of oxSWNHs/TBP–NHBP as a drug-delivery system to stimulate osteogenesis and bone regeneration. Therefore, our findings suggested that surface modification is a useful approach for designing medical biomaterials with desirable properties and possible practical applications.
Acknowledgments

We thank Dr T Imanura for kindly providing the MC3T3-E1 cells and ST2 cells, Dr K-I Sano for valuable discussion, and Ms T Minamisawa for technical assistance.

This work was supported by a JSPS KAKENHI (grant no 21791959, 16K11524).

Disclosure

The authors report no conflicts of interest in this work.

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