Title:
Amine modification of calcium phosphates by low-pressure CH₄/N₂/He plasma for bone regeneration

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Abstract

Calcium phosphates are promising materials for artificial bone but lack of satisfied osteogenic ability on their surfaces. In the present study, we applied a low-pressure plasma technology to chemically (amine) modify the surface of calcium phosphates (hydroxyapatite or beta-tricalcium phosphate) using a CH₄/N₂/He plasm gas mixture to improve their osteogenic ability. The CH₄/N₂/He plasma treatment produced a thin, stable amine-rich carbon polymer on the surface of the calcium phosphates, and enhanced hydrophilicity, deep infiltration of cells into porous calcium phosphates, cell adhesion and osteogenic differentiation on the surface of calcium phosphates. In a rat calvarial defect model, the CH₄/N₂/He plasma treatment afforded calcium phosphates a significant higher bone regeneration capacity. Together, these results suggest that surface modification of calcium phosphates with CH₄/N₂/He plasma might improve osteogenic ability of calcium phosphates in vitro and in vivo.
Introduction

Autogenous iliac bone grafting remains the gold standard for repairing large bone defects caused by trauma or tumors and spinal fusion surgeries. However, the amount of autograft that can be harvested is limited and the harvesting procedure can cause donor site morbidity 1. To overcome these limitations, the use of artificial bones, in combination with autograft is prevalent. However, the widespread use of artificial bone is hampered by its lack of satisfied osteogenic ability, despite their superiority in terms of bone conduction and availability 2.

Calcium phosphates are representative materials for artificial bone. To enhance the osteogenic ability of calcium phosphates, surface modification mediated by plasma technology has gained considerable attention. Functional groups created by plasma polymerization (i.e., polymer formation via plasma discharges) can provide selected surface properties such as hydrophilicity/hydrophobicity, cytocompatibility, and bacterial resistance to meet different clinical needs 3. We previously reported that plasma treatment of hydroxyapatite (HA, a typical calcium phosphate used for artificial bone) with O₂/He gas could improve surface hydrophilicity and promote the osteogenic differentiation of rat bone marrow stromal cells (BMSCs). However, the O₂/He plasma-treated HA exerted only minimal effects in vivo, possibly owing to the instability of the generated hydroxyl (–OH) groups 4-5. In the present study, we focused on the addition of amino groups or amines (−NHₓ with x = 0–2) to the surface of HA and beta-tricalcium phosphate (β-TCP, another typical calcium phosphate used for artificial bone), which has been suggested to promote cell attachment 6-7. Using a mixture gas of CH₄/N₂/He as the precursor, we
successfully generated a thin, stable amine-rich carbon polymer on and inside the surfaces of the HA and β-TCP through a low-pressure plasma treatment, and enhanced cell infiltration, migration, cell adhesion, osteogenic differentiation in vitro, and bone regeneration with a rat calvarial defect model *in vivo*.

**Materials and methods**

**Materials and reagents**

Both dense disks and porous disks of HA \([\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2]\) and β-TCP \([\text{Ca}_3(\text{PO}_4)_2]\) were kindly provided by Coors Tek KK (Tokyo, Japan). Porous disks of both HA and β-TCP had well-organized interconnected structure \(^8\), and contained interconnected pores (porosity; 72–78%, average pore diameter; 150 μm, average diameter of interconnected passages; 40 μm).

The following culture media were used for *in vitro* experiments: (1) growth medium (GM) comprising α Eagle’s minimal essential medium (α-MEM, Gibco, ThermoFisher Scientific, Waltham, MA, USA) supplemented with 10% fetal bovine serum (Sigma-Aldrich, St. Louis, MO, USA) and 1% antibiotic-antimycotic solution (Sigma-Aldrich); (2) osteogenic differentiation medium (ODM) consisting of GM supplemented with 50 μg/ml L-ascorbic acid 2-phosphate (Sigma-Aldrich), 10 mM β-glycerol phosphate (Merck KGaA, Frankfurt, Germany), and 10 nM dexamethasone (Sigma-Aldrich).

Cell Counting Kit-8 (CCK-8, Dojindo Molecular Technologies, Kumamoto, Japan) was used for the cell proliferation assay. LabAssay ALP (FUJIFILM Wako Pure Chemical Corporation) was used for the evaluation of alkaline phosphatase (ALP) activity and BCIP/NBT Color Development Substrate (Promega Corp.,
Madison, WI, USA) was used for ALP staining. M-PER and Pierce Rapid Gold BCA Protein Assay Kits (Thermo Fisher Scientific) were used for total protein extraction and quantification, respectively. K-CX AT solution (Falma Co., Tokyo, Japan) was used for decalcification of in vivo specimens.

Plasma polymerization of HA and β-TCP disks

Plasma polymerization using a bipolar pulsed-plasma deposition system (also known as an inverter plasma system) was as shown in Fig. 1A. The details of the system are described elsewhere. Briefly, HA and β-TCP disks were placed on the bottom metal (molybdenum) electrode, 190 mm in diameter, connected to the bipolar high-voltage power supply. The applied bipolar pulse voltages were 1.3 and -1.1 kV (peak-to-peak voltage of 2.4 kV) and the pulse duration was 1 μs for each positive or negative pulse. The power, repetition frequency, and duty cycle were 15 W, 5 kHz, and 1%, respectively, such that the time elapse between a positive pulse and the subsequent negative pulse was 100 μs. The upper metal (aluminum) electrode, 80 mm in diameter, was grounded and the distance between the two electrodes was 38 mm. For plasma polymer deposition, the discharge was generated in a CH₄/N₂/He gas mixture with flow rates of 10, 20, and 10 sccm, respectively, at 70 Pa.

Dense disks were treated on single side for 30 min, which were used for in vitro experiments; porous disks of HA was treated on single side for 30 min, which were used for cross section analysis; porous disks of β-TCP were treated on both sides for 60 min (30 min for each side), which were used for in vivo experiments.
Plasma polymer film characterization

X-ray photoelectron spectroscopy (XPS) was carried out to determine the chemical compositions of the surfaces of dense disks using the Shimadzu ESCA-850 with a non-monochromatized Mg-Kα (1253.6 eV) X-ray source at Osaka University, Japan. The pass energy was 75 eV and the spot diameter was 8 mm (90% uniformity).

A high-resolution XPS Axis Supra (Kratos Analytical, Manchester, UK) spectrometer at the Central European Institute of Technology (CEITEC), Brno, Czech Republic was used to analyze chemical compositions of plasma polymers, especially those on the inner surfaces of interconnected pores of plasma-treated porous disks. This high-resolution XPS uses an X-ray source of monochromatic Al-Kα 1486.6 eV with an emission current of 15 mA. The spectra were acquired with a pass energy of 20 eV. The analysis area provided by the larger slot aperture was approximately 700 × 300 μm².

For cross section analysis, the single side treated porous disk were cut in half at the center and the cross sections were scanned using high-resolution XPS. The measurements were performed on 21 spots with a diameter of 110 μm that were separated by an equal distance of 100 μm along the center axis of the cross-section (Fig. 1B). During the analysis, the pressure of the XPS chamber was maintained at < 10⁻⁷ Pa. The peaks of XPS signals used for the analyses in this study were C 1s, Ca 2p, N 1s, O 1s, and P 2p.

Detection of primary amines (–NH₂) on CH₄/N₂/He plasma-treated disk surfaces was performed using the standard derivatization with 4-trifluoromethyl-
benzaldehyde (TFBA) \(^{11}\). The derivatization reactions of TFBA vapors with the CH\(_4\)/N\(_2\)/He plasma-treated disk surfaces were allowed to occur in an Ar atmosphere (Ar flow rate of 60 sccm at atmospheric pressure) at room temperature (approximately 25 °C) for 4 h inside the glove box. Relative concentrations of primary amines on the surfaces of the HA and β-TCP were analyzed with ESCA-850 XPS following the derivatization reactions.

To examine the stability of the deposited plasma polymers, we stored CH\(_4\)/N\(_2\)/He plasma-treated disks in a sterilized dish at room temperature (approximately 24 °C), and then examined relative concentrations of nitrogen and primary amines on the disk surfaces were using ESCA-850 XPS at several time points over 65 days.

**Scanning electron microscopy (SEM)**

The surfaces of untreated and plasma treated porous HA disks (φ 5 mm × h 2 mm) were observed with a scanning electron microscope (S-4800, Hitachi, Ltd., Tokyo, Japan).

**Ethic declarations**

All animal work was approved by the Animal Experimentation Committee of our institution (01-070-000) and restrictedly followed ARRIVE guidelines and the National Institutes of Health Guide for the Care and Use of Laboratory Animals \(^{12}\).
Rat BMSCs were obtained from the bone shafts of the femora of four 3-week-old green fluorescent protein (GFP)-transgenic male Sprague-Dawley rats (SD-Tg (CAG-EGFP) rat, Japan SLC, Hamamatsu, Japan). Following sacrifice using CO₂ inhalation, both ends of the femur were removed from the epiphysis; the marrow was flushed out using 10 ml of GM expelled from a syringe through a 21-gauge needle according to the previously described method 13. The released cells were collected in two 100 mm culture dishes containing 15 ml of GM. The medium was changed after 24 h to remove hematopoietic cells and renewed twice weekly. Cultures were maintained in a humidified atmosphere of 95% air with 5% CO₂ at 37 °C. When the cells reached 80–90% confluency they were washed with phosphate-buffered saline (PBS) and trypsinized with 1% trypsin-ethylenediaminetetraacetic acid (EDTA). Following centrifugation for 5 min at 400 g, the cells were resuspended and plated at a density of 3.6 × 10⁴/cm². After again reaching confluency, cells were collected and stored at -80 °C (Passage 1). Prior to in vitro experiments, stocked cells were thawed and resuspended in 15 ml of GM, then plated in a 100 mm dish and cultured for three days to reach 80-90% confluency (Passage 2).

Morphology analysis of adhered cells

Cell suspension (5 × 10³ cells/35 μl GM) was gently dropped on dense β-TCP disks (φ 5 mm × h 2 mm) and incubated for 3 h in a 24-well culture plate to initiate adhesion. Then, wells were slowly filled with 1 ml of GM and incubation was continued for another 24 h. Macro fluorescence photos of random areas were obtained. The open source software CellProfiler (www.cellprofiler.org) 14 was
used for sorting cells and cell morphology analysis. In particular, the cell area was measured to quantify the spreading of attached BMSCs. Two cell-shape descriptors (circularity and solidity) were investigated; circularity indicates the closeness of the cell shape to a perfect circle, and solidity is an index to quantify the amount and size of concavities of the cell:

\[ Circularity = \frac{4\pi A}{P^2} \]

where \( A \) is the cell area and \( P \) is the perimeter.

\[ Solidity = \frac{A}{\text{Convex}A} \]

where Convex\( A \) is the area of the smallest convex hull that contains the cell.

**Cell adhesion assay**

Cell suspensions (5 x 10^3 cells/35 μl GM) were gently dropped on dense β-TCP disks (φ 5 mm x h 2 mm) to form centroclinal water drops on the disks. After incubating for 30 min, a centrifugation cell adhesion assay was performed. Briefly, 1 ml of PBS was gently added into the wells and fluorescent macro-photos were obtained to quantify the initial adherent cells. β-TCP disks on which cells were attached were then embedded into a 48-well culture plate containing 100 μl of Vaselin. After filling each well with PBS, the plate was sealed with sealing tape. Then, the culture plate was set upside-down on a centrifuge and centrifuged at 10 g for 5 min to detach weakly adherent cells. PBS and detached cells were slowly aspirated, then the wells were carefully filled with 200 μl PBS. Macro fluorescence photos were obtained using the same conditions as prior to the centrifugation. Automatic cell counting was performed using ImageJ software
The adhesion rate was calculated as follows:

\[
\text{Adhesion Rate} = \frac{\text{Cell Number}_{\text{before centrifugation}}}{\text{Cell Number}_{\text{after centrifugation}}} \times 100\%
\]

**Cell proliferation assay**

To balance the initial cell count of adherent cells on plasma-treated and untreated $\beta$-TCP disks, two different concentrations ($5 \times 10^3$ cells/$35 \mu l$ GM for the untreated group and $3 \times 10^3$ cells/$35 \mu l$ GM for the plasma-treated group) of cell suspension were dropped on dense $\beta$-TCP disks ($\varphi 5 \text{ mm} \times h 2 \text{ mm}$) in a 48-well plate. After incubating for 30 min, 500 $\mu l$ of GM was slowly added into each well. At day 1, 3, 5, 7, 11, and 14, 50 $\mu l$ of CCK-8 solution was added to each well and the plate was incubated for 2 h, then 100 $\mu l$ of GM from each well was transferred into a 96-well plate. Optical density at 450 nm was measured using a spectrophotometer (Multiskan GO, Thermo Fisher Scientific).

**Osteogenic differentiation assay**

A total of $2 \times 10^4$ cells/$35 \mu l$ GM were dropped on dense $\beta$-TCP disks ($\varphi 5 \text{ mm} \times h 2 \text{ mm}$) for 30 min in a 48-well plate to initiate adhesion. Then, 500 $\mu l$ of GM was slowly added into the wells, incubated for 24 h, and the culture medium substituted with ODM for osteogenic differentiation. The subculture was maintained for another four days. After washing the disks twice with PBS, the attached cells were fixed using 500 $\mu l$ of 4% PFA for ALP staining according to the manufacturer's instructions. For ALP activity, 60 $\mu l$ of M-PER was added to each well and the cells were detached from the disks using a mini scraper. After lysing the cells for 5 min, supernatant was collected for ALP and total protein
assays. An ALP activity unit was defined as the release of 1 nmol p-nitrophenol per min of incubation at 37 °C for one β-TCP disk. The total protein content of the samples was measured to standardize the ALP activity values.

Rat calvarial defect model

A total of 20, 8-week-old male Sprague-Dawley rats (Charles River Laboratories Japan, Yokohama, Japan) were used to generate the calvarial defect model. Anesthesia was maintained by intraperitoneal injection of a mixture of 0.15 mg/kg medetomidine, 2.0 mg/kg midazolam, and 2.5 mg/kg butorphanol after introducing anesthesia by inhalation of 5% isoflurane. A 1.5 cm longitudinal incision was made at the center of the vertex and two full-thickness bone defects with a diameter of 5 mm were then carefully created using a high-speed trephine burr under constant irrigation with saline to avoid heat injury of the surrounding tissue. The CH₄/N₂/He plasma-treated and untreated β-TCP disks were implanted in the right and left defects, respectively. The rats were given free access to water and food. Rats were sacrificed at 3 (n = 5) and 6 (n = 15) weeks by CO₂ inhalation. Microfocus computed tomography (micro-CT) and histological analyses were performed to evaluate new bone formation in the inner pores of β-TCP disks at 3 and 6 weeks.

Microfocus computed tomography (micro-CT)

Harvested specimens were fixed in 10% buffered formalin, dehydrated and degreased using a graded ethanol series, and stored in 70% ethanol at 4 °C for micro-CT scanning (Skyscan 1272 micro-CT, Bruker, Kontich, Belgium) with the
following parameters: Camera binning = 2x2, Source voltage (kV) = 80, Source current (μA) = 125, Image pixel size (μm) = 4, Rotation step (degree) = 0.6, Filter = Al 1 mm. Image analysis was performed using CTAN software (Version 1.18.8.0+, Brucker). The micro-CT images corresponding to the histological sections were compared and the intensities of the newly formed bone and the residual β-TCP were determined and the respective areas extracted (Fig. 1C).

Figure 1. Plasma generation system, methodology of cross-section XPS analysis and micro CT evaluation. (A) Schematic diagram of the pulsed-plasma deposition system used in this study. Samples were placed on the bottom electrode. The discharge chamber was filled with a gas mixture of CH₄, N₂, and He at 70 Pa and bipolar high-
voltage short pulses were applied to the bottom electrode periodically to generate the plasma. (B) Schematic diagram of cross-section XPS analysis. A porous disk was treated with the plasma on a single side and cut in half to expose the cross section. Then, the chemical compositions along the center axis were measured. (C) H&E-stained histological sections and corresponding micro-CT slices were compared to identify $\beta$-TCP (blue area) and newly formed bone (yellow area). Unstained areas in histological sections constitute the areas containing remaining $\beta$-TCP, which were decalcified. Pink areas stained by eosin reflect newly formed bone inside the $\beta$-TCP, which correspond to the gray areas in micro-CT slices. An original algorithm was designed to extract and calculate the volume of $\beta$-TCP and newly formed bone separately.

Histological evaluation

After micro-CT scanning, specimens were demineralized using K-CX AT solution at 4 °C, cleared in xylene, and embedded in paraffin. Then, sections of 4 μm thickness were created from the center of specimens and stained with hematoxylin and eosin (H&E).

Statistical analysis

Statistical analysis was performed using GraphPad Prism version 7.04 for Windows (GraphPad Software, San Diego, CA, USA) applying the Mann–Whitney U test for non-parametric data, student’s t-test for parametric data, and Wilcoxon matched-pairs signed rank test for in vivo results. The values are presented as mean ± standard deviation (SD). The differences were considered
statistically significant for p value < 0.05.

Results

Plasma polymerization on calcium phosphate porous disks

The atomic percentages of calcium (Ca), phosphorus (P), oxygen (O), and carbon (C) was 27.5 ± 0.8, 13.9 ± 0.5, 46.4 ± 0.4, and 12.2 ± 1.2 at.% on the surface of the untreated porous HA, and 28.5 ± 0.2, 14.0 ± 1.5, 44.0 ± 0.6, and 13.4 ± 2.2 at.% on the surface of the untreated dense β-TCP, respectively. After excluding C, the atomic percentages were Ca:P:O = 31.3:15.8:52.8 for the porous HA surface and 32.9:16.2:50.9 for the dense β-TCP surface, which exhibited a Ca-rich stoichiometry compared with the theoretically ideal HA (Ca:P:O = 25.0:10.0:65.0), and β-TCP (Ca:P:O = 23.1:15.4:61.5).

The atomic percentages of Ca, P, O, C, and N, on the surfaces of CH4/N2/He plasma-treated porous HA disks was 0.0 ± 0.0, 0.0 ± 0.0, 8.2 ± 3.6, 78.1 ± 3.5, and 13.6 ± 1.9 at.%. A separate measurement performed using a Si substrate exposed to the same CH4/N2/He plasma revealed that the deposited plasma polymer film thickness was approximately 100 nm, which is consistent with our XPS measurements.

Fig. 2A shows the change in atomic percentage of N atoms in the polymer film on porous HA disks from the day of polymerization (day 0) to day 65. It was observed that percentage of N atoms in the deposited polymer film remained in the range of 12–15 at.%.

Fig. 2B shows the stability of primary amines in the polymer film on porous HA disks in ambient air up to 65 days from the day of plasma polymerization. The
The chemical compositions at the 21 spots along the center axis on the cross section of porous HA disk, as shown in Fig. 1B, are presented in Fig. 3C. It was observed that, in the region very close to the top surface, the atomic percentage closely resembled that of the top surface and the C and N atomic percentages decreased along with the depth. The observed atomic percentages of N and O of 3-5 and 20-35 at.% indicated that plasma polymerization occurred even on the surfaces of the inner pores with a penetration depth of at least 2 mm.

SEM photos of outer surfaces of untreated and plasma treated porous HA are shown in Fig. 2D. Plasma polymerization did not cause structural changes of the porous disks. At x1000 magnification, interconnecting channels are clearly visible in both groups. The deposited polymer was not visible with SEM since its thickness was about 100 nm.
Effect of plasma treatment on cell behaviors

β-TCP are originally hydrophobic. CH₄/N₂/He plasma treatment in this study enhanced the hydrophilicity of the β-TCP (Supplementary Fig. S1A, B, Supplementary Movie S2) and cell adhesion to it.

Both cell counts and adhesion rate were all significantly higher on CH₄/N₂/He plasma-treated than the untreated β-TCP disks (Fig. 3A, 3B). The morphology of attached cells in the CH₄/N₂/He plasma-treated group following 24h incubation, presented a significant wider spreading cytoplasm (larger cell area) and showed a decreased circularity and solidity, which emphasized the presence of membrane protrusions (Fig. 3C, 3D)
Figure 3. Cell adhesion and morphology. (A) Fluorescence photos of BMSCs from GFP rats attached onto dense β-TCP disks before and after centrifugation. (B) Quantitative analysis of attached cells before and after centrifugation, and adhesion rate. Data are expressed as mean ± SD (n = 5). *p < 0.05, **p < 0.01. Mann-Whitney U test. (C) Fluorescence photos of attached cells on untreated or plasma-treated dense β-TCP disks. Scale bars = 500 μm. (D) Quantitative analysis of area, which represents the size, and solidity which represents the shape of attached cells. Data are expressed as mean ± SD with n = 1860 for “untreated” and n = 1440 for “plasma”. ****p < 0.0001. T-test.

CH₄/N₂/He plasma treatment did not significantly influence cell proliferation (Fig. 4A), but also significantly improved osteogenic differentiation as demonstrated by ALP staining (Fig. 4B, 4C) and ALP activity assay (Fig. 4D).
Figure 4. *In vitro* cell proliferation and osteogenic differentiation of GFP rat BMSCs.

Cell proliferation curve (A). Macro photos of ALP-stained BMSCs from GFP rats subcultured on untreated (B) and CH₄/N₂/He plasma-treated (C) dense β-TCP disks. Scale bars = 1 mm. ALP activity assay (D). ALP activity was normalized by total protein content. Data are expressed as mean ± SD (n = 3). *p < 0.05. T-test.
CH$_4$/N$_2$/He plasma treatment enhances new bone formation in vivo

Figure 5 shows micro-CT slices and corresponding histological sections (Fig. 5A, 5B: “Untreated”; 5E, 5F: “Plasma-treated”). Magnified histological images of the center and interface regions demonstrated more abundant bone formation and solid interface union in the CH$_4$/N$_2$/He plasma-treated (Fig. 5C, 5D) than in the untreated group (Fig. 5G, 5H) at postoperative 6 weeks.

At 3 weeks, volumes of new bones were observed inside the CH$_4$/N$_2$/He plasma-treated porous β-TCP disks and did not differ from the untreated porous β-TCP disks. At 6 weeks, the new bone inside the CH$_4$/N$_2$/He plasma-treated porous β-TCP disks was significantly higher than the untreated β-TCP disks (Fig. 5I). The 3D-reconstructed image of extracted new bone along with the host cranial bone revealed the abundant new bone formation in the CH$_4$/N$_2$/He plasma-treated β-TCP disk (Fig. 5J). In both groups, nearly one-third of the β-TCP was resorbed during 3 to 6 weeks (Fig. 5K) in similar resorption speed.
Figure 5. CH₄/N₂/He plasma treatment promotes bone formation inside β-TCP. Micro-CT slices and the corresponding histological sections stained with H&E of the untreated (A, B) and the plasma-treated group (E, F. Scale bars for D–G = 1 mm) at postoperative 6 weeks. Magnified images of the center and interface regions (C, D: “Untreated” group; G, H: “Plasma-treated” group. Scale bars = 100 μm). (I) Micro-CT
analysis of newly formed bone volumes between the plasma-treated and untreated group at 6 weeks. (J) 3D reconstruction of calvarial bone implanted with treated and untreated β-TCP (6 weeks, left: “Untreated”, right: “Plasma-treated”. Scale bar = 1 mm). Resorption rates of β-TCP between plasma-treated and untreated groups (K). Data are expressed as mean ± SD (n = 5 for 3 weeks, n = 15 for 6 weeks). **p < 0.01, ****p < 0.0001. Wilcoxon matched-pairs signed rank test.

**Discussion**

In this study, we generated a thin, stable amine-rich carbon polymer on the surface of calcium phosphates (HA and β-TCP) by low-pressure plasma technology using a mixed gas containing CH₄/N₂/He as precursors.

Low-pressure CH₄/N₂/He plasma allows chemically reactive gaseous species to penetrate into small pores of the porous HA and β-TCP disks during the plasma treatment and induces plasma polymerization on the inner surfaces of interconnected pores. In the present study, the top surface was completely covered with the amine-rich carbon polymers, and about 72–78% of the cross-section area distant from the top surface was covered with the amine-rich carbon polymers.

Through XPS measurements, we confirmed that under the same plasma conditions, plasma polymerization on the outer surface of both HA and β-TCP disks occurs in a similar manner regardless of disk type or composition. Therefore, extensive and systematic surface analyses of plasma-treated artificial bone were performed only with porous HA disks, with the expectation that the obtained results should also apply to plasma polymerization on β-TCP disks. Considering
the increasing use of β-TCP as the clinical product of bone substitute, cell and animal studies were performed with the CH₄/N₂/He plasma treated β-TCP disks. The results showed that CH₄/N₂/He plasma treatment had enhanced the hydrophilicity, *in vitro* cell migration, adhesion, osteogenic differentiation and *in vivo* bone regeneration of the β-TCP.

Stability of the amine-rich carbon polymer and its enhancement of osteogenic differentiation are considered to be the main mechanisms underlying this enhanced bone regeneration capacity.

As the surface N concentration did not substantially change over the first two months, the primary amines were considered to have been converted to secondary or ternary amines, mostly likely by oxidation. Notably, the reduction of primary amines in plasma polymers does not necessarily indicate the reduction of their biological effects as secondary or even ternary amines embedded in plasma polymers may have similar biological effects. However, determining how different amino groups interact within biological systems is beyond the scope of the current study.

Amine constitutes a hydrophilic functional group. The present study demonstrated that amine modification of calcium phosphates could enhance its hydrophilicity, and fasten the infiltration of cell suspension dropped on the calcium phosphate surface. Notably, the CH₄/N₂/He plasma-treated β-TCP resorbed tissue fluid and rapidly became wet, in contrast to non-treated β-TCP, which remained almost completely dry during the implantation (Supplementary Fig. S3). Moreover, histological sections at early phase (postoperative 3 weeks) revealed denser tissue formation in the CH₄/N₂/He plasma-treated disks.
these findings suggest that the amine-rich carbon polymers can efficiently promote early tissue integration into the deep center of the disks, which makes preliminary preparations for later bone regeneration.

β-TCP coated with amine-rich carbon polymers enhanced in vitro osteoblastic differentiation and in vivo new bone formation. Several effects provided by amine modification must likely contributed to this enhanced bone formation.

Firstly, amine modification strengthens cell adhesion by enhancing integrin binding, which has been shown to be required for osteoblastic differentiation\(^{18-20}\). In detail, cell adhesion is mainly mediated by the binding of cellular integrins and adhesive proteins such as fibronectin in the extra-cellular matrix. Hydrophilic and positively charged amines can increase the density of fibronectin and change its conformation\(^{21}\). These changes in fibronectin strengthen the cell adhesion via integrin\(^{22,23}\) and trigger rapid phosphorylation of focal adhesion-associated tyrosine kinase (FAK)\(^{24}\), subsequently triggering ERK/MAPK signaling to upregulate Runt-related transcription factor 2 (Runx2), which is a master regulator of osteoblastic differentiation\(^{25-31}\).

In addition, amine modification was found to modulate the cell morphology (e.g., larger cell area and decreased circularity and solidity), which has been shown to be relevant to the osteogenic differentiation of rat mesenchymal stem cells\(^ {32}\). Moreover, human mesenchymal stem cells exhibiting a spreading rather than a round shape appear inclined toward an osteogenic lineage as well\(^ {33}\). The underlying mechanism is considered to involve upregulation of Ras homolog family member A (RHOA), a transcription factor that regulates the actin cytoskeleton, along with increased osteogenesis\(^ {32-34}\).
Lastly, positively charged amines can also improve osteogenesis by affecting pH \(^{35}\). In an aqueous environment, amine is protonated and becomes positively charged (–NH\(_3^+\)), which can increase the interfacial pH. A high pH environment around implant materials has been reported to enhance osteoblastic differentiation \(^{35-36}\).

In summary, our results support that amine modification of calcium phosphates with low-pressure CH\(_4\)/N\(_2\)/He plasma influences cell adhesion, cell spreading, and possibly the interfacial environment of calcium phosphates (here \(\beta\)-TCP). We consider that these factors in combination underlie the high ability of the modified \(\beta\)-TCP to improve bone regeneration.

There are limitations to this study. Firstly, the stability of the amine modification was observed for only two months, which should be elongated in future study for confirmation of longer off-the-shelf use. Secondly, we investigated the bone regeneration capacity only by the calvarial defect model. The calvaria has a rich blood supply and therefore is easy for bone regeneration. Further investigations of the bone regeneration effects in harsh environment such as malunions of fractures is needed. Thirdly, as the volume of newly formed bone inside the pores of the \(\beta\)-TCP disks was tiny, we could only evaluate the bone volume even with even a high-resolution micro-CT, but could not evaluate other bone parameters such as trabecular numbers and thicknesses. Finally, lacking biomechanical tests, we do not exactly know the real strength of the regenerated bone.

In treatment of large bone defects with artificial bone, the successful introduction of cells and blood vessels at a substantial distance from the host bone remains challenging \(^{37-38}\). To treat “critical size” bone defects, pre-loading of
bone or vascular-forming cells or vascular transplantation inside artificial bones has been attempted \(^{39-42}\), which usually require long and costly pre-treatments. In the present study, we have reported a “chemical modification” approach as a next-generation surface treatment and processing strategy to form a stable amine-rich carbon polymer on the surfaces of calcium phosphates (representative material of artificial bone) to improve their bone regeneration ability in vitro and in vivo.

**Author contributions**

J. Kodama, A.H.H., S.H., and T. Kaito designed the study; A.H.H., T.I. S.S., and S.H. performed the material development and plasma polymerization; A.H.H., M.M., T.I., D.N., L.Z., and S.H. performed the surface analyses of materials. J. Kodama and T. Kaito performed the *in vitro* and *in vivo* experiments; Y.U., J. Kushioka, R.O., K.H., T. Kamatani, D.T., H.T., S.N., S.T., T.M., and Y.S. contributed to the data analysis and interpretation; J. Kodama, A.A.H., S.H., and T. Kaito wrote the manuscript.

**Data availability**

The datasets generated and/or analysed during the current study are available from the corresponding authors on reasonable request.

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relevant to this article.

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