Efficient induction of antimicrobial activity with vancomycin nanoparticle-loaded poly(trimethylene carbonate) localized drug delivery system

Abstract: Surgery and the local placement of an antibiotic are the predominant therapies to treat chronic osteomyelitis. Vancomycin-loaded N-trimethyl chitosan nanoparticles (VCM/TMC NPs) as a potential drug delivery system have high intracellular penetration and effective intracellular antibacterial activity. This study investigated the effects of a biocompatible material, poly(trimethylene carbonate) (PTMC), to increase the sustained effectiveness of an intracellular antibiotic and its potential application in antibiotic delivery. VCM/TMC NP-PTMC was characterized using scanning electron microscopy and Fourier transform infrared spectroscopy to determine the morphology, stability and chemical interaction of the drug with the polymer. Further, the biodegradation, antibacterial activity, protein adsorption, cell proliferation and drug release characteristics were evaluated. In addition, a Staphylococcus aureus-induced osteomyelitis rabbit model was used to investigate the antibiotic activity and bone repair capability of VCM/TMC NP-PTMC. The results showed that the composite beads of VCM/TMC NPs followed a sustained and slow release pattern and had excellent antibacterial activity and a higher protein adsorption and cell proliferation rate than the VCM-PTMC in vitro. Furthermore, VCM/TMC NP-PTMC inhibits bacteria and promotes bone repair in vivo. Thus, VCM/TMC NP-PTMC might be beneficial in periodontal management to reduce the bacterial load at the infection site and promote bone repair.

Keywords: vancomycin-loaded N-trimethyl chitosan nanoparticles, poly(trimethylene carbonate), sustained release, intracellular antibacterial effect, bone infection

Introduction
Chronic osteomyelitis is caused by pathogenic microorganisms, which can adhere to – and even invade – mammalian cells. The most common causative organism in osteomyelitis is Staphylococcus aureus. Drug-resistant pathogens such as methicillin-resistant S. aureus have developed the ability to persist in intracellular locations following drug treatment. Vancomycin (VCM) is a glycopeptide antibiotic primarily active as an inhibitor of cell wall synthesis in susceptible organisms.

At present, the predominant therapies for chronic osteomyelitis are surgery and the local placement of VCM-loaded calcium sulfate beads. Nevertheless, our study on patient samples showed that the local VCM concentration reached a peak at the fourth day postimplantation and then decreased rapidly, indicating unsatisfactory control over the sustained release of drug. Therefore, a new delivery system with controlled and sustained release is crucial. Nanocarriers have attracted prominent interest in the past few decades. Chitosan (CS), a natural and plentiful biopolymer, possesses various biological characteristics; however, its application has been hampered...
largely by its solubility only in acidic environments.\textsuperscript{13} As a result, CS derivatives were synthesized to overcome this defect. Carboxymethyl chitosan (CMC) is a biocompatible material and it is soluble in water without acid.\textsuperscript{14} There were negative charges on the surface of antibiotic-loaded CMC NPs, and burst release of the drug in the initial stage was observed.\textsuperscript{15}

In our study, CS was derivatized by N-methylation to N,N,N-trimethyl chitosan (TMC), a water-soluble compound, to form nanocomplexes with anionic compounds, such as drugs, proteins and DNA, through ionotropic gelation as opposed to a pH-dependent charge.\textsuperscript{16} VCM-loaded TMC NPs (VCM/TMC NPs), prepared in our previous study,\textsuperscript{17} proved to be an effective intracellular mode of drug administration. Additionally, the VCM/TMC NPs had an excellent probability of VCM release in the chronic osteomyelitis rabbit model. However, a burst release phenomenon of VCM was observed. Poly(trimethylene carbonate) (PTMC),\textsuperscript{18,19} a surface-eroding biodegradable material that is fully biocompatible,\textsuperscript{20} exhibits an ideal sustained, zero-order release profile and promotes bone regeneration\textsuperscript{21} due to its unique degradation behavior.\textsuperscript{22}

In this study, with the aim to increase the sustained effectiveness of intracellular antibiotics and the probability of biocompatibility, VCM/TMC NP-PTMC was prepared using a composite of VCM/TMC NPs and PTMC. Scanning electron microscopy (SEM) was performed to characterize the surface of the beads before and after erosion. Dialysis was performed to evaluate the drug release, which was measured by reverse-phase high-performance liquid chromatography (HPLC). In vitro cytology examination and the effect on antibacterial activity were analyzed for the NPs. Additionally, the VCM/TMC NP-PTMC was implanted into a chronic osteomyelitis rabbit model to evaluate its antibiotic activity, biocompatibility and bone regeneration ability.

**Materials and methods**

**Materials**

PTMC was purchased from Shanghai Leon Chemical Co., Ltd (Shanghai, China). VCM-HCl was supplied by Dalian Meilun Biology Technology Co., Ltd (Dalian, China). Lipase solution from *Thermomyces lanuginosus* (100,000 U/g) was obtained from Sigma-Aldrich (St Louis, MO, USA) and used as received. Tetrahydrofuran (THF) and anhydrous hexane were purchased from Tianjin Shield Specialty Chemical Co., Ltd (Zhejiang, China). Beyotime Biotechnology Ltd Co (Shanghai, China) supplied the Cytotoxicity Assay Kit and all cell culture reagents. Methanol (HPLC grade) was purchased from Fisher Chemical (Suzhou, China). Other reagents and chemicals were of analytical reagent grade, unless indicated otherwise.

**VCM analysis**

The antibiotic concentration of VCM was determined using our established method\textsuperscript{23} in order for calculating the antibiotic efficiency of VCM/TMC NP-PTMC.

**Preparation and characterization of VCM/TMC NP-PTMC**

VCM/TMC NPs were prepared as described previously.\textsuperscript{17} VCM/TMC NP-PTMC was prepared according to the reported literature with a slight improvement.\textsuperscript{22} In short, PTMC solution was prepared in a glass dish sealed by a cover by dissolving approximately 0.67 g of polymer in 25 mL THF while stirring at room temperature until complete dissolution. Then, under continuous stirring, 128 mg of lyophilized VCM/TMC NPs was slowly added and aged for 24 h to obtain a relatively uniform paste. The obtained suspension was precipitated with an excess of anhydrous hexane and then immediately dried in vacuum overnight. For preparing the test sample of the drug delivery system, the collected dried material was further cut into cuboids – with a square shape of 10 mm length, 10 mm breadth and a height of 2 mm – and stored in a desiccator. VCM-PTMC was prepared in the same way as the experimental control group, and the drug loading was calculated.

The surface topography and morphology of VCM/TMC NP-PTMC and VCM-PTMC (used as a control) were examined by SEM (S-3400N; Hitachi, Tokyo, Japan). The resulting samples were cut into smaller pieces, placed on the copper cylinders and coated with gold by an ion sputter. The presence of VCM or VCM/TMC NPs in the PTMC was confirmed through X-ray diffraction (XRD) (X’Pert PRO X-ray diffractometer; PANalytical, Mainz, Germany) and Fourier transform infrared (FTIR) spectroscopy analysis.

**In vitro biodegradation study**

The biodegradation behavior of VCM/TMC NP-PTMC and VCM-PTMC (used as a control) was examined in an incubator shaker under sink conditions at 37°C. In brief, the prepared specimens were suspended in a vial of 5 mL phosphate-buffered saline (PBS) pH 7.4 (NaCl: 8 g/L, KCl: 0.4 g/L, KH\textsubscript{2}PO\textsubscript{4}: 0.4 g/L and Na\textsubscript{2}HPO\textsubscript{4}: 2.86 g/L) or lipase aqueous solution (containing 2% enzyme concentrate, 0.5% CaCl\textsubscript{2}, and 25% propylene glycol, w/w), closed tightly and rotated at 100 rpm. At predetermined time points, square
specimens were collected and vacuum-dried for 0.5 h. Afterward, the masses were measured in triplicate, and the changes to the surface morphology during the degradation process were examined by SEM. The relative mass represents the degradation ratio and was calculated as Relative mass = (Wf - Wi) / Wf, where Wf and Wi represent the weight of dried specimen before and after incubation in medium.

**In vitro release profile**

A dialysis technique with two kinds of release media was used to evaluate the in vitro release profile of VCM from VCM/TMC NP-PTMC. One randomly chosen prepared specimen (8.39 ± 1.15 mg) was placed in 1 mL of release medium and then transferred into a dialysis bag (molecular weight cutoff MWCO: 2 kDa), which was dialyzed against 4 mL of the corresponding medium (PBS 7.4 or lipase aqueous) at 100 rpm in a 37°C water bath. On days 2, 5, 7, 12, 18, 27, 32 and 36, 400 μL of the outside solution was harvested, and an equal amount of PBS 7.4 or lipase aqueous solution was supplemented to maintain a constant volume. The amount of released drug was then determined and calculated as the cumulative percentage release.

**Antibacterial activity assay**

**Bacterial culture**

Gram-positive *S. aureus* (American Type Culture Collection [ATCC] 6538; Shanghai BioRc Co., Ltd, Shanghai, China) was purchased and cultured in Luria-Bertani (LB) broth (in triplicate) were tested for osteoblast proliferation by measuring the DIZ around the disk. The DIZ values were taken in four perpendicular directions and measured by a Vernier caliper. All tests were performed in triplicate.

**Antibacterial activity on the infected rat calvarial osteoblasts (ROBs)**

ROBs were isolated from neonatal (2-day-old) Sprague Dawley rats;25 cells at passage four were used. To identify the infected ROBs, *S. aureus* was labeled with fluorescein isothiocyanate (FITC; 100 μg/mL) for 4 h. The nuclei of the ROBs were labeled with 4',6-diamidino-2-phenylindole, and the cytoskeleton was labeled with rhodamine phalloidin, following a modified previous protocol.6 An *S. aureus* suspension (bacterial cells corresponded to approximately 1×10⁶ CFU/mL) was added to induce intracellular infection. The cells were incubated for 2 h in a CO₂-free incubator, and then VCM/TMC NP-PTMC or VCM-PTMC (the VCM concentration was 50 μg/mL) was added. Untreated ROBs were the control group. The number of bacterial colonies was quantified using a medical image management system (Image-Pro Plus, IPP6.0; Media Cybernetics, Inc., Bethesda, MD, USA).

**Protein adsorption**

Bicinchoninic acid (BCA) assay was performed to quantify the total protein adsorption onto the VCM/TMC NP-PTMC (5×5×2 mm with similar weight), as reported by Shalumon et al,26 with small change. Briefly, VCM/TMC NP-PTMC and VCM-PTMC (used as a control) were placed in a 24-well plate and incubated at 37°C with Dulbecco’s Modified Eagle’s Medium (DMEM, pH 7.4) supplemented with 10% (v/v) heat-inactivated fetal bovine serum (FBS) for 24 h and 48 h. The total amount of protein in the medium was determined prior to adding the materials. At the preestablished time, according to the manufacturer’s guidelines (Beyotime Biotechnology Ltd Co.), the supernatants were collected to determine the remains in the medium using the following equation. Protein adsorption was expressed as a percentage content of the control cells.

\[
R(\% \text{ w/w}) = \frac{(M_p - S_p)}{M_p} \times 100
\]

where R, Mp and Sp represent the protein adsorption ratio, total amount of protein in medium, and amount of protein in supernatants, respectively.

**Cell proliferation**

The EtO-sterilized VCM/TMC NP-PTMC and VCM-PTMC (in triplicate) were tested for osteoblast proliferation by
rabbits were euthanized with overdosages of pentobarbital sodium. The marrow in the tibia was spread on top of sheep blood agar plates, incubated overnight at 37°C and the number of colonies counted. After incubation, 100 μL of formazan dissolution buffer was added to each well to dissolve the MTT formazan crystals. The plates were shaken for 5 min and the absorbance was measured at 490 nm using a microplate reader. The results of five repeated experiments were expressed as a percentage viability of the control cells.

In vivo treatment in a chronic osteomyelitis rabbit model

Animal groups and treatment

All the animal experiments were performed in accordance with the Guide for the Care and Use of Laboratory Animals and were approved by the Bioethics Committee of Zhejiang Academy of Traditional Chinese Medicine. Male New Zealand White rabbits (SCXK 2014-0001), aged 3 months and weighing 3.0±0.2 kg, were housed in individual cages under air-controlled conditions (20°C±1°C and 12 h/12 h light–dark illumination cycles). The model was prepared according to a previous report, and five rabbits were verified by microbiological and histological examination at the fourth week after infection. The model rabbits were divided into the model, VCM/TMC NP-PTMC and VCM-PTMC groups. The two test groups were implanted with VCM/TMC NP-PTMC or VCM-PTMC beads (10 mg VCM per rabbit) after thoroughly cleaning the infected tissue. The rabbits in the model group were housed under the same conditions but not treated.

Antibacterial activity evaluation

At the end of the fourth week after infection and at the second, fourth and sixth weeks after treatment, inflammatory indexes in serum, such as highly sensitive C-reactive protein (CRP) and white blood cell count (WBC), were determined. At the end of the sixth week after treatment, the rabbits were euthanized with overdosages of pentobarbital sodium. The marrow in the tibia was spread on top of sheep blood agar plates, incubated overnight at 37°C and the number of colonies counted.

Bone repair capability evaluation

At the end of the fourth and eighth weeks after treatment, rabbits were euthanized with overdosages of pentobarbital
Figure 1 The characterization of VCM-PTMC and VCM/TMC NP-PTMC.

Notes: (A) SEM images of the surfaces of VCM-PTMC (left) and VCM/TMC NP-PTMC (right) before and after incubation, as seen after incubation in PBS 7.4 and lipase aqueous medium at 37°C. The top images show the surfaces of the beads before incubation, the middle images show the surfaces of the beads after 5 days of incubation in PBS 7.4, and the bottom images show the exposure to the surfaces of the beads after 3 days of incubation in lipase aqueous medium. (B) XRD spectra of VCM/PTMC and VCM/TMC NP-PTMC. (C) FTIR spectra of VCM-PTMC and VCM/TMC NP-PTMC. (1) VCM-TMC NPs; (2) PTMC; and (3) VCM/TMC NP-PTMC.

Abbreviations: FTIR, Fourier transform infrared spectroscopy; NP, nanoparticle; PBS, phosphate-buffered saline; PTMC, poly(trimethylene carbonate); SEM, scanning electron microscopy; TMC, N-trimethyl chitosan; VCM, vancomycin; XRD, X-ray diffraction.
appeared. Compared to the IR spectrum of VCM-TMC NPs, a new band at 1747 cm\(^{-1}\), attributed to the C–O stretching region, appeared in the spectrum of the PTMC. The region between 1500 cm\(^{-1}\) and 1360 cm\(^{-1}\) is characterized by the 1475 cm\(^{-1}\) CH\(_2\) band, which is attributed to the asymmetrical stretching of C–H in the methyl groups on the quaternary ammonium, and this appeared in the spectrum of TMC. The 1380 cm\(^{-1}\) and 1360 cm\(^{-1}\) peaks could be ascribed to the C–H deformation and asymmetric bands (Figure 1C).

**In vitro biodegradation study**

The loading efficiency (LE) of VCM/TMC NP-PTMC was calculated according to the following equation:

\[
\text{LE}(\%, \text{w/w}) = \frac{Aa}{(Aa + Ap)} \times 100
\]

\[= \frac{0.01(0.01 + 0.67)}{1.47}
\]

where Aa and Ap represent the total amount of antibiotic and total amount of polymer material, respectively.

PBS at pH 7.4 and lipase aqueous media were applied to determine the in vitro biodegradation. Figure 2A depicts that on days 2–6, the VCM-PTMC in the lipase aqueous medium showed a slightly higher relative degradation mass in comparison to the VCM/TMC NP-PTMC, and on the sixth day, 0.19±0.01 relative degradation mass was observed for the VCM/TMC NP-PTMC. However, the VCM-PTMC showed 0.35±0.03 relative degradation mass in the lipase aqueous medium. In addition, in PBS 7.4, the VCM-PTMC showed higher relative degradation mass compared to the VCM/TMC NP-PTMC, and on the sixth day, 0.59±0.11 relative degradation mass was observed for the VCM-PTMC. At the end of the 36th day, the VCM/TMC NP-PTMC showed 0.53±0.05 relative degradation mass.

**In vitro drug release study**

The cumulative release curves (Figure 2B) showed the release behavior of VCM from VCM-PTMC and VCM/TMC NP-PTMC over 36 days in PBS 7.4 and lipase aqueous medium. The cumulative release of VCM followed a steady, continued-release pattern, without a burst release phenomenon. The cumulative release rates of VCM from VCM-PTMC and VCM/TMC NP-PTMC were 90.79±0.06% and 83.57±0.02%, respectively, with 36-day incubation in the lipase aqueous medium. The cumulative release rates of VCM from VCM-PTMC and VCM/TMC NP-PTMC were 24.21±0.02% and 18.71±0.04% with 36-day incubation in the PBS 7.4 medium, respectively.

**Antibacterial activity**

DIZ results (Figure 2C) showed that the DIZ values of the release solution from VCM-PTMC and VCM/TMC NP-PTMC were 10.84±0.70 mm and 11.18±0.66 mm on the 18th day of incubation in lipase aqueous solution. On the 32nd day, the DIZ value of VCM-PTMC was reduced to 6.23±0.69 mm but was 9.93±0.41 mm for VCM/TMC NP-PTMC. The DIZ values of release solution from VCM-PTMC and VCM/TMC NP-PTMC incubated in PBS 7.4 were 10.30±0.36 mm and 11.06±0.30 mm on the 27th day and 9.45±0.37 mm and 9.46±0.25 mm on the 36th day, respectively.

As shown in Figure 2F and G, following 2 h of infection with *S. aureus*, labeled bacteria became visible inside the ROBs, while after 24 h incubation of infected ROBs with VCM/TMC NP-PTMC, the amount of *S. aureus* both inside and outside the cells was drastically reduced. In addition, in infected ROBs incubated with VCM-PTMC, *S. aureus* visible outside the ROBs was drastically reduced, but *S. aureus* inside the ROBs showed strong density of fluorescence. The results indicated that VCM/TMC NPs inhibited *S. aureus* intracellularly and extracellularly.

**Protein adsorption**

The protein adsorption studies were carried out for 24 h and 48 h, respectively, wherein the VCM/TMC NP-PTMC showed a significantly higher protein adsorption compared to the control beads at both the time periods (\(P<0.01\)) (Figure 2D). Additionally, the maximum protein adsorption on the VCM/TMC NP-PTMC was achieved in the initial 48 h period. The VCM-PTMC showed a significantly higher protein adsorption in the initial 48 h period than at 24 h (\(P<0.05\)).

**Cell proliferation**

Cell proliferation activity was tested by the MTT assay. Figure 2E showed that after 4-day incubation, VCM/TMC NP-PTMC elicited significantly higher cell proliferation activity (\(P<0.05\)) compared to VCM-PTMC. Similar results were obtained with significance (\(P<0.01\)) after 7 days of incubation.

**In vivo antibacterial activity**

Most model rabbits had a large number of infected sinuses, and white and yellow pus overflowed from the wounds (Figure 3A), similar to the representative symptom of chronic osteomyelitis in clinic. Streaking the marrow lysates on agar plates resulted in a concordantly decreased number of colonies for the VCM/TMC NP-PTMC and VCM-PTMC.
groups, which even reduced down to zero for the VCM/TMC NP-PTMC group after 6-week treatment (Figure 3B). Compared to the control group, the levels of CRP and WBC were higher in the model group. At the end of the fourth week and eighth week after treatment with VCM/TMC NP-PTMC or VCM-PTMC, both antibiotic delivery systems significantly decreased the CRP and WBC (Figure 3D).

Bone repair capability
After 8 weeks of treatment, the tibial plateau of the VCM/TMC NP-PTMC group seemed flatter than that of the VCM-PTMC group (Figure 3C). The images of the 3D reconstruction in Figure 4A indicated a progressive increase in bone volume at the end of eighth week after treatment with VCM/TMC NP-PTMC and VCM-PTMC while it tended
toward bone loss in the model groups. The micro-CT indexes
(Figure 4B) showed that the BV/TV and BMD decreased
further and were significantly lower in the model group than
in the VCM/TMC NP-PTMC and VCM-PTMC groups.
In addition, the BV/TV and BMD were markedly higher in the
VCM/TMC NP-PTMC group than in the VCM-PTMC group.
A histomorphometric analysis showed a significant difference
between the VCM/TMC NP-PTMC and VCM-PTMC groups,
demonstrated by increased MAR and BFR (Figure 4C and D).
These results suggested the effect of VCM/TMC NP-PTMC
on promoting bone healing. Infection caused a decrease in the
number of bone trabeculae, but the VCM/TMC NP-PTMC
and VCM-PTMC groups showed less pronounced decreases
than the model group. A pronounced decrease in number of
osteoblasts per trabecular bone area (NOb/TAr) and number
of osteoblasts per bone perimeter (NOb/Bpm) was induced
by \textit{S. aureus}, but this decrease was mitigated by VCM/TMC
NP-PTMC and VCM-PTMC treatment (Figure 5A and B).

As shown in Figure 5C, significantly decreased rates of
integrated optical density/area (IOD/area) for osteocalcin
(OC) were observed in the model group than in both VCM/
TMC NP-PTMC and VCM-PTMC groups, which was more
pronounced in the VCM/TMC NP-PTMC group.

**Discussion**

In our previous study,\textsuperscript{17} the geometric mean diameter of
VCM/TMC NPs obtained from atomic force microscopy
was 200–325 nm, mean size distribution was 220–230 nm
with a fairly monodisperse polydispersity index, which was
almost <0.2; meanwhile, VCM/TMC NPs carried a positive
zeta potential of 14.6±0.8 mV at physiological pH, suggest-
ing that these NPs are probably ionically stabilized against
particle agglomeration due to the positive charges on their
surface. In this study, the SEM results illustrated the pres-
ence of spherical VCM/TMC NPs with a smooth surface
and compact structure. The X-ray and IR spectrum results

![Figure 2](https://www.dovepress.com/)

**Figure 2** The biodegradation, antibacterial activity, protein adsorption, cell proliferation and drug release characteristics.

**Notes:** (A) Relative biodegradation mass of VCM-PTMC (\(\pi\)) and VCM/TMC NP-PTMC (\(\theta\)) in PBS 7.4 and lipase aqueous media. (B) The curves of cumulative release of VCM from VCM-PTMC (\(\pi\)) and VCM/TMC NP-PTMC (\(\theta\)) in the PBS 7.4 and in the lipase aqueous media. (C) The graphical representation of the antibacterial activity of VCM-PTMC and VCM/TMC NP-PTMC solution released from PBS 7.4 and lipase aqueous media. (C1) The antibacterial activity of VCM/TMC NP-PTMC solution released from PBS 7.4 and lipase media. (C2) The antibacterial activity of VCM/TMC NP-PTMC solution released from PBS 7.4 and lipase media. (D) The graphical representation of cell proliferation and drug release ability on the surface of VCM-PTMC and VCM/
TMC NP-PTMC. (E) The graphical representation of attachment and proliferation ability on the surface of VCM-PTMC and VCM/TMC NP-PTMC. (F) Fluorescence optical images of fluorescently stained osteoblastic cell nuclei (blue arrow) and cytoskeletal F-actin (pink arrow), invaded by \textit{Staphylococcus aureus} (green arrow). (G) The bacterial count after 24 h and 48 h incubation of infection ROBs with VCM-PTMC and VCM/TMC NP-PTMC. **P < 0.05.***

**Abbreviations:** NP, nanoparticle; OD, optical density; PBS, phosphate-buffered saline; PTMC, poly(trimethylene carbonate); ROBs, rat calvarial osteoblasts; TMC, \(N\)-trimethyl chitosan; VCM, vancomycin.
showed that VCM/TMC NPs are successfully introduced to the PTMC.

VCM-PTMC had more rapid biodegradation than VCM/TMC NP-PTMC, which is due to the slow degradation of VCM/TMC NPs in the VCM/TMC NP-PTMC. Biodegradable polymers can be arbitrarily classified into two groups: bulk eroding (homogeneous) and surface eroding (heterogeneous) polymers. 22 PTMC biodegradation only occurs on the surface; therefore, its degradation process is slower than materials degraded through bulk erosion, which prevents the bulk erosion of the VCM/TMC NPs due to immediate contact with the solvent and premature release. 31 In addition, compared to the PBS pH 7.4 medium, both delivery systems had obvious swelling phenomenon in the lipase aqueous medium, which is associated with the conformation of the adsorbed enzymes on the surface. As the PTMC
Figure 4 The bone repair capability of VCM-PTMC and VCM/TMC NP-PTMC.

Notes: (A) Micro-CT morphometry of bone defect of the tibia in chronic osteomyelitis model rabbits that underwent 8 weeks of treatment with VCM-PTMC and VCM/TMC NP-PTMC. (B) The columns represent the scores of the BV/TV and BMD from five rabbits per group. (C) Typical photographs of fluoroscope observation on tibia with double-fluorescent labeling. (D) The columns represent the scores of MAR and BFR. *P<0.05.

Abbreviations: BFR, bone formation rate; BMD, bone mineral density; BV/TV, bone volume/tissue volume; CT, computed tomography; MAR, mineral appositional rate; NP, nanoparticle; PTMC, poly(trimethylene carbonate); TMC, N-trimethyl chitosan; VCM, vancomycin.
Figure 5 The histopathology of tibia and immunohistochemical analysis.

Notes: (A) Typical histopathology of the rabbit tibia. (B) The columns represent the scores of osteoblast number/trabecular bone area (NOb/TAr); osteoblast number/bone perimeter (NOb/Bpm). (C) The columns represent the IOD/area rate of immunohistochemical staining for OC from six rabbits per group. Data are presented as the mean ± SD. *P < 0.05.

Abbreviations: IOD, integrated optical density; NP, nanoparticle; OC, osteocalcin; PTMC, poly(trimethylene carbonate); SD, standard deviation; TMC, N-trimethyl chitosan; VCM, vancomycin.
possesses more hydroxyl termini, it is liable to absorb water and is not as subject to enzymatic degradation. Overall, the release of VCM/TMC NP-PTMC may be better for the prolonged administration of antibiotics.

Protein adsorption was enhanced in the VCM/TMC NP-PTMC, mainly due to 1) the presence of NPs that increase the total surface area and binding sites for proteins or 2) an electrostatic interaction between the material surface and proteins. Protein adsorption further aids in cellular adhesion, infiltration and proliferation by the adsorption of adhesion molecules such as fibronectin and/or vitronectin. The adhesion and proliferation of ROBs could be influenced by varying the nanostructured surfaces. The fixed surface structure in the nanometer-size scale could be useful for designing novel scaffolds in tissue engineering applications. Therefore, it is reasonable to conclude that the VCM/TMC NP-PTMC are not only cytocompatible but also significantly effective on ROB proliferation to induce bone regeneration. PTMC is recognized as a cell-friendly material as well, with high levels of cell viability and cell adhesion being observed on the PTMC surface. As a result, VCM/TMC NP-PTMC can be useful nanocarriers for the intracellular delivery of various antibiotics.

The different antibacterial activities in vitro were due to the presence of VCM/TMC NPs in the VCM/TMC NP-PTMC, possessing the slow biodegradation and sustained drug release characteristics of VCM/TMC NPs. Moreover, in our previous study, the synthesis mechanism and cell uptake evaluation proved that the VCM/TMC NPs with a positive zeta potential were responsible for the effective attachment to the negatively charged cell membrane of bacteria, which caused the leakage of proteinaceous and intracellular components, leading to bacterial rupture and death. In this study, the VCM/TMC NP-PTMC could improve delivery of water-soluble VCM across the cytomembrane, due to its positive surface charge, facilitating the movement of antibiotics across the ROB membrane. Therefore, antibiotic treatment of the infected osteoblasts indicated that VCM/TMC NP-PTMC inhibited intracellular *S. aureus* efficiently, which can explain the excellent antibacterial activity of VCM/TMC NP-PTMC in the chronic osteomyelitis rabbits.

In chronic osteomyelitis, although the necrotic tissue has been adequately debrided, a large bone defect is inevitable. As a result, the appropriate reconstruction of bone defect is needed. VCM-CaSO₄ was used for local antibiotic treatment for many years in clinic; however, the overdosage of VCM administered locally caused the inhibition of osteoblast activity and proliferation. NPs have been shown to promote cell attachment and proliferation ability. Actually, the VCM/TMC NP-PTMC in our study exhibited a sustained and slow release that can eliminate the inhibition of high dose of VCM and possessed excellent antibiotic activity and bone repair capability. Meanwhile, the PTMC is best known as a surface-eroding polymer and is fully biocompatible, which appears to fully degrade in bone and assists in promoting bone regeneration. Generally, in the clinic, patients are commonly supplied therapy for at least 4 weeks. Therefore, VCM/TMC NP-PTMC could meet the requirements.

Although, the proliferative probability of ROBs and the antibacterial activities of VCM/TMC NPs and VCM/TMC NP-PTMC in vivo and in vitro were compared and evaluated in this article, the molecular mechanism of effect of VCM/TMC NPs and VCM/TMC NP-PTMC on bone regeneration and bone repair has not been stated. The relevant studies will be performed in our further research.

**Conclusion**

This work has demonstrated that rationally designed VCM/TMC NP-PTMC can be readily prepared using a composite method with a high drug-loading capacity and a steady, biodegradable, cytocompatible and antibacterial nature. An in vivo study showed that implantation of VCM/TMC NP-PTMC was an efficient method of antibiotic treatment in chronic osteomyelitis, and VCM/TMC NP-PTMC had excellent probability of promoting bone healing. These NPs thus can be a promising strategy to treat intracellular infections such as chronic osteomyelitis in the clinic.

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**Disclosure**

The authors report no conflicts of interest in this work.

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