Evaluation of interbody fusion efficacy and biocompatibility of a polyetheretherketone/calcium silicate/porous tantalum cage in a goat model

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ABSTRACT

Objective: To evaluate the interbody fusion efficacy and biocompatibility of a graft-free cage made of polyetheretherketone/calcium silicate composite/porous tantalum (PEEK/CS/pTa cage) compared with a PEEK/CS cage with an autogenous bone graft in a goat model.

Methods: PEEK/CS/pTa and PEEK/CS cages were prepared through an injection-moulding method. The PEEK/CS composites and porous tantalum were characterized by Fourier transform infrared spectroscopy (FTIR), X-ray photoelectron spectroscopy (XPS), X-ray diffraction (XRD), scanning electron microscopy (SEM), and energy-dispersive spectroscopy (EDS) mapping. Then, adult goats were chosen for C2/C3 and C3/C4 discectomy via the anterior cervical approach and randomly implanted with PEEK/CS/pTa and PEEK/CS/cages with autogenous bone grafts. The fusion performance and osseointegration of the cages were evaluated by X-ray imaging, magnetic resonance imaging (MRI) scanning, and bone histomorphometry analysis. Moreover, the concentrations of Ca and Si in urine, serum, tissue around the fusion segments and major organs of the goats were determined by inductively coupled plasma–optical emission spectrometry (ICP–OES). Histological observation of major organs of the goats was used to evaluate the biosafety of PEEK/CS/pTa and PEEK/CS cages.

Results: X-ray and MRI imaging suggested that both PEEK/CS/pTa cages and PEEK/CS cages maintained similar average intervertebral space heights. The tissue volumes in the fusion area were comparable between the two groups of cages at 26 weeks after surgery. Histological morphometric data showed that PEEK/CS/pTa cages and PEEK/CS cages with autogenous bone grafts had similar bone contact and osseointegration via the anterior cervical approach and randomly implanted with PEEK/CS/pTa and PEEK/CS/cages with autogenous bone grafts. The fusion performance and osseointegration of the cages were evaluated by X-ray imaging, magnetic resonance imaging (MRI) scanning, and bone histomorphometry analysis. Moreover, the concentrations of Ca and Si in urine, serum, tissue around the fusion segments and major organs of the goats were determined by inductively coupled plasma–optical emission spectrometry (ICP–OES). Histological observation of major organs of the goats was used to evaluate the biosafety of PEEK/CS/pTa and PEEK/CS cages.

Conclusion: Overall, these results suggested that the graft-free PEEK/CS/pTa cage showed similar bony fusion performance to the PEEK/CS cages with autogenous bone grafts. The cages releasing calcium and silicon had good biological safety in vivo. The translational potential of this article: This study provided a new graft-free interbody fusion solution to patients with degenerative disc diseases, which could avert potential donor-site complications. This study also

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1. Introduction

Discectomy and fusion are effective surgical procedures to treat degenerative disk diseases and rebuild spinal stability. Most surgeons apply interbody fusion cages to enhance spinal mechanical endurance and augment spine function [1]. Autogenous bone grafts are usually packed with interbody fusion cages and act as osteogenesis substrates to achieve bony fusion in the long term [2]. For decades, autogenous bone grafts have been widely regarded as the gold standard for interbody fusion due to their strong osteogenic potential and complete histocompatibility [3-5]. However, the acquisition of autogenous bone grafts inevitably causes trauma in the donor site and could potentially lead to multiple kinds of donor-site complications, such as persistent donor-site pain, deep haematoma formation, deep infection, incisional hernia, and donor-site fracture [6,7]. According to previous reports, 2%-37.9% of patients suffered donor-site pain 6 months after the operation, and 2.5% of patients suffered deep infection at the donor site [8-11]. The limited bone availability and donor-site complications drove the development of alternative materials [12-14].

Porous tantalum, recognized as an effective material for artificial metal trabecular bone, has been used in spine surgery for years [15-17]. Porous tantalum has a high volumetric porosity, which allows ingrowth of nearby tissue and subsequent osteogenesis [18]. In addition, the Young’s modulus of porous tantalum was approximately 3-4 GPa, which was relatively similar to that of cancellous bone (approximately 10.4 GPa) [19] and could avert the stress-shielding effect of traditional metal implants [18,20]. In addition, porous tantalum has strong corrosion-resistant properties and good biocompatibility [18,21,22]. In previous studies, porous tantalum has been proven to show considerably good osteoconductive and osteoinductive performance in a cervical interbody model, with evident trabecular bone ingrowth and osseointegration at the fusion interface [23]. Therefore, porous tantalum is an ideal alternative material to autogenous bone grafts.

Among the various kinds of materials used for interbody cages, polyetheretherketone (PEEK) accounts for a large percentage because of its good biocompatibility and moderate elastic modulus, similar to cancellous bone [24-26]. PEEK cages were biologically inert, and this property led to relatively high biosafety as well as some disadvantages, including fibrous healing and low osseointegration at the implant surface [27]. With the efforts of recent years, various methods have been developed to improve the biological activity of PEEK, such as surface coatings made of titanium or hydroxyapatite and other incorporated composites [28-31]. In our previous studies, we designed and developed a compounding and injection-moulding technique to incorporate calcium silicate (CS) into PEEK to manufacture a PEEK/CS composite material, which was optimized for osteogenesis with a CS ratio of 40% by weight (wt%) [32,33]. A comparative study showed that the PEEK/CS interbody fusion cage had a better pro-ostegenic effect in vitro and better fusion and osseointegration performances in vivo than the pure PEEK interbody fusion cage in a goat cervical interbody fusion model [34]. Based on these findings, we step further to replace the packed-in autogenous bone graft with porous tantalum and designed a new PEEK/CS/pTa cage that could potentially avert trauma and complications at the donor site and achieve similar fusion performance and osseointegration. This study was designed to compare the interbody fusion performance of PEEK/CS/pTa cages and PEEK/CS cages packed with autogenous bone grafts and investigate the calcium (Ca) and silicon (Si) distribution in the body, with the aim of laying a foundation for biosafety assessment and clinical translation.

2. Materials and methods

2.1. Preparation of interbody fusion cage

The PEEK/CS composite interbody fusion cage was manufactured as previously described. In brief, PEEK powder (Solvay Specialty Polymers, GA, USA) and micron-sized calcium silicate powders were blended in a ball mill (QM-3B, T-Bota Ltd., China) with a weight ratio of 6/4 for 1 h. Then, the mixed powders were dried at 150 °C for 24 h and heated to 380 °C in an injection-moulding machine (BA-300/050CD, Awans, Belgium) and fabricated into PEEK/CS composites, which were further processed through a turning lathe (C616-1, Jinan First Machine Tool Group Co. Ltd., China) and cut into interbody fusion cage rings. For PEEK/CS/pTa interbody fusion cage preparation, the inner void space of the PEEK/CS interbody fusion cage was filled with a pre-cut porous tantalum metal block (Trabecular MetalTM VBR-21, Zimmer Biomet, USA) with a size of 5 mm × 5 mm × 3 mm. Otherwise, the inner void space was filled with the autogenous bone graft, which was harvested from the iliac crest during the operation. The specific shape and size of these interbody fusion cages are shown in Fig. 1.

2.2. Characterization of PEEK/CS/pTa interbody fusion cage

The surface morphology, element composition and distribution of the PEEK/CS interbody fusion cage and porous tantalum metal block were characterized by field-emission scanning electron microscopy (SEM, S4800, HITACHI, Japan) and energy-dispersive spectrometry (EDS, S4800, HITACHI, Japan). The composition of the PEEK/CS interbody fusion cage was determined by X-ray photoelectron spectroscopy (XPS, ESCALAB 250, Thermo Scientific, USA) and Fourier transform infrared spectroscopy (FTIR, Nicolet iN10, Thermo Scientific, USA), and the phase structure of the porous tantalum metal block was determined by X-ray diffraction (XRD, Rigaku Ultima IV, Rigaku Corporation, Japan).

2.3. Animal model and surgical procedures

An anterior cervical discectomy and fusion model in goats was carried out according to our previous study [34]. All animal procedures and experiments were reviewed and approved by the Animal Ethical Committee of Shanghai Ninth People’s Hospital. The approval number is SH9H-2021-A649-1. In brief, 15 female goats (2 years old, 40–50 kg) were purchased from Shanghai Jiagan Biotechnology Company and accustomed to the new environment for one week before the surgery. The PEEK/CS cages and PEEK/CS/pTa cages used for surgery were sterilized by ethylene oxide sterilization. On the day of surgery, the goats were anaesthetized with gas anaesthesia (isoflurane) and fixed on the operation table in a supine position with tracheal intubation inserted to maintain an unobstructed airflow. Then, the skin of the neck area was disinfected with an iodophor and 75% ethanol scrub. After this, a longitudinal skin incision was made 1.5 cm to the left of the midline of the neck. Then, we fully exposed the operation field of the ventral C2/C3 and C3/C4 disc surfaces for discectomy, which made enough intervertebral space for interbody cage implantation. The PEEK/CS and PEEK/CS/pTa interbody fusion cages were randomly implanted in the intervertebral space of C2/C3 and C3/C4 in a goat. The inner holes of the PEEK/CS interbody fusion cage were filled with autogenous bone grafts harvested from the iliac crest. Additionally, a titanium plate was utilized to fix the cervical vertebrae. Finally, routine closure was performed, and 4 million
Fig. 1. Design and characterization of PEEK/CS and PEEK/CS/pTa interbody fusion cages. (A) Schematic design of the PEEK/CS (upper) and PEEK/CS/pTa (lower) interbody fusion cages. (B) Photographs of the actual PEEK/CS (upper) and PEEK/CS/pTa (lower) interbody fusion cages. (C) FTIR spectra of the PEEK/CS composites. (D) Full XPS spectra of the PEEK/CS composites. (E) XRD pattern of the porous tantalum scaffold. (F) SEM images and EDS mapping of the PEEK/CS composites and porous tantalum scaffold.
units of penicillin sodium were intramuscularly injected into each goat for 5 days after surgery. Goats that had undergone surgery were carefully raised and monitored for any adverse reactions and complications until they reached corresponding endpoints. Goats were intramuscularly injected with alizarin red solution (20 mg/kg) and calcein solution (20 mg/kg) 2 weeks and 3 days before being sacrificed. As indicated in Fig. 2A, at each endpoint, goats were euthanized, and the main organ tissues, including heart, liver, spleen, lung, kidney, and cervical spinal segments, were immediately collected. Some of the main organ tissues were immersed in 4% formalin solution, and the remaining organ tissues and spinal segments were stored at −20 °C for further analysis.

2.4. X-ray analysis

At predetermined time points (Fig. 2A), goats were anaesthetized, and lateral cervical X-ray images were harvested with an X-ray DR system (Ultimax, TOSHIBA, Japan). The average interbody disc space height (DSH) was determined to evaluate the fusion performance as previously described. Briefly, the average DSH was measured from the lateral cervical X-ray images according to the following formula: The average DSH = (anterior DSH + middle DSH + posterior DSH)/3 [34,35].

2.5. Magnetic resonance imaging (MRI) scanning and analysis

A high-field MRI scanning system (BRUKER BIOSPEC 70/30 MRI, BRUKER BIOSPEC, Germany) was used to detect and analyse the tissue formation and fusion performance of interbody cages. In brief, experimental spinal segments were sawed to a proper length. After this, the samples were subjected to a high-field MRI scan with a field strength of 7 T and a slice thickness of 0.5 mm. The tissue volume was selected with a global threshold in the inner space of the interbody fusion cage in T1-weighted imaging; the signals were further reconstructed to 3D images and used to quantify the tissue volume. MRI data analysis and reconstruction were performed with 3D Slicer software (Version 4.11).

2.6. Undecalciﬁed bone histomorphometry

Experimental spinal segment samples at two time points were subjected to undecalciﬁed bone histomorphometry analysis to determine the bone formation rate in the fusion area and bone-implant contact as previously reported [34]. In brief, spinal segment samples were dehydrated in a graded ethanol series. The samples were then embedded in methyl methacrylate (MMA, Merck, Schuchardt, Germany) monomer for 2 weeks. After this, the samples were transferred into an embedded device with colloidal MMA and stored at room temperature until the embedded samples totally solidiﬁed. Then, the solidiﬁed samples were cut into thin 100 μm slices using a microtome (SP1600, Leica, Germany). The slices were ﬁxed on plastic slides and polished to approximately 50 μm, and Van Gieson staining was performed according to the manufacturer’s instructions. Finally, the volume of new bone in the fusion area was measured with Image-Pro Plus 6.0 software (Media Cybernetics, United States). Three sections of each sample were randomly chosen for analysis.

2.7. Fluorescence microscopy analysis

Dynamic bone formation during the observation period was tracked by double labelling with alizarin red and calcein, which have already been widely applied in labelling bone tissue. Image-Pro Plus 6.0 software was used to measure the bone formation rate and bone-implant contact.

Fig. 2. Experimental design of the goat interbody fusion model and DSH analysis based on X-ray radiography. (A) Schematic illustration of the surgery and analysis procedures of the goat interbody fusion model. (B) Representative X-ray images at 0, 4, 8, 12, and 26 weeks. (C) DSH analysis at 0 (n = 15), 4 (n = 15), 12 (n = 15), and 26 weeks (n = 8).
rate by observing the fluorescence labelling on slides with confocal laser scanning microscopy (CLSM, Leica, Germany).

2.8. Analysis of elemental Ca and Si

The metabolism of calcium and silicon released from PEEK/CS and PEEK/CS/pTa cages was monitored by determining the concentrations of Ca and Si in the serum and urine of the goats before and after the operation. The accumulation of Ca and Si in goats’ major organs, including the heart, liver, spleen, lung, and kidney, was determined and compared with that of normal healthy female goats. The element concentrations of Ca and Si in bone, dura mater, and spinal cord from operation segments and distal segments (C7) were also measured and compared. Serum and urine samples from goats were collected at 0, 4, 8, 12, 18, and 26 weeks after the operation, and major organ samples of goats and cervical spine segments were harvested immediately after goats were sacrificed. All samples were stored at −80 °C until being analysed. The samples were digested with aqua regia supplemented with 0.25% hydrofluoric acid for 1 h at 180 °C using a microwave digester (ultraWAVE, Milestone, USA) and further analysed for Ca and Si concentrations by inductively coupled plasma optical emission spectroscopy (ICP-OES, Agilent 720 ES, Agilent, USA). To prevent potential Si contamination during sample processing, urine samples were collected with catheters made of polyvinyl chloride (PVC), and all serum and urine samples were stored in polypropylene tubes and chemically digested in Teflon vessels [36,37].

2.9. Histological analysis of major organs

After the goats were sacrificed at predetermined time points, the major organs, including the heart, liver, spleen, lung, and kidney, were collected from goats. Some organ tissues were separated and fixed in 4% paraformaldehyde. The specimens were then embedded in paraffin embedding and cut into 4 µm slices. The changes in the histology of major organs were observed by microscopy with H&E staining.

2.10. Statistical analysis

In this experiment, all data were analysed with SPSS 13.0 software (Statistical Package for the Social Sciences, USA) and presented as the means ± standard deviation. We used two-way analysis of variance (two-way ANOVA) to analyse the data of average interbody disc space height. Multigroup parametric data of element concentrations of Si and Ca in urine, serum, major organs, bone, dura mater, spinal cord and degradation rate of PEEK/CS interbody fusion cage were compared with one-way analysis of variance (ANOVA) followed by Tukey’s test. The tissue formation data determined by MRI, bone-implant contact rate, and bone volume in the fusion area were analysed with the t test.

3. Results

3.1. Characterisation of PEEK/CS/pTa interbody fusion cage

The design draft and actual products of PEEK/CS and PEEK/CS/pTa cages are shown in Fig. 1A-B. FTIR analysis (Fig. 1C) of the PEEK/CS composites showed the characteristic peaks of PEEK. XPS analysis (Fig. 1D) of the PEEK/CS composites revealed the surface composition of calcium silicate. The surface element of tantalum was validated by the signal dots of tantalum of the EDS mapping (Fig. 1F) and characteristic peaks of tantalum in the XRD result (Fig. 1D). SEM and EDS spectrum images (Fig. 1F) demonstrated the rough surface of PEEK/CS composites and porous tantalum scaffolds. The signal dots of Ca and Si had a relatively even distribution pattern on the surface of PEEK/CS composites.

3.2. X-ray imaging and DSH measurement

To monitor interbody fusion, X-ray photographs were performed at predetermined time points (Fig. 2A). As shown in Fig. 2B, in both fusion segments of PEEK/CS cages with autogenous bone grafts and PEEK/CS/pTa cages, an increased density of upper and lower endplates and high-density bony ingrowth were observed at 12 weeks. At 26 weeks, bridging bone was formed between the fusion segments of PEEK/CS cages with autogenous bone grafts. Quantitative measurement (Fig. 2C) revealed that DSH in segments of PEEK/CS cages with autogenous bone grafts or those with PEEK/CS/pTa cages gradually decreased during the follow-up period from 0 to 12 weeks, and no significant difference was observed between the two groups, which indicated that PEEK/CS/pTa cages had fusion stability similar to that of PEEK/CS cages with autogenous bone grafts.

3.3. MRI scanning and analysis

To quantify and visualize the tissue ingrowth and 3D structure in the fusion segments, MRI scanning and tissue volume analysis were performed. As shown in Fig. 3A, obvious tissue ingrowth and bridging tissue formation were observed in both fusion segments of PEEK/CS cages with autogenous bone grafts and PEEK/CS/pTa cages at 12 weeks and 26 weeks. Quantitative analysis of tissue volume revealed that PEEK/CS cages with autogenous bone grafts supported a higher volume of new tissue growth than PEEK/CS/pTa cages at 12 weeks (p < 0.01). The tissue volume in the fusion area of PEEK/CS cages with autogenous bone grafts remained slightly higher than that of PEEK/CS/pTa cages at 26 weeks. However, no obvious significant difference was found between these two groups, which indicated that both the PEEK/CS/pTa and PEEK/CS cages with autogenous bone grafts had satisfactory fusion performance.

3.4. Osseointegration assessment by histomorphometry analysis

Undecalciﬁed bone histomorphometry analysis served as the gold standard for assessing osseointegration and bony fusion. To evaluate the osseointegration performance of PEEK/CS cages with autogenous bone grafts and PEEK/CS/pTa cages, newly formed bone around the cage was double-labelled with ﬂuorescence. As shown in Fig. 4A, ﬂuorescence-labelled mineralized bone was found at the contact surfaces of both cages at 12 weeks. At 26 weeks, denser ﬂuorescence labelling was found at the contact surfaces of both cages. Bone-implant contact analysis (Fig. 4B) revealed that osseointegration of PEEK/CS cages with autogenous bone grafts increased from 33.99 ± 6.57% (12 weeks) to 62.29 ± 6.47% (26 weeks), while the bone-implant contact ratio at the contact interface of PEEK/CS/pTa increased from 31.75 ± 5.20% (12 weeks) to 62.27 ± 13.69% (26 weeks). There were no significant differences between the two groups at either time point. As shown in Fig. 4C, bone tissue formed in the fusion area inside the cages was stained with Van Gieson staining. At 12 weeks, continuously mineralized bone ingrowth was found in PEEK/CS cages with autogenous bone grafts, and deposition of newly formed bone was observed in the porous tantalum scaffolds of PEEK/CS/pTa cages. At 26 weeks, continuously mineralized bone tissue was found in both PEEK/CS cages with autogenous bone grafts and PEEK/CS/pTa cages. Bone volume in the fusion area was quantitatively analysed, as shown in Fig. 4D, which indicated that PEEK/CS cages with autogenous bone grafts supported more mature mineralized bone tissue formation than PEEK/CS/pTa cages at 12 weeks (p < 0.05) and 26 weeks (p < 0.01).

3.5. Determination of Ca and Si levels in serum and urine

To monitor the levels of elemental Ca and Si in the serum and urine of goats, serum and urine samples were collected before and after surgery at predetermined time points and were further analysed by ICP–OES. As shown in Fig. 5A, the Ca concentration in the urine of goats showed a
relatively steady pattern during the whole observation period. The Ca concentration in the serum of goats (Fig. 5B) remained relatively steady at 0, 4, and 8 weeks but showed a significant decrease at 12 weeks and subsequently rose to normal levels (0 weeks) at 18 and 26 weeks. For Si concentration in the urine of goats, Fig. 5C demonstrates that there was no difference in Si level in urine during the observation period except for 26 weeks, at which time it was significantly increased compared with 0 weeks. Si concentration in serum (Fig. 5D) at 8 and 12 weeks significantly decreased compared with 0 weeks, while no difference in the serum Si concentration was observed at 4, 12, or 26 weeks compared with 0 weeks.

3.6. Determination of Ca and Si levels in tissues

To detect the accumulation of Ca and Si in major organs, the heart, liver, spleen, lungs and kidneys were collected and subjected to element analysis. The results showed that the Ca content in the liver and kidneys of surgically manipulated goats was not different from that of normal goats (Fig. 6B and E). The Ca contents in the liver, spleen, and lungs of surgically manipulated goats at 12 weeks were significantly lower than those of normal goats (Fig. 6A, C, 6D). However, these differences diminished at 26 weeks. At 12 and 26 weeks, the Si content in the heart, liver, spleen, lungs, and kidneys of surgically manipulated goats showed no difference compared with those of normal goats (Fig. 6G–J). These results demonstrated that no obvious Ca or Si accumulation was found in surgically manipulated goats implanted with cages made of PEEK/CS composite. Although the Ca content in some organs (heart, spleen, and lung) of surgical goats was lower than that of normal goats, this transient fluctuation was restored to normal levels at 26 weeks. Element analysis of Ca and Si in local tissues, including the spinal cord, dura mater, vertebral bone at the fusion segments and distal segments and new bone formed at the fusion segments, was also carried out to determine the element deposition in tissues around the interbody fusion cages. Compared with distal segments, there was no difference in Ca and Si content in the spinal cord and dura mater of fusion segments (Fig. S1A-B, D-E). In terms of bone tissues, the results (Fig. S1F) of Si content in vertebral bone showed no difference among distal segments, fusion segments and new bone at 12 weeks and 26 weeks. The Ca content in new bone was significantly higher than that of vertebral bone of fusion segments at 12 weeks and 26 weeks (Fig. S1C). The overall results implied that Ca and Si content in tissues was maintained at a normal level.

3.7. Histological observation of major organs

Histological sections and H&E staining of major organs of goats were performed to determine whether pathological changes existed in goats implanted with cages. As shown in Fig. 7, in all goats at 12 weeks and 26 weeks, no obvious histologic abnormalities were observed, indicating that PEEK/CS and PEEK/CS/pTa cages and their degradation products are non-toxic to the major organs of goats.

4. Discussion

Recent years have witnessed the rapid development of artificial and synthetic graft materials, including ceramics, porous metals, polymers, bioactive glass, demineralized bone matrix (DBM) and other composite
materials consisting of pro-osteogenic factors \cite{38-41}. Unlike degradable materials such as ceramic material, bioactive glass and DBM, porous tantalum scaffolds serve as permanent implants, exhibiting strong mechanical properties and enabling a rapid postoperative return to movement and exercise \cite{16,42}.

Interbody fusion cages made of porous tantalum have been applied in clinical practice for years; however, porous tantalum alone has some limitations in application. Due to the relatively high brittleness \cite{43,44}, plastic deformation and cracking of porous tantalum easily occur during the knocking process of the implant surgery, which ultimately reduces the mechanical stability of fusion segments. In most cases of porous tantalum cage implantation, autogenous bone grafts were also used \cite{45,46}. This study demonstrated that the PEEK/CS cage combined with porous tantalum can serve as a new graft-free interbody fusion cage, which is expected to diminish the donor-site complications of autogenous bone grafts.

To examine whether PEEK/CS/pTa cages could achieve fusion performances similar to those of PEEK/CS cages with autogenous bone grafts, two cages were randomly implanted and fixed in segments of C2/C3 and C3/C4 within a goat. Restoration of DSH after implanting an interbody fusion cage was one of the main goals of interbody fusion. According to the DSH measurement results, there was no significant difference between the PEEK/CS/pTa cages and PEEK/CS cages with autogenous bone grafts at 4, 8, 12, and 26 weeks, which reflected the similar fusion stability of these two cages.

Owing to the radiopacity, micro-CT and X-ray cannot show the structure of new bone inside the porous tantalum. According to previous reports, however, artefacts from tantalum metal can be avoided in MRI.
scanning. Zhou et al. utilized MRI scanning to evaluate the fusion progress of porous tantalum interbody fusion devices in a porcine spinal arthrodesis model and proved that T1-weighted spin-echo MRI scanning was an effective and non-invasive way of assessing spinal interbody fusion with porous tantalum devices [23]. MRI scanning was also reported to be compatible with PEEK devices in vivo [47]. Therefore, in this study, we used high-field T1-weighted spin-echo MRI scanning and 3D reconstruction of MRI images to evaluate the tissue formation and fusion performances of the cages. The MRI results showed that both PEEK/CS/pTa cages and PEEK/CS cages with autogenous bone grafts formed continuous bridging tissue at 12 weeks and 26 weeks. Although the newly formed tissue volume of PEEK/CS/pTa was lower than that of PEEK/CS cages with autogenous bone grafts, there was no significant difference between the tissue volume of these two cages at 26 weeks.

Although the tissue formation assessed by MRI scanning revealed that the tissue volume formed in the fusion area was similar between PEEK/CS/pTa cages and PEEK/CS cages with autogenous bone grafts, the bone volume formed in these two cages was unclear. To investigate bone tissue formation, we utilized undecalciﬁed bone sections and Van Gieson staining to determine the bone volume at the fusion area. The results indicated that the bone volume formed in the fusion area of PEEK/CS cages with autogenous bone grafts was signiﬁcantly higher than that of PEEK/CS/pTa cages at 12 weeks and 26 weeks. In fact, the existence of porous tantalum took up much space in the fusion area, which may partly account for the lower bone tissue formation inside the PEEK/CS/pTa cages. Other studies focusing on the osteogenic performance of pTa also conﬁrmed bone ingrowth and occupation of new bone within porous tantalum scaffolds [48,49].

The results of the bone-implant contact rate reﬂect the osseointegration performance of the cages. Mineralized new bone labelled with calcein and alizarin red was observed at the interfaces between the cages and the ingrowing bone. The bone contact rate of these two cages was similar at 12 weeks and 26 weeks, with no signiﬁcant difference being found. Porous tantalum exhibits satisfactory osteoconductivity and mechanical properties due to its highly porous structure and lack of osteoinductive inorganic mineral ions and proteins when compared with ceramic and DBM graft materials. Our previous studies demonstrated that PEEK/CS composites facilitated osteogenic differentiation through continuous surface decomposition of PEEK/CS composites and the release of Ca2⁺ and Si4⁺ in the microenvironment, which could facilitate osteoblastic differentiation and osseointegration [32,33]. Ca2⁺ and Si4⁺ supplements from the PEEK/CS composites, to some extent, are expected to improve the osteogenic performance of porous tantalum.

Evaluation of biocompatibility is a crucial step for clinical translation of any new medical device [50–52]. Ca is widely involved in various biochemical reactions and life activities, and the concentration of Ca in blood is precisely maintained in a narrow range to ensure the normal processes of life [53]. As for silicon, an appropriate rate of silicon release contributes greatly to bone repair [54–56]. On the other hand, dysfunction of silicon excretion and abnormal accumulation in organs would lead to structural damage and dysfunction of important organs and death [57,58]. For interbody fusion cages made of PEEK/CS composites, detailed analysis of the metabolism of Ca and Si in vivo is needed to validate their biosafety. In this regard, we continuously monitored the element levels of Ca and Si in the serum and urine of goats before and after implantation of cages.
after surgery. The results of element concentrations of Ca and Si in serum and urine demonstrated that the overall element levels of Ca and Si maintained a steady state. The decreased Si content in serum at 8 and 18 weeks and decreased Ca content at 18 weeks may be attributed to increased demands of Ca and Si in tissue repair from 8 to 18 weeks postoperatively, during which bone formation was active. The concentration of Si in urine increased at 26 weeks, which may be associated with decreased demand for Si in the fusion area after active bone regeneration. The level of Si in urine ranges from 3.56 to 81.41 mg/L (ppm), which is far less than the saturation level (180 ppm) of Si in urine [59]. Previous studies on Si excretion from implants demonstrated that Si was constantly dissolved in the circulating interstitial fluid at the implant site and further diffused into the blood, and Si in the blood was mainly excreted in the form of soluble silicate by the kidney [59-61]. In particular, implants placed in bony sites exhibited a slower dissolution rate of Si than implants placed in soft tissue such as muscle [59,60]. The excretion rate of Si by the kidney was estimated to far exceed the dissolution rate of Si at the implant sites, which enables the maintenance of normal levels of Si in the body and avoids the abnormal accumulation of Si in organs [59].

Element content in goat tissues was also measured to examine whether abnormal element deposition and structural damage occurred. The results showed no obvious deposition of Ca or Si in the spinal cord, dura mater, or vertebral bone around the fusion segments except in the newly formed bone. In major organs of goats, including the heart, liver, spleen, lung, and kidney, there was no obvious accumulation of Ca and
damage or dysfunction in major organs [54, 59].

Histological observation of major organs of goats further validated the biosafety and biocompatibility of cage implantation. No obvious structural dysfunction or damage was found in the major organs of goats at 12 weeks and 26 weeks after surgery. Several experiments have validated that Si dissolved from bioactive glass and ceramics containing Si. We noticed that the calcium content in the heart, spleen, and lung of goats implanted with interbody fusion cages made of PEEK/CS composites. All the data lay a strong foundation for the clinical translation of bioactive cages made of PEEK/CS composites in spine surgery.

5. Conclusion

Overall, this study indicated that both PEEK/CS/pTa cages and PEEK/CS cages with autogenous bone grafts supported bone ingrowth and osseointegration in the fusion area and obtained stable fusion performance. Application of PEEK/CS/pTa could potentially avoid autogenous bone graft harvest and donor-site complications, while achieving similar fusion efficacy compared with PEEK/CS cages with autogenous bone graft. The results of element metabolism of Ca and Si in vivo, as well as histological observation of major organs, proved that the PEEK/CS/pTa cage and PEEK/CS cages had credible biosafety and biocompatibility.

Author contributions

Conception and study design: Tingting Tang, Jie Zhao, Zhifen Yu, Jie Wei, Kai Yuan. Data acquisition and analysis: Kai Yuan, Kai Zhang, Yiqi Yang, Xiyuan Feng, Jie Zhou, Jingtian Mei, Hanjun Li. Drafting the manuscript: Kai Yuan and Kai Zhang. Critical revision of the manuscript for intellectual content: Tingting Tang, Jie Zhao, and Zhifen Yu. All authors agreed to this submission. Kai Yuan and Kai Zhang contributed equally to this study.

Declaration of competing interest

The authors report no conflicts of interest related to this work.

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