Fluoride-coated high-purity magnesium cage promotes bone fusion in goat models

Luchao Yu1,2#, Yu Sun3#, Mingfei Wang2, Yingshan Wu1, Xiaonong Zhang3, Jianguang Xu1

1Department of Orthopedic Surgery, Shanghai Jiao Tong University Affiliated Sixth People's Hospital, Shanghai, China; 2Department of Orthopedic Surgery, Putuo Hospital, Shanghai University of Traditional Chinese Medicine, Shanghai, China; 3State Key Laboratory of Metal Matrix Composites, School of Materials Science and Engineering, Shanghai Jiao Tong University, Shanghai, China

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*These authors contributed equally to this work.

Background: Cervical fusion devices made by polyether ether ketone (PEEK) cause concomitant effects which decompress the spinal cord and nerve roots. Magnesium has good biocompatibility and bioactivity as a biodegradable orthopedic implant material; however, its fusion rate is low. In this paper, we aimed to improve interbody fusion rate of high-purity magnesium (HP-Mg) by coating it with fluoride.

Methods: Fluoride-coated HP-Mg (F-HP-Mg) cages were prepared, and HP-Mg cages served as controls. We tested hydrogen release in phosphate-buffered saline (PBS) and weight loss in chromic acid. Anterior cervical discectomy and bone graft fusion (ACDF) was performed at the C2-C3 segment on goats with F-HP-Mg and HP-Mg cages to evaluate fusion score.

Results: Hydrogen release of F-HP-Mg cages was significantly lower than that of HP-Mg cages. Weight was significantly decreased in both types of cages after rinsing with chromic acid, while F-HP-Mg cages were more resistant to corrosion compared to HP-Mg cages. There were no significant differences in disc space height (DSH) and remaining cage volume between the two groups in computed tomography (CT) images of goat cervical spine, while cavities were found at postoperative 12 weeks and confirmed by histological staining. No complications were found, while serum aspartate aminotransferase (AST) level was significantly higher in the HP-Mg group compared to the F-HP-Mg group. Fusion rate at 24 weeks after ACDF was significantly higher with F-HP-Mg cages.

Conclusions: The use of F-HP-Mg improved histological fusion in the cervical intervertebral space of goats compared to HP-Mg and showed good biosafety.

Keywords: Fluoride; high-purity magnesium (HP-Mg); bone fusion

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standard” for clinical interbody fusion; however, common complications of donor site, such as pain, bleeding, fracture, and nerve injury have limited widespread use of the iliac crest as an intervertebral bone graft material. The cervical fusion device made by polyether ether ketone (PEEK), which is widely used in clinical practice, can cause a strong non-specific inflammatory response in the surrounding bone tissue, triggering fibroblast and macrophage infiltration as well as bone resorption and osteolysis, mimicking the local reaction caused by ultra-high molecular weight polyethylene powder after joint replacement.

Magnesium (Mg) has attracted attention in biodegradable orthopedic implant material research, displaying characteristics of good biocompatibility, bioactivity, and desirable elastic modulus similar to bone (3). Success has been reported in high-purity Mg (HP-Mg) (99.99 wt.%) screw in in vivo studies with femoral intracondylar fractured rabbit models (4) and HP-Mg interference screws can promote fibrocartilaginous entheses regeneration in the anterior cruciate ligament reconstruction rabbit model (5). However, the results of in vivo ACDF experiments on animal models of Mg-based cervical cages have been unsatisfactory (6,7). Potential reasons include that co-implantation of titanium and Mg accelerates degradation of Mg cages, and hydrogen and excessively increasing pH value produced by rapid Mg corrosion negatively influence surrounding tissues healing. Recently, Guo et al. claimed that they achieved histological fusion of HP-Mg cages in a goat model, although total fusion area was less than 30% (8).

Coating is an effective way to regulate degradation rate of Mg fusion cages (9). Some studies have suggested that fluoride-coated Mg alloy could improve corrosion resistance and promote osteogenic differentiation (10,11). In this study, we tried to improve interbody fusion of HP-Mg cages by coating fluoride on HP-Mg (F-HP-Mg). We present the following article in accordance with the ARRIVE reporting checklist (available at https://atm.amegroups.com/article/view/10.21037/atm-22-2098/rc).

Methods

Sample and materials

The HP-Mg cages were built with 99.982 wt.% Mg (Suzhou Origin Medical Technology, Jiangsu, China), 0.0178 wt.% silicon (Si), <0.001 wt.% iron (Fe), and <0.001 wt.% aluminum (Al) in the State Key Laboratory of Metal Matrix Composites of Shanghai Jiao Tong University. Cages were designed and fabricated for ACDF and possessed dimension parameters similar to PEEK cage (Medtronic Cornerstone-SR PEEK; Medtronic, Parkway Minneapolis, MN, USA), which was modified with specification of 14 mm × 11 mm × 4 mm and a 4-degree wedge angle.

The F-HP-Mg cages (mainly composed of MgO and MgF₂) were made through immersion of HP-Mg cages in a hydrofluoric acid solution (9). Untreated HP-Mg cages of the same size were used as controls in goat models. An Mg sheet (size 12 mm × 10 mm × 1 mm) was prepared for in vitro testing.

Surface modification

The fluoridation process was operated in fume hood hydrofluoric acid. The Mg sheet samples that had been polished and ultrasonically cleaned and dried were removed and placed into a plastic beaker reaction vessel with a plastic agitator at the bottom and a suspended yarn net. The samples were placed into each beaker on the yarn net, and then 250 mL 40% hydrofluoric acid reagent was poured into the beaker. It was observed that all Mg samples were immersed in liquid. The beaker was then sealed with plastic wrap and the opening was secured with a rubber band. The beaker was placed on the magnetic stirrer at low speed, with the sample fully immersed in the acid flow at room temperature for 96 hours. Finally, all samples were rinsed in ethanol and dried. The untreated pure Mg group was used as the control.

Immersion test

The method of the immersion test was employed as described in our previous study, which was in accordance to ASTM-G31-72 (12). The F-HP-Mg and HP-Mg sheets were incubated in PBS (pH =7.40, 1 cm²/20 mL) separately at 37 °C for 2 weeks, with 3 samples in each group. The pH value of PBS was measured across time by pH meter (FE20; Mettler Toledo, Columbus, OH, USA). After 14 days, 1 side of the sheets was ultrasonically cleaned with 100% ethanol, and the other side was rinsed with 180 g/L chromic acid solution and distilled water, air dried, and weighed.

Animals and surgery

Experiments were performed under a project license (No. SYXK2011-0128) granted by ethics board of Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, in
compliance with institutional guidelines for the care and use of animals. A protocol was prepared before the study without registration. Goats have similar cervical spines to human, especially at the C2-C3 level, which makes them ideal experimental models to test cervical spine fusion (13,14). A total of 10 healthy 1.5-year-old goats weighing 38.78±3.72 kg (range, 33 to 46 kg) were purchased from Shanghai Jiagan Biotechnology Co., Ltd. (Shanghai, China) and given adequate access to sterilized water and food. All animals were randomly assigned to either the experimental or control group for assessment at post-operative 4, 12, and 24 weeks.

Goats were anesthetized with 2.5% pentobarbital sodium, followed by ACDF surgery at C2-C3 segment, which was implanted with a HP-Mg or F-HP-Mg cage, a titanium plate, and fixed with screws. Cage holes were filled with autogenous bone from the anterior spinous process of cervical vertebra.

**Radiographical analysis**

Anteroposterior and lateral radiographs were performed between post-surgery and sacrifice to determine implant location, gas accumulation, disc space height (DSH), and implant settlement. The DSH is the distance between upper and lower anterior edge of the vertebral body fixed by screws of the titanium plate.

Micro-CT scan and three-dimensional (3D) reconstruction were performed on sacrificed specimens, and quantitative analysis of the fusion device was carried out. Interbody fusion could be scored with CT images (15), as follows: level 0, no new bone or even vertebral endplate destruction; level 1, new bone formation but not continuous between C2/3; and levels 2 and 3, continuous bridging new bone and comprises <30% and >30% of fusion area, respectively.

**Histological analysis**

Sacrificed goat vertebrae were fixed in paraformaldehyde, dehydrated in acetone, embedded in methyl methacrylate, cut into 300 µm sections and ground to 20 µm. Staining was performed on 4 sections with hematoxylin-eosin (HE), toluidine blue, Van Gieson, and Masson to determine osteogenesis and inflammation.

**Statistical analysis**

No data was excluded. Data were presented as mean ± standard deviation (SD). A two-tailed t-test was used to compare CT fusion scores between the HP-Mg cage group and F-HP-Mg cage group. The two-way analysis of variance (ANOVA) was used to compare weight of cages, Mg concentrations, alanine aminotransferase (ALT), aspartate aminotransferase (AST), and creatinine (CREA). Statistical analysis was conducted with the software SPSS 20 (IBM Corp., Armonk, NY, USA), and P<0.05 was statistically significant.

**Results**

**In vitro test**

**Gross characteristics**

The MgF$_2$ films formed in 40% hydrofluoric acid after 96 hours appeared dark black. The MgF$_2$ layer did not change macro sizes of samples (Figure 1A). No significant difference was detected in weight between the two groups.

**Hydrogen release**

Hydrogen release in the buffer was measured for 336 hours. With the prolongation of immersing time, hydrogen (H$_2$) release was increased from both HP-Mg and F-HP-Mg cages (Figure 1B). The H$_2$ concentration was significantly higher in HP-Mg cages compared to F-HP-Mg cages.

**Weight loss**

Weight of HP-Mg cages and F-HP-Mg cages was similar before the weight loss experiment. After rinsing with 180 g/L chromic acid, both the HP-Mg cages and F-HP-Mg cages lost a significant volume of weight (Figure 1C). The F-HP-Mg cages were significantly heavier than the HP-Mg cages after chromic acid, indicating that the former were more resistant to corrosion than the latter.

**In vivo test**

**Radiological findings**

The interbody fusion was evaluated with 64-slice-spiral CT. The DSH measurement is shown in Figure 2A. At 4 weeks postoperatively, there was no significant difference in DSH, but a significant difference could be observed in CT images between the two groups, with a small amount of gas observed in the anterior tissue of the cervical spine in HP-Mg cages (Figure 2B). We found that HP-Mg cages caused damage to the upper and lower endplates and surrounding bone, forming cavities at 12 weeks postoperatively. The
cavity was further enlarged at 24 weeks, and the autogenous bone filling in the middle hole of the cage was completely lost (Figure 2C).

No significant DSH difference and implant settlement was found in HP-Mg cages and F-HP-Mg cages at postoperative 4, 12, and 24 weeks (Figure 2D). The remaining volume of both kinds of cages was significantly lower after 24 weeks implantation, while not significantly different between HP-Mg and F-HP-Mg cages (Figure 2D).

Histological results
Bone fusion was observed in F-HP-Mg cages with a continuous bone found between endplates (Figure 3A-3E). However, osteolytic phenomenon with destroyed bone tissues was found in HP-Mg cages.

Mg concentrations, ALT, AST, and CREA in serum
Serum Mg²⁺ concentration did not markedly change in both groups (Figure 4A). No complications related to hypermagnesemia were found. Serum ALT level was similar between the 2 goat groups (Figure 4B), while AST level was statistically higher in the HP-Mg group compared to the F-HP-Mg group at postoperative weeks 12 and 24 (Figure 4C). Serum CREA was not statistically different (Figure 4D).

Fusion results
During the experiment, CT fusion score of F-HP-Mg cages increased over time, but the score of HP-Mg cage decreased at weeks 12 and 24. The fusion score of segments with F-HP-Mg cages was remarkably higher than HP-Mg cages at 24 weeks, with HP-Mg as (0.2±0.45) and F-HP-Mg as (2.8±0.45) (P<0.01) (Figure 5A). After 24 weeks, new bone tissue between endplates in graft segments was found in CT 3D reconstruction images (Figure 5B), while no fusion with a cage was observed (Figure 5C).

Discussion
Guo et al. reported the use of an HP-Mg cage to perform
Figure 2 Radiological characterization of HP-Mg cages and F-HP-Mg cages in goat. (A) Measurement of DSH. (B) CT scan images of goats C2-C3 spine in the two groups after 4 weeks of operation. (C) CT images of goat cervical spine in the two groups after 4, 12, and 24 weeks of operation. (D) DSH and cages volume right after 0 and 24 weeks after operation in the two groups. N=5. Data are presented as mean ± SEM. ****, P<0.0001. HP-Mg, high-purity magnesium; F-HP-Mg, fluoride-coated HP-Mg; DSH, disc space height; CT, computed tomography; SEM, standard error of the mean.
Figure 3 Histological staining of goat cervical spine after 24 weeks implantation of HP-Mg cages and F-HP-Mg cages. HE staining (A), Masson staining (B), Van Gieson staining (C), toluidine blue staining (D), and CT scan (E) of goat spine in the two groups. Magnification in A-D: ×10. HP-Mg, high-purity magnesium; F-HP-Mg, fluoride-coated HP-Mg; HE, hematoxylin and eosin; CT, computed tomography.

Figure 4 Safety evaluation of HP-Mg cages and F-HP-Mg cages in goats. Serum magnesium (A), ALT (B), AST (C), and CREA (D) in the two groups. N=5. Data are presented as mean ± SEM. *, P<0.05. HP-Mg, high-purity magnesium; F-HP-Mg, fluoride-coated HP-Mg; ALT, alanine transaminase; AST, aspartate aminotransferase; CREA, creatinine; SEM, standard error of the mean.
ACDF surgery on a goat model, and achieved the first successful interbody fusion, indicating the possibility of Mg-based cage application in ACDF (8). However, this study also reported that the total area of interbody fusion was less than 30%. There was a 300–400 µm gap between the cage and the bone tissue, which was filled by hyperplastic fibrous tissue. The bone was not tightly bound to the cage interface.

The results of this experiment revealed that in the experimental group using an HP-Mg cage, because the cage caused osteolysis to the surrounding bone tissue, the CT sagittal view showed the formation of a cavity around the cage, and the pathological section showed that there was a space between the cage and the bone tissue. With infiltration of fibrous tissue, the effect of vertebral fusion was not ideal. However, after the surface modification of HP-Mg with hydrofluoric acid, the degradation rate of HP-Mg slowed down, continuous bone tissue formation was seen on a CT sagittal view, the gap between the cage and the bone tissue was reduced, and new trabecular bone formation was seen surrounding the cage.

For the cage material, we chose to use HP-Mg (99.98 wt.%) for 2 reasons: first, Mg-based implants had been revealed as osteoinductive in in vivo studies in unstressed bones, leading to successful fusion. The mechanism may be related to the release of Mg$^{2+}$ from Mg-based implants in neuronal calcitonin gene-related polypeptide (CGRP)-mediated osteogenic differentiation, which plays an important role in induction of osteogenesis (16). Secondly, compared with Mg alloys, HP-Mg has better corrosion resistance. Preliminary research shows that the higher the purity, the stronger the corrosion resistance, and the slower the degradation rate. This experiment required Mg-based cages in the vertebral column which takes about 6 months, so the purity of HP-Mg needs to be improved as much as possible to obtain sufficient support time.

It has been shown that Mg-based materials immersed in hydrofluoric acid can form a fluoride coating on the metal surface, which can improve the corrosion resistance and mechanical properties of the material. This modification process is simple and cost-effective, allowing for widespread application in spinal fusion surgery.
surface (10), which can effectively reduce the degradation rate of Mg, thereby slowing the release rate of Mg\(^{2+}\) and increased deposition of localized calcium and phosphate. The results of this study showed that fluoride-coated Mg alloy had good osteogenic activity and biocompatibility, and the fluoride coating significantly upregulated expressions of type I collagen and bone morphogenetic protein 2 (BMP-2). Therefore, in this experiment, we modified the surface of the HP-Mg cage in hydrofluoric acid to form a MgF\(_2\) coating, and the thickness of the MgF\(_2\) coating depended on the reaction time of the MgF\(_2\) cage in hydrofluoric acid, which we continued for 96 hours.

The results of this experiment showed that after the HP-Mg cage was implanted into the goat cervical spine, galvanic corrosion occurred with the fixed anterior cervical vertebrae titanium plate, which accelerated degradation of the HP-Mg cage and damaged the upper and lower vertebral body endplates. As a result, the fusion failed. Despite this, the prevailing view is that locally high Mg ion concentrations do not produce significant cytotoxicity and are safe for topical application as a degradable material (17). However, the accelerated corrosion rate of Mg can lead to loss of mechanical properties of orthopedic implants and failure (18). If Mg cages are coated with MgF\(_2\), the degradation rate can be reduced, galvanic corrosion can be isolated, and osteogenic activity and biocompatibility can be improved.

In this study, serological tests were performed at 4, 12, and 24 weeks after the operation. Serum levels of Mg\(^{2+}\), AST, ALT, and CREA were examined, and no obvious liver and kidney function damage was found. In addition, studies involving pathological sectioning on animal organs found no obvious lesions, which supports the good biosafety of HP-Mg fusion in vivo. Excessive Mg ions can be excreted in urine and not absorbed by the body.

The results of this study suggest that the application of the HP-Mg interbody fusion cage with MgF\(_2\) coating can achieve cervical vertebral body fusion, and thus may be used in ACDF surgery. However, more in vivo and in vitro experiments and longer observation times are needed. In addition, the fusion effect of the HP-Mg cage group in our study was not satisfactory, the rapid degradation of Mg\(^{2+}\) would destroy the bone of the upper and lower vertebral body endplates and form a cavity in the intervertebral space, although it was possible to form a bone bridge connection at the anterior and posterior borders of the vertebral body. Partial fusion was achieved, which was different from the related research results of using the same material to make cages (8), and might be the result of difference of surgical operation, animal breeding environment, and other factors.

The in vivo experiment of cervical degradable cage is time consuming, requires complicated operation, and has big individual differences. Previous researchers used cages with β-calcium triphosphate as the core and polyactic acid as the shell to conduct a phase III clinical trial of cervical interbody fusion (19). After an average follow-up of 27 months, the fusion rate was 96% (26/27). However, in another study, the results of cages with the same specifications were disappointing. Although the experiment planned to include 50 participants, it had to terminate early due to the high dislocation and complication rates after recruiting only 33 participants (20). Therefore, animal experiments should be carried out to evaluate safety of degradable cages before the implementation of clinical trials.

Conclusions

The HP-Mg cage with fluoride coating can successfully achieve histological fusion in the cervical intervertebral space of goat: that is, the new bone is connected between the upper and lower vertebral bodies through the middle hole of the cage. It is worth noting that the fusion area of the HP-Mg cage with fluoride coating was greater than 30%, the fusion results of the fluoride coated group at 24 weeks were significantly better than that of the uncoated group, and the degradation rate was also significantly lower than that of the uncoated group. The F-HP-Mg cage can play a stable supporting role in the early post-implantation period, and then degrade steadily during the observation period. The cage has sufficient mechanical support for intervertebral applications. In addition, the F-HP-Mg cage has good biocompatibility, no adverse reactions for vital organs. More studies are needed to evaluate the long-term fusion effect and degradation behavior of this cage, which is critical for potential use in ACDF in the future.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. Experiments were performed under a project license (No. SYXK2011-0128) granted by ethics board of Shanghai Jiao Tong University Affiliated Sixth People's Hospital, in compliance with institutional guidelines for the care and use of animals.

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