Research Article

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Safety study of malapposition of the bio-corrodible nitrided iron stent in vivo

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Abstract: To evaluate the safety of stent malapposition of corrodible nitride iron stent as biodegradable cardiovascular implants, a total of 108 stents were implanted into the abdominal aortas, iliac arteries, and iliac artery bifurcations of 36 New Zealand white rabbits separately. Each rabbit was implanted with three stents. After a follow-up period of 3 months, no thrombus and embolism were found in local and downstream vessels. And no other adverse events occurred either. Stent strut covered by endothelial layer started to show signs of degradation, while struts exposed to bifurcated blood flow covered by a layer of tissue and no rust particle was found on the surface. Also, there were no traces of thrombosis and traces of excess inflammation. The authors conclude that the risk brought by stent malapposition in less than 9 months is acceptable.

Keywords: bio-absorb iron stent, corrosion, iron overload, stent malapposition, thrombosis

1 Introduction

Coronary stent implantation is the most common and effective treatment means for coronary artery disease, which works by opening the narrowed arteries thus improving blood flow [1–3]. Coronary stents have undergone enormous development since their first introduction in 1987 [4]. Bio-absorbable coronary stents have been widely acknowledged to be the 4th revolution in vascular intervention, following Percutaneous Transluminal Coronary Angiography, Bare Metal Stent, and Drug-Eluting Stent (DES), getting over the adverse event of late stent thrombosis occurred in the third generation DES, providing mechanical support and anti-restenosis properties in short term and then metabolized physiologically, thus avoiding the permanent metallic enclosing of the treated coronary artery. Therefore, bio-absorbable coronary stents have extensive research significance.

Iron-based stents are a promising candidate for bio-absorbable stents, due to their good biocompatibility and outstanding mechanical performances like 316L stainless steel [5–10]. According to reported studies [8–10], it has been demonstrated that safety of iron stents implantation without the significant obstruction of the stented vessel as a result of inflammation, thrombotic events, or neointimal proliferation, but a faster degradation rate is demanded theoretically; an ideal bio-absorbable iron stent should provide enough support to ensure that preventing vascular retraction in the initial stage after implantation, after 3–6 months of action, most of the loss of support caused by the corrosion process is beneficial to tissue regeneration. However, some studies demonstrated that the corrosion rate of the pure iron stents was slower than required. For the past few years, further researches focused on modifying the composition, surface, and microstructure of pure iron stents to increase their corrosion rate [11–14]. Here vacuum plasma nitriding was employed to enhance its strength and corrosion rate by incorporating nitrogen into the iron matrix to form dispersive iron nitride precipitation phase [9,15]. But along with the increase of corrosion rate of the iron stent, the release of iron ions will also increase during the short term after implantation, especially the stage when the struts were not completely covered by endothelium cells. And too much iron in the body is at risk [16,17]. Some studies show that iron overload has
a direct relationship with heart and liver disease, diabetes, and other relevant diseases [18,19]. Conversely, after the stent strut was completely covered by endothelium cells, the human body has a series of iron control mechanisms (an iron regulation mechanism) to control the systematic or local iron overload risk [20]. Hence, it is very necessary to evaluate the iron overload risk in the short term after the implantation of the nitrided iron stent.

Another safety concern about bio-corrodi ble stent is late thrombogenicity or embolism of corrosion products when exposed to blood flow for a long time, probably occurs in stent malapposition, which has not been reported yet. The stent malapposition is usually caused by inadequate expansion or plaque existed at the vascular wall [21]. And this phenomenon will eventually cause delayed endothelialization, stent thrombosis complications, and other diseases [22–25]. In vitro studies cannot fully simulate the real physical and chemical environment in vivo. Therefore, appropriate animal models are necessary to further study the implantation of bio-corrosive and degradable iron nitride stents.

Given these considerations, this study adopted implanting the bio-absorbable stent in the iliac artery bifurcation position of the New Zealand rabbit, to simulate the stent implantation state in a vessel. And through monitoring the thrombus and embolism of supported vessels and downstream vessels, the topography of nitrided iron stent after implantation for 1, 3, and 9 months separately, the corrosion rate and the late risk brought by the implanted stents were evaluated.

2 Materials and methods

2.1 Stents

The degradable stents, with a pattern similar to a commercially available permanent coronary stent of a nominal inflated outer diameter of 3.0 mm and a nominal length of 18 mm, were laser cut from iron tubes (>99.8% iron, made by Biotyx Medical (Shenzhen) Co., Ltd). Vacuum plasma nitriding technique is added into the manufacturing process after laser cutting using a self-designed vacuum nitriding furnace. Vacuum plasma nitriding in an engineering field is originally a surface modification technology. However, when applied to small coronary stents, it becomes a bulk alloying method since the original strut thickness is 100 µm or less. As a result of dispersion strengthening and solution strengthening, the hardness, radial strength, and stiffness of the nitrided iron stent were significantly increased. After the treating process of vacuum nitriding, the nitrided iron stents were polished to achieve a strut thickness of 70 ± 5 µm. The stent weighs 10–15 mg. The stents were sterilized with ethylene oxide and stored in vacuumed packages before implantation operation.

2.2 Animals

The study was conducted with the approval of the local governmental authorities and adhered to the Guide for the care and use of laboratory animals (NIH publication 85–23, 1985). Thirty six adult New Zealand white rabbits (mean weight 2.2 kg, range 2.0–2.5 kg) were purchased from Guangdong province medical animal test center (China).

2.3 Procedure

Anesthesia was induced by pentobarbitone (30 mg/kg) intravenously. Antibiotic prophylaxis was administered intramuscularly. The right carotid artery was surgically exposed and a 5 French sheath was introduced over a 0.018-inch guidewire after being immersed in the heparinized saline for 0.5 h. Quantitative angiography of the descending aorta was performed to determine the luminal diameter of the descending aorta at the site of the implantation (5 French Berman angiography catheter, Arrow, Reading, Pennsylvania, USA). Balloon catheters with a diameter of 3 mm (Savy, Cordis, Miami, Florida, USA) were chosen to achieve a balloon to vessel ratio of about 1.2 ± 0.1. The degradable iron stent was manually crimped to the balloon catheter. Under fluoroscopic control, the stent was introduced and positioned at the predetermined implantation site. The stents were implanted with 8 atm for 10 s. After removal of the catheter and sheath, the carotid artery was ligated and the incision of the skin was closed with sutures. The animals were returned to the recovery area. Figure 1 shows the implantation position of a testing rabbit sacrificed after 3 months of implantation.

2.4 Follow-up study

All animals were followed up and results were documented throughout the study. Vessel morphology was
observed after 1 month \((n = 15)\), 3 months \((n = 15)\), and 9 months \((n = 6)\).

### 2.5 Morphological investigation

After angiography for the downstream vessels (using an x-ray machine with the assistant of diatrizoate), 1,000 U of heparin were injected into the rabbits, and then the animals were sacrificed by an intravenous injection of a lethal dose of pentobarbital. Immediately after death, the infra diaphragmatic descending aorta was exposed and the stented segment with stents was identified and harvested. The segments were placed in 3.5% neutral buffered formalin and kept for histopathological investigation. The stents were cut longitudinally and the lumen of the stented vessel was evaluated for signs of adherent thrombi or overt neointimal obstruction of the vessel. One specimen of each study group was viewed under a scanning electron microscope after critical point drying and gold sputtering to assess the endothelialization of the stent struts.

### 3 Results

The SEM and EDS of the nitride iron material surface were shown in Figure S1 (Supporting Information). After nitride, the atom ratio of Nitrogen is about 5.27%, with a weight percent of 1.37%. The degradable iron stents were implanted in the predetermined segment of the iliac arteries in 36 rabbits without major complications. No
abnormalities on macroscopic inspection of the implantation site were observed. During the 3 months of the follow-up period, no cases of animal death or other obvious symptoms of pathological changes occurred. After the animals were put to dissection, no thrombosis or angiomphraxis is observed in the downstream vessels. The segment of the iliac arteries was separated and cut down for the next study of degradation. There was also no blockage or thrombosis observed at the implant position. Figure 2 shows the rabbit iliac arteries with degradable iron stent after 1 month of implantation. As seen from Figure 2a, the strut in the orifice of vascular branch position was not covered by endothelial cells, and the other struts contacted with the vascular wall were covered by endothelial cells completely. In addition, the nitrided iron stent was corroded and there was no obvious corrosion product formed on the surface of the strut. Compared with the strut contacted with the vascular wall (position C in Figure 2b) and the transitional strut (incompletely covered by endothelial cells, position A in Figure 2b), the strut in the branch position is relatively smooth. In other words, the corrosion level in position B is relatively slight. SEM image gives information that the strut in the vascular branch position is covered by a layer of uniform complete fibrous tissue membrane, which effectively prevented the direct contact between the strut and the blood flow. In short, it increased safety when there was a stent malapposition state occurred.

Figure 3 shows rabbit iliac artery with degradable iron stent after 3 months implantation. As seen from Figure 3, the corrosion level increased compared with the case of 1 month. Iron transportation was observed from the implantation site towards the reticuloendothelial system after 1 month, and the level became more serious after 3 months. There are obvious black corrosion products stacked on the surface of the struts which are located at the transitional position. However, the strut which is located at the vascular branch position is still shining. No corrosion products could be detected on the strut surface. Figure 3c and d show the SEM images of rabbit iliac artery with degradable iron stent after 3 months of implantation. We can see that the fiber tissue membranes covered on the strut located at the branch position became thicker than the ones after 1 month of implantation. The thickness of the fiber tissue membrane is about several microns. The fiber tissue membrane shown in Figure 3d is deliberately broken for the aim of evaluating the thickness change.

Figure 3: Rabbit iliac artery with degradable iron stent after 3 months implantation. (b) is the scale-up diagram in (a); (d) is the scale-up diagram in (c).
Figure 4 shows the strut located at the vascular branch position with a degradable iron stent after 9 months of implantation. As shown, the surface of the strut is relatively smooth, and also no obvious corrosion product was found. To evaluate the risk of thrombosis brought by the stent malposition, the rabbits’ lower limbs vessels’ X-ray morphology was performed before the rabbit was sacrificed at 3 and 9 months separately (Figure 5). As seen, there were no obstacles found and there was no abnormal blood flow of the downstream vessels.

4 Discussions

Stent malapposition refers to improper contact between the struts and vascular walls [26]. And it includes two kinds: early acute and late chronic malapposition [27]. Early acute stent malapposition cases are mainly caused by the inexperienced operation, partially caused by some vascular diseases like calcified lesions [28]. Late stent malapposition is caused by the positive reconstruction of vessels or by the chronic rebound of the stent. Research has shown that late stent malapposition may lead to the formation of stent thrombosis [29,30]. Once the iron stent was implanted, it corrodes gradually; when there the stent malapposition happens, the struts’ endothelialization process would be delayed. At the same time, the corroded strut may produce degradation product particles. And the particles may flow away and even enter into the downstream small vessels. This hazard could cause complications like thrombosis and obstruction. However, the follow-up animal study of 1 and 3 months found that there are no attached degradation products found on the surface of the strut which is located at the vascular branch position. In addition, there was a layer of fiber tissue membranes covered on the strut. This kind of fiber layer tissue membrane isolated the stent strut and the flowing blood [31]. The strut at the branch position remained shining for 3 months. Corrosion mechanism is a major difference between the degradation iron metal stent and the permanent stent, thus the extensive research attention in the
The degradation process of the nitrided iron stent in vivo could be explained as follows: First, the iron was oxidized to metal ions following equation (1). The electrons from the anodic reaction were consumed by a corresponding cathodic reaction and the reduction of oxygen dissolved in water, following equation (2). These reactions occurred randomly over the entire surface where a difference existed, at grain boundaries or the interface between different phases.

Anodic reaction:

$$\text{Fe} \rightarrow \text{Fe}^{2+} + 2\text{e}.$$  \hspace{1cm} (1)

Cathodic reaction:

$$\frac{1}{2}\text{O}_2 + \text{H}_2\text{O} + 2\text{e} \rightarrow 2\text{OH}^-.$$  \hspace{1cm} (2)

Then, the released metal ions reacted with the hydroxyl ion (OH⁻) released from the cathodic reaction to form insoluble hydroxides (hydrous metal oxides) according to equations (3) and (4).

$$2\text{Fe}^{2+} + 4\text{OH}^- \rightarrow \text{Fe(OH)}_2 \text{ or } 2\text{FeO} \cdot \text{H}_2\text{O},$$  \hspace{1cm} (3)

$$\text{Fe}_3\text{Fe(OH)}_2 + \text{O}_2 \rightarrow 4\text{FeOOH} + 2\text{H}_2\text{O}.$$  \hspace{1cm} (4)

However, the corrosion conditions are differences between the strut at the branch position and the position adjacent to the vascular wall. As for the strut at the branch position, the membrane covered on the strut is thinner; the iron ion released from the strut is easy to pass through the membrane and taken away by the blood flow. So, there was no corrosion product stacked on the surface of strut. Conversely, as for the strut at the position adjacent to vascular wall, 3 months after implantation it was revealed that it was covered by a relatively thicker and denser layer mainly composed of endothelium. Hence, the iron ion released from the strut is hard to pass through and chemical reactions (3) and (4) occurred. To some extent, this could explain the result we see from Figure 3. To further distinguish the difference of degradation conditions, SEM analysis was performed to the surface topography of strut at different implant positions after cleaning the corrosion product. As seen in Figure 6, the corrosion depth of the strut adjacent to the vascular wall is slightly deeper than the one at the branch position (6 months after implantation). The corrosion extent of the strut at the transitional position looks more serious (Figure 3b). This phenomenon may cause by the difference in oxygen diffusion rate. In another word, the oxygen concentration of the tissue fluid around the strut at the two positions is different. In addition, the difference in biochemical environment between the two positions also (galvanic corrosion) accelerated the corrosion rate at the transitional position.

Figure 7 shows the corrosion speed between different existed states of an iron element. In a real degradation situation in vivo environment, corrosion products are always produced on the surface of an iron stent. The amount of rust is determined by the difference between V1 and V2, while V3 is not so important as V1 and V2 since rust will stay much longer than intact iron during the lifespan of the iron stent. And the final degradation time of the stent is mainly determined by V2 since V2 is much lower than V1. V2 becomes a bottleneck procedure in all degradation activities of the iron stent. The
exception occurred before the iron stent is covered by tissue when the bloodstream flows on the surface of the iron stent, which will expedite the V2 extremely to the level faster than V1. So the struts of the stent at the branch position keep free rust on.

Currently adopted in vivo experiments are using bare nitride iron stents, without being coated with drug-eluting coatings. After being covered with coatings, the corrosion rate should be slower, especially at the initial contacting stage, which ultimately causes less corrosion products accumulation and thus lead to a safe performance at the implantable site. However, the endothelialization process might be affected by the released anti-hyperplasia drugs, as drug-releasing would possibly cause delayed healing of endothelium cell layer [38–41]. Future study will focus on this consideration. Overall, the data presented in this work properly supported the safety of the malapposition of the bio-corrodible nitrided iron stent.

5 Conclusion

After 3 months of implantation of the nitrided iron stents, no thromboembolic complications and no adverse events occurred; iron stent strut covered by endothelial layer started to show signs of degradation without evidence of stent particle embolization, thrombosis traces, and traces of excess inflammation. This animal study together with corrosion mechanism calculation concludes that the risk brought by stent malapposition within 9 months is acceptable.

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