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Proof of concept of FOLDAVALVE, a novel 14 Fr totally repositionable and retrievable transcatheter aortic valve

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Abstract

Aims: Transcatheter aortic valve implantation (TAVI) is an emerging field with technological challenges. One major challenge is to minimise delivery catheter size to reduce vascular trauma, while maintaining features such as repositionability and retrievability. This study was designed as proof of concept of FOLDAVALVE, using a short-term ovine model.

Methods and results: FOLDAVALVE is a fully repositionable and retrievable transcatheter aortic valve with a 14 Fr delivery system whose leaflets are excluded from the stent during crimping. A short-term ovine model (n=3) was used for transfemoral implantations at the Institut Mutualiste Montsouris in Paris. There was a smooth transition of the valve over the aortic arch followed by seamless delivery into the native ovine aortic position. Implantation was followed by repositioning, resheathing and retrieval, then implantation of a fresh prosthesis in the same animal. Transoesophageal echocardiography and fluoroscopy were used to monitor the delivery and implantation processes and to demonstrate valve functionality. Animal sacrifice and direct visualisation after explant confirmed excellent final position.

Conclusions: FOLDAVALVE was successfully implanted in the aortic position of an ovine model. FOLDAVALVE is a promising technology which has been shown to be feasible in this preclinical TAVI study.

KEYWORDS
- aortic stenosis
- crimping
- low profile
- self-expandable nitinol stent
- transcatheter aortic valve implantation

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Abbreviations

AS    aortic stenosis
LV    left ventricle
PARTNER    Placement of AoRTic TranNscathetER valve trial
SURTA VI    Surgical Replacement and Transcatheter Aortic Valve Implantation
TAVI    transcatheter aortic valve implantation
TEE    transoesophageal echocardiogram
TTE    transthoracic echocardiogram

Introduction

Transcatheter therapies for severe aortic stenosis (AS) have progressed rapidly over the past few years. Based on the success of the PARTNER trial and the CoreValve U.S. Pivotal trial, multiple valves are in use for patients at high or extreme surgical risk. Trials such as PARTNER II and SURTA VI are underway to evaluate the efficacy and risks of transcatheter aortic valve implantation (TAVI) in patients with intermediate surgical risk.

Some of the most common complications of TAVI are major and minor vascular complications, which can be due to a number of factors, including high sheath to femoral artery ratio. This can be controlled, as the sheath to femoral artery ratio is entirely dependent on the device which is being used and its sheath size. Another important factor to consider in valve design is durability. Patients with an intermediate surgical risk tend to be younger, have fewer comorbid medical issues and thus a longer life expectancy. Therefore, valve durability has become of paramount concern. Crimping has been shown to damage the microstructure of pericardial leaflets permanently, and may lead to decreased durability, although this has not been definitively shown up to this point. In lower-profile valves, increased crimping induces more damage to the leaflets than larger devices. As the trend towards lower-profile devices continues, this damage should be somehow mitigated. Finally, the ability to reposition a valve prior to final deployment and to retrieve it once it is deployed is critical due to the negative consequences of malpositioning.

FOLDA VALVE (FOLDA LLC, Rancho Santa Margarita, CA, USA) is a self-expanding transcatheter aortic valve (TAV) which is deliverable through the femoral artery (Figure 1, Figure 2). Due to its unique delivery system, INGENUITY (FOLDA LLC) (Figure 3), the valve is fully repositionable and retrievable. It utilises a novel mechanism to exclude the valve leaflets from the stent crimping process. Here we describe the preclinical assessment of FOLDA VALVE in a short-term ovine model.

Methods

VALVE CHARACTERISTICS

FOLDA VALVE is composed of a self-expanding nitinol stent and three bovine pericardial leaflets. When crimped, the leaflets are folded outside of the nitinol frame and are pulled into the expanding stent using a drawstring mechanism during deployment. Formation of the trileaflet valve occurs simultaneously with the stent expansion. These unique design features allow the valve’s leaflets to be spared from damage that occurs during stent crimping and valve delivery. During delivery and positioning, the valve is fully repositionable and retrievable through the INGENUITY delivery system. The INGENUITY system allows for six degrees of freedom in valve positioning and repositioning, and can be used for a femoral, subclavian or direct aortic approach.

The radiopaque nitinol frame, in addition to markers on the delivery system, is used to guide implantation of the valve in the aortic position. Valve compression, steering and release are controlled.
by the operator to attain a satisfactory position for implantation (Figure 3). Repositioning occurs by partially compressing the stent back into the delivery system by means of force fibres which temporarily connect the stent to the delivery system. Compressing the stent to a size less than the native valve’s annulus allows for seamless repositioning of the valve in six degrees of freedom (Moving image 2). The retrieval process, which can be performed any time before the full release, is initiated by collapsing the stent with the leaflets inside by means of the force fibres. This results in complete resheathing and facilitates retrieval. Once fully resheathed, the valve is retrieved via the INGENUITY system and will not be used for reimplantation; a new valve system will be introduced to ensure the leaflet quality. At any time before full release and separation from the delivery system, the valve can be fully retrieved.

**SHORT-TERM ANIMAL MODEL**

To determine the feasibility and safety of FOLDAVALVE implantation, a short-term animal study was designed to test the capacity of the valve to be implanted, repositioned and retrieved. Three healthy domestic sheep (mean weight 45 kg) were used with a follow-up of 90 minutes post implantation. During the post-procedure monitoring, a complete transthoracic echocardiogram was obtained with haemodynamic assessment of the implanted valve. The animals were then euthanised after the 90-minute follow-up, and their hearts and aorta were harvested for gross macroscopic examination.

**FOLDAVALVE IMPLANTATION**

The animal protocols were developed in conjunction with and approved by the animal care and use committee of the Institut Mutualiste Montsouris (IMM), Paris, France. Transfemoral implantation of the FOLDAVALVE was performed by a multidisciplinary team. The procedure was undertaken in the hybrid operating room after pre-screening the animals for proper aortic root size.

Under general anaesthesia and mechanical ventilation, a 15 Fr introducer sheath was placed in one femoral artery for ease of manipulation. The INGENUITY delivery sheath was inserted into one femoral artery, while a standard 6 Fr introducer sheath was placed in the contralateral femoral artery. A 6 Fr pigtail catheter was then inserted and positioned in the aortic root. Following an aortic root angiogram, a 0.035” J-tipped guidewire was inserted and advanced into the LV after which it was exchanged for a stiff pre-shaped guidewire. FOLDAVALVE was then loaded onto the delivery system and advanced into the LV, then pulled back to the aortic root (Figure 4).

Under angiographic guidance, the valve was deployed, repositioned, properly positioned, and finally released (Figure 5, Moving image 3). After initial deployment, the valve was partially recaptured by the delivery system, repositioned and then redeployed into the native aortic valve followed by repeated assessment. The valve was then fully released (Moving image 3). In one of the cases, the fully deployed valve was recaptured and fully removed from the animal, and a new valve system was then introduced and implanted in the same animal to test FOLDAVALVE’s retrieval capacity (Moving image 4). A final evaluation was performed that consisted of angiographic, transthoracic echocardiographic (TEE) and haemodynamic evaluation of the FOLDAVALVE. Patency of the coronary arteries was assessed and prosthesis function was evaluated including valvular and paravalvular regurgitation. LV function was monitored and transvalvular gradients were measured.

**Results**

**IMPLANTATION**

FOLDAVALVES were successfully implanted in three domestic sheep under fluoroscopic and transoesophageal echocardiographic (TEE) guidance. The prostheses were easily able to traverse the aortic arch and be implanted in the aortic position (Figure 4, Figure 5). After initial implantation, the valves were partially recaptured then
repositioned and reimplanted. Retrieval, repositioning and reim-
plantation were successful in all studied cases. Procedure time var-
ied from five minutes to seven minutes for initial implantations.
Repositioning took five minutes and retrieval took two minutes.

**FLUOROSCOPIC EVALUATION**

The implanted valves were closely investigated using fluoroscopy.
Central prosthetic regurgitation was not present in any of the three 
animals. Paravalvular leak was present in two of the three cases: 
in one animal the paravalvular leak was mild by fluoroscopy, and 
in the other case paravalvular leak was mild to moderate. Three-
dimensional evaluation of the stent was also undertaken using 
fluoroscopy, and the stents were found to be well apposed with no 
evidence of stent disruption or fracture in any of the cases. Two-
dimensional measurements showed the valves were properly sized.
Finally, aortic root angiography revealed patent coronary arteries in 
all three cases (Figure 5).

**TTE EVALUATION**

In each studied case, echocardiographic evaluation of FOLDA VALVE 
indicated proper valve function and alignment with no evidence of 
perforation or pericardial effusion in both 2D and 3D. Pulse wave 
Doppler of the valves yielded no evidence of stenosis with the max-
imum instantaneous velocity of 102 cm/s and a mean velocity of 
70.2 cm/s in one case which corresponded to a maximum gradi-
ent of 4 mmHg and a mean gradient of 2 mmHg across the valve.

Paravalvular leak was again confirmed in two cases, with one case 
being mild and one being mild to moderate according to colour 
Doppler studies (Online Figure 1). There was no evidence of central 
aortic regurgitation in any of the cases. Dobutamine was infused and 
a modified apical five-chamber view demonstrated no valve migra-
tion during or after infusion (Online Figure 2).

**CLINICAL AND ANATOMICAL ASSESSMENT**

The animals were clinically and haemodynamically stable throughout 
the procedure and through follow-up. None of the animals demon-
strated evidence of heart block, significant bradycardia, atrial or ven-
tricular arrhythmias on telemetry during or after implantation. None 
of the valves migrated or embolised during implantation or post-
implantation monitoring. Macroscopic evaluation of the explanted 
hearts demonstrated that the valves were secure and in a stable posi-
tion (Figure 6). There was no evidence of traumatic injury to the aor-
tic root, left ventricular outflow tract or the thoracic aorta.

**Discussion**

Here we describe the proof of concept study of FOLDAVALVE, 
a novel self-expanding 25 mm transcatheter aortic valve with 
a 14 Fr delivery system. FOLDAVALVE is fully repositionable 
and retrievable, and spares the leaflets from the damage induced 
by crimping by means of excluding them from the stent during the 
crimping process (Figure 2, Moving image 1). The INGENUITY 
delivery system provides exceptional flexibility and control in

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**Figure 4.** FOLDAVALVE being easily passed over the aortic arch and into the left ventricular cavity immediately before staged deployment 
and positioning in the aortic root. A) The valve passes up the descending thoracic aorta. B) This valve smoothly passes over the aortic arch 
and across the native aortic valve. C) The valve is being positioned from the left ventricle into the native aortic valve.

**Figure 5.** Positioning and release of the FOLDAVALVE. FOLDAVALVE is pulled back into the aortic root and positioned properly (A). 
The valve is released from the delivery system (B & C). The valve is fully expanded and functional at the aortic valve position (D).
addition to full repositioning and retrieval capabilities. Procedural success was also shown with successful animal implantations, no short-term complications, and multimodality imaging showing good valve function.

FOLDAVALVE addresses many of the challenges that will be faced in the quest to improve upon the currently available TAVs. In this short-term animal study, there were no instances of valve embolisation, coronary obstruction or heart block. The mild to moderate degree of paravalvular leak present in one case is believed to be due to the animal prosthesis size mismatch (Online Figure 1).

Through the use of a short-term animal model it was demonstrated that the valve and its delivery system were highly functional and provided the operator with meticulous control. By using a streamlined delivery system that excludes the leaflets from stent crimping and capitalising on a design with a physiologic flow profile, FOLDAVALVE holds the promise of being a durable valve with excellent haemodynamics and, due to its low profile, a reduced rate of procedure-associated bleeding.

**Impact on daily practice**

TAVI is rapidly becoming a treatment option for more and more patients with severe AS. However, many TAVI delivery systems are bulky and induce a great deal of vascular trauma. The work presented here describes a TAV with advanced features such as repositioning and retrievability while maintaining a low-profile delivery system (14 Fr) and sparing the leaflets from crimping inside the stent. We have shown that implantation of this valve is feasible and that the functionality of the valve is good in a short-term animal model.

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**Conflict of interest statement**

A. Kheradvar is a co-founder and is associated financially and scientifically with FOLDA LLC, the parent company of FOLDAVALVE. The other authors have no conflicts of interest to declare.

**References**


**Online data supplement**

**Online Figure 1.** Colour Doppler echocardiogram of the valve in long axis.

**Online Figure 2.** 2D and 3D echocardiograms of the implanted FOLDAVALVE.

**Moving image 1.** Schematic animation showing how leaflets are turning inside the stent via a drawstring mechanism to form a trileaflet heart valve.

**Moving image 2.** The six degrees of freedom in positioning when the valve is still attached to the delivery system.

**Moving image 3.** Transfemoral delivery, staged deployment, repositioning and complete implantation of FOLDAVALVE in sheep under fluoroscopy.

**Moving image 4.** Transfemoral delivery, staged deployment, implantation, retrieval and fresh implantation of FOLDAVALVE in sheep under fluoroscopy.
Online data supplement

Online Figure 1. Colour Doppler echocardiogram of the valve in long axis with the paravalvular leak highlighted by the arrow. The leak is probably due to animal prosthesis size mismatch.

Online Figure 2. Two- and three-dimensional echocardiograms of the implanted FOLDAVALVE. A) & B) 2D echocardiograms of FOLDAVALVE in the long and short axis, respectively. The stent is well positioned and symmetrically expanded. C) 3D echocardiogram of the valve in short axis.