LIFE IS DIFFERENT

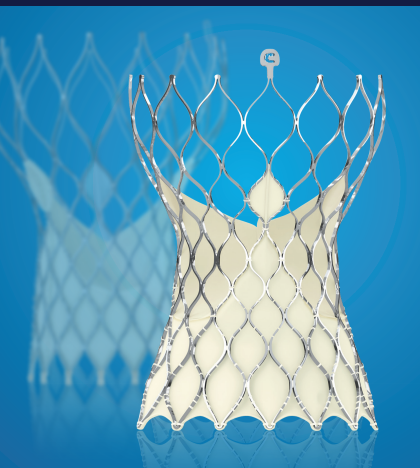


CoreValve™ Evolut™ R

Transcatheter Aortic Valve Replacement (TAVR) Platform

Medtronic

BUILT ON A PROVEN FOUNDATION

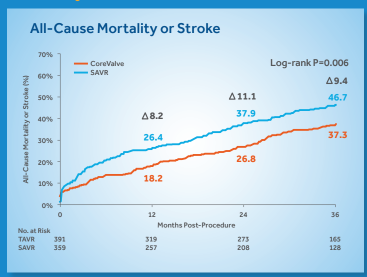


The CoreValve™ System continues to demonstrate exceptional outcomes — and we've taken what we've learned from the design of that platform and applied it to the Evolut™ R System.

Supra-annular Valve Design
Self-expanding Nitinol Frame
Porcine Pericardial Tissue
Low Delivery Profile

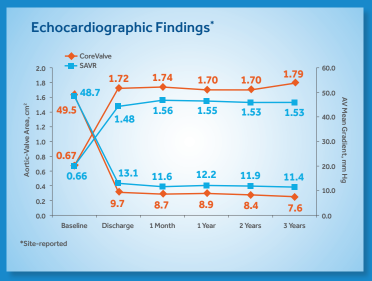
SUPERIOR LONG-TERM CLINICAL OUTCOMES

Lower Rate of Mortality or Stroke



The CoreValve™ Platform shows superior outcomes vs. surgery.¹

Unsurpassed Sustained Hemodynamic Performance



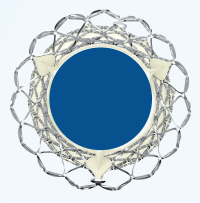
CoreValve[™] system had significantly better valve performance over SAVR at all follow-up visits (P<0.001)¹

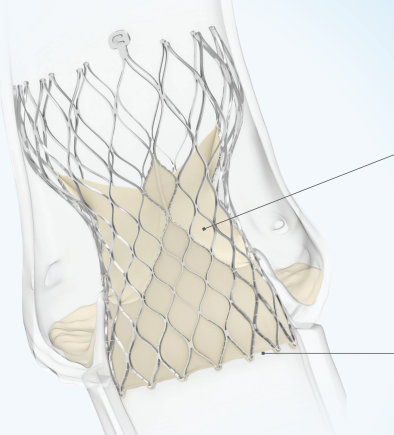
UNSURPASSED HEMODYNAMICS

Supra-annular valve design maximizes leaflet coaptation and promotes single digit gradients and large EOA's.

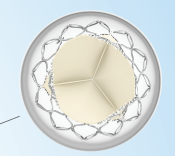
7.5 mm Hg single digit gradients

2.0 cm² Large EOA





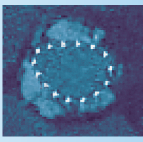






Supra-annular Valve | Optimizes coaptation in non-circular anatomy with supra-annular valve position





Annulus Conforms to the native annulus

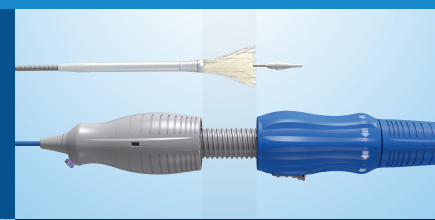
Exceptional Survival

98.8%

CONTROL DURING DEPLOYMENT

ACCURATE POSITIONING

1:1 response provides immediate feedback between the deployment knob and the movement of the capsule



Tactile Indicator ~ 2/3 Deployment Just Prior to Point of No Recapture†

RECAPTURE AND REPOSITION

EnVeo™ R provides option to recapture and reposition for accurate placement.

†Up to 80% deployment.

ACCESS MORE PATIENTS



BROADEST ANNULUS RANGE ON THE MARKET**

The only TAVR platform indicated to treat annulus up to 30 mm

17/18







29 mm Valve



30 mm

LOWEST DELIVERY PROFILE

The only TAVR system with a vessel indication down to 5.0 mm***



[&]quot;Broadest annulus range based on CT derived diameters

[‡]Measurement for TAV-in-SAV only.

^{***}Evolut™ R 23, 26 and 29 mm valves. 34 mm valve minimum vessel indication ≥ 5.5 mm

INDICATIONS The Medtronic CoreValve and CoreValve Evolut R systems are indicated for use in patients with symptomatic heart diseased due to either severe native calcific aortic stenosis or failure (stenosed, insuficient, or combile) of a surgical bioprosthetic aortic valve who are judged by a heart team, including a cardiac surgeon, to be at high or greater risk for open surgical therapy (ie. Society of Thoracic Surgeons predicted risk of operative mortality score 28% or at a 215% risk of traiting at 31 days.

CONTRAINDICATIONS The CoreValve and CoreValve Evolut R systems are contraindicated for patients presenting with any of the following conditions: known hypersensitivity or contraindication to aspirin, heparin (HIT/HITTS) and bivalirudin, ticlopidine, clopidogrel, Nitinol (Titanium or Nickel), or sensitivity to contrast media, which cannot be adequately premedicated; ongoing sepsis, including active endocarditis; preexisting mechanical heart valve in aortic position.

WARNINGS General Implantation of the CoreValve and CoreValve Evolut R systems should be performed only by physicians who have received Meditonic CoreValve training. This procedure should only be performed where emergency a ortic valve surgery can be performed promptly. Mechanical failure of the delivery catheter system and/or accessories may result in patient complications. Transcatheter Aortic Valve (Bioprosthesis) Accelerated deterioration of the bioprosthesis may occur in patients presenting with an altered calcium metabolism.

PRECAUTIONS General The safety and effectiveness of the CoreValve and CoreValve Evolut R systems have not been evaluated in the pediatric population. The safety and effectiveness of the bioprosthesis for a ortic valve replacement have not been evaluated in the following patient populations: patients who do not meet the criteria for symptomatic severe native aortic stenosis as defined. (1) symptomatic severe high gradient aortic stenosis – aortic valve area ≤1 0cm² or aortic valve area index ≤0.6 cm²/m², a mean aortic valve gradient ≥40 mmHg, or a peak aortic-jet velocity ≥4.0 m/s. (2) symptomatic severe low-flow/low-gradient aortic stenosis – aortic valve area ≤1.0cm² or aortic valve area index ≤0.6 cm²/m², a mean aortic valve gradient <40 mmHg, and a peak aortic-jet velocity < 4.0 m/s; who are at moderate or low surgical risk (predicted perioperative mortality risk of < 15%); with untreated, clinically significant coronary artery disease requiring revascularization; with a preexisting prosthetic heart valve with and any significant client that the state of hemodynamic support. The safety and effectiveness of a CoreValve or CoreValve Evolut R bioprosthesis implanted within a failed preexisting transcatheter bioprosthesis has not been demonstrated. Implanting a CoreValve or CoreValve Evolut R bioprosthesis in a degenerated surgical bioprosthesis [transcatheter agrtic valve in surgical agrtic valve (TAV in SAV)] should be avoided in the following conditions. The degenerated surgical bioprosthesis presents with a significant concomitant perivalvular leak (between the prosthesis and the native annulus), is not securely fixed in the native annulus, or is not structurally intact (eq. wireform frame fracture); partially detached leaflet that in the aortic position may obstruct a coronary ostium, stent frame with a manufacturer's labeled inner diameter < 17 mm. The safety and effectiveness of the bioprosthesis for aortic valve replacement have not been evaluated in patient populations presenting with the following: blood dyscrasias as defined: leukopenia (WBC <1000 cells/mm²), thrombocytopenia (platelet count <50,000 cells/mm²), history of bleeding diathesis or coagulopathy, or hypercoagulable states; congenital bicuspid or unicuspid valve; mixed aortic valve disease (aortic stenosis and aortic regurgitation with predominant aortic regurgitation [3-4+1]); moderate to severe (3-4+) or severe (4+) mitral or severe (4+) tricuspid regurgitation; hypertrophic obstructive cardiomyopathy; new or untreated echocardiographic evidence of intracardiac mass, thrombus, or vegetation; native aortic annulus size <18 mm or >29 mm for CoreValve and <18 mm or >30 mm for CoreValve Evolut R per the baseline diagnostic imaging or surgical bioprosthetic aortic annulus size <17 mm or >29 mm for CoreValve and <17 mm or >30 mm for CoreValve Evolut R; transarterial access not able to accommodate an 18 Fr introducer sheath or the 14 Fr equivalent EnVeo R InLine sheath when using Model ENVEOR-US or transarterial access not able to accommodate a 20 Fr introducer sheath or the 16 Fr equivalent EnVeo R InLine sheath when using Model ENVEOR-N-US; sinus of valsalva anatomy that would prevent adequate coronary perfusion; moderate to severe mitral stenosis; severe ventricular dysfunction with left ventricular ejection fraction (LVEF) <20%; symptomatic carotid or vertebral artery disease; severe basal septal hypertrophy with an outflow gradient

Prior to Use Exposure to glutaraldehyde may cause irritation of the skin, eyes, nose, and throat. Avoid prolonged or repeated exposure to the vapors. Damage may result from forceful handling of the catheter. Prevent kinking the catheter when removing it from the packaging. This device was designed for single patient use only. Do not reuse, reprocess, or resterilize this product. Reuse, reprocessing, or resterilization may compromise the structuralintegrity of the device at Orac reater airsk for contamination of the device, which could result in patient injury, illness, or death. The bioprosthesis size must be appropriate to fit the patient's nantomy. Proper sizing of the device is the responsibility of the physician. Refer to Instructions for Use for available sizes. Failure to implant a device within the sizing matrix could lead to adverse effects such as those listed below. Patients must present with access vessel diameters of 26 mm for the CoreValve Evolut R system when using Model ENVEOR-US, or 25.5 mm when using Model ENVEOR-US, or 35.5 mm when using Model ENVEOR-US, or patients must present with an ascending aortic (direct aortic) access site 260 mm from the Dasal plane for both systems. Implantation of the bioprosthesis should be avoided in patients with aortic root angulation (angle between plane of aortic valve annulus and horizontal plane/vertebrae) of 350° for right subclavian/axillary access or >70° for femoral and left subclavian/axillary access. Use caution when using the subclavian/axillary acreas or patient RIMA graft. For direct aortic access, ensure the access site and trajectory are free of patent RIMA or a preexisting patent RIMA graft.

During Use For direct aortic and subclavian access procedures, care must be exercised when using the tip-retrieval mechanism to ensure adequate clearance to avoid advancement of the catheter tip through the bioprosthesis leaflets during device closure. For direct aortic access procedures, use a separate introducer sheath, do not use the EnVeo R InLine sheath. Adequate rinsing of the bioprosthesis with sterile saline, as described in the Instructions for Use, is mandatory before implantation. During rinsing, do not touch the leaflets or squeeze the bioprosthesis. If a capsule becomes damaged during loading or the capsule fails to close, replace the entire system (bioprosthesis, catheter, and CLS). Do not use a catheter with a damaged capsule. After a bioprosthesis has been inserted into a patient, do not attempt to reload that bioprosthesis on the same or any other catheter. AccuTrak DCS Only. During implantation, if resistance to deployment is encountered (e.g., the micro knob starts clicking or it sight or stuck), apply upward pressure to the macro silder while turning the micro knob. If the bioprosthesis still does not deploy, remove it from the patient and use another system. AccuTrak DCS Only. Once deployment is initiated, retrieval of the bioprosthesis from the patient (e.g., use of the catheter) is not recommended. Retrieval of a partially deployed valve using the catheter may cause mechanical

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CardioVascular Technical Support

Tel: (877) 526-7890 Tel: (763) 526-7890 Fax: (763) 526-7888 failure of the delivery catheter system, aortic root damage, coronary artery damage, myocardial damage, vascular complications, prosthetic valve dysfunction (including device malposition), embolization, stroke, and/or emergent surgery. AccuTrak DCS Only: During deployment, the bioprosthesis can be advanced or withdrawn as long as annular contact has not been made. Once annular contact is made, the bioprosthesis cannot be advanced in the retrograde direction; if necessary, and the frame has only been deployed \leq 2/3 of its length, the bioprosthesis can be withdrawn (repositioned) in the antegrade direction. However, use caution when moving the bioprosthesis in the antegrade direction. EnVeo R DCS Only: If a misload is detected, unsheath the bioprosthesis and examine the bioprosthesis for damage (for example, permanent frame deformation, frayed sutures, or valve damage). Do not attempt to reload a damaged bioprosthesis. Do not load the bioprosthesis onto the catheter more than 2 times or after it has been inserted into a patient. EnVeo R DCS Only: Use the deployment knob to deploy and recapture the bioprosthesis. Do not use the trigger for deploying or recapturing because it could cause inaccurate placement of the bioprosthesis. EnVeo R DCS Only: Once the radiopaque capsule marker band reaches the distal end of the radiopaque paddle attachment (point of no recapture), retrieval of the bioprosthesis from the patient is not recommended. Retrieval after the point of no recapture may cause mechanical failure of the delivery catheter system, aortic root damage, coronary artery damage, myocardial damage, vascular complications, prosthetic valve dysfunction (including device malposition), embolization, stroke, and/or emergent surgery. EnVeo R DCS Only: During deployment, the bioprosthesis can be advanced or withdrawn as long as annular contact has not been made. Once annular contact is made, the bioprosthesis cannot be advanced in the retrograde direction; recapture until the bioprosthesis is free from annular contact, and then reposition in the retrograde direction. If necessary, and the radiopaque capsule marker band has not yet reached the distal end of the radiopaque paddle attachment, the bioprosthesis can be withdrawn (repositioned) in has not yet reactive the distail each of the realizable and a tractiment, the bioprostress can be withdrawn repositioned in the antegrade direction. However, use caution when moving the bioprostress in the antegrade direction. While the catheter is in the patient, ensure the guidewire from the tip. Do not remove the guidewire from the catheter while the catheter is inserted in the patient. Use the handle of the delivery system to reposition the bioprosthesis. Do not use the outer catheter sheath. Once deployment is complete, repositioning of the bioprosthesis (e.g., use of a snare and/or forceps) is not recommended. Repositioning of a deployed valve may cause aortic root damage, coronary artery damage, myocardial damage, vascular complications, prosthetic valve dysfunction (including device malposition), embolization, stroke, and/or emergent surgery. Do not attempt to retrieve or to recapture (EnVeo DCS only) a bioprosthesis if any one of the outflow struts is protruding from the capsule. If any one of the outflow struts has deployed from the capsule, the bioprosthesis must be released from the catheter before the catheter can be withdrawn. Ensure the capsule is 6 losed before catheter removal. When using a separate introducer sheath, if increased resistance is encountered when removing the catheter through the introducer sheath, do not force passage. Increased resistance assumed a not included any other and order assage may result in damage to the device and/or harm to the patient. Increased resistance may indicate a problem and forced passage may result in damage to the device and/or harm to the patient. If the cause of resistance cannot be determined or corrected, remove the catheter and indirected as as a single unit over the guidewine, and inspect the catheter and confirm that it is complete. Clinical long-term durability has not been established. for the bioprosthesis. Evaluate bioprosthesis performance as needed during patient follow-up. Postprocedure, administer appropriate antibiotic prophylaxis as needed for patients at risk for prosthetic valve infection and endocarditis. Postprocedure, administer anticoaquiation and/or antiplatelet therapy per physician/clinical judgment. Excessive contrast media may cause arminister anticognistion amount an impacted thereby be impossible in the imposition of the imposition bioprosthesis must be implanted within a transcatheter bioprosthesis to improve valve function, valve size and patient anatomy In the event that valve function or sealing is impaired due to excessive calcification or incomplete expansion, a postimplant balloon dilatation of the bioprosthesis may improve valve function and sealing. To ensure patient safety, valve size and patient anatomy must be considered when selecting the size of the balloon used for dilatation. The balloon size chosen for dilatation should not exceed the diameter of the native sortic annulus or, for surgical bioprosthetic valves, the manufacturer's labeled inner diameter. Refer to the specific balloon catheter manufacturer's labeling for proper instruction on the use of balloon catheter devices. Note: Bench testing has only been conducted to confirm compatibility with NuMED Z-MED II™ Balloon Aortic Valvuloplasty catheters where CoreValve or CoreValve Evolut R bioprosthesis device performance was maintained after dilation. Data on File

For EnVeo R DCS: For transfemoral access, use caution in patients who present with multiplianar curvature of the aorta acute angulation of the aortic arch, an ascending aortic aneurysm, or severe calcification in the aorta and/or vasculature. If 82 of these factors are present, consider an alternative access route to prevent vascular complications. There will be some resistance when the catheter is advanced through the vasculature. If there is a significant increase in resistance, stop advancement and investigate the cause of the resistance (for example, magnify the area of resistance) before proceeding. Do not force passage. Forcing passage could increase the risk of vascular complications (for example, vessel dissection or rupture). Persistent force on the catheter can cause the catheter to kink, which could increase the risk of vascular complications (for example, vessel dissection or rupture).

POTENTIAL ADVERSE EVENTS Potential risks associated with the implantation of the CoreValve or CoreValve Evolut R transcatheter aortic valve may include, but are not limited to, the following: death - myocardial infarction, cardiac arrest, cardiogenic shock, cardiac tamponade - coronary occlusion, obstruction, or vessels pasam (including acute coronary closure) - cardiovascular injury (including rupture, perforation, tissue erosion, or dissection of vessels, ascending aorta trauma, ventricle, myocardium, or valvular structures that may require intervention) - emergent surgical or transcatheter intervention (for example, coronary artery bypass, heart valve replacement, valve explant, percutaneous coronary intervention [PCI], balloon valvuloplasty) - prosthetic valve dysfunction (regurgitation or stensois) due to fracture; bending out-or-fround configuration) of the valve frame; underexpansion of the valve frame; calcification; pannus; leaflet wear, tear, prolapse, or retraction; por valve coaptation, suture breaks or disruption; leaks, mal-sixing (prosthesis) - patient mismatch); malposition (either too high or too lowly) malplacement - prosthetic valve migration/embolization - prosthetic valve endocarditis - prosthetic valve thrombosis - delivery catheter system component migration/embolization - stroke (ischemic or hemorrhagic), transient ischemic attack (TIA), or other neurological deficits - heart failure - cardiac radiac output - andilar or lowly or disabling bleeding) - vascular access-related complications (e.g. dissection, perforation, pain, bleeding, hematoma, pseudoaneurysm, irreversible nerve injury, compartment syndrome, arteriovenous fistula, stensosis) - mitral valve regurgitation or injury - conduction system disturbances (for example, atrioventricular node block, left-bundle branch block, asystole), which may require a permanent pacemaker - infection (including electrolyte imbalance) - allergic reaction to antiplatelet agents, contrast medium, or anesthesia - exposure to radiation through fluorosc

 $Please reference the CoreValve \ and \ CoreValve \ Evolut \ R \ Instructions for \ Use for more information regarding indications, warnings, precautions and potential adverse events.$

CAUTION Federal law (USA) restricts this device to sale by or on the order of a physician.

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