

Dear Imaging Center:

This letter is in response to your inquiry concerning the safety of performing magnetic resonance (MR) procedures in patients who have been implanted with Edwards Lifesciences LLC (formerly Baxter Healthcare Corporation, CardioVascular Group) products.

MR Information:

MR procedures have been performed on numerous occasions on patients with Edwards' implantable products without reported problems. The products listed below are made from non-ferromagnetic, weakly ferromagnetic materials or paramagnetic materials. For all products, the *in vivo* forces are greater than those pertaining to the magnetic field interactions (i.e., the forces associated with translational attraction and torque are less than those associated with gravitational forces). Thus, these products are considered safe for patients undergoing magnetic resonance imaging (MRI) procedures using MR systems operating under the conditions described in the following pages.

Product Information:

Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Carpentier-Edwards aortic and mitral bioprostheses	2625, 6625	12, 20, 21
Carpentier-Edwards S.A.V. aortic bioprosthesis	2650	12, 20, 21
Carpentier-Edwards Duraflex low pressure porcine mitral bioprosthesis	6625LP	12, 20, 21
Carpentier-Edwards Duraflex low pressure porcine mitral bioprosthesis with extended sewing ring	6625-ESR-LP	12, 20, 21
Carpentier-Edwards bioprosthetic valved conduit	4300	12, 20, 21



MR Conditional

Non-clinical testing has demonstrated that these devices are MR Conditional. A patient with these devices can be scanned safely immediately after placement of the implant under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of 3000 gauss/cm or less.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode.

Under the scan conditions defined above these devices are expected to produce a maximum temperature rise of 3 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by these devices extends approximately as far as 30 mm from the devices when imaged with a gradient echo pulse sequence and approximately as far as 14 mm from the devices when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of these devices. Optimization of MR imaging parameters is recommended.

The valve wireform stent is composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

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Cobalt	Chromium	omium Nickel Molybden		Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Carpentier-Edwards PERIMOUNT pericardial aortic bioprostheses	2700, 2700TFX	18, 19, 20, 21, 22
Carpentier-Edwards PERIMOUNT RSR pericardial aortic bioprostheses	2800, 2800TFX	
Carpentier-Edwards PERIMOUNT Magna pericardial aortic bioprostheses	3000, 3000TFX	



Non-clinical testing has demonstrated that these devices are MR Conditional. A patient with these valves can be scanned safely, immediately after placement of this valve under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of less than 3000 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above these devices are expected to produce a maximum temperature rise of 2.3 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by these devices extends approximately as far as 27.5 mm from the bioprostheses when imaged with a gradient echo pulse sequence and approximately as far as 8.5 mm from the valves when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the bioprostheses. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobaltchromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
Carpentier-Edwards PERIMOUNT Magna Ease pericardial aortic	3300TFX	19, 20, 21
bioprosthesis		



Non-clinical testing has demonstrated that this device is MR Conditional. A patient with this valve can be scanned safely, immediately after placement of this implant under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of less than 3000 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above this device is expected to produce a maximum temperature rise of 2.3 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately as far as 25.5 mm from the bioprosthesis when imaged with a gradient echo pulse sequence and approximately as far as 12.5 mm from the valve when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the bioprosthesis. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Carpentier-Edwards PERIMOUNT pericardial mitral bioprosthesis	6900	19, 20, 21
Carpentier-Edwards PERIMOUNT Plus pericardial mitral bioprosthesis	6900P	
Carpentier-Edwards PERIMOUNT Theon mitral pericardial bioprosthesis	6900PTFX	



Non-clinical testing has demonstrated that these devices are MR Conditional. A patient with these valves can be scanned safely, immediately after placement of these implants under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of less than 3000 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above these devices are expected to produce a maximum temperature rise of 2.3 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately as far as 33 mm from the bioprostheses when imaged with a gradient echo pulse sequence and approximately as far as 12.5 mm from the valves when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of these bioprostheses. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
Carpentier-Edwards PERIMOUNT Magna Mitral pericardial bioprostheses	7000, 7000TFX	19, 20, 21
Carpentier-Edwards PERIMOUNT Magna Mitral Ease pericardial bioprostheses	7200TFX, 7300TFX	



Non-clinical testing has demonstrated that these devices are MR Conditional. A patient with these valves can be scanned safely immediately after placement of these implants under the following conditions:

- Static magnetic field of 3 tesla or less.
- Maximum spatial gradient field of 3000 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above these devices are expected to produce a maximum temperature rise of 2.3 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately as far as 36 mm from the bioprostheses when imaged with a gradient echo pulse sequence and approximately as far as 11.5 mm from the valves when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of these bioprostheses. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Ī	Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
Ī	40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
EDWARDS INTUITY Elite aortic valve	8300AB	14



Non-clinical testing has demonstrated that this device is MR Conditional. A patient with this valve can be scanned safely, immediately after placement of this implant under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of less than 2670 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above this device is expected to produce a maximum temperature rise of 0.8 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately as far as 40 mm from the bioprosthesis when imaged with a gradient echo pulse sequence and approximately as far as 40 mm from the valve when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the bioprosthesis. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

The expandable frame is composed of a stainless steel alloy that is commonly used in implantable devices. The nominal composition (wt. percent) of the stainless steel material used is as follows:

Chromium	Nickel	Molybdenum	Manganese	Silicon	Carbon	Phosphorus	Sulfur	Copper	Iron
18%	14%	2.6%	< 2.0%	< 0.75%	< 0.03%	< 0.025%	< 0.01%	< 0.5%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
INSPIRIS RESILIA aortic valve	11500A	23



Non-clinical testing has demonstrated that this device is MR Conditional. A patient with this valve can be scanned safely, immediately after placement of this implant under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of less than 3000 gauss/cm.
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of continuous scanning per sequence in the normal operating mode

Under the scan conditions defined above this device is expected to produce a maximum temperature rise of 2.5 °C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately as far as 17 mm from the bioprosthesis when imaged with a gradient echo pulse sequence and approximately as far as 10 mm from the valve when imaged with a spin echo pulse sequence in a 3 T MRI system. The lumen is partially to fully obscured under these conditions. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the bioprosthesis. Optimization of MR imaging parameters is recommended.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
KONECT RESILIA aortic valved conduit	11060A	19, 20, 21, 26



Non-clinical testing has demonstrated that the KONECT RESILIA aortic valved conduit (AVC), Model 11060A, is MR Conditional. A patient with the Model 11060A AVC can be scanned safely immediately after placement of this implant, under the following conditions:

- Static magnetic field of 3 tesla or less
- Spatial gradient field of less than 3000 gauss/cm (30 T/m)
- Maximum MR system-reported whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg in the normal operating mode

Under the scan conditions defined above, KONECT RESILIA AVC Model 11060A is expected to produce a maximum in vivo temperature rise of less than 2 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact extends approximately 12.5 mm from the Model 11060A valve when imaged with a spin echo pulse sequence, and 25.5 mm from the device when imaged with a gradient echo pulse sequence and a 3 tesla MRI system. The artifact obscures the device lumen.

The valve wireform stent and orifice-stiffening band are composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal composition (wt. percent) is as follows:

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Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
MITRIS RESILIA mitral valve	11400M	27



Non-clinical testing demonstrated that the Model 11400M valve is MR Conditional. A patient with this device can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5T and 3.0T only
- Maximum spatial gradient field of 3000 gauss/cm (30 T/m) or less
- Maximum MR system-reported, whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg per 15 minutes of scanning (i.e. per pulse sequence)
- Normal mode operation of the MR system for both SAR and gradients.

Under the scan conditions above, the Model 11400M valve is expected to produce a maximum temperature rise of 2 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 20 mm from the Model 11400M valve when imaged with a gradient echo pulse sequence and a 3.0 tesla MRI system. Optimization of MR imaging parameters is recommended.

The valve wireform stent is composed of a corrosion-resistant, nickel-titanium superelastic alloy that is commonly used in implantable devices. The valve orifice-stiffening band is composed of a corrosion-resistant cobalt-chromium spring alloy that is commonly used in implantable devices. The nominal compositions (wt. percent) are as follows:

Component	Cobalt	Chromium	Nickel		Carbon	Iron	Nic	obium	Tita	anium	C	Copper
Wireform	<0.05%	<0.01%	55.8% <0.04%		<0.05%	<0.025%			Bal	3al <0.01%		
Component	Cobalt	Chromium	Nickel	Мо	lybdenum	Mangane	ese	Carbo	n	Berylliu	m	Iron
Band	40%	20%	15%		7%	2%		< 0.10	%	< 0.10%	6	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Cribier-Edwards aortic bioprosthesis (PHV)(Caution: Investigational	9000, 9000PHV,	N/A
device. Limited by Federal law to investigational use.)		

Non-clinical testing has demonstrated that the Cribier-Edwards aortic bioprosthesis (PHV) is MR Conditional. It can be scanned safely under the following conditions:

- Static magnetic field of 3 tesla or less.
- Spatial gradient field of 720 gauss/cm or less.
- Maximum whole-body-averaged specific absorption rate (SAR) of 3.0 W/kg for 15 minutes of scanning.

In non-clinical testing, the device produced a maximum temperature increase of 0.5 °C at a maximum whole body averaged specific absorption rate (SAR) of 3.0 W/kg for 15 minutes of MRI.

MR image quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the device. Optimization of MR imaging parameters is recommended.

The valve's stent frame is composed of stainless steel material. The nominal composition (wt. percent) of the stainless steel material as follows:

	Chromium	Nickel	Molybdenum	Manganese	Silicon	Copper	Carbon	Phosphorus	Sulfur	Iron
I	17.3%	14.4%	2.53%	1.74%	0.54%	0.093%	0.026%	0.017%	0.001%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards SAPIEN transcatheter heart valve	9000TFX	N/A



Non-clinical testing has demonstrated that the Edwards SAPIEN transcatheter heart valve is MR Conditional. It can be scanned safely under the following conditions:

- Static magnetic field of 1.5 tesla (T) or 3 tesla.
- Spatial gradient field of 2500 gauss/cm or less.
- Maximum whole-body-averaged specific absorption rate (WB-SAR) of 2 W/kg for 15 minutes of scanning
- Normal mode operation, as defined in IEC 60601-2-33 Ed. 3.0, of the MR system.

In non-clinical testing and analysis, the implant was determined to produce a temperature rise of less than 1.1 °C above background for a whole body SAR of 2W/kg for 15 minutes of MR scanning in a 1.5 T and 3.0 T cylindrical whole body MR system.

The image artifact extended as far as 15 mm from the device for spin echo images and 40 mm for gradient images when scanned in non-clinical testing in a 3 T GE Signa HDx MR system. The artifact obscures the device lumen in gradient echo images. The implant has not been evaluated in MR systems other than 1.5 or 3.0 T.

MR image quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the device.

The valve's stent frame is composed of stainless steel material. The nominal composition (wt. percent) of the stainless steel material used is as follows:

Chromium	Nickel	Molybdenum	Manganese	Silicon	Copper	Carbon	Phosphorus	Sulfur	Iron
17.3%	14.4%	2.53%	1.74%	0.54%	0.093%	0.026%	0.017%	0.001%	Bal

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards SAPIEN XT transcatheter heart valve (THV)	9300TFX	N/A



Non-clinical testing has demonstrated that the Edwards SAPIEN XT transcatheter heart valve is MR Conditional. A patient with this device can be scanned safely, immediately after placement of this device under the following conditions:

- Static magnetic field of 1.5 tesla or 3 tesla
- Maximum spatial gradient field of 2500 gauss/cm (25 T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg (Normal Operating Mode)

Under the scan conditions defined above, the SAPIEN XT transcatheter heart valve is expected to produce a maximum temperature rise of 2.6 $^{\circ}$ C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends as far as 14.5 mm from the implant for spin echo images and 30 mm for gradient echo images when scanned in a 3.0 T MRI system. The artifact obscures the device lumen in gradient echo images. The implant has not been evaluated in MR systems other than 1.5 or 3.0 T.

For valve-in-surgical valve implantation or in the presence of other implants, please refer to the MRI safety information for the surgical valve or other devices prior to MR imaging.

The frame of the implant is composed of MP35N alloy with the chemical constituents listed below:

Carbon	max. 0.025 wt%
Silicon	max. 0.15 wt%
Manganese	max. 0.15 wt%
Phosphorus	max. 0.015 wt%
Sulfur	max. 0.010 wt%
Chromium	19.0 – 21.0 wt%
Nickel	33.0 – 37.0 wt%
Iron	max. 1.0 wt%
Molybdenum	9 – 10.5 wt%
Titanium	max. 1.0 wt%
Boron	max. 0.015 wt%
Cobalt	balance

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards SAPIEN 3 transcatheter heart valve (THV)	9600TFX	N/A
Edwards SAPIEN 3 Ultra transcatheter heart valve (THV)	9750TFX	
Edwards SAPIEN 3 Ultra RESILIA transcatheter heart valve (THV)	9755RSL	



Non-clinical testing has demonstrated that the Edwards SAPIEN 3 transcatheter heart valve, Edwards SAPIEN 3 Ultra transcatheter heart valve, and the Edwards SAPIEN 3 Ultra RESILIA transcatheter heart valve are MR Conditional. A patient with this device can be scanned safely, immediately after placement of this device under the following conditions:

- Static magnetic field of 1.5 tesla or 3 tesla.
- Maximum spatial gradient field of 2500 gauss/cm (25 T/m) or less.
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg (Normal Operating Mode)

Under the scan conditions defined above, the Edwards SAPIEN 3 transcatheter heart valve and the Edwards SAPIEN 3 Ultra transcatheter heart valve are expected to produce a maximum temperature rise of 3.0 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends as far as 14.5 mm from the implant for spin echo images and 30 mm for gradient echo images when scanned in a 3.0 T MRI system. The artifact obscures the device lumen in gradient echo images. The implant has not been evaluated in MR systems other than 1.5 or 3.0 T.

For valve-in-valve implantation or in the presence of other implants, please refer to the MRI safety information for the surgical valve or other devices prior to MR imaging.

The frame of the implant is composed of MP35N alloy with the chemical constituents listed below:

Carbon	max. 0.025 wt%
Silicon	max. 0.15 wt%
Manganese	max. 0.15 wt%
Phosphorus	max. 0.015 wt%
Sulfur	max. 0.010 wt%
Chromium	19.0 – 21.0 wt%
Nickel	33.0 – 37.0 wt%
Iron	max. 1.0 wt%
Molybdenum	9 – 10.5 wt%
Titanium	max. 1.0 wt%
Boron	max. 0.015 wt%
Cobalt	balance

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards SAPIEN X4 transcatheter heart valve (THV)	14000RSL	N/A



Non-clinical testing has demonstrated that the Edwards SAPIEN X4 transcatheter heart valve is MR Conditional. A patient with this device can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5 Tesla or 3 T
- Maximum spatial gradient field of 2500 Gauss/cm (25 T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2.0 W/kg (Normal Operating Mode).

Under the scan conditions defined above, the SAPIEN X4 transcatheter heart valve is expected to produce a maximum temperature rise of 3.0 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends as far as 14.5 mm from the implant for spin echo images and 30 mm for gradient echo images when scanned in a 3.0 T MRI system. The artifact obscures the device lumen in gradient echo images.

For valve in valve implantation or in the presence of other implants, please refer to the MRI safety information for the surgical valve or other devices prior to MR imaging.

The frame of the implant is composed of MP35N allow with the chemical constituents listed below:

The hame of the implant is com	posed of MF 33N alloy with the chemical constituents listed below.
Carbon	max. 0.025 wt%
Silicon	max. 0.15 wt%
Manganese	max. 0.15 wt%
Phosphorus	max. 0.015 wt%
Sulfur	max. 0.010 wt%
Chromium	19.0 – 21.0 wt%
Nickel	33.0 – 37.0 wt%
Iron	max. 1.0 wt%
Molybdenum	9 – 10.5 wt%
Titanium	max. 1.0 wt%
Boron	max. 0.015 wt%
Cobalt	balance

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The radiopaque markers on the implant is composed of electron-beam or vacuum-arc cast Tantalum (R05200) with the chemical constituents listed below in accordance with ASTM F560-17:

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Carbon	max. 0.01%
Oxygen	max. 0.015%
Nitrogen	max. 0.01%
Hydrogen	max. 0.0015%
Niobium	max. 0.1%
Iron	max. 0.01%
Titanium	max. 0.01%
Tungsten	max. 0.05%
Molybdenum	max. 0.02%
Silicon	max. 0.005%
Nickel	max. 0.01%
Tantalum	balance

CAUTION: investigational device. Limited by Federal (USA) law to investigational use only. To be used by qualified investigators only.

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
CardiAQ-Edwards transcatheter mitral valve (TMV)	TMV3040B,	N/A
	9650TMV	



Non-clinical testing has demonstrated that the TMV is MR Conditional. A patient with this device can be scanned safely in an MR system meeting the following conditions:

- Static magnetic field of 1.5 tesla or 3.0 tesla only
- Maximum spatial gradient field of 4,000 gauss/cm (40 T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg

Under the scan conditions defined above, the TMV is expected to produce a maximum temperature rise of 1.7 °C in a 1.5 tesla system and 1.8 °C in a 3.0 tesla system after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 10 mm from the TMV when imaged with a gradient echo and spin echo pulse sequence and a 3.0 tesla MRI system. MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the TMV. Therefore, optimization of MR imaging parameters to compensate for the presence of this device may be necessary.

The frame of the implant is composed of Nitinol alloy with the chemical constituents listed below in accordance with ASTM F2063-12.

Nickel	54.5 to 57.0 wt%
Titanium	Balance
Nitrogen plus Oxygen	0.05 wt%
Carbon	<0.05 wt%

*INVESTIGATIONAL DEVICES. CAUTION: The CardiAQ-Edwards transcatheter mitral valve is an investigational device. Limited by Federal (USA) law to investigational use only. Exclusively for clinical investigations. To be used by qualified investigators only. See instructions for use for full information, including indications, contraindications, warnings, precautions and adverse events.

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards EVOQUE Transcatheter Mitral Valve (TMV)	9850TMV	N/A



Non-clinical testing has demonstrated that the Edwards 9850TMV valve is MR Conditional. A patient with the valve can be scanned safely, immediately after placement of this valve under the following conditions:

- Static magnetic field of 3.0 tesla or less
- Spatial magnetic gradient field of less than 3000 gauss/cm
- Maximum MR system-reported, whole body averaged SAR of 2.0 W/kg
- Normal operating mode of the MR system for both gradients and SAR

Based on worst-case non-clinical testing and calculated SAR in the patient during MRI, the 9850TMV valve was determined to produce a temperature rise of less than 3 °C at a maximum MR system reported, whole-body-averaged specific absorption rate (SAR) of 2.0 W/kg, for 15 minutes of MR scanning at 1.5 T and a rise of less than 4 °C at a background local specific absorption rate (SAR) of 2.0 W/kg, for 15 minutes of MR scanning at 3.0 T.

Image artifact was measured non-clinically in a GE Signa 3T Discovery 750 MR system according to ASTM F2119-07 using the spin echo and gradient echo sequences specified therein. The spin echo images had artifacts that extended as far as 4 mm from the implant. The gradient echo images had artifacts that extended as far as 5.85 mm from the valve. The lumen of the valve was partially to fully obscured.

The frame of the implant is composed of Nitinol alloy with the chemical constituents listed below in accordance with ASTM F2063-12.

Nickel	55.8 wt%
Titanium	Balance
Nitrogen plus Oxygen	<0.04 wt%
Carbon	<0.04 wt%

*INVESTIGATIONAL DEVICES. CAUTION: The Edwards EVOQUE Transcatheter Mitral Valve is an investigational device. Limited by Federal (USA) law to investigational use only. Exclusively for clinical investigations. To be used by qualified investigators only. See instructions for use for full information, including indications, contraindications, warnings, precautions and adverse events.

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Replacement Heart Valve Product Description (Stented Tissue)	Models	Reference
Edwards EVOQUE Transcatheter Tricuspid Valve	9850EV44 9850EV48 9850EV52 9850EV56*	N/A



Non-clinical testing has demonstrated the Edwards EVOQUE valve, Model 9850EV, is MR Conditional. A patient with the valve can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5 and 3 T only
- Maximum spatial gradient magnetic field of 3000 gauss/cm (30.0 T/m) or less
- Maximum MR system-reported, whole body averaged specific absorption rate (SAR) of 2.0 W/kg
- Normal operating mode of the MR system for both gradients and SAR

Under the scan conditions defined above, the EVOQUE valve is expected to produce a maximum temperature rise of 4 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the EVOQUE valve extends approximately 0.8 cm from the device when imaged with a gradient echo or spin echo pulse sequence and a 3 T MRI system.

The frame of the implant is composed of Nitinol alloy with the chemical constituents listed below in accordance with ASTM F2063-12.

Nickel	55.8 wt%
Titanium	Balance
Nitrogen plus Oxygen	<0.04 wt%
Carbon	<0.04 wt%

*CAUTION INVESTIGATIONAL DEVICE: The Edwards EVOQUE Tricuspid Valve Replacement System Model 9850EV56 is an investigational device. Limited by Federal (USA) law to investigational use only. Exclusively for clinical investigations. To be used by qualified investigators only. See instructions for use for full information, including indications, contraindications, warnings, precautions and adverse events.

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Transcatheter Valve Repair Product Description	Models	Reference
Edwards PASCAL Precision Transcatheter Valve Repair System	20000IS	N/A
Edwards PASCAL Precision Transcatheter Valve Repair System	20000ISM	N/A



Non-clinical testing demonstrated that the PASCAL and PASCAL Ace implants are MR Conditional. A patient with this device can be safely scanned in a MR system meeting the following conditions:

- Static magnetic field of 1.5 T and 3.0 T
- Maximum spatial field gradient of 3,000 gauss/cm (30 T/m)
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 4 W/kg (First Level Controlled Operating Mode)

Under the scan condition defined above, the implant is expected to produce a maximum temperature rise of less than 4°C after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device in a worst-case multiple implant configuration extends up to 15 mm from the implant when imaged in the worst-case gradient echo pulse sequence in a 3.0 T MRI system.

The PASCAL (Model 20000IS) and PASCAL Ace (Model 20000ISM) implants are primarily composed of Nitinol spacer, paddles and clasps (in accordance with ASTM F2063). The nominal composition (wt. percent) of the materials are as follows:

Nickel	54.5 to 57.0%
Carbon	Max 0.040%
Cobalt	Max 0.050%
Copper	Max 0.010%
Chromium	Max 0.010%
Hydrogen	Max 0.005%
Iron	Max 0.050%
Niobium	Max 0.025%
Nitrogen + Oxygen	Max 0.040%
Titanium	Balance

The 20000IS implant also comprises a titanium nut and bolt. The 20000ISM implant comprises a titanium nut, bolt, proximal plate and distal plate (in accordance with ASTM F136). The nominal composition (wt. percent) of the materials are as follows:

Nitrogen	Max 0.05%	
Carbon	Max 0.08%	
Hydrogen	Max 0.012%	
Iron	Max 0.25%	
Oxygen	Max 0.13%	
Aluminum	5.5-06.50%	
Vanadium	3.5-4.5%	
Titanium	Balance	

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
Edwards CENTERA transcatheter heart valve	9551S	25



The Edwards CENTERA THV has been determined to be MR Conditional. A patient with this device can be immediately scanned safely in an MR system meeting the following conditions:

- Static magnetic fields of 1.5 tesla (T) or 3.0 T.
- Maximum spatial gradient field of 3000 Gauss/cm (30 T/m).
- Maximum MR System reported, whole-body-averaged specific absorption rate (WB-SAR) of 2.0 W/kg (Normal Operating Mode).

Under the scan conditions defined above, the CENTERA valve is expected to produce a maximum temperature rise of less than 2.0 °C after 15 minutes of continuous scanning.

Image artifact was measured non-clinically in a GE Signa 3T HDx MR system according to ASTM F2119-07 using the spin echo and gradient echosequences specified therein. The spin echo images had artifacts that extended as far as 4 mm from the implant and partially to fully obscured the lumen. The gradient echo images had artifacts that extended as far as 5 mm from the valve.

The THV has not been evaluated in MR systems other than 1.5 T or 3.0 T. The delivery system has not been evaluated for MR compatibility and is considered MR unsafe.

The frame of the implant is composed of Nitinol alloy with the chemical constituents listed below in accordance with ASTM F2063-12

accordance with ASTM F2003-12.		
Nickel	54.5% - 57.0%	
Cobalt	max. 0.05%	
Iron	max. 0.05%	
Carbon	max. 0.05%	
Niobium	max. 0.025%	
Copper	max. 0.01%	
Chromium	max. 0.01%	
Oxygen	max. 0.04%	
Oxygen + Nitrogen	max. 0.05%	
Hydrogen	max. 0.005%	
Titanium	Balance	

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
Edwards Alterra adaptive prestent in conjunction with	29AP4045,	N/A
Edwards SAPIEN 3 transcatheter heart valve	9600TFX	



Non-clinical testing has demonstrated that the Edwards Alterra adaptive prestent, alone or with a deployed SAPIEN 3 transcatheter heart valve, is MR Conditional. A patient can be scanned safely immediately after placement of this implant in an MR system meeting the following conditions:

- Static magnetic fields of 1.5 Tesla or 3.0 Tesla
- Spatial magnetic gradient field of 3000 Gauss/cm (30 T/m) or less
- Maximum MR system-reported, whole body averaged specific absorption rate (SAR) of 2.0 W/kg (normal operating mode) scanning per sequence
- · Gradient system is in normal operating mode

Under the scan conditions defined above, the Edwards Alterra adaptive prestent is expected to produce a maximum temperature rise of 4.0 °C or less after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends as far as 6.6 mm for gradient echo images when scanned using a 3.0 T MRI system. The artifact obscures the device lumen in spin and gradient echo images.

The frame of the valve implant is composed of MP35N alloy with the chemical constituents listed below:

Carbon	max. 0.025 wt%
Silicon	max. 0.15 wt%
Manganese	max. 0.15 wt%
Phosphorus	max. 0.015 wt%
Sulfur	max. 0.010 wt%
Chromium	19.0 – 21.0 wt%
Nickel	33.0 – 37.0 wt%
Iron	max. 1.0 wt%
Molybdenum	9 – 10.5 wt%
Titanium	max. 1.0 wt%
Boron	max. 0.015 wt%
Cobalt	balance

(continues on next page)

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
Edwards Alterra adaptive prestent in conjunction with	29AP4045,	N/A
Edwards SAPIEN 3 transcatheter heart valve	9600TFX	

(continued from previous page)

The frame of the prestent implant is composed of Nitinol alloy with the chemical constituents listed below in accordance with ASTM F2063-12:

Nickel	54.5% - 57.0%
Cobalt	max. 0.05%
Iron	max. 0.05%
Carbon	max. 0.05%
Niobium	max. 0.025%
Copper	max. 0.01%
Chromium	max. 0.01%
Oxygen	max. 0.04%
Oxygen + Nitrogen	max. 0.05%
Hydrogen	max. 0.005%
Titanium	Balance

The radiopaque markers on the prestent implant is composed of electron-beam or vacuum-arc cast Tantalum (R05200) with the chemical constituents listed below in accordance with ASTM F560-17:

Carbon	max. 0.01%
Oxygen	max. 0.015%
Nitrogen	max. 0.01%
Hydrogen	max. 0.0015%
Niobium	max. 0.1%
Iron	max. 0.01%
Titanium	max. 0.01%
Tungsten	max. 0.05%
Molybdenum	max. 0.02%
Silicon	max. 0.005%
Nickel	max. 0.01%
Tantalum	balance

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
SAPIEN M3 dock in conjunction with the SAPIEN M3 valve	9770DDS/9780DDS/9680DSC with 9680TFX29M	N/A



Non-clinical testing has demonstrated that the Edwards SAPIEN M3 dock implant, with a deployed SAPIEN M3 valve, is MR Conditional. A patient can be scanned safely immediately after placement of these devices under the following conditions:

- Static magnetic field of 1.5 tesla or 3.0 tesla only
- Maximum spatial gradient field of 3,000 gauss/cm (30 T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg (Normal Operating Mode)

Under the scan conditions defined above, the Edwards SAPIEN M3 implants are expected to produce a maximum temperature rise of 2 °C or less after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 8 mm from the implant when imaged with spin echo pulse sequence and a 3.0 tesla MRI system. The lumen of the valve inside the dock was partially to fully obscured in spin and echo gradient images.

Reduction in artifact may be possible with sequences designed for reduction of metal artifact.

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Replacement Heart Valve Product Description (Stented Tissue)	Model	Reference
SAPIEN M3 dock in conjunction with the	9880DDS with	N/A
SAPIEN M3 valve	9880TFX29M	



Non-clinical testing has demonstrated that the Edwards SAPIEN M3 dock implant, with a deployed SAPIEN M3 valve, is MR Conditional. A patient can be scanned safely immediately after placement of this valve under the following conditions:

- Static magnetic field of 1.5 Tesla (T) or 3.0 T.
- Spatial magnetic gradient field of 4,000 Gauss/cm (40 T/m) or less.
- Maximum MR system-reported, whole body averaged specific absorption rate (WB-SAR) of 2.0 W/kg (Normal Operating Mode)

Under the scan conditions defined above, the Edwards SAPIEN M3 implants are expected to produce a temperature rise of 2 °C or less after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 1.1 cm from the dock when imaged with a gradient echo pulse sequence and a 3.0 T MR system. The lumen of the valve inside the dock was partially to fully obscured in spin and gradient echo images.

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Replacement Heart Valve Product Description (Stentless Tissue)	Models				
Edwards Prima aortic stentless bioprosthesis	2500				
Edwards Prima Plus aortic stentless bioprosthesis	2500P				
These valves are made of porcine aortic valves and there are no metallic components. Therefore there					
are no MRI issues for these implants, and they may be considered as MR safe.					

	nt Heart Valve	e Product Desc cal)		Models		Refe	erence	
Starr-Edward	ds aortic and ı	mitral prosthese	2400, 60	200, 2300, 2 200, 6120, 6 320, 640	300,	2	2, 3	
		n a static magne I.5 tesla or less					during N	⁄IR
Starr-Edward	ds prostheses	,	Pre-1000, Pre-6000, 1260, 2320, 6520 (plastic disk)			4, 5		
		n a static magne 2.35 tesla or les					e during	MR
2300, 2310,	2320, 2400, 6	d of Stellite 21. 3300, 6310, 632 of Stellite 21 is a	0 and 6400) a					
Cobalt	Carbon	Manganese	Silicon	Chromium	Nickel	Molybo	lenum	Iron
61.5%	<0.35%	< 1.0	1.0%	28.5%	<1.0%	69	%	0.75%

Replacement (Bileaflet Mec	Heart Valve Pr hanical)	oduct Descript	Models			Reference		
Edwards-Duro	medics aortic ai	nd mitral bileafle	et prosth	eses	31	60, 9	120	2
procedures pe leaflets are gra commercially p	se devices in a s rformed at 1.5 to aphite substrate oure titanium gra ellite 25 is as fo	esla or less. Val coated with pyr ade II. The stiffe	lve hous olytic ca	ings are irbon. Ti	e composed he retainer	d of so	olid pyrolytic in the sewir	carbon and the g ring are
Cobalt	Chromium	Tungsten	Nic	kel	Iron		Manganese	Carbon
50%	20%	15%	10	%	< 3%		1.5%	0.1%
The nominal co	omposition (wt.	percent) for con	nmercial	lly pure	titanium gr	ade II	is as follows	3:
Nitrogen	Nitrogen Carbon		gen Ir		ron	(Oxygen	Titanium
< 0.03%	< 0.10%	< 0.01	2%	< 0	.30%	, , ,		99%

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Replacement Heart Valve Product Description (Bileaflet Mechanical)	Models	Reference
Edwards MIRA aortic and mitral mechanical valves	3600, 3600f, 3600u, 9600	1
(Caution: Investigational device. Limited by Federal		
law to investigational use.)		

Testing of these devices in a magnetic field of 1.5, 3.0, and 8.0 tesla has shown that these devices are safe and compatible during MRI (magnetic resonance imaging) procedures. Valve housing is composed of ASTM B348 Grade 5 Ti-6Al-4V titanium alloy coated with turbostatic carbon. Leaflets are composed of graphite substrate coated with pyrolytic carbon. The nominal composition for Ti-6Al-4V titanium alloy is as follows:

Nitrogen	Carbon	Hydrogen	Iron	Oxygen	Aluminum	Vanadium	Titanium
< 0.03%	< 0.10%	< 0.0125%	< 0.40%	< 0.20%	5.5 to	3.5 to	Balance
					6.75%	4.5%	(~90%)

Valve Repa	ir Product De	Mod	els F	Reference				
Carpentier-Edwards Classic annuloplasty mitral and tricuspid rings						4500	1	
Carpentier-Edwards Classic annuloplasty mitral and tricuspid rings						4525	1	
with Duraflo	treatment							
Edwards MC3 Tricuspid annuloplasty ring						0	1	
Testing of these devices in a magnetic field of 1.5 tesla has shown that these devices are safe and compatible during MRI (magnetic resonance imaging) procedures. Rings have titanium alloy cores. The nominal composition (wt. percent) of the titanium alloy is as follows:								
Nitrogen	Carbon	Hydrogen	Iron	Oxygen	Aluminum	Vanadium	Titanium	
< 0.05%	< 0.08%	< 0.012%	< 0.25%	< 0.13%	6%	4%	89%	

Exceptions:

Carpentier-Edwards annuloplasty rings, Models 4400 and 4500, marketed from 1980 to 1983, were made of stainless steel. Therefore we are unable to advise on the safety of MR procedures for patients with these particular annuloplasty rings. These older rings were labeled with lot numbers (not serial numbers) that had the following format: 1C005 (i.e., where the first character was numeric, the second character was a letter from A to L and the last three or four characters were numeric).

Valve Repair Product Description	Models	Reference
Carpentier-McCarthy-Adams IMR ETlogix mitral annuloplasty ring	4100	1
GeoForm mitral annuloplasty ring	4200	1

The device has been shown not to have magnetic interactions at up to 8 tesla. It is also safe with respect to RF heating at 1.2 W/kg for up to 15 minutes. Artifacts have been determined at 1.5 tesla. Optimization of MR imaging parameters is recommended.

Rings have titanium alloy cores. The nominal composition (wt. percent) of the titanium alloy is as follows:

3	g							
Nitrogen	Carbon	Hydrogen	Iron	Oxygen	Aluminum	Vanadium	Titanium	
< 0.05%	< 0.08%	< 0.012%	< 0.25%	< 0.13%	6%	4%	89%	

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Valve Repair Product Description	Models
Cosgrove-Edwards annuloplasty mitral and tricuspid band	4600
Cosgrove-Edwards annuloplasty mitral and tricuspid band with Duraflo treatment	4625

These bands are composed of a silicone rubber strip impregnated with barium sulfate covered with a knit polyester cloth and there are no metallic components. Therefore, there are no MRI issues for these implants, and they may be considered as MR safe.

Valve Repair Product Description	Models	Reference
Carpentier-Edwards Physio mitral annuloplasty ring	4450	1
Carpentier-Edwards Physio mitral annuloplasty ring with Duraflo	4475	1
Treatment		

Testing of these devices indicates that MR procedures may be conducted safely with static fields of 1.5 tesla and 3.0 tesla. Rings have corrosion-resistant cobalt-chromium spring alloy bands separated by polyester film strips covered by silicone rubber and a knit polyester covering. The nominal composition (wt. percent) of the cobalt-chromium alloy is as follows:

Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	< 0.10%	< 0.10%	16.0%

Valve Repair Product Description	Model	Reference
Carpentier-Edwards Physio II mitral annuloplasty ring	5200	1



MR Conditional

Non-clinical testing has demonstrated that the Carpentier-Edwards Physio II annuloplasty ring, model 5200, is MR Conditional. A patient with this annuloplasty ring can be scanned safely, immediately after placement of this implant under the following conditions:

- Static magnetic field of 3 tesla or less
- Spatial gradient field of 720 gauss/cm or less
- Maximum MR system reported whole-body-averaged specific absorption rate (SAR) of 3 W/kg for 15 minutes of scanning

In non-clinical testing, the Carpentier-Edwards Physio II annuloplasty ring produced a temperature rise of less than or equal to 1.8 °C at a maximum MR system reported whole-body-averaged specific absorption rate (SAR) of 3 W/kg for 15 minutes of MR scanning in a 3 tesla MR System.

MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the device. Optimization of MR imaging parameters is recommended.

Rings have metal alloy bands separated by polyester film strips covered by silicone rubber and a woven polyester covering. The nominal composition (wt. percent) of the metal alloy is as follows:

peryester esterning. The normal composition (was percently of the motor and) to do remove.							
Cobalt	Chromium	Nickel	Molybdenum	Manganese	Carbon	Beryllium	Iron
40%	20%	15%	7%	2%	<0.10%	<0.10%	16%

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Valve Repair Product Description	Model	Reference
Physio Flex annuloplasty ring	5300	24



Non-clinical testing demonstrated that the Physio Flex annuloplasty ring, model 5300, is MR Conditional. A patient with this device can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5T and 3.0T only
- Maximum MR spatial gradient field of 3000 gauss/cm (30 T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of < 2 W/kg (Normal Operating Mode)

Under the scan conditions above, the Physio Flex annuloplasty ring, model 5300 is expected to produce a maximum temperature rise of 2 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device extends approximately 6 mm from the Physio Flex annuloplasty ring when imaged with a gradient echo pulse sequence and a 3.0 tesla MRI system. Optimization of MR imaging parameters is recommended.

The Physio Flex annuloplasty rings have nitinol cores. The nominal composition (wt. percent) of the nitinol is as follows:

Nickel	Carbon	Cobalt	Copper	Chromium	Hydrogen	Iron	Niobium	Nitrogen + Oxygen	Titanium
55.8%	<0.04%	<0.05%	<0.01%	<0.01%	<0.005%	<0.05%	<0.025%	<0.04%	Bal

Valve Repai	r Product De	escription	Model	Refere	ence			
Carpentier-E	dwards Phys	io Tricuspid ar	6200 11					
compatible d	Testing of these devices in a magnetic field of 3.0 tesla has shown that these devices are safe and compatible during MRI (magnetic resonance imaging) procedures. Rings have titanium alloy cores. The nominal composition (wt. percent) of the titanium alloy is as follows:							
Nitrogen	Carbon	Hydrogen	Iron	Oxygen	Aluminum	Vanadium	Titanium	
< 0.05%	< 0.08%	< 0.012%	< 0.25%	< 0.13%	6%	4%	89%	

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Valve Repair Product Description	Model	Reference
dETlogix mitral annuloplasty ring	5100	1



Non-clinical testing has demonstrated that the dETlogix annuloplasty ring, model 5100, is MR Conditional. A patient with the dETlogix annuloplasty ring can be scanned safely, immediately after placement of this implant under the following conditions:

- Static magnetic field of 3 tesla or less
- Spatial gradient field of 720 gauss/cm or less
- Maximum MR system reported whole-body-averaged specific absorption rate (SAR) of 3 W/kg for 15 minutes of scanning

In non-clinical testing, the dETlogix annuloplasty ring produced a temperature rise of less than or equal to 0.6 °C at a maximum MR system reported whole-body-averaged specific absorption rate (SAR) of 3 W/kg for 15 minutes of MR scanning in a 3 tesla MR System.

MR image quality may be compromised if the area of interest is in the same area or relatively close to the position of the device. Optimization of MR imaging parameters is recommended.

The ring has a titanium alloy core. The nominal composition (wt. percent) of the titanium alloy is as follows:

Nitrogen	Carbon	Hydrogen	Iron	Oxygen	Aluminum	Vanadium	Titanium
< 0.05%	< 0.08%	< 0.012%	< 0.25%	< 0.13%	6%	4%	89%

Bovine Pericardial Patch	Model		
Bovine Pericardial Patch	4700		
These patches are constructed from bovine pericardial tissue and there are no metallic components.			
Therefore this device is MR Safe.			

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Product Description	Model	Reference	
Atrial Shunt	9700AS07 (part of 9770ASDC)	28	



Non-clinical testing of Edwards Lifesciences Atrial Shunt is MR Conditional. A patient with the Atrial Shunt can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5T and 3.0T
- Maximum spatial field gradient of 3,000 Gauss/cm (30T/m)
- Maximum MR system-reported whole body average specific absorption rate (SAR) of 2.0W/kg for normal mode of the MR system

Under the scan condition defined above, the Atrial Shunt is expected to produce a maximum temperature rise of 2°C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the device may extend up to 5mm from the Atrial Shunt when imaged in the worst-case gradient pulse sequence and a 3.0T MRI system.

The implant is composed (weight %) of Nitinol alloy with the chemical constituents listed below:

Nickle	Titanium	Oxygen + Nitrogen	Carbon		
55.8%	Balance	< 0.04%	< 0.04%		
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Product Description	Model	Reference
APTURE transcatheter shunt	N/A (part of 9771AIS)	29



MRI Safety Information. A person with the APTURE transcatheter shunt may be safely scanned at 1.5T or 3.0T under the following conditions. Failure to follow these conditions may result in injury.

Parameter	Condition		
Device name	APTURE transcatheter shunt		
Static Magnetic Field Strength (B0)	1.5 and 3T		
MR Scanner Type	Cylindrical		
B0 Field Orientation	Horizontal		
Maximal Spatial Field Gradient	3,000 gauss/cm (30T/m)		
RF Excitation	Circularly Polarized (CP)		
RF Transmit Coil Type	Integrated Whole Body Transmit Coil		
Operating Mode	Normal Operating Mode		
Maximum Whole-Body SAR	2 W/kg (Normal Operating Mode)		
Maximum Head SAR	3.2 W/kg (Normal Operating Mode)		
Temperature Rise	Under the scan conditions defined above, the APTURE shunt is expected to produce a temperature rise of less than 3°C after 15 minutes of continuous scanning.		
Scan Duration	Up to 1 hour of continuous RF (a sequence or back-to-back series/scan without breaks).		
Image Artifact	Non-clinical testing of the APTURE shunt in a 3T environment resulted in an image artifact of 5mm. Some manipulation of scan parameters may be needed to compensate for the artifact.		

The implant is composed (weight %) of Nitinol alloy with the chemical constituents listed below:

	(1101911170)		
Nickle	Titanium	Oxygen + Nitrogen	Carbon
55.8%	Balance	< 0.04%	< 0.04%

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Contact us in the USA at 800-424-3278 or outside the USA at 949-250-2500 if you have any questions.

Sincerely, Technical Support

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