#### Cardiac Resynchronisation Therapy (CRT) Devices

### Allure<sup>™</sup> RF

#### Merlin@home™ Transmitter Compatible

#### Cardiac Resynchronisation Therapy Pacemaker

#### **Product Highlights**

- Angled header and physiologic tear drop shape provide better lead wrap
- CorVue™ Congestion Monitoring feature monitors the intrathoracic impedance and provides the option for both patient and physician alerts
- The DirectTrend™ Report provides a summary of three month daily, one year weekly or one year daily diagnostic trends
- Better patient unitilization from Day 1 when paired with the Merlin@home™ transmitter at point of care
- AT/AF Alerts can be programmed to notify patients and their clinics when a programmed AT/AF threshold or continuous episode duration has been exceeded, or when a high ventricular rate accompanies the AT/AF episode
- Exclusive AF Suppression™ algorithm is clinically proven to suppress episodes of paroxysmal and persistent AF
- AT/AF burden trend provides a graphical representation of the percentage of time in AT/AF and the number of AT/AF episodes in the previous 52 weeks
- Up to 14 minutes of stored electrograms help identify key intrinsic and pacemaker-related events and simplify the diagnosis of complex ECG rhythms associated with heart failure
- Industry-leading longevity offers 8.2 years of service life supported by a 6 year warranty\*

#### Ordering Information

Contents: Cardiac pulse generator

Model Number	Dimensions (H x W x T, mm)	Weight (g)	Volume (cc)	Connector
PM3222	55 x 59 x 6	24	14	IS-1

Indications: Implantation of Allure and Allure RF devices is indicated for: maintaining synchrony of the left Indications: Implantation of Allure and Allure Rr devices is indicated for: maintaining synchrony of the left and right ventricles in patients who have undergone an AV nodal ablation for chronic atrial fibrillation and have NYHA Class III or IVI) in those patients who remain symptomatic despite stable, optimal medical therapy, and have a left ventricular ejection fraction < 35% and a prolonged ORS duration, implantation of Assurity, Endurity and Allure family of devices is indicated in one or more of the following permanent conditions: Endurity and Allure family of devices is indicated in one or more of the following permanent conditions: syncope, presyncope, fatigue, disorientation, or any combination of those symptoms. Rate-Modulated Pacing is indicated for patients with chronotropic incompetence, and for those who would benefit from increased stimulation rates concurrent with physical activity. Pual-Chamber Pacing is indicated for those patients exhibiting: sick sinus syndrome, chronic, symptomatic second- and third-degree AV block, recurrent Adams-Stokes syndrome, symptomatic bilateral bundle branch block when tachyarrhythmia and other causes have been ruled out. Atrial Pacing is indicated for patients with sinus node dysfunction and normal AV and intraventricular conduction systems. Ventricular Pacing is indicated for patients with significant bradycard and normal sinus rhythm with only rare episodes of A-V block or sinus arrest, chronic atrial fibrillation, severe physical disability. AF Suppression algorithm is indicated for suppression of paroxysmal or persistent atrial fibrillation episodes in patients with one or more of the above pacing indications.

Contraindications: Implanted Cardioverter-Defibrillator (ICD). Devices are contraindicated in patients with an implanted cardioverter-defibrillator. Rate-Adaptive Pacing may be inappropriate for patients who experience angina or other symptoms of myocardial dysfunction at higher sensor-driven rates. An appropriate Maximum Sensor Rate should be selected based on assessment of the highest stimulation rate tolerated by the patient. AF Suppression stimulation is not recommended in patients who cannot tolerate high atrial-rate stimulation. Dual-Chamber Pacing, though not contraindicated for patients with chomic atrial flutric, chronic atrial flutric, and provide no benefit beyond that of single-chamber pacing in such

patients. Single-Chamber Ventricular Demand Pacing is relatively contraindicated in patients who have patients. Single-Innumber Ventricular Demand Pacing is relatively contraindicated in patients who have demonstrated pacemaker syndrome, have retrograde VA conduction, or suffer a drop in arterial blood pressure with the onset of ventricular pacing. Single-Chamber Atrial Pacing is relatively contraindicated in patients who have demonstrated compromise of AV conduction. Atrial Fibrillation. Anthem devices are contraindicated in patients having chronic atrial fibrillation or intermittent atrial fibrillation that does not terminate. For specific contraindications associated with individual modes, refer to the programmer's on-screen help.

specific contraindications associated with individual modes, refer to the programmer's on-screen help.

Potential Adverse Events: The following are potential complications associated with the use of any pacing system: air embolism, body rejection phenomena, cardiac tamponade or perforation, hematoma, bleeding hematoma, seroma, formation of fibrotic tissue, local tissue reaction, inability to interrogate or program due to programmer or device malfunction, infection/erosion, interruption of desired pulse generator function due to electrical interference, either electromyogenic or electromagnetic, lead malfunction due to conductor fracture or insulation degradation, loss of capture or sensing due to lead displacement, body reaction at electrode interface, or lead malfunction (fracture or damage to insulation), loss of normal device function due to battery failure or component malfunction, pacemaker migration or pocket erosion, pectoral muscle or diaphragmatic stimulation, phrenic nerve stimulation, pneumothorax/hemothorax, endocarditis, excessive bleeding induced atrial or ventricular arrhythmias, myocardial irritability, pericardial effusion, pericardial rub, pulmonary edema, rise in threshold and exit block, valve damage, cardiac/coronary sinus dissection, cardiac/coronary sinus or cardiac vein thrombosis. sinus perforation, coronary sinus or cardiac vein thrombosis

Refer to the User's Manual for detailed indications, contraindications, warnings, precautions and potential

#### Customer Support: 46-8-474-4756

Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of Birds Jummary: Prior to using these devices, please review the instructions for use for a complete listing or indications, contraindications, contraindications, contraindications, contraindications, contraindications, contraindications, swarnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country. Unless otherwise noted, Mindicates that the name is a trademark of, or licensed to, St. Jude Medical or noe of its subsidiaries. ST. JUDE MEDICAL, the nine-squares symbol and MORE CONTROL. LESS RISK, are registered and unregistered trademarks and service marks of St. Jude Medical, Inc. and its related companies.

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<sup>\*</sup>Longevity calculated based on the following settings: 2.5 V, 500 Ohm, 60 BPM, 100% DDD-BiV Pacing, 0.4ms, Cap Confirm Off, and Stored EGM On

## Allure<sup>™</sup> RF

#### Cardiac Resynchronisation Therapy Pacemaker

#### **Product Specifications**

PHYSICAL SPECIFICATIONS	
Model	PM3222
Telemetry	RF
Dimensions (mm)	55 x 59 x 6
Weight (g)	24
Volume (cc)1	14
Connector	IS-1
PARAMETER	SETTINGS

#### Resynchronisation Therapy

QuickOpt™ Timing Cycle RV and LV Pulse Width (ms) RV and LV Pulse Amplitude (V) RV Pulse Configuration LV Pulse Configuration Ventricular Sense Configuration

Ventricular Pacing Chamber First Chamber Paced Interventricular Pace Delay (ms) Sensed/Paced AV Delay; Interventricular Paced Delay 0,05; 0,1–1,5 in steps of 0,1

0,25–4,0 in steps of 0,25; 4,5–7,5 in steps of 0,5 Unipolar; Bipolar Unipolar; Bipolar Unipolar; Bipolar; LV Tip-RV Ring; LV Ring-RV Ring BV Unipolar Tip; BV Bipolar; RV Unipolar Tip; RV Bipolar;

LV Unipolar Tip; LV Bipolar; RV Unipolar Ring; LV Tip-RV Tip BV; RV only; LV only (temporary mode) Simultaneous2: RV: LV

#### Output/Sensing Negative AV

Hysteresis Search (ms) Shortest AV/PV Delay (ms) Atrial ACap™ Confirm Primary Pulse Confirmation Backup Pulse Confirmation Backup Pulse Amplitude (V) Searchable Intervals (hrs) Atrial Pulse Configuration Atrial Sense Configuration Atrial Sensitivity<sup>3,4</sup> (Fixed) (mV) Atrial Pulse Amplitude (V) Atrial Pulse Width (ms) RVCap™ Confirm Searchable Interval (hrs)

LVCap™ Confirm Searchable Interval (hrs)
SenseAbility™ Technology A Max Sensitivity (mV) V Max Sensitivity (mV)

Threshold Start

Decay Delay (ms)

Ventricular Sensitivity (fixed) (mV)

Off; -10 to -120 in steps of 10  $25{-}50$  in steps of 5;  $60{-}120$  in steps of 10 On; Off; Monitor

Rinolar Bipolar 5,0 8: 24

8: 24 Unipolar (tip-case); Bipolar (tip-ring) Unipolar (tip-case); Bipolar (tip-ring); Unipolar Ring (ring-case) 0,1—0,5 in steps of 0,1; 0,75–2,0 in steps of 0,25; 2,5–5,0 in steps of 0,5 0,25–4,0 in steps of 0,25; 4,5–7,5 in steps of 0,5 0,05; 0,1–1,5 in steps of 0,1

On; Off; Monitor 8; 24 On; Off; Monitor

8; 24 Off; On (Automatic Sensitivity Control adjustment for atrial and ventricular events)

0,2-1,0 in steps of 0,1 0,2-2,0 in steps of 0,1

(Atrial and Ventricular Post-Sense) 50; 62,5; 75; 100% (Atrial Post-Pace) 0,2-3,0 in steps of 0,1 mV (Ventricular Post-Pace) Auto; 0,2-3,0 in steps of 0,1 mV (Atrial and Ventricular Post-Sense) 0, 30; 60; 95; 125; 160; 190; 220 (Atrial Post-Pace) 0, 30; 60; 95; 125; 160; 190; 220 (Ventricular Post-Pace) 0, 30; 60; 95; 125; 160; 190; 220 (9-12,5) in steps of 0,5<sup>14</sup>

#### Rate/Timing

Mode DDT Trigger<sup>5</sup> DDT Timing<sup>5</sup> Base Rate (min<sup>-1</sup>) Hysteresis Rate (min<sup>-1</sup>) Search Interval (min) Intervention Rate (min-1)

Intervention Duration (min-1) Recovery Time Recovery Time Rest Rate (min<sup>-1</sup>) Maximum Tracking Rate (min<sup>-1</sup>) Sensed AV Delay (ms) Paced AV Delay (ms) Ventricular Pace/Sense Refractory<sup>7</sup> (Fixed) (ms) Atrial Pace Refractory Atrial Sense Refractory PVARP (ms)

Atrial Protection Interval (ms)<sup>5</sup> Far-Field Protection Interval (ms)<sup>5</sup>

A00(R): AAI(R): AAT(R): VOO(R): VVI(R): VVT(R): DOO(R): ...o(m), AAN(N); AAI(K); VUU(R); VVI(R); VVT(R); DOD DVI(R); DDI(R); DDD(R); Pacing Off R wave

30–130 in steps of 5; 140–170 in steps of 10 Off; 30-150 in steps of 5<sup>6</sup>

Off; 1; 5; 10; 15; 30

Fast; Medium; Slow; Very Slow

rast; medium; slow; very slow off; 30-150 in steps of 5 90–130 in steps of 5; 140–180 in steps of 10 25; 30–200 in steps of 10; 225–325 in steps of 25 25; 30–200 in steps of 10; 225–300 in steps of 25; 350

125; 160-400 in steps of 30; 440; 4708 190-400 in steps of 30; 440; 4708 93; 125; 157; 190-400 in steps of 30; 440; 4708 125-500 in steps of 25

1 ± 0,5 cc
2 LV first with 10 ms interventricular delay.
3 Sensitivity is with respect to a 20 ms haversine test signal.
4 Values 0,1-0,4 not available in a Unipolar Sense Configuration.
5 This parameter is not programmable resis rate is 5 min-¹ below the programmed base rate.
7 in dual-chamber modes, the maximum Ventricular Refractory Period is 325 ms.
8 Programming options dependent on pacing mode.
9 During ratin MINS in dual-chamber modes, the maximum Ventricular Refractory Vendo is 325 ms.
10 SI Burst Cycle is applied at the preprogrammed SI cycle length.

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Rate Responsive AV/PV Delay Rate Responsive PVARP/VREF Shortest PVARP/VRFF

Max Sensor Rate (min-1) Threshold

Slope Reaction Time Recovery Time

AF Suppression™ Algorithm Lower Rate Overdrive (min<sup>-1</sup>)<sup>5</sup> Upper Rate Overdrive (min<sup>-1</sup>)<sup>5</sup> No. of Overdrive Pacing Cycles

Rate Recovery (ms) Auto Mode Switch

AMS Base Rate (min-1)

Off; Low; Medium; High Off; Low; Medium; High 125-475 in steps of 25

On; Off; Passive 80–150 in steps of 5; 160-180 in steps of 10 Auto (-0,5); Auto (+0,0); Auto (+0,5); Auto (+1,0); Auto (+1,5); Auto (+2,0); 1-7 in steps of 0,5 Auto (-1); Auto (+0); Auto (+1); Auto (+2); Auto (+3); 1-16

Very Fast: Fast: Medium: Slow Fast; Medium; Slow; Very Slow

#### AF Management

Off; On

5 15–40 in steps of 5 8-12

Off; Low; High

Off; Low; High

Off; Low; High

Off; Low; High

Off; Low; High Off; Low; High 125-300 in steps of 25 2; 3; 4; 5; 10; 15; 20 Off; Low; High 125-300 in steps of 25

2; 3; 4; 5; 10; 15; 20 Off; Low; High

Off: Low: High

Off; Low; High

2-3-4-5

o; 12
Off; DDD(R) to DDI(R); DDD(R) to DDT(R); DDD(R) to VVI(R);
DDD(R) to VVT(R); VDD(R) to VVI(R); VDD(R) to VVT(R)
40-170 in steps of 5

#### **Stored Electrograms**

Priority Options Channel Triggers Advanced Hysteresis AMS Entry/AMS Exit/ AMS Entry and Exit AT/AF Detection Magnet Response High Atrial Rate Rate (min-1) No. of Consecutive Cycles High Ventricular Rate Rate (min<sup>-1</sup>) No. of Consecutive Cycles PMT Termination Consecutive PVCs

Magnet Response Ventricular Intrinsic Preference, VIP™ (ms) VIP Search Interval VIP Search Cycles of the Atrial Tachycardia Detection Rate (min-1) Post Vent. Atrial Blanking (PVAB) (ms)

No. of Consecutive PVCs Noise Reversion

Other

Ventricular Safety Standby PVC Response PMT Options PMT Detection Rate (min-1) Lead Type NIPS Options

Stimulation Chamber Coupling Interval<sup>®</sup> (ms) S1 Count

S1<sup>10</sup>; S2; S3 and S4 Cycle (ms) Right Ventricular Support Rate (min<sup>-1</sup>) Sinus Node Recovery Delay (s) Diagnostic Trends

CorVue Congestion Trigger

CorVue™ Congestion Monitoring

Off: Battery Test

Off; 50-150 in steps of 25; 160-200 in steps of 1030 sec.; 1; 3; 5; 10; 30 min.

110-200 in steps of 10; 225-300 in steps of 25

60-200 in steps of 10; 225; 250 Off: On Off; Atrial Pace<sup>8</sup> Off; Passive; Atrial Pace<sup>8</sup> 90-180 in steps of 5 Uncoded; Unipolar; Bipolar

Atrial Right Ventricular 200-800 in steps of 10 2-25 in steps of 1

Off; 100-800 in steps of 10 (Fixed or Adaptive)

Off: 30-95 in steps of 5

2; 4; 6; 8; 10; 12; 14; 16

17.5 in steps of 1 AT/AF Activity; Exercise; Lead Impedance; P and R Wave; A and V threshold, CorVue™ Congestion Monitoring

8-18 days

#### **Patient Notifiers**

Programmable Notifiers (On; Off)

Device at ERI; Atrial Lead Impedance Out of Range; Ventricular Lead Impedance Out of Range; LV Lead Impedance Out of Range; AT/AF Burden; AT/AF Episode Duration; High V Rate During AT/AF; High V Rate; Percent BIV/RV Pacing Alert; CorVue Alert

Device Reset Entry into Backup VVI Mode Audible Duration (sec) Number of Audible Alerts per Notification Number of Notifications

Time Between Notifications (hours)

1-16



Item GMCRM1094EN

## Tendril<sup>™</sup> STS

#### **Pacing Lead**

#### Product Highlights - Pacing Lead

- The Tendril STS lead allows patients to undergo MRI scans when used in conjunction with a MRI Ready pacemaker from St. Jude Medical
  - Allows MRI scans (See Parameter Settings for scan exclusion zone)
  - Permits a maximum whole body averaged specific absorption rate (SAR) of 2 watts per kilogram (W/kg)
- Soft silicone tip offers more compliance and less tip pressure at the lead tip-endocardium interface
- Small diameter lead offers improved ease of venous passage, reduced risk of venous thrombosis or rib-clavicle crush and ability to accommodate additional leads more easily
- Optim<sup>™</sup> lead insulation—a chemical co-polymer that blends the best features of polyurethane and silicone for improved handling and increased durability
- Titanium nitride (TiN) fractal coating on the tip and ring electrodes is designed to promote precise sensing and to provide improved contact with the myocardium
- Lubricious Fast-Pass™ coating facilitates lead insertion through the introducer and veins to ease implantation
- Fits through a 6 F introducer



Model Number	Description	Insulation	Fixation	Min. Introducer (F)	Connector	Length (cm)
2088TC	Tendril™ STS Pacing Lead	Optim™	Ext/Ret helix	6	IS-1 bipolar	46*; 52*; 58*; 65; 100

<sup>\*</sup> Indicates lead lengths that are MRI conditional with a scan exclusion zone.

Model Number	Description	Dimensions (H x W x T, mm)	Weight (g)	Volume (cc)	Connector
PM1140	Endurity <sup>™</sup> Core Pacemaker	41 x 50 x 6	19	9,7 (± 0,5)	IS-1
PM2140	Endurity Core Pacemaker	46 x 50 x 6	19	10,4 (± 0,5)	IS-1
PM1152	Endurity Core Pacemaker	41 x 50 x 6	19	9,7 (± 0,5)	IS-1
PM2152	Endurity Core Pacemaker	46 x 50 x 6	19	10,4 (± 0,5)	IS-1
PM1162	Endurity Pacemaker	41 x 50 x 6	19	9,7 (±0,5)	IS-1
PM2162	Endurity Pacemaker	46 x 50 x 6	19	$10,4 (\pm 0,5)$	IS-1
PM1172	Endurity MRI <sup>™</sup> Pacemaker	41 x 50 x 6	19	9,7 (± 0,5)	IS-1
PM2172	Endurity MRI Pacemaker	46 x 50 x 6	19	$10,4 (\pm 0,5)$	IS-1
PM1272	Assurity MRI <sup>™</sup> Pacemaker	47 x 50 x 6	20	10,4 (±0,5)	IS-1
PM2272	Assurity MRI Pacemaker	47 x 50 x 6	20	10,4 (±0,5)	IS-1

Indications: Tendril™ STS lead is designed for permanent sensing and pacing in either the right atrium or the right ventricle, in combination with a compatible device. Active leads such as the Tendril STS lead may be indicated for patients where permanent fixation of passive leads is suspected to be unstable.

In atrial applications, the use of screw-in leads such as Tendril STS lead may be indicated in the presence of an abnormal, surgically altered or excised atrial appendage.

**Contraindications:** Tendril STS lead is contraindicated: in the presence of tricuspid atresia, for patients with mechanical tricuspid valves, in patients who are expected to be hypersensitive to a single dose of one milligram of dexamethasone sodium phosphate.

Adverse Events: Potential complications associated with the use of Tendril STS lead are the same as with the use of other active fixation leads and include: cardiac tamponade, diaphragmatic stimulation, embolism, excessive bleeding, induced ventricular ectopy, infection, loss of pacing and/or sensing due to dislogment or mechanical malfunction of the pacing lead, phrenic nerve stimulation, thrombosis. Complications reported with direct subclavian venipuncture include pneumothorax, hemothorax, laceration of the subclavian artery, arteriovenous fistula, neural damage, thoracic duct injury, cannulation of other vessels, massive hemorrhage and, rarely, death.

Refer to the User's Manual for detailed indications, contraindications, warnings, precautions and potential adverse events.





## Tendril<sup>™</sup> STS

#### **Pacing Lead**

### Product Specifications - Pacing Leads

#### PHYSICAL SPECIFICATIONS 2088TC Model Minimum Introducer Size Type of Lead Active-fixation, bipolar, steroid-eluting, endocardial, pacing lead Lead Connector IS-1 bipolar Lead Lengths 46; 52; 58; 65; 100 cm Fixation Mechanism Extendable/Retractable helix Typical Number of Rotations for Helix Extension 6-11 (straight stylet) Lead Body Diameter 1,9 mm (max) Tip-to-Ring Spacing 10 mm Lead Tip Electrode (Cathode) Active titanium-nitride-coated Pt/Ir helix (2,0 mm extension) Tip Electrode Surface Area $6,9 \text{ mm}^2$ Ring Electrode (Anode) Titanium-nitride-coated Pt/Ir Ring Electrode Surface Area 16 mm<sup>2</sup> Capable with titanium-nitride-coated Pt/Ir helix Mapping Steroid $< 1~{\rm mg}$ dexamethasone sodium phosphate Inner Conductor/Outer Conductor MP35N™\* coil Inner Insulation Silicone rubber Outer Insulation $\text{Optim}^{\scriptscriptstyle\mathsf{TM}} \text{ lead insulation}$ Lead Body Coating Fast-Pass<sup>™</sup> coating

In Pack

Straight stylets 1 x-soft in lead; 1 x-soft; 1 soft

J-curved stylets 2 soft Helix extension/retraction clip-on tools 2 clip-on tools

#### Accessory Kits

Available Separately	Model Number	Compatible Lengths	Description
Stylet Kit	DS06002 with appropriate length designation	46; 52; 58; 65; 100 cm	1 fixation tool; 1 clip-on tool; 1 J-shaped soft; 1 x-soft; 1 soft; 1 firm; 1 x-firm
	DS06003 with appropriate length designation	46; 52; 58; 65; 100 cm	1 clip-on tool; 1 J-shaped soft; 1 x-soft; 1 soft; 1 firm; 1 x-firm
Locator™ Plus Deflectable Stylet	1281 with appropriate length designation	46; 52; 58; 65 cm	Disposable implant tool to facilitate precise lead positioning
	1292 with appropriate	46; 52; 58; 65 cm	and manipulation with one hand

#### MRI Conditional Parameters

Magnet strength: 1.5 Tesla

 $SAR: \le 2 W/kg$ 

Scan region: Isocenter must be inferior to L4 or 10 cm superior to C1



\*MP35N is a trademark of SPS Technologies, Inc.

 $\textbf{Customer Support:}\ 46\text{-}8\text{-}474\text{-}4756$ 

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## QuickFlex<sup>™</sup> µ

4F Bipolar, Optim™ Insulation-Insulated, Left Ventricular Pacing Lead

#### MODEL 1258T



#### **SPECIFICATIONS**

St. Jude Medical's innovative QuickFlex™µ lead—the latest in lead technology—is designed to provide predictable outcomes through superb access, delivery and fixation

Based on the QuickFlex™ lead family, this 4 F bipolar lead features a narrow ring electrode for lead tip flexibility, and a steerable tip for control and deliverability. The large S-curve provides superior fixation for this small diameter lead. The 4.3 F lead body diameter allows Direct-to-Target™ placement through a sub 5 F inner catheter to deliver leads to small, acute venous anatomies that may have been unreachable in the past.

The addition of the QuickFlex  $\mu$  lead to the QuickFlex CRT family of leads provides implanters with even more lead options to enable predictable placement and stability for varied patient needs.

#### **DESIGNED TO DELIVER**

- Optim<sup>™</sup> Lead Insulation
  - Optim insulation is a hybrid insulation material—the first of its kind developed specifically for cardiac lead use. It blends the biostability and flexibility of high-performance silicone rubber with the strength, tear resistance and abrasion resistance of polyurethane. This insulation allows for an abrasion resistant, thin diameter lead.
- Low Profile
  - Entire lead body: 4,3 F
  - Lead tip: 4,0 F
- Flexible Lead Body
  - Tip-to-ring electrode spacing of 20 mm and reduced lengths of rigid portions (tip and ring) create superb flexibility.
- Steerable Tip
  - Distal tip angle can be controlled to maneuver the lead through venous anatomy.
- Over-the-Wire or Stylet-Approach Compatibility
  - Specially designed leads give the implanting physician the option of using either approach during the same procedure.
- Fast-Pass™ Lubricious Coating
  - Enables multiple leads to easily slide against one another, possibly reducing inadvertent dislodgement.

#### **EXCEPTIONAL STABILITY AND PERFORMANCE**

- S-Shaped for Stability
  - The S-curve shape is designed to provide enhanced lead stability in a wide variety of vein sizes.
- Options for Any Anatomy
  - The complete family of QuickFlex leads, including QuickFlex μ, QuickFlex and QuickFlex XL is a comprehensive suite of CRT leads with varying diameters and S-Shape sizes, providing options to enable predictable procedures regardless of the patient's venous anatomy.
- Suture Sleeve
  - The new suture sleeve has been designed with silicone ridges to secure a thinner lead body.
- Titanium Nitride Coating (TiN)
  - TiN coating on the tip and ring electrodes has been shown to improve stimulation efficiency and lower polarisation.
- Steroid Elution
  - Steroid elution minimizes inflammatory reaction at the electrode-tissue interface and provides lower acute and chronic thresholds than non-steroideluting leads.



MODEL 1258T

#### SPECIFICATIONS

Parameter	Description
Connector	IS-1 Bipolar
Lead Length	75 cm, 86 cm, 92 cm
Lead Body Size	4,3 F (1,42 mm/0.056")
Tip Electrode Size	4,0 F (1,33 mm/0.052")
LV Lead Delivery System Introducer Size	Minimum 5 F ID
Minimum S-Curve Height	16 mm
Tip Electrode	Pt/Ir, TiN coated, ring-shaped, two grooves
Steroid	Dexamethasone sodium phosphate
Tip Electrode Surface Area	5,0 mm <sup>2</sup>
Ring Electrode Surface Area	7,4 mm <sup>2</sup>
Tip-to-Ring Electrode Spacing	20 mm
Lead Body Insulation	Optim <sup>™</sup> insulation
Lead Body Coating	Fast-Pass™ coating
Conductors	
Distal (coil)	MP35N™
Proximal (cables)	MP35N™
Suture Sleeve	Attached

#### Indications and Usage

The QuickFlex lead has application as part of a St. Jude Medical™ biventricular system.

#### Contraindications

The use of QuickFlex leads is contraindicated in patients who:

- Are expected to be hypersensitive to a single dose of 1,0 mg of dexamethasone sodium phosphate.
- Are unable to undergo an emergency thoracotomy procedure.
- Have coronary venous vasculature that is inadequate for lead placement, as indicated by venogram

Global Headquarters One Lillehei Plaza St. Paul, Minnesota 55117

+1 651 483 2000 +1 651 490 4310 Fax

St. Jude Medical Europe, Inc. The Corporate Village Figueras Building Avenue Da Vinci Iaan, 11 Box F1 B-1935 Zaventem

Belgium +32 2 774 68 11 +32 2 772 83 84 Fax Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, California 91342

+1 818 362 6822 +1 818 364 5814 Fax

St. Jude Medical Brasil Ltda. Rua Frei Caneca, 1380 7° ao 9° andares 01307-002 - São Paulo (SP) Brazil +55 11 5080 5400

+55 11 5080 5423 Fax

St. Jude Medical AB Veddestavägen 19 175 84 Järfälla Sweden

+46 8 474 4000 +46 8 760 9542 Fax

St. Jude Medical (Hong Kong) Ltd. Unit 2701-07 27/F, COSCO Tower

Grand Millennium Plaza 183 Queen's Road Central, Hong Kong +852 2996 7688 +852 2956 0622 Fax

St. Jude Medical Japan Co., Ltd. 3-1-30, Minami-Aoyama Minato-ku Tokyo 107 0062

Japan +81 3 3423 6450 +81 3 3402 5586 Fax



#### www.sjm.com

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## CPS Direct<sup>™</sup> Universal

Slittable Outer Guide Catheter

MODELS DS2C018, DS2C019, DS2C020, DS2C021, DS2C022, DS2C023, DS2C025, DS2C026, DS2C027, DS2C028, DS2C029



#### **SPECIFICATIONS**

- The CPS Direct Universal family of outer guide catheters is designed to facilitate left heart lead delivery. They are compatible with other products in the St. Jude Medical™ Cardiac Positioning System (CPS™) family an inter-compatible system of tools designed to give you more control to efficiently and predictably deliver the left-heart lead to your vein of first choice.
- Enables Direct-To-Target<sup>™</sup> delivery to the desired vein:
  - Soft, atraumatic tip with multi-durometer PEBAX<sup>TM</sup> shaft is designed to provide flexibility to allow advancement of the catheter deep into the coronary venous system.
  - CPS Direct<sup>™</sup> Universal catheter is compatible with CPS Aim<sup>™</sup> Universal inner catheters, designed to assist with branch vein subselection and left ventricular lead delivery , including delivery of the Quartet<sup>™</sup> quadripolar LV lead.
- Designed to reduce procedural steps during implant:
  - Slittable hub and integrated shaft provide smooth transition during slitting of catheter.
  - U-channel valve bypass tool simplifies lead delivery.
  - Ergonomic slitter facilitates smooth slitting.

- Designed to provide reliable coronary sinus access:
  - Excellent torque transmission and soft, atraumatic tip due to braid-reinforced, multi-durometer PEBAX™\* material design.
  - Unique SiteMark™ 3D markers provide fluoroscopic visibility to determine anterior/posterior location and verify torque transfer.
  - Six curve options to satisfy needs of various anatomies and different implanter techniques.
  - Compatible with CPS Aim<sup>™</sup> Universal cannulators and CPS Luminary<sup>™</sup> bideflectable catheter with lumen to modify shape and extend reach if necessary.
- Designed for worry-free removal:
- Catheter design features Smooth-Slit<sup>™</sup> braiding technology and ergonomic slitter, designed to allow effortless, best-in-class cutting, minimizing the risk of lead dislodgement upon catheter removal.

PEBAX is a trademark of Arkema Inc.



#### PHYSICAL SPECIFICATIONS

Slittable Outer Guide Catheter

Models	CURVE SHAPE	AVAILABLE LENGTH	OVERALL LENGTH	INNER DIAMETER	OUTER DIAMETER
DS2C018	Straight	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C019	115°	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C020	135°	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C021	Wide	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C022	X-Wide	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C023	Right Side	47 cm	50.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C025	Straight	54 cm	57.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C026	115°	54 cm	57.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C027	135°	54 cm	57.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C028	Wide	54 cm	57.6 cm	8F (2.67mm)	10F (3.34mm)
DS2C029	X-Wide	54 cm	57.6 cm	8F (2.67mm)	10F (3.34mm)

Multi-durometer PEBAX™ material reinforced with stainless steel braid wire for a kink-resistant catheter shaft and soft distal tip. Material

Lubricious coating on inner and outer surface.

Marker Three gold marker bands and two tungsten stripes on distal tip.

#### ACCESSORIES

INCLUDED Dilator 2 Valve bypass tools SEPARATELY AVAILABLE

CPS™ Universal Slitter CPS Direct™ Valve Bypass Tool

Global Headquarters

One St. Jude Medical Drive St. Paul, Minnesota 55117

+1 651 756 2000 +1 651 756 3301 Fax

SJM Coordination Center BVBA

The Corporate Village Da Vincilaan 11 Box F1 1935 Zaventem, Belgium +32 2 774 68 11 +32 2 772 83 84 Fax

St. Jude Medical Cardiovascular & **Ablation Technologies** 5050 Nathan Lane North

Plymouth, Minnesota 55442

+1 651 756 5400 +1 651 756 5470 Fax

St. Jude Medical Brasil Ltda.

Rua Itapeva, 538 5° ao 8° andar 01332-000 – São Paulo – SP

+55 11 5080 5400 +55 11 5080 5423 Fax

St. Jude Medical Implantable

Electronic Systems 15900 Valley View Court Sylmar, California 91342

+1 818 362 6822

+1 818 364 5814 Fax

St. Jude Medical (Hong Kong) Ltd.

Suite 1608, 16/F Exchange Tower 33 Wang Chiu Road Kowloon Bay, Kowloon Hong Kong SAR +852 2996 7688 +852 2956 0622 Fax

U.S. Division

6300 Bee Cave Road Bldg. Two, Suite 100 Austin, TX 78746 USA +1 512 286 4000

+1 512 732 2418 Fax

St. Jude Medical Australia Pty, Ltd.

17 Orion Road Lane Cove, NSW 2066 Australia +61 2 9936 1200 +61 2 9936 1222 Fax

#### SJMprofessional.com



Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country.

#### Accessories-Miscellaneous

Model Number	Receptacle (for adapting from)
AC-0160	Test Magnet 90 gauss at 1"
60007717-001	Vein Pick
442-2	Torque Wrench (#2)
437-246	Set of "L" Hex Wrenches (#2, #4, #6)
4033A	DF4/IS-1/DF-1 Lead Terminal Cap
6201	FasTac™ Flex Epicardial Lead Implant Tool
4080	Lead Removal Tool
DS0A001	Suture Sleeve (radiopaque 7.0 F)
AC-0130	Silicone Oil
424	Medical Adhesive
FL-1056	Lead Flushing Tool
4071	Torque Tool and Tip Introducer
AC-IP-2	IS-1 Port Plug
AC-DP-3	DF-1 Port Plug
AC-IS4PP	IS4/DF4 Port Plug
4078G	Custom Floppy Firm Guidewire, Straight, 5 cm Floppy Tip, 180 cm, 0.014", PTFE Coated
EX3151	IS4/DF4 Connector Sleeve

FasTac is a trademark of Greatbatch Medical.

**Customer Support:** 46-8-474-4756

**Brief Summary:** Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country.









EC Design-Examination Certificate
Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4)
(Other devices than custom made or intended for clinical investigation)

No. I7 014607 0234 Rev. 00

Manufacturer: St. Jude Medical

**Cardiac Rhythm Management** 

**Division** 

15900 Valley View Court Sylmar CA 91342

USA

EC-Representative: St. Jude Medical Coordination Center BVBA

The Corporate Village, Da Vincilaan 11 Box F1, 1935 Zaventem,

BELGIUM

#### Product: Implantable Pacemakers

The Certification Body of TÜV SÜD Product Service GmbH declares that a design examination has been carried out on the respective devices in accordance with AIMDD Annex 2 (4). This design of the devices conforms to the requirements of this Directive. For marketing of these devices an additional Annex 2 certificate is mandatory. See also notes overleaf.

**Report no.:** 713149860

 Valid from:
 2019-06-15

 Valid until:
 2024-05-26

Date, 2019-06-14

Stefan Preiß

1. Pumil





EC Design-Examination Certificate Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4) (Other devices than custom made or intended for clinical investigation)

No. I7 014607 0234 Rev. 00

Model(s): see below

St. Jude Medical Cardiac Rhythm Management Division Facility(ies):

15900 Valley View Court, Sylmar CA 91342, USA

St. Jude Medical Puerto Rico LLC

Lot A Interior - #2 Rd Km. 67.5, Santana Industrial Park, Arecibo

PR 00612, USA

St. Jude Medical Operations (M) Sdn.Bhd.

Plot 102, Lebuhraya Kampung Jawa, Bayan Lepas Industrial Zone,

11900 Penang, MALAYSIA

**Parameters** ./.

Design St. Jude Medical Cardiac Rhythm Management Division

Facility(ies): 15900 Valley View Court, Sylmar, CA 91342, USA

Product: Implantable Pacemakers

Test Report No.: 70069297

Model: Model No.: Variant:

Microny™ II SR+ 2525T

Test Report No.: 70110810

Model No.: Variant: Model:

Zephvr™ SR 5620 Zephyr™ DR 5820 5826 Zephyr™ XL DR

Page 2 of 4

TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123



EC Design-Examination Certificate
Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4)
(Other devices than custom made or intended for clinical investigation)

No. I7 014607 0234 Rev. 00

Test Report No.: 71321436

Model: Model No.: Variant:

Zephyr™ XL SR 5626

Test Report No.: 713017309\_1

Model: Model No.: Variant:

 Assurity™
 PM1240

 Assurity™
 PM2240

 Endurity™
 PM1160

 Endurity™
 PM2160

 Allure™
 PM3120

 Allure™ RF
 PM3222

 Allure Quadra™ RF
 PM3242

Test Report No.: 713028360

Model: Model No.: Variant

Quadra Allure MP™RF PM3262

Test Report No.: 713043621

Variant: Model No.: Model: MR Conditional PM1272 Assurity MRI™ MR Conditional Assurity MRI™ PM2272 MR Conditional Endurity MRI™ PM1172 MR Conditional Endurity MRI™ PM2172 MR Conditional PM1162 Endurity™ MR Conditional PM2162 Endurity™

Page 3 of 4

TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123



EC Design-Examination Certificate Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4) (Other devices than custom made or intended for clinical investigation)

No. I7 014607 0234 Rev. 00

Endurity™ Core PM2140 MR Conditional Endurity™ Core PM1152 MR Conditional	Model:	Model No.:	Variant:
Endulity Cole PM2152 MR Conditional	Endurity <sup>™</sup> Core	PM2140	

Test Report No.: 713084189

Model:	Model No.:	Variant:
Quadra Allure™	PM3542	MR Conditional
Quadra Allure MP™	PM3562	MR Conditional

Model No.:

Variant:

Test Report No.: 713130819

Model:

Zenex™	PM1250	
Zenex™	PM2250	
Zenus™	PM1170	
Zenus™	PM2170	
Zenex MRI™	PM1282	MR Conditional
Zenex MRI™	PM2282	MR Conditional
Zenus MRI™	PM1182	MR Conditional
Zenus MRI™	PM2182	MR Conditional

Page 4 of 4 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123



St. Jude Medical (SJM) hereby declares that the following SJM facilities and products conform to the applicable provisions of Annex 2 the European Union's Active Implantable Medical Devices Directive, AIMDD, 90/385/EEC. with the

any other notified body for the same products.	e premises of SJM. We declare no application has been lodged with the sole responsibility of eclaration is issued under the sole responsibility of eclaration issued previously for the same product(s).	
Manufacturer Address:	St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, CA 91342	
European Representative:	St. Jude Medical Coordination Center BVBA The Corporate Village Da Vincilaan 11 Box F1 1935 Zaventem, Belgium	
Product Type:	Implantable Pacemakers	
Product Name(s):	See Attachment	
Model Number(s):	See Attachment	
Classification:	AIMD	
GMDN Code(s):	See Attachment	
Original CE Mark Date:	See Attachment	
(FQA or EC as appropriate) Certificate No and expiration date:	EC Certification No: I7 014607 0234 Rev. 00 Expiration Date: 2024-05-26	
	FQA Certificate No: I1 16 12 14607 211 Expiration Date: 2021-07-25	
	ISO13485 Certificate No: Q1N 17 09 14607 217 Expiration Date: 2020-10-31	

Signature:

Manager Regulatory Affairs

**Applicable Quality System Standards:** 

**Notified Body:** 

**Notified Body Number:** 

Manufacturing Facilities:

Fulfills the requirements of Annex 2 of the European Union's Active Implantable Medical Devices Directive, AIMDD, 90/385/EEC and corresponding national legislation.

Fulfills applicable requirements including CE marking and the Essential Requirements of the AIMDD, 90/385/EEC and corresponding national legislation.

TÜV SÜD Product Service GmbH Zertifizierstelle Ridlerstraße 65, 80339, Münich, Germany

0123

St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, CA 91342 USA

St. Jude Medical Puerto Rico LLC Lot A Interior - #2 Rd Km. 67.5, Santana Industrial Park, Arecibo PR 00612, USA

St. Jude Medical Operations (M) Sdn. Bhd Plot 102, Lebuhraya Kampung Jawa, Bayan Lepas Industrial Zone, 11900 Penang, MALAYSIA

Signature:

Manager Regulatory Affairs

Issue Date



The following product(s) is/are approved under EC-certificate number I7 014607 0230 Rev. 00:

Product Name	Model No.	GMDN Codes	First Date of CE Marking
Microny™ II SR+	2525T	47267	1999-9-17
Zephyr™ XL DR	5826	47265	2006-5-9
Zephyr™ DR	5820	47265	2006-5-9
Zephyr™ SR	5620	47267	2006-5-9
Zephyr™ XL SR	5626	47267	2007-6-13
Assurity™	PM1240	47267	2013-3-7
Assurity™	PM2240	47265	2013-3-7
Endurity™	PM1160	47267	2013-3-7
Endurity™	PM2160	47265	2013-3-7
Allure™ Allure™ RF	PM3120	47263	2013-3-7
Allure Quadra™ RF	PM3222	47263	2013-3-7
	PM3242	47263	2013-3-7
Quadra Allure MP ™ RF	PM3262	47263	2014-7-31
Assurity MRI ™	PM1272 (MR Conditional)	47267	2014-12-18
Assurity MRI™	PM2272 (MR Conditional)	47265	2014-12-18
Endurity MRI ™	PM1172 (MR Conditional)	47267	2014-12-18
Endurity MRI ™	PM2172 (MR Conditional)	47265	2014-12-18
Endurity ™	PM1162 (MR Conditional)	47267	2014-12-18
Endurity ™	PM2162 (MR Conditional)	47265	2014-12-18
Endurity <sup>™</sup> Core	PM1140 (MR Conditional)	47267	2015-7-24
Endurity <sup>™</sup> Core	PM2140 (MR Conditional)	47265	2015-7-24
Endurity <sup>™</sup> Core	PM1152 (MR Conditional)	47267	2015-7-24
Endurity <sup>™</sup> Core	PM2152 (MR Conditional)	47265	2015-7-24
Quadra Allure ™	PM3542 (MR Conditional)	47263	2016-10-21
Quadra Allure MP TM	PM3562 (MR Conditional)	47263	2016-10-21
Zenex ™	PM1250	47267	2018-10-12
Zenex ™	PM2250	47265	2018-10-12
Zenus ™	PM1170	47267	2018-10-12
Zenus ™	PM2170	47265	2018-10-12
Zenex MRI™	PM1282 (MR Conditional)	47267	2018-10-12

Signature:	
Kothy Ocks	14Jun 2019
Kathy Berg	Issue Date
Manager Regulatory Affairs	



Product Name	Model No.	GMDN Codes	First Date of CE Marking
Zenex MRI TM	PM2282 (MR Conditional)	47265	2018-10-12
Zenus MRI TM	PM1182 (MR Conditional)	47267	2018-10-12
Zenus MRI TM	PM2182 (MR Conditional)	47265	2018-10-12

Signature:

Manager Regulatory Affairs

Issue Date

Page 4 of 4







## **Certificate**

No. Q5 014607 0231 Rev. 03

**Holder of Certificate: Abbott Medical** 

15900 Valley View Court Sylmar CA 91342

USA

**Certification Mark:** 



Scope of Certificate: Design and Development, Production and

Distribution of Implantable Pulse Generators and Implantable Cardioverter Defibrillators, Implantable

Leads for AIMDs, Programmers for AIMDs,

Application Software (external), Cardiac Rhythm Management Device Accessories (adapters,

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stylets, guidewires, tools, etc.)

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 014607 0231 Rev. 03

**Report No.:** 713237689

 Valid from:
 2022-08-12

 Valid until:
 2025-03-31

Date, 2022-08-12 Christoph Dicks

Head of Certification/Notified Body





## **Certificate**

No. Q5 014607 0231 Rev. 03

EN ISO 13485:2016 Applied Standard(s):

Medical devices - Quality management systems -

Requirements for regulatory purposes

(ISO 13485:2016) DIN EN ISO 13485:2016

**Abbott Medical** Facility(ies):

15900 Valley View Court, Sylmar CA 91342, USA

Design and Development, Production and Distribution of Implantable Pulse Generators and Implantable Cardioverter Defibrillators, Implantable Monitoring and Recording Systems, Implantable Leads for AIMDs, Programmers for AIMDs, Application Software (external), Cardiac Rhythm Management Device, Accessories (adapters, stylets, guidewires, tools, etc)

Abbott Medical

645 Almanor Avenue, Sunnyvale CA 94085, USA

Design and Development of Implantable Pulse Generators and Implantable Cardioverter Defibrillators, Implantable Monitoring and Recording Systems, Implantable Leads for AIMDs, Programmers for AIMDs, Application Software (external), Cardiac Rhythm Management Device Accessories (adapters, stylets, guidewires, tools, etc.); and returned product analysis of Implantable Cardioverter Defibrillators, Implantable Monitoring and Recording Systems and Cardiac Rhythm Management Device Accessories





## **CERTIFICATE**



This is to certify that



### **SANTE INTERNATIONAL S.A.**

Str. Mantuleasa nr. 33, Sector 2 023961 Bucuresti Romania

has implemented and maintains a Quality Management System.

#### Scope:

Import, trade and storage of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Through an audit, documented in a report, it was verified that the management system fulfills the requirements of the following standard:

ISO 9001: 2015

Certificate registration no. 497269 QM15

Valid from 2021-06-16

Valid until 2024-06-15

Date of certification 2021-06-16





**DQS GmbH** 

Markus Bleher Managing Director







### Annex to certificate Registration No. 497269 QM15

#### SANTE INTERNATIONAL S.A.

Str. Mantuleasa nr. 33, Sector 2 023961 Bucuresti Romania

#### Location

#### 075906 Sante International SA Sos. Mihai Bravu nr. 7, bl. P37-P37A, sector 2 021303 Bucuresti Romania

#### 497270

Sante International SA Str. Pupitrului, nr. 81, sect. 3 033036 Bucuresti Romania

#### 31050285

Sante International SA Calea Ghirodei, nr. 36 300327 Timisoara Romania

#### 31050284

Sante International SA Calea Dorobantilor, nr. 111 400609 Cluj-Napoca Romania

#### 31050283

Sante International SA Str. Lascar Catargi, nr. 37 700107 Iasi Romania

#### Scope

Import, trade and storage of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices. Consulting for state and private medical units.

# Storage of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

