**ACCOLADE™ EL Pacing System**

Model L321

- Extended longevity device labeled to last up to 16.7 years
- Automatic Daily Monitoring with the LATITUDE™ NXT Patient Management System
- RightRate™ MV sensor is the only sensor clinically proven to restore chronotropic competence
- Post-Operative System Test (POST) function to facilitate patient follow-up with a fully automatic device and lead check

### Mechanical Specifications

<table>
<thead>
<tr>
<th>Model</th>
<th>Type</th>
<th>Size (cm) (W x H x D)</th>
<th>Mass (g)</th>
<th>Volume (cc)</th>
<th>Connector Type (RA RV LV)</th>
<th>C-Codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>L321</td>
<td>DR</td>
<td>4.45 x 5.88 x 0.75</td>
<td>29.1</td>
<td>14.2</td>
<td>RA: IS1 – RV: IS1</td>
<td>C1785</td>
</tr>
</tbody>
</table>

### Additional Longevity Information

- Settings: pacing pulse width 0.4ms, Impedance 500Ω, LRL 60bpm, Sensor On, EGM Onset On. These calculations also assume that the pulse generator spends 6 months in Storage mode during shipping and storage, the ZIP™ telemetry use for 1 hour at implant time and for 40 minutes annually for in-clinic follow-up checks.
- The following LATITUDE usage will decrease longevity by approximately 10 months: Daily device check on, monthly full interrogations (scheduled remote follow-ups, and quarterly patient-initiated interrogations). Daily device checks and quarterly full interrogations will decrease longevity by approximately 9 months.
- Power Supply: lithium-carbon monofluoride cell; Boston Scientific; 402294.
ACCOLADE™ EL Pacing System
Model L321

Pacing Therapy

| Brady Modes          | Normal: DDD(R)/DDI(R)/VDD(R)/V-V(R)-A(AI)/R-DOO-VOO- AOO-Off  
|                     | Temporary: DDD(R)/DDI(V)-VVI(AI)-DOO-VOO-AOO-Off  

| AT/AF Management     | ATR Mode Switch, Ventricular Rate Regulation (VRR), Atrial Flutter Response (AFR), Rate Smoothing  
|                     | Right Atrial Automatic Threshold (RAAT)  
|                     | Right Ventricular Automatic Capture (RVAC)  

| Automaticity         | Automatic Gain Control (AGC) for sensitivity  
|                     | Rate Adaptive Pacing  

| Rate Adaptive Pacing| Accelerometer, RightRate™ (Minute Ventilation) or blended sensors with sensor trending function  

| RV Pacing Reduction | AV Search +, RYTHMIQ™, AV Delay to 400 ms, Rate Hysteresis  

| Rate Management     | Sudden Brady Response (SSB), PMT Termination, PVARP after PVC, Dynamic PVARP  

| Pace/Sense          | Unipolar, Bipolar, Bipolar/Unipolar, Unipolar/Bipolar, Unipolar/Off, Bipolar/Off, Lead Safety Switch  

Patient Diagnostics

| Arrhythmia Logbook  | Event Summary, Stored Electrograms with Annotation Markers (Intervals and approximately 14 minutes all multi channel EGM, always with 10 seconds Onset and event storage prioritization). Implant activation of all available EGMs. On screen measurements of all stored signal, amplitudes and timing. Snapshot Function (up to 12 seconds trace of ECG/EGM display stored).  

| Histograms & Counters| Ventricular Tachy Counter, Brady Counter, Histograms, Intrinsic Promotion (Rate Hysteresis % successful and AVSH+ % successful).  

| Therapy/Diagnostics | Heart Rate Variability (HRV) with SDANN and ABM, Respiratory Rate Trend, Signal Artifact Monitor, AT/AF Burden, Activity Level, A & V Arrhythmias, Weight and Blood Pressure*  

| Atrial Arrhythmia Report | AT/AF% and Total Time in AT/AF, AT/AF Burden Trend, RV Rate during AT/AF Trend, Pacing Percent Trend, Heart Rate Trend, Activity Level and Respiratory Rate Trends, RV Rate during AT/AF Histogram. Timeline history of interrogations, programming, and counter resets for one year. Longest AT/AF, Fastest RVs rate in AT/AF, and most recent episode.  

| DAILY TREND for last 365 Days | Events, Activity Level, AT/AF Burden, Pacing Percent, Respiratory Rate, Heart Rate, SDANN, HRV Footprint, ABM, Lead Impedance and Amplitude, RAAT Trend, RVAC Trend  

Implant/In-Clinic Follow-Up

| Implant Communication Mode | Programmable values: Enable use of ZIP™ telemetry (MICS) (Requires initial use of wand for device ID) or use wand for all telemetry  
|                           | Nominal: Enable use of ZIP telemetry (Requires initial use of wand for device ID)  

| In-Clinic Follow-Up | Snapshot Function: 12 seconds trace of ECG/EGM display stored  
|                     | POST (Post-Operative System Test): provides an automatic device/lead check at a pre-determined time post-implant to help document proper system functionality without requiring manual system testing  

| Indications-Based Programming (IBP) | Tool that provides specific programming recommendations based on the patient’s clinical needs and primary indications  

Remote Follow-Up

| Remote Monitoring | This device is designed to be LATITUDE™ NXT enabled. LATITUDE NXT availability varies by region  

| Thresholds | Automatic storage of last successful daily PaceSafe** threshold test for all active chambers  

| Wireless | Remote follow-up for all devices (MICS)  

| Patient-Triggered Monitor (PTM) | Triggers the storage of two minutes onset and one minute post – EGMs, intervals, and annotated marker data during a symptomatic episode by placing a magnet over the device  

Safety Functions**

| Safety Core | Is intended to provide life-sustaining therapy if certain non-recoverable or repeat fault conditions occur. Safety Core operates independently and acts as a backup to these components  

| Electrocautery Protection Mode | Provides asynchronous pacing at the programmed outputs and LRL when commanded by the programmer  

**The Safety Functions do not have programmable parameters.

*Weight and Blood Pressure are only available via LATITUDE NXT.

---


All trademarks are the property of their respective owners.
Caution:

Patients may develop psychological intolerance to a pulse generator system and may experience the following:
- Dependency
- Depression
- Fear of premature battery depletion
- Fear of device malfunction
- Low cardiac output or congestive heart failure secondary to bradycardia

For a list of potential adverse events associated with MRI scanning, refer to the MRI Technical Guide.

Venous trauma (e.g., perforation; dissection, erosion); Worsening heart failure.

Current during defibrillation with internal or external paddles; Syncope; Tachyarrhythmias, which include acceleration of atrial tachycardia; Thrombosis/thromboemboli; Valve damage; Vasovagal response; Venous occlusion; (e.g., tissue damage, valve damage); Myopotential sensing; Oversensing/undersensing; Pacemaker-mediated tachycardia (PMT) (applies to dual-chamber devices only); Pericardial rub, effusion; Pneumothorax; Pulse generator migration; Shunting endocarditis; Lead dislodgement; Lead fracture; Lead insulation breakage or abrasion; Lead perforation; Lead tip deformation and/or breakage; Local tissue reaction; Loss of capture; Myocardial infarction (MI); Myocardial necrosis; Myocardial trauma (e.g., tissue damage, valve damage); Myopotential sensing; Oversensing/undersensing; Pacemaker-mediated tachycardia (PMT) (applies to dual-chamber devices only); Precordial rub, effusion; Pneumothorax; Pulse generator migration; Shunting current during defibrillation with internal or external paddles; Syncope; Tachyarrhythmias, which include acceleration of atrial tachycardia and early, recurrent atrial fibrillation; Thrombosis/thromboemboli; Valve damage; Vasovagal response; Venous occlusion; Venous trauma (e.g., perforation; dissection, erosion); Worsening heart failure.

These Boston Scientific pacemakers are contraindicated in patients who have a separate implanted cardioverter defibrillator (ICD) with transvenous leads.

Use of certain pacing modes and/or features available in these Boston Scientific pacemakers is contraindicated for the following patients under the circumstances listed:
- Unipolar pacing or use of the My Sensor with a Subcutaneous Implantable Cardioverter Defibrillator (S-ICD) because it may cause inappropriate therapy or inhibition of appropriate S-ICD therapy.

Warnings

Read this manual thoroughly before implantation to avoid damage to the pulse generator and/or lead. Such damage can result in patient injury or death. For single patient use only. Do not re-use, re-process, or re-sterilize. Always have external defibrillation equipment available during implant and electrophysiologic testing. Using multiple pulse generators could cause pulse generator interaction, resulting in patient injury or a lack of therapy delivery. In response to applicable non-recoverable or repeat fault conditions, the pulse generator will switch irreversibly to Safety Core operation. Do not kink, twist, or braid the lead with other leads. Do not use atrial tracking modes in patients with chronic refractory atrial tachyarrhythmias. Lead Safety Switch should be programmed Off for patients with an ICD. Unipolar pacing due to Lead Safety Switch is contraindicated for patients with an ICD. Automatic Lead Recognition should be programmed to Off before implant for patients with an ICD. Unipolar pacing due to RAAT is contraindicated and should be programmed off for patients with an ICD. If programmed to a fixed atrial Sensitivity value of 0.15 mV or a fixed sensitivity value of 3.0 mV or less in a unipolar lead configuration in any chamber, the pulse generator may be more susceptible to electromagnetic interference. Advise patients to seek medical guidance before entering environments that could adversely affect the operation of the active implantable medical device. ACCOLADE MRI, PROPONENT MRI, ESSENTIO MRI, FORMIO MRI, VITALIO MRI and INGENIO MRI devices are considered MRI Conditional. For these devices, unless all of the MRI Conditions of Use are met, MRI scanning of the patient does not meet MRI Conditional requirements for the implanted system. Significant harm to or death of the patient and/or damage to the implanted system may result. All other devices covered by this statement are not MR Conditional. For potential adverse events applicable when the Conditions of Use are met or not met, refer to the MRI Technical Guide. Do not subject a patient with an implanted pulse generator and/or lead to diathermy.

Precautions

For specific information on precautions, refer to the following sections of the product labeling: clinical considerations, sterilization and storage, implantation, device programming, environmental and medical therapy hazards, hospital and medical environments, home and occupational environments, follow up testing, explant and disposal, supplemental precautionary information.

Potential Adverse Events

Based on the literature and on pulse generator and/or lead implant experience, the following alphabetical list includes the possible adverse events associated with implantation of the included devices: Air embolism; Allergic reaction; Bleeding; Bradycardia; Cardiac tamponade; Chronic nerve damage; Component failure; Conductor coil fracture; Death; Elevated thresholds; Erosion; Excessive fibrotic tissue growth; Extracardiac stimulation (muscle/nerve stimulation); Fluid accumulation; Foreign body rejection phenomena; Formation of hematomas or seromas; Heart block; Heart failure following chronic RV apical pacing; Inappropriate pacing; Infrasonic pain; Incomplete lead connection with pulse generator; Infection including endocarditis; Lead dislodgement; Lead fracture; Lead insulation breakage or abrasion; Lead perforation; Lead tip deformation and/or breakage; Local tissue reaction; Loss of capture; Myocardial infarction (MI); Myocardial necrosis; Myocardial trauma (e.g., tissue damage, valve damage); Myopotential sensing; Oversensing/undersensing; Pacemaker-mediated tachycardia (PMT) (applies to dual-chamber devices only); Precordial rub, effusion; Pneumothorax; Pulse generator migration; Shunting current during defibrillation with internal or external paddles; Syncope; Tachyarrhythmias, which include acceleration of atrial tachycardia and early, recurrent atrial fibrillation; Thrombosis/thromboemboli; Valve damage; Vasovagal response; Venous occlusion; Venous trauma (e.g., perforation; dissection, erosion); Worsening heart failure.

For a list of potential adverse events associated with MRI scanning, refer to the MRI Technical Guide.

Patients may develop psychological intolerance to a pulse generator system and may experience the following: Dependency; Depression; Fear of premature battery depletion; Fear of device malfunction.

Caution: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete "Directions for Use" for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator’s Instructions.