OPERATOR’S MANUAL

Pacing System Analyzer (PSA)

Application for use with the LATITUDE™ Programming System

Model 3922 Pacing System Analyzer (PSA)

Model 3300 LATITUDE™ Programming System

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician trained or experienced in device implant and follow-up procedures.
# TABLE OF CONTENTS

INFORMATION FOR USE ................................................................................................................. 1
  Trademark Statement .................................................................................................................... 1
  Description and Use ...................................................................................................................... 1
    Intended Use ................................................................................................................................. 1
    Intended Audience ...................................................................................................................... 1
  Required Expertise and Knowledge .............................................................................................. 1
    Physician Supervision ................................................................................................................. 1
  Essential Performance .................................................................................................................... 2
  Contraindications ........................................................................................................................... 2

WARNINGS ........................................................................................................................................ 3

PRECAUTIONS ................................................................................................................................. 6
  General ........................................................................................................................................... 6
  Adverse Effects ............................................................................................................................... 7

PSA FEATURES .................................................................................................................................. 7

PSA APPLICATION OVERVIEW .................................................................................................... 9

SYSTEM ACCESSORIES .................................................................................................................. 9
  Optional External Equipment ......................................................................................................... 10

PSA SETUP AND CONNECTION ..................................................................................................... 10
  Connect the PSA cable to the Programmer and Leads ................................................................... 11
    PSA Dual Chamber Brady Leads Hookup, example .................................................................... 13
    PSA Quadripolar Lead Hookup, example .................................................................................. 14

NAVIGATING THE PSA SETTINGS .................................................................................................. 15
  Ventricular Sensing ....................................................................................................................... 15
  LV Quadripolar Support ............................................................................................................... 15
  Current of Injury (COI) Support ..................................................................................................... 16
  Screen Layout and options .............................................................................................................. 17
    PSA Main Screen panels ............................................................................................................. 17
    Lead Traces panel ......................................................................................................................... 18
    Pacing and Amplitude panel ......................................................................................................... 19
    PSA Settings panel ....................................................................................................................... 19
    Test Chamber Measurements panel ............................................................................................ 20

LEAD IMPLANT EVALUATION STEPS ............................................................................................ 22
  1. Preparation ................................................................................................................................. 22
  2. Measure P/R wave amplitude and Current of Injury ................................................................. 22
  3. Complete a Pacing Threshold Test ............................................................................................ 23
  4. Store and Save Lead Evaluation Data ....................................................................................... 24

PSA - MORE TESTS ......................................................................................................................... 25
  Conduction Test Support .............................................................................................................. 25
  Antegrade Conduction Test .......................................................................................................... 25
  Retrograde Conduction Test ......................................................................................................... 26
  Burst Pacing ..................................................................................................................................... 26

PSA - Test Results ............................................................................................................................ 27

STAT BUTTON .................................................................................................................................. 28

REAL-TIME LOGS ............................................................................................................................ 31
  PSA Real-time Log .......................................................................................................................... 32
  Electronic Calipers .......................................................................................................................... 32
Real-time Log Tools .................................................................................. 32
TEST REPORTS .......................................................................................... 33
PSA Reports ............................................................................................. 33
End Session ............................................................................................. 34
PSA EVENTS, NOISE DETECTION, PARAMETERS, AND SPECIFICATIONS .................................................. 35
Noise Detection ....................................................................................... 35
Programmable Parameters .................................................................... 35
MAINTENANCE, TROUBLESHOOTING, SERVICE, AND STANDARDS .... 39
WARRANTY INFORMATION ..................................................................... 39
INFORMATION FOR USE

Trademark Statement
The following are trademarks of Boston Scientific or its affiliates: LATITUDE, and Quick Start.
DisplayPort is a trademark of the Video Electronics Standards Association (VESA).

Description and Use
The Pacing System Analyzer (PSA) is an application of the LATITUDE™ Programming System, Model 3300, which is a portable cardiac rhythm management system designed to be used with specific Boston Scientific (BSC) systems, i.e. implantable pulse generators (PGs) and leads.
The PSA application provides the ability to:
• Assess electrical performance and placement of cardiac lead systems during implant of cardiac rhythm management devices and provide other diagnostic information.

NOTE: The screen images used in this manual are representative and may not exactly match your screens.

Intended Use
The LATITUDE Programming System, Model 3300 is intended for use in hospital and clinical environments to communicate with Boston Scientific implantable systems. The PSA application is intended to be used during the implantation of pacemakers and defibrillators (including Cardiac Resynchronization Therapy or CRT devices) to evaluate the placement of pacing and defibrillation leads.

Intended Audience
This literature is intended for use by professionals trained or experienced in device implant procedures.

Required Expertise and Knowledge
Users must be thoroughly familiar with electrotherapy of the heart. Only qualified medical specialists and professionals having the special knowledge required for the proper use of the device are permitted to use it.

Physician Supervision
The LATITUDE Programming System may only be operated under the constant supervision of a physician. During a procedure, the patient must be continuously monitored by medical personnel with the aid of a surface ECG monitor.
Essential Performance

In order for the LATITUDE Programming System to meet its intended use, it must communicate with Boston Scientific implantable pulse generators. Therefore those functions that pertain to communications with the implanted pulse generators using telemetry wands are considered essential performance.

LATITUDE Programming System performance determined to be essential by Boston Scientific for electromagnetic compatibility (EMC) testing, as per IEC 60601-1-2, is the ability to:

- Initiate a STAT PACE, PSA STAT PACE, STAT SHOCK, or DIVERT THERAPY command to a PG where supported
- Display real-time intracardiac electrograms
- Supports touchscreen tap and button press interactions
- Deliver pacing and perform impedance lead measurements with the Pacing System Analyzer (PSA) function

Boston Scientific hereby declares that this device is in compliance with the essential requirements and other relevant provisions of Directive 1999/5/EC for Radio and Telecommunications Terminal Equipment (RTTE). To obtain a full text Declaration of Conformity, contact Boston Scientific using the information on the back cover of this manual.

**NOTE:** No recurring calibration of the LATITUDE Programming System or its applications is required or needed.

Contraindications

The LATITUDE Programming System is contraindicated for use with any pulse generator other than a Boston Scientific pulse generator. For contraindications for use related to the pulse generator, refer to the associated product literature for the pulse generator being interrogated.

The PSA application is contraindicated for use with any programming system other than the Boston Scientific Model 3300 LATITUDE™ Programming System.

The following uses of the PSA are contraindicated:

- With AV conduction disorders; atrial single-chamber pacing
- With competing intrinsic rhythms; asynchronous modes
- With chronic atrial tachycardia as well as chronic atrial fibrillation or flutter; modes with atrial control (DDD, VDD)
- With poor tolerance of high ventricular rates (e.g., with angina pectoris); tracking modes (i.e., atrial control modes) and propensity for atrial tachycardia
- Use as an external pacemaker

a. During the duration of the implantation, the Programmer PSA application is suitable for temporary external pacing while the patient is being continuously monitored by medical personnel.
WARNINGS

Refer to the LATITUDE Programming System Operator’s Manual, Model 3300 for additional Warnings.

⚠️ **Use of unspecified cables and accessories.** The use of any cables or accessories with the LATITUDE Programming System other than those provided by or specified by Boston Scientific could result in increased electromagnetic emissions, decreased electromagnetic immunity, or electrical shock of the LATITUDE Programming System. Anyone connecting such cables or accessories to the LATITUDE Programming System, including the use of MSOs (Multiple Socket Outlets), may be configuring a medical system and is responsible to ensure that the system complies with the requirements of IEC/EN 60601-1, Clause 16 for medical electrical systems.

⚠️ **Radio Frequency (RF) communications equipment.** Keep all RF communications equipment (including peripherals such as antennas, wands, and cables) at least 30 cm (12 inches) away from the Model 3300 Programmer, including cables specified by Boston Scientific, to avoid degradation of the performance of this equipment.

⚠️ **Connector contacts.** Do not simultaneously touch the patient and any accessible LATITUDE Programming System connector or exposed conductor.

⚠️ **Electric shock.** To avoid the risk of electric shock, only connect the Programmer to a grounded/earthed power source.

⚠️ **Electrostatic charges.** The PSA lead system is in electrical contact with the patients’ heart and blood.

- Do not touch the metal clips on the PSA cable or the pacing lead. Electrical currents can be dangerous to the patient and the user.
- Discharge any electrical static charge on your person by touching a grounded metal surface before touching the patient, the PSA cable or the device.

⚠️ **Electrical currents.** Unused PSA cable connections can induce electrical currents into the patient’s heart.

- Attach unused cable connections to surgical draping near the patient or disconnect the unused cables from the system.

⚠️ **Electrocautery.** The LATITUDE Programming System is designed and tested to be electrocautery safe.

- While the device is designed and tested to be electrocautery safe, electrocautery can induce electrical currents in the PSA cables that can be conducted into the patient’s heart.
Whenever possible disconnect the PSA cables from the pacing leads when performing an electrocautery procedure.

- If the Programmer is connected to the patient during an electrocautery procedure, check its operation afterwards.
- If there is an electrical overload, the Programmer will reset and reboot. During the reset and reboot, which takes about one minute, there will be no pacing support. A backup PSA/pace resource must be available in case electrocautery is applied.

**Programming System location.** Use of the PSA application on the Model 3300 Programmer adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.

**Programming System must remain outside sterile field.** The Model 3300 Programmer is non-sterile and cannot be sterilized. Do not allow the device to enter a sterile zone in an implant environment.

**Physiological signals.** Operation of the LATITUDE Programming System with physiological signals that are lower than the minimum detectable amplitude may cause inaccurate results.

**Programming System is MR Unsafe.** The LATITUDE Programming System is MR Unsafe and must remain outside the MRI site Zone III (and higher) as defined by the American College of Radiology Guidance Document for Safe MR Practices. Under no circumstances should the LATITUDE Programming System be brought into the MRI scanner room, the control room, or the MRI site Zone III or IV areas.

**Induction.** When activating PSA Burst Pacing, which may cause unpredictable arrhythmias, always have cardiac emergency equipment (e.g., external pacemaker, external defibrillator) in an operational status available for immediate life support.

- Consider additional preemptive measures in patients where acceleration or a loss of rhythm could cause life threatening danger.

**External Defibrillation.** The LATITUDE Programming System is designed and tested to be defibrillation safe.

- While the Programmer is designed and tested to be defibrillation safe, the patient can be endangered and the Programmer can be damaged.
- The PSA cable must be disconnected from the lead(s) before using external defibrillation.

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Whenever possible disconnect all cables from the patient when using external defibrillation equipment.

If the LATITUDE Programming System is connected to the patient during defibrillation, verify that the Programmer is operating after defibrillation.

**External pacing equipment.** If the patient is pacer dependent and the Programmer encounters a fault condition, pacing operation continues unless the fault was in the PSA component itself. For this reason, always have external pacing equipment available for patient back-up.

**Loss of Power.** Operating the Programmer with a depleted internal battery or no battery can suspend Programmer function if AC power is temporarily interrupted.

- If an optional battery is used, do not use a depleted or unapproved battery. For additional patient safety, when the battery level indicator shows 25% or less remaining, connect the AC power to the Programmer.
- When operating on battery power, do not attempt to replace the battery.
- An attention message displays on the Programmer screen when the battery reaches 25% depletion. When the battery reaches 10% depletion or less, an additional warning displays. At 5%, there is a warning dialog followed by a 60 second automatic shutdown.

**Loss of pacing support.** Always have external cardiac pacing equipment in an operational status available for immediate life support.

- Initially, when the Programmer is switched on, the pacing functions are switched off while a self-test is conducted. No pacing is possible during the self-test, which can take up to one minute.
- Connecting the PSA cable to the wrong lead may result in ineffective sensing and pacing behavior and loss of pacing support.
- If the user manually restarts the Programmer, pacing support is lost until the system completes its self-test, which can take up to one minute and the user must restart PSA manually if desired.
- If there is no battery installed, pacing support will be lost if AC power is lost.
- Consider additional preemptive measures in patients where loss of pacing could cause life threatening danger.

**Impaired AV conduction.** Single chamber atrial modes are contraindicated for patients with impaired AV conduction.

- If the patient has impaired AV conduction, AAI programming and antegrade conduction tests must not be performed.
Abruptly terminating pacing. Abruptly terminating pacing may result in extended periods of asystole in some patients.

- Gradually decrease the pacing rate until the patient’s intrinsic rate is detected for a controlled transition from pacing to intrinsic action.

Loss of capture. Pacing threshold testing implies loss of capture. At loss of capture, asystole and pacing during vulnerable periods can occur.

- Consider the health of the patient prior to performing a pacing threshold test.

Use of protective sleeves. Incorrect positioning of the protective silicone rubber sleeves over the PSA cable clips can cause unintended electrical connections that can impair cable function and endanger the patient.

- Before connecting cables, ensure correct position of protective sleeves.

Do not use wet cables. Moisture on wet cables can impair cable function and endanger the patient.

Equipment Modifications. No modification of this equipment is allowed unless approved by Boston Scientific.

PRECAUTIONS

Refer to the LATITUDE Programming System Operator’s Manual, Model 3300 for additional Precautions.

General

- **Functional impairment due to external damage.** Mechanical impact, for example dropping the Model 3300 Programmer unpackaged can permanently impair the function of the system. Do not use the device if there is apparent damage. If damage has occurred, contact Boston Scientific using the information on the back cover of this manual to return the unit.

- **Programming System.** Use only the appropriate Boston Scientific LATITUDE Programming System equipped with the appropriate software to perform PSA functions.

- **Stylus use.** If you want to use a stylus, ensure that it is a projected capacitance stylus. The use of any other object could damage the touchscreen.

- **Electrocautery cables.** Keep all electrocautery cables at least 30 cm (12 in.) away from the LATITUDE Programming System to avoid false signals due to electrocautery energy.
• **Leakage current.** Although optional external equipment connected to the Model 3300 Programmer meets leakage-current requirements for commercial products, it may not meet the more stringent leakage requirements for medical products. Consequently, all external equipment must be kept outside the patient environment.
  - Never touch the electrical contacts on the side panels of the Model 3300 Programmer and the patient, the telemetry wand, or any cable at the same time.

• **PSA connections.** Ensure leads are connected appropriately for desired use; incorrect setup can result in pacing/sensing events, which display under a different chamber on the screen. The PSA application user interface (UI) associates specific lead connections with the RA, RV and LV chambers on screen to support testing all three chambers with minimal change of physical connections. Saved PSA measurements are also labeled automatically based upon the chamber in use on the screen. These labels can later be adjusted by the user if the decision is made to use one physical connection to test other chambers (for example, using only the RV connection to test RA, RV and LV leads).

• **PSA connector clips.** Do not clip any PSA connector directly to the skin, pocket, or other tissue of the patient.

• **Ventricular Sensing.** During a PSA session, ventricular sensing behavior is driven by the most recently selected ventricular pacing configuration: RV-only, LV-only, or Bi-V.
  - At system startup, the PSA mode is set to ODO (non-pacing) and the effective ventricular pacing configuration is Bi-V.
  - When a non-pacing mode (ODO or OVO) is selected from the Mode panel, sensing is set to Bi-V to ensure sensing is enabled on both leads regardless of any prior configuration.

**Adverse Effects**

None known.

**PSA FEATURES**

The Pacing System Analyzer application determines in-situ lead characteristics of impedance, capture threshold, P/R wave amplitude, P/R wave, and slew rate. It supports three chambers (RA, RV, and LV) and provides the following features and functions:

• Real-time surface ECG
• Real-time intrinsic E-grams (EGM)
• Real-time Brady event markers (Pace, Sense, Noise)
• Brady settings (programmable modes are ODO, OAO, OVO, AOO, VOO, DOO, AAI, VVI, VDI, DDI, VDD, and DDD)
• Real-time heart rate display
• Intrinsic amplitude(s)
• Intrinsic P/R Interval
• Slew rate
• Pace impedances
• Pace threshold tests (amplitude and pulse-width)
• PSA STAT pacing operation
• Conduction testing (Antegrade and Retrograde)
• Burst pacing
• High output pacing (10V @ 2ms) for evaluating Phrenic Nerve Stimulation (PNS)
• RV-LV interval
• Detailed EGM viewing for Current of Injury (COI) diagnostics
• LV quadripolar testing support
• Noise filter for 50Hz and 60Hz frequencies
• Real-time Log storage and viewing
• Test results screen

The Pacing System Analyzer application performs the following functions:
• Display real time lead signals for testing RA, RV and LV Leads (including quadripolar leads) that are appropriately connected to the Programmer via PSA cables
• Display real-time signals for surface ECG and telemetered PG EGM signals (if in session with implanted device)
• Capture, annotate and review Real-time Log recordings of lead signal traces and markers
• Provide PSA configuration parameters for pacing and sensing, including burst pacing therapy
• Provide ability to perform and (as applicable) record lead evaluation results: intrinsic amplitude, slew rate, impedance, threshold and timing
• Provide ability to review recorded results, and save (to a USB pen drive or the Programmer hard drive) or print the PSA results

The Programming System supports PSA operation by:
• Displaying the PSA user interface on an external display during implant
• Exporting saved patient data from the Programmer hard drive to a removable USB pen drive
• Providing the option to encrypt patient data prior to exporting to a USB pen drive
• Transferring final measured data to the implanted PG (if in session with the implanted device)

Refer to the LATITUDE Programming System Operator’s Manual, Model 3300 for details on the operation of other features.

PSA APPLICATION OVERVIEW

The Pacing System Analyzer application is used to assess electrical performance and placement of cardiac lead systems during implant of cardiac rhythm management devices.

The following will help integrate the data, organize the behavior, and provide optimal flexibility to switch between the PSA and the PG application during implant. When these steps are used, all saved data is organized together and associated with the PG Model/Serial being implanted.

1. Identify the PG to be implanted and start a session/interrogate using the Quick Start button.
2. Access the PSA application from within the PG application once the PG session has started.
3. Switch back and forth between PSA and PG applications as desired during the procedure.

**NOTE:** Even if the PSA application is switched to a PG session, PSA operation (pacing and sensing) continues until the Programmer is turned off.

**NOTE:** Boston Scientific recommends using PSA within the PG session because the data can be easily transferred to the pulse generator.

SYSTEM ACCESSORIES

The Pacing System Analyzer application of the Programming System supports use of the following accessories:

• Model 6763 PSA Cable, re-sterilizable and re-usable; the cable clip protective covers contain Elastosil R 401, (silicone rubber)
• Model 6697 (Remington Model S-101-97) PSA Disposable Cable, single use only and requires a Model 6133 Safety Adapter
• Model 6133 (Remington Model ADAP-2R) Safety Adapter
WARNING: The use of any cables or accessories with the LATITUDE Programming System other than those provided by or specified by Boston Scientific could result in increased electromagnetic emissions, decreased electromagnetic immunity, or electrical shock of the LATITUDE Programming System. Anyone connecting such cables or accessories to the LATITUDE Programming System, including the use of MSOs (Multiple Socket Outlets), may be configuring a medical system and is responsible to ensure that the system complies with the requirements of IEC/EN 60601-1, Clause 16 for medical electrical systems.

Optional External Equipment

For information about optional external equipment, refer to the LATITUDE Programming System Operator's Manual, Model 3300.

PSA SETUP AND CONNECTION

Before starting a PSA session, the LATITUDE Programming System must be started and the PG should be interrogated.

1. Ensure that the PSA cable(s) are sterile.

   The Model 6763 PSA cable is shipped non-sterile. If this cable is being used in a sterile procedure, then follow the sterilization procedures in the Instructions For Use (IFU) for this PSA cable.

2. Select the PSA button to turn on the PSA function (Figure 2 on page 11).

   NOTE: Once the PSA application starts it continues to operate until the Programmer is turned off.

3. Continue with “Connect the PSA cable to the Programmer and Leads” on page 11.
**NOTE:** Manually powering Off the Programmer and powering it back On resets all PSA parameters to the nominal values.

**CAUTION:** If you want to use a stylus, ensure that it is a projected capacitance stylus. The use of any other object could damage the touchscreen.

**NOTE:** The screen images in this manual are representative and may not exactly match the screens displayed.

[1] PSA application button

**Figure 2.** Main PG screen after Quick Start

**Connect the PSA cable to the Programmer and Leads**

For the PSA cable connection, refer to the illustration of the Programming System right side (Figure 3 on page 12).

For an example of a dual lead PSA connection, refer to Figure 4 on page 13.
For an example of a quadripolar PSA connection, refer to Figure 5 on page 14.
1. Connect the PSA cable to the appropriate connector (LV or A/RV) on the right side panel of the Programming System.

2. For cables with protective sleeves (e.g. Model 6763 PSA cable), position the protective sleeves so that they cover the cable clips.

   **NOTE:** The Model 6763 PSA cable protective sleeves must cover the clips when in use.

3. Connect the PSA cable clips to the lead(s) and consider the following:
   
   a. Clips of cable and leads.
      
      - Do not allow yourself or others to touch the metal clips on the PSA cable or the pacing lead. The device is in electrical contact with the patient’s heart and blood via the implanted leads.
      
      - Touching the metal clips on the PSA cable or the pacing lead may expose the patient’s heart to dangerous electrical currents.

   b. Connecting PSA cable to leads.
      
      - Verify that the PSA cable clips are attached to the correct lead(s).
      
      - Connecting the PSA cable clips to the wrong lead may result in ineffective sensing and pacing behavior and loss of pacing support.

      **NOTE:** Refer to Figure 1 on page 10 for PSA cable connector identification.

      **NOTE:** Refer to Figure 13 on page 22 for a lead connection example.
c. Keep PSA cable dry.
   - Do not use wet cables.

d. Unused PSA cable connections.
   - Attach unused cable connections to surgical draping near the patient.

**CAUTION:** Ensure the left side of the unit is accessible at all times so that the power cord can be disconnected.

⚠️ **WARNING:** The PSA cable **must** be disconnected from the lead(s) before using external defibrillation.

⚠️ **WARNING:** Do not use the Programming System adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, check the Programming System for normal operation in that configuration.

This completes the PSA setup section. Continue with “Navigating the PSA Settings” on page 15.

**PSA Dual Chamber Brady Leads Hookup, example**

Figure 4 illustrates the proper PSA cable hookup for dual chamber Brady leads.

![PSA Dual Chamber Lead hookup, example using Model 6763 PSA cable](image)


**Figure 4.** PSA Dual Chamber Lead hookup, example using Model 6763 PSA cable

**NOTE:** See Model 6697/S-101-97 IFU for cable connection information.
PSA Quadripolar Lead Hookup, example

When using a quadripolar lead, Figure 5 on page 14 illustrates the proper PSA cable hookup for a unipolar configuration.

If you want a unipolar configuration using the Can as a vector, use any LV lead electrode as a cathode, move the A+ connector from the atrial lead to a temporary, indifferent electrode (e.g. hemostat, pocket spreader) placed at the patient’s implant site to serve as the anode. Check the box for the “Use the A+ connection ...,” then check the desired Can button (see Figure 13 on page 22).

To end a unipolar configuration you must deselect the Can button and deselect the “Use the A+ connection ...” box to return to normal operation of the atrial lead anode (see Figure 13 on page 22).

CAUTION: Do not clip any PSA connector directly to the skin, pocket, or other tissue of the patient.

Note that the A+ connector on the PSA cable may be attached to the patient’s implant site to simulate the PG device connection.

NOTE: For unipolar configurations, attach the A+ connector on the PSA cable to a temporary, indifferent electrode (e.g. hemostat, pocket spreader) placed at the patient’s implant site to simulate the PG device connection.


Figure 5. PSA Quad Lead Hookup using Model 6763 PSA cable

NOTE: See Model 6697/S-101-97 IFU for cable connection information.
NAVIgATING THE PSA SETTinGS

When the PSA application is started, a check of the battery level is performed. The user is warned that PSA support may be terminated if the programmer’s optional internal battery is at a low level (or missing) in the event that AC power is lost.

**NOTE:** The PSA performs a self-test prior to each use. If the self-test fails, the PSA considers this a non-recoverable fault. The PSA monitors for non-recoverable faults while active and indicates to the user when it has experienced one.

**NOTE:** If the PSA is programmed in a Brady Pacing mode when a non-recoverable fault occurs, PSA falls back to the nominal set of pacing parameters in DOO pacing mode with the LV vector maintained as previously programmed.

**NOTE:** Once the PSA application starts it continues to operate until the Programmer is turned off.

**Ventricular Sensing**

During a PSA session, ventricular sensing behavior is driven by the most recently selected ventricular pacing configuration: RV-only, LV-only, or Bi-V.

At system startup, the PSA mode is always set to ODO Bi-V, which is the default setting. Sensing chamber options include:

- BiV enabled: sensing (and pacing if in a pacing mode) in both the RV and LV
- RV-only enabled: sensing (and pacing if in a pacing mode) in the RV but not LV
- LV-only enabled: sensing (and pacing if in a pacing mode) in the LV but not RV

**LV Quadripolar Support**

CRT can improve survival and symptoms in patients with heart failure and LBBB. However, lead location, phrenic nerve stimulation, timing between RV and LV, and high capture thresholds can impact benefit. Quadripolar LV leads, compared to bipolar leads, maybe associated with improved survival and decreased risk of replacement and deactivation. Ongoing follow-up and vector configuration of the quadripolar lead may be essential to sustain its potential benefits.

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The LV Quadripolar feature supports the implant assessment of left-ventricle leads. It allows additional vectors to be used when evaluating and configuring the location for lead operation.

The LV Quadripolar feature provides organized control of the LV pacing/sensing vectors, thus avoiding manual repositioning of the pacing cable clips by the user for each vector test. It provides measurement of the time between the RV and LV signals and displays that measurement to the user as a surrogate replacement measure for a QLV interval measurement.

The role the PSA plays in LV Quadripolar support is to:

- Provide an electrical/mechanical interface that does not require manual repositioning of the PSA cable clips to test each vector
- Support programmatic control of the LV pacing/sensing vector

This feature is designed to allow evaluation convenience and operational efficiency for the user.

**NOTE:** PSA uses the same sensing vector as the pacing vector for LV leads.

**Current of Injury (COI) Support**

Selecting the Current of Injury button (callout [4] in Figure 6 on page 17) provides information that can be used in addition to the measured pacing information (i.e. pacing threshold, sensing) and may assist in determining adequate lead position.

The COI feature displays the injury to the myocardium at the site of anchoring the active-fixation lead. The COI manifests itself as an increase in the duration of the intracardiac electrogram and elevation of the ST-segment compared to the baseline. COI may be recorded during placement of passive leads as well, presumably because of focally injured cell membranes consequent to the trauma of electrode pressure against the endocardium. With active-fixation leads, the ST-segment elevation is expected to be even more pronounced. It has been shown that magnitude of ST-segment elevation can predict adequate performance in the acute phase during implant of active-fixation leads. Studies have suggested ‘adequate values of COI measured’ in order to predict good midterm performance of the lead. It should be noted that COI is readily visualized when present without the need for specific measurements. Boston Scientific does not make recommendations of ST segment elevation measurements that are representative of an adequate COI.

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a. References:
The role the PSA plays in COI is to minimize filtering of displayed EGM signals in order to preserve signal morphology, and to isolate the most recent EGM cycle to promote visual detection and measurement of changes in morphology. This feature is an enhancement (for user convenience) that allows the user to zoom into a single real-time waveform for the chamber selected. The waveform display is updated each time a pace or sense event is detected by the PSA in the selected chamber. The view allows a high resolution view of each waveform in order to allow observable changes to the intrinsic waveform in real-time.

**Screen Layout and options**

**PSA Main Screen panels**

This section provides details on each of the three PSA main screen panels:

1. Lead Traces (page 18)
2. PSA Pacing and Amplitude (page 19)
3. PSA Test Chamber Measurements (page 20)

![PSA Main Screen Layout](image)


**Figure 6.** PSA Main Screen Layout
Figure 7.  PSA Lead trace selection examples, low voltage PG (Lead-I and PSA A)

Lead Traces panel

The PSA displays real-time surface ECG, EGM traces, and event markers for each enabled channel (lead) including a heart rate indicator.

**NOTE:** Before evaluating the lead(s), confirm that the PSA EGMs are selected using the Lead trace selections (Figure 7 on page 18).

**NOTE:** PSA Test Results and Real-time Logs should be saved before powering off the Programmer in order to avoid losing the PSA data.

Event markers generated by the PSA may overlap on the Real-time EGM display based upon the selected display speed and event intervals. If overlap occurs, the most recent marker information will be visible as the top layer. To reduce/remove overlap, the Real-time display speed can be adjusted. Also, a Snapshot or Real-time Log can be captured for review at an appropriate display speed.

- Up to four real time traces can be displayed (see callout [1] in Figure 6 on page 17). Selecting a lead trace button displays the Real-time Trace Selection panel. Figure 7 lists two of the lead trace names (Lead-I and PSA A) for a low voltage PG. Other selections display when high voltage PGs are interrogated.

- The Magnify Traces button enlarges the lead trace area to fill the display window and provides additional information at the bottom of the traces display. See Figure 8.
  - The Calibrate button transmits a 1 mV calibration pulse so the user has a reference point to evaluate amplitudes.
  - The Baseline button forces the trace back to the baseline, and is normally used after a defibrillation shock.

Figure 8.  Lead Traces panel example (lower portion, magnified)
**Pacing and Amplitude panel**

In preparing for PSA tests, verify the settings on the PSA Pacing and Amplitude panel and the PSA Settings panel.

From the PSA Pacing and Amplitude panel, verify the Mode, Lower Rate, Pacing chamber, and Amplitude. Make modifications as necessary.

**NOTES:**

- The Settings magnify button provides for additional PSA settings (see “PSA Settings panel” on page 19)
- The More Tests magnify button provides for More Tests (see “PSA - More Tests” on page 25)
- The Test Results magnify button provides for Test Results (see “PSA - Test Results” on page 27)

![PSA Pacing and Amplitude panel](image)

**Figure 9. PSA Pacing and Amplitude panel**

**PSA Settings panel**

From the PSA Pacing and Amplitude panel, click on the Settings button to display the PSA Settings panel. Verify the Parameters and Pacing and Sensing settings prior to beginning a lead testing session. Make modifications as necessary.

![PSA Settings panel](image)

**Figure 10. PSA Settings panel**
**Test Chamber Measurements panel**

The information for each lead (P/R Wave, Slew, Impedance, and RV-LV Interval) updates on a beat by beat basis when the PSA clips are attached to the corresponding lead.

From the Test Chamber Measurements panel (Figure 11), use the magnify buttons (A, RV, and LV) to select the chamber to test.

![Magnify buttons for A, RV, and LV lead Thresholds](image)

**Figure 11. PSA Test Chamber Measurements panel**

Use the Magnify button for each lead threshold to be tested to start pacing and adjust the threshold parameters on the Threshold panel (Figure 12).

Once threshold is determined, click the Save Threshold button to store the result in Test Results.
On the PSA LV Threshold panel, select the LV Pace/Sense Vector button to configure the desired cathode/anode pacing and sensing configuration (Figure 13 on page 22).

Be sure to check the “Use the A+ connection...” when a configuration including the Can vector is desired, and ensure that the A+ PSA clip uses an indifferent electrode to make electrical contact with the patient in the sterile field.

CAUTION: Do not clip any PSA connector directly to the skin, pocket, or other tissue of the patient.
LEAD IMPLANT EVALUATION STEPS

1. Preparation
   
   1. Interrogate the PG.
   
   2. Select the PSA button in the upper right of the screen.
   
   3. Change the Real-time lead trace selection(s) to view the PSA lead trace(s). Refer to callout [1] in Figure 6 on page 17 and the PSA Lead Trace selections in Figure 7 on page 18).
   
   4. Use the PSA Settings button (callout [8] in Figure 6 on page 17) to open the PSA Settings panel (Figure 10 on page 19). Then, select/confirm the desired parameters for the PSA settings. Select the Close button to close the panel and continue the session.

2. Measure P/R wave amplitude and Current of Injury

   1. Use the Test Chamber Measurements panel (Figure 11 on page 20) to assess the P wave, R wave, and Slew rate, for the attached lead(s). The RV-LV interval may also be assessed.

   **NOTE:** If the signal is noisy, first try to remove the source of interference. If noise is still evident on your electrogram
trace, consider turning on the filter for 50/60Hz to reduce the noise on the electrogram.

2. To assess current of injury morphology select the Current of Injury button (callout [6] in Figure 6 on page 17).

3. Complete a Pacing Threshold Test

For the following steps, refer to:
- PSA Pacing and Amplitude panel (Figure 9 on page 19)
- PSA Threshold panels (Figure 12 on page 21)
- Test Chamber Measurements panel (Figure 11 on page 20).

1. Adjust the Lower Rate to overdrive the intrinsic rate (e.g. 10 bpm above intrinsic rate) from the PSA Pacing and Amplitude panel.

2. Turn on Pacing for the lead to be threshold tested (A, RV, or LV) from the PSA Pacing and Amplitude panel. This will automatically adjust the mode setting to the appropriate value (AAI, VVI, or DDD) based on the lead(s) selected. If desired, the mode can be manually changed.

3. Check the impedance from the Test Chamber Measurements panel.
   
   **NOTE:** Impedance also displays in the current calculation box (Figure 12 on page 21).

4. From the Test Chamber Measurements panel, select the Magnify button for the desired lead (A, RV, or LV) to display the PSA Threshold panel for that lead.

5. Determine the Pacing Threshold by decrementing Amplitude or Pulse Width.

6. Press the Save Threshold button to save the data for P/R Wave, Slew, Impedance, and Threshold.
   - The most recent sensed settings are retained and, upon press of the “Save Threshold” button, are saved with the pace threshold results. So for a given lead placement, the lead’s sensing values are checked first, then pacing characteristics are checked. The settings, although not from the same instant in time are from the same lead placement location. Therefore checking sensing, then relocating or moving the lead and proceeding immediately to pacing tests would result in an inconsistent measurement.
   - This data will be saved in the PSA Test Results, and PSA report (which is accessed by pressing the Data button at the bottom of the screen to display the Data Management panel) during the active session.

   **NOTE:** A Real-time Log event is captured automatically (each time the Save Threshold button is pressed), which can
be reviewed later, saved, or printed as a PDF while in the current session.

7. Check for extracardiac stimulation by pressing and holding the “Hold for 10V @ 2ms” button from the PSA Threshold panel (see Figure 12 on page 21).
   a. If there is no stimulation, continue with the next step.
   b. If there is stimulation, adjust the amplitude and/or pulse width and check again for extracardiac stimulation. Press the PNS button to store the amplitude and pulse width where the Phrenic Nerve Stimulation (PNS) occurred.

   **NOTE:** The PNS button simply stores the most recent amplitude and pulse width in Test Results at the time the button is pressed. It does not perform a PNS test.

4. Store and Save Lead Evaluation Data

PSA results are stored in Test Results (Figure 6 on page 17) and in the PSA report. Press the Data button at the bottom of the screen to display the Data Management panel.

1. Review the Real-time Logs. Save and/or print as desired (see Figure 23 on page 31).
2. Review the PSA Test Results. Save and/or print as desired (see Figure 19 on page 28).

   **NOTE:** PSA test results and Real-time Logs should be saved or printed before powering off the Programmer in order to avoid losing the PSA data.

   **NOTE:** PSA functional state (pacing/sensing configuration) is retained when transitioning into a PG session if the PSA was used prior to interrogating a device. PSA will power off once a PG session is ended or the Programmer is powered off (manually, or loss of power). Any recorded thresholds/results, snapshots or Real-time Logs will be lost upon any transition into or out of a PG session.

   **NOTE:** If PSA is not used within a PG session, the user must manually re-enter the PSA data to the PG during the PG session.

   **NOTE:** If during implant testing, the physician changes to another PG; the PSA data must be manually entered to the new PG.
PSA - MORE TESTS

The More Tests button (see Figure 6 on page 17) is available as clinically needed. More Tests include antegrade and retrograde conduction tests and Burst Pacing as illustrated in Figure 14 on page 25.

Conduction Test Support

It has been demonstrated that 45% of patients who require dual chamber system implantation for any indication have retrograde conduction at some paced rate if paced from the ventricle. Even patients who have had AV block for many years may retain retrograde conduction.

The mean range of VA time is 110 - 450 ms. The existence of retrograde conduction via the natural pathway, and antegrade conduction via the implanted dual-chamber system provides a reentry circuit. Measurement of antegrade and retrograde conduction intervals allows evaluation of the state of AV and VA conduction as supporting evidence for device system implantation and to allow the setting of the atrial refractory interval after the ventricular event to avoid retrograde conduction and the onset of the endless loop tachycardia.

Figure 14. PSA More Tests (Antegrade and Retrograde Conduction and Burst Pacing)

When you push a conduction test button you see beat by beat conduction measurements for the selected test.

NOTE: No automatic Real-time Logs are captured for Antegrade or Retrograde conduction tests. If desired, these tests must be manually recorded using a Snapshot or the Real-time recorder. Burst Pacing does automatically capture a Real-time Log of this event.

Antegrade Conduction Test

The Antegrade Conduction Test measurement uses the AAI Brady mode with ventricular sensing enabled to measure the patient’s A-V conduction times based on either a paced or sensed atrial event.

NOTE: If there is no conduction in A, ventricular sensing continues.

Retrograde Conduction Test

The Retrograde Conduction Test measurement uses the VDI Brady mode to measure the patient’s V-A conduction times based on either a paced or sensed ventricular event.

Burst Pacing

Burst Pacing is used to induce or terminate arrhythmias when delivered to the desired chamber. Only the selected chamber receives Burst Pacing.

Burst Pacing can be enabled for an A, RV, or LV lead as illustrated in Figure 17.
To deliver Burst Pacing, perform the following steps:

**NOTE:** Before you begin Burst Pacing, ensure that pacing is active in the chamber where you will deliver the burst pacing.

1. Select the chamber (A, RV, or LV).
2. Select a Pacing Interval.
3. Check the Enable box.
4. A warning displays indicating burst pacing will be activated (Figure 18 on page 27).
5. Press and hold the “Hold for Burst” button. (There is a timeout maximum at 45 seconds for A and 30 seconds for RV and LV.)
6. If PSA Pacing is on before the burst test, PSA Pacing will resume after Burst Pacing completes.
7. An automatic Real-time recording is triggered when Burst Pacing stops.

**NOTE:** Pacing resumes (as needed) at the PSA lower rate limit and Mode (if programmed on) when Burst Pacing ends.

### PSA - Test Results

This screen presents the list of test results from the current PSA application session, including the Threshold Test Panel lead/chamber (Right Atrium, Right Ventricle or Left Ventricle) where the result was documented, the time the result was captured, as well as the Amplitude and Pulse Width captured for the result. The Notes column can be edited. The LV results, by default, contain the LV Pace/Sense Vector configured at the time of the result.

The user can edit the Lead location of a result to any of the three chambers; this supports the use case where leads in multiple chambers were tested using a single physical connection/chamber on the Programmer and PSA Application.

Check-boxes allow the user to select any and all valid and desired set(s) of results to print or save to PDF. If the PSA used within a PG application session
is Saved, the most recent selected results for each chamber\textsuperscript{a} automatically transfer to the PG application\textsuperscript{b} for storage in the PG upon a subsequent Program operation. This provides a set of data from the implant PSA session to the implanted device for future reference; it is recommended this data be captured in the PG and this functionality provides an automated replacement for a previously manual entry.

![Figure 19. PSA - Test Results](image)

### STAT BUTTON

The red STAT button, \(\text{△}\), is at the top right of the Model 3300 Programmer to provide a rescue shock or pacing. The STAT function is available in the same location at all times to initiate a PSA STAT PACE or deliver a rescue pace or shock. Pressing the STAT button displays the Emergency Function screen as shown in Figure 20 and Figure 22. Check the pulse generator labeling for specific details of the STAT parameters.

**NOTE:** Ensure there is effective connection between the PSA cable and the lead(s) before using the STAT button.

1. Press the STAT button.
   The following conditions determine the actions available when the STAT button is pressed:
   - When the PG is in “Storage,” “Off,” or “Monitor Only” mode, a STAT SHOCK / STAT PACE is delivered. If the STAT SHOCK / STAT PACE is delivered in storage, the Tachy mode changes to “Off.”
   - When in telemetry communication with a high voltage (ICD or CRT-D) PG, a pop-up displays allowing the user to initiate a STAT PACE, STAT SHOCK, or DIVERT THERAPY command. If a PSA session is in

\begin{itemize}
\item a. Maximum of 3 chambers total, one for RA, RV and LV.
\item b. The transfer is into the Patient Implant Data.
\end{itemize}
progress, a PSA STAT PACE option also displays as shown in Figure 20 on page 29.

- When in telemetry communication with a low voltage PG, a pop-up displays allowing the user to initiate a STAT PACE or DIVERT THERAPY command. If a PSA session is in progress a PSA STAT PACE option also displays.

- When not in communication with a PG, an Interrogate button displays with text prompting the user to perform Quick Start to attempt to identify the device (see Figure 22 on page 31). Once in a session with an implanted transvenous device, press the red STAT button again to display available options.

2. Select the desired action.

After the STAT button is pressed, the following occurs when an action is clicked:

- PSA STAT PACE—when a PSA session is active, it configures the PSA with STAT PACE settings and functionality.
- STAT PACE—initiates PG pace functionality specific to the supported transvenous device (ICD, CRT-D, Pacemaker/CRT-P).

**NOTE:** When selected, STAT PACE or PSA STAT PACE remain active until the Brady settings in the PG or PSA are changed.

- STAT SHOCK—initiates PG shock functionality specific to the high-voltage transvenous ICD and CRT-D pulse generators.
- DIVERT THERAPY—initiates PG divert therapy for any supported transvenous device (ICD, CRT-D, Pacemaker/CRT-P) and, while in a PG session, stops the pending therapy.

Figure 20. Red STAT Button pop-up in a High Voltage transvenous PG session with
the PSA application running

![EMERGENCY FUNCTIONS](image)

**Figure 21. Red STAT Button pop-up in a Low Voltage transvenous PG session with the PSA application running**

In Figure 20, the top row buttons (STAT PACE, DIVERT THERAPY, and STAT SHOCK) display only during a High Voltage PG session. PSA STAT PACE displays when the PSA feature is active.

In Figure 21, the top row button (STAT PACE) displays only during a Low Voltage PG session. PSA STAT PACE displays when the PSA feature is active.

If in a PSA session only (no PG interrogated), then the dialog in Figure 22 displays along with the PSA STAT PACE button.

If not in a transvenous PG session, pressing the STAT button displays the following dialog with no buttons - “There is no active device session. - Press interrogate to initiate the Quick Start function. - PSA Stat Pace is available below.”
REAL-TIME LOGS

Use the two buttons, Snapshot and Real-time Recorder, to record real-time lead traces. Examples of recorded events and a sample snapshot are shown in the following two figures.
Using the buttons on the left side of the screen, events can be selected/deselected and saved, printed or deleted. Events may be saved to the Programmer’s hard drive or to a USB pen drive.

**NOTE:** PSA events are not automatically saved when a PSA session ends. Use the Real-time Log to save, print, or delete these events before ending the PSA session.

**PSA Real-time Log**

![Real-time Log diagram](image)


Figure 24. PSA Real-time Log example

**Electronic Calipers**

Use the electronic calipers (slide bar) to measure the time span within the event. The time frame measured between the calipers is measured in seconds. A caliper can be repositioned by selecting it and then dragging it to expand or collapse the time frame. For detailed instructions on using the electronic calipers, refer to the associated product literature for the pulse generator being interrogated.

**Real-time Log Tools**

Select any part of the Real-time Log event display and the tool pop-up displays as in Figure 24. At the top center of the pop-up is an arrow and a target icon. When a tool is selected, the tool action occurs at that target point on the screen. A new tool pop-up displays each time you select another part of the Real-time Log event display, so that you can use multiple tools anywhere on the display as shown in Figure 24 on page 32.
The five tools are:

- **Circle tool** - places a circle on the display at the target point.
- **Line tool** - places a dashed vertical line on the display at the target point.
- **Left scissor tool** - creates a copy of the Real-time Log and removes the entire portion of the recording to the left of the target point. The original recording is retained.
- **Right scissor tool** - creates a copy of the Real-time Log and removes the entire portion of the recording to the right of the target point. The original recording is retained.
- **Note tool** - displays a keyboard to type in any notes, which will then appear at the bottom of Real-time Log horizontally aligned with the target point.

**TEST REPORTS**

The following information can be saved to a PSA Test Report for each lead:

- Date/Time Stamp
- Intrinsic Amplitude
- Lead Impedance
- Slew Rate
- Pace Threshold Amplitude
- Pace Threshold Pulse Width
- LV Vector (LV lead only)
- PNS (phrenic nerve stimulation)
- RV-LV Interval (LV lead only)
- Notes

**PSA Reports**

The following is an example of a PSA Report created in PDF format.
End Session

**NOTE:** PSA test results and Real-time Logs should be saved before powering off the Programmer in order to avoid losing the PSA data.

The only way to end a PSA session is to power off the Programmer. There is no Off button for the PSA application.
PSA EVENTS, NOISE DETECTION, PARAMETERS, AND SPECIFICATIONS

Table 1. PSA Events

<table>
<thead>
<tr>
<th>Event Type</th>
<th>Trigger Event</th>
<th>Duration of Recording (seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSA PACE THRESHOLD TEST (A, RV, and LV)</td>
<td>PSA Save Threshold button press</td>
<td>12</td>
</tr>
<tr>
<td>PSA BURST PACING</td>
<td>PSA Burst button release</td>
<td>24</td>
</tr>
<tr>
<td>PSA STAT PACE</td>
<td>PSA STAT PACE (commanded)</td>
<td>12</td>
</tr>
</tbody>
</table>

Table 2. Burst Pacing Programmable Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Programmable Values</th>
<th>Increment</th>
<th>Nominal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pacing Interval</td>
<td>100 - 750 ms</td>
<td>10 ms</td>
<td>240 ms</td>
</tr>
<tr>
<td>Chamber</td>
<td>A, RV, LV</td>
<td>n/a</td>
<td>n/a</td>
</tr>
</tbody>
</table>

Noise Detection

When noise is detected, the PSA switches to asynchronous pacing at the lower rate limit. The following table defines the PSA noise response:

Table 3. Noise Response

<table>
<thead>
<tr>
<th>Brady Mode</th>
<th>Noise Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>AAI</td>
<td>AOO</td>
</tr>
<tr>
<td>VVI, VDI, VDD</td>
<td>VOO</td>
</tr>
<tr>
<td>DDI, DDD</td>
<td>DOO</td>
</tr>
</tbody>
</table>

Programmable Parameters

**NOTE:** Manually powering Off the Programmer and powering it back On resets all PSA parameters to the nominal values.

Table 4. Programmable Parameter Nominals

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Nominal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brady Mode</td>
<td>ODO</td>
</tr>
<tr>
<td>Lower Rate Limit (LRL)</td>
<td>60 ppm</td>
</tr>
<tr>
<td>Parameter</td>
<td>Value</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Maximum Tracking Rate (MTR)</td>
<td>120 ppm</td>
</tr>
<tr>
<td>Ventricular Pacing Chamber</td>
<td>RV</td>
</tr>
<tr>
<td>Ventricular Sensing</td>
<td>Baseline to Peak</td>
</tr>
<tr>
<td>LV Offset</td>
<td>0 ms</td>
</tr>
<tr>
<td>AV Delay</td>
<td>120 ms</td>
</tr>
<tr>
<td>PVARP/ARP Interval</td>
<td>250 ms</td>
</tr>
<tr>
<td>VRP Interval</td>
<td>240 ms</td>
</tr>
<tr>
<td>LVRP Interval</td>
<td>250 ms</td>
</tr>
<tr>
<td>Atrial/RV/LV Pace Amplitude</td>
<td>5.0 V</td>
</tr>
<tr>
<td>Atrial/RV/LV Pulse Width</td>
<td>0.5 ms</td>
</tr>
<tr>
<td>Atrial Sensitivity</td>
<td>0.6 mV</td>
</tr>
<tr>
<td>RV Sensitivity</td>
<td>2.5 mV</td>
</tr>
<tr>
<td>LV Sensitivity</td>
<td>2.5 mV</td>
</tr>
<tr>
<td>LV Vector</td>
<td>(LV1)→(LV2)</td>
</tr>
<tr>
<td>Noise Filter</td>
<td>OFF</td>
</tr>
</tbody>
</table>

**Table 5. PSA STAT PACE Parameters**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brady Mode</td>
<td>VVI</td>
</tr>
<tr>
<td>Lower Rate Limit (LRL)</td>
<td>60 ppm</td>
</tr>
<tr>
<td>Ventricular Pacing Chamber</td>
<td>BiV</td>
</tr>
<tr>
<td>LV Offset</td>
<td>0 ms</td>
</tr>
<tr>
<td>RV/LV Pace Amplitude</td>
<td>7.5 V</td>
</tr>
<tr>
<td>RV/LV pulse width</td>
<td>1.0 ms</td>
</tr>
<tr>
<td>RV/LV Sensitivity</td>
<td>2.5 mV</td>
</tr>
<tr>
<td>LV Vector</td>
<td>(LV1) - (LV2)</td>
</tr>
</tbody>
</table>

**Table 6. Parameter Ranges**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSA parameters</td>
<td></td>
</tr>
<tr>
<td>Mode</td>
<td>ODO, OAO, OVO, AOO, VOO, DOO, AAI, VVI, VDI, DDI, VDD, and DDD</td>
</tr>
<tr>
<td>Lower Rate Limit (LRL)</td>
<td>30 - 175 ppm in increments of 5 ppm</td>
</tr>
<tr>
<td>Maximum Tracking Rate (MTR)</td>
<td>50 - 175 ppm in increments of 5 ppm</td>
</tr>
<tr>
<td>Parameter</td>
<td>Range</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>AV delay</td>
<td>30 - 300 ms in increments of 10 ms</td>
</tr>
<tr>
<td>LV Offset</td>
<td>± 100 ms in increments of 10 ms</td>
</tr>
<tr>
<td>PVARP/ARP Interval</td>
<td>150 - 500 ms in increments of 10 ms</td>
</tr>
<tr>
<td>Paced VRP interval</td>
<td>150 - 500 ms in increments of 10 ms</td>
</tr>
<tr>
<td>Paced LVRP interval</td>
<td>150 - 500 ms in increments of 10 ms</td>
</tr>
<tr>
<td>Filter values</td>
<td>Off, 50 Hz, 60 Hz</td>
</tr>
<tr>
<td>Ventricular Pacing Chamber</td>
<td>BiV, RV, or LV</td>
</tr>
<tr>
<td>LV Pace /Sense vector</td>
<td>E1 to E2/E3/E4/Coil/Can</td>
</tr>
<tr>
<td></td>
<td>E2 to E3/E4/Coil/Can</td>
</tr>
<tr>
<td></td>
<td>E3 to E2/E4/Coil/Can</td>
</tr>
<tr>
<td></td>
<td>E4 to E2/E3/Coil/Can</td>
</tr>
<tr>
<td>PSA EGM channel gain</td>
<td>0.5, 1.0, 2.0, 5.0, and 10.0 mm/mV</td>
</tr>
<tr>
<td>Burst Pacing Interval</td>
<td>100 - 750 ms in increments of 10 ms</td>
</tr>
<tr>
<td></td>
<td>80 - 600 ppm in various increments</td>
</tr>
<tr>
<td></td>
<td>(maximum duration of 45 seconds for A and 30 seconds for RV and LV)</td>
</tr>
<tr>
<td>Noise Filter</td>
<td>Off, 50 Hz, 60 Hz</td>
</tr>
<tr>
<td>Atrial, LV, or RV pacing amplitude</td>
<td>0.1 - 5.0 V in increments of 0.1 V and</td>
</tr>
<tr>
<td></td>
<td>between 5.0 - 10.0 V in increments of 0.5 V</td>
</tr>
<tr>
<td>Atrial, LV, or RV pulse width</td>
<td>0.1 - 2.0 ms in increments of 0.1 ms</td>
</tr>
<tr>
<td>Atrial, RV, or LV sensitivity</td>
<td>0.2 - 1.0 mV in increments of 0.2 mV</td>
</tr>
<tr>
<td></td>
<td>1.0 - 8.0 mV in increments of 0.5 mV</td>
</tr>
<tr>
<td></td>
<td>8.0 - 10.0 mV in increments of 1.0 mV</td>
</tr>
<tr>
<td>Traces</td>
<td>Lead-I, Lead-II, Lead-III, Lead-aVR,</td>
</tr>
<tr>
<td></td>
<td>Lead-aVL, Lead-aVF, Lead-V</td>
</tr>
<tr>
<td>Surface Gain</td>
<td>Auto, 0.5, 1, 2, 5, 10, 20 mm/mV</td>
</tr>
<tr>
<td>Trace Speed</td>
<td>0, 25, 50 mm/s</td>
</tr>
<tr>
<td>Show PSA Markers</td>
<td>Off, On</td>
</tr>
<tr>
<td>Enable Surface Filter</td>
<td>Off, On</td>
</tr>
<tr>
<td>Display Pacing Spikes</td>
<td>Off, On</td>
</tr>
<tr>
<td>P/R wave amplitude</td>
<td>0.25 - 30 mV with an accuracy of ± 10%</td>
</tr>
<tr>
<td></td>
<td>or ± 0.2 mV</td>
</tr>
<tr>
<td>P/R wave interval</td>
<td>0 - 500 ms</td>
</tr>
</tbody>
</table>
Parameter | Range
---|---
Lead Impedance | For pacing amplitudes ≥ 0.5 V and ≤ 7.5 V and pulse widths ≥ 400 μs, the accuracy is ± 10%.
| For impedances 100 - 2499 Ω and 2500 - 3000 Ω, the accuracy is ± 20% for pacing voltages ≤ 1.1 V and ± 15% for pacing voltages > 1.1 V.
Conduction Rate | 30 - 175 ppm in increments of 5 ppm
Conduction Amplitude | 0.1 - 5.0 V in increments of 0.1 V and between 5.0 - 10.0 V in increments of 0.5 V
Slew Rate | 0.5 - 4.0 V/s with an accuracy of ± 0.2 V/s or ± 20% whichever is greater

Table 7. PSA Markers

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>AS</td>
<td>Atrial Sense after refractory</td>
</tr>
<tr>
<td>[AS]</td>
<td>Atrial Sense during noise window</td>
</tr>
<tr>
<td>(AS)</td>
<td>Atrial Sense during refractory</td>
</tr>
<tr>
<td>AN</td>
<td>Atrial Rate Noise</td>
</tr>
<tr>
<td>AP</td>
<td>Atrial Pace</td>
</tr>
<tr>
<td>AP-Ns</td>
<td>Atrial Pace - Noise (asynchronous pacing)</td>
</tr>
<tr>
<td>RVS</td>
<td>Right Ventricular Sense after refractory</td>
</tr>
<tr>
<td>[RVS]</td>
<td>Right Ventricular during noise window</td>
</tr>
<tr>
<td>RVN</td>
<td>Right Ventricular rate noise</td>
</tr>
<tr>
<td>RVP</td>
<td>Right Ventricular Pace</td>
</tr>
<tr>
<td>RVP-Ns</td>
<td>Right Ventricular Pace - Noise (asynchronous pacing)</td>
</tr>
<tr>
<td>LVS</td>
<td>Left Ventricular Sense after refractory</td>
</tr>
<tr>
<td>[LVS]</td>
<td>Left Ventricular during noise window</td>
</tr>
<tr>
<td>LVN</td>
<td>Left Ventricular rate noise</td>
</tr>
<tr>
<td>LVP</td>
<td>Left Ventricular Pace</td>
</tr>
<tr>
<td>LVP-Ns</td>
<td>Left Ventricular Pace - Noise (asynchronous pacing)</td>
</tr>
<tr>
<td>RPVC</td>
<td>RV Premature Ventricular Contraction (PVC)</td>
</tr>
<tr>
<td>LPVC</td>
<td>LV Premature Ventricular Contraction (PVC)</td>
</tr>
<tr>
<td>&gt; 2s</td>
<td>Large interval greater than 2 seconds</td>
</tr>
</tbody>
</table>
MAINTENANCE, TROUBLESHOOTING, SERVICE, AND STANDARDS

Refer to the LATITUDE Programming System Operator’s Manual, Model 3300 for maintenance, troubleshooting, handling (including symbols on devices and packaging), standards, and specifications information.

WARRANTY INFORMATION

For all warranty information, refer to the LATITUDE Programming System Operator’s Manual, Model 3300.