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.
### LIST OF ACRONYMS

<table>
<thead>
<tr>
<th>AC</th>
<th>Alternating current</th>
<th>LCD</th>
<th>Liquid crystal display</th>
</tr>
</thead>
<tbody>
<tr>
<td>AF</td>
<td>Atrial Fibrillation</td>
<td>MRI</td>
<td>Magnetic resonance imaging</td>
</tr>
<tr>
<td>ATP</td>
<td>Anti-tachycardia pacing</td>
<td>NSR</td>
<td>Normal sinus rhythm</td>
</tr>
<tr>
<td>CRT</td>
<td>Cardiac resynchronization therapy</td>
<td>RF</td>
<td>Radio Frequency</td>
</tr>
<tr>
<td>CPR</td>
<td>Cardiopulmonary resuscitation</td>
<td>RFI</td>
<td>Radio Frequency Interference</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiogram</td>
<td>RFID</td>
<td>Radio Frequency Identification</td>
</tr>
<tr>
<td>EMI</td>
<td>Electromagnetic interference</td>
<td>S-ECG</td>
<td>Subcutaneous electrocardiogram</td>
</tr>
<tr>
<td>EOL</td>
<td>End of life</td>
<td>S-ICD</td>
<td>Subcutaneous implantable cardioverter defibrillator</td>
</tr>
<tr>
<td>ERI</td>
<td>Elective replacement indicator</td>
<td>USB</td>
<td>Universal serial bus</td>
</tr>
<tr>
<td>ESD</td>
<td>Electrostatic discharge</td>
<td>VAC</td>
<td>Voltage alternating current</td>
</tr>
<tr>
<td>FCC</td>
<td>Federal Communications Commission</td>
<td>VF</td>
<td>Ventricular fibrillation</td>
</tr>
<tr>
<td>GUI</td>
<td>Graphic user interface</td>
<td>VT</td>
<td>Ventricular tachycardia</td>
</tr>
<tr>
<td>IC</td>
<td>Industry Canada</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This literature is intended for use by professionals trained or experienced in device implant and/or follow-up procedures.

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This product may be protected by one or more patents. Patent information can be obtained at http://www.bostonscientific.com/patents.

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Description

The EMBLEM S-ICD programmer (the “programmer”) is a component of Boston Scientific’s subcutaneous implantable cardioverter defibrillator system (the S-ICD System) which is prescribed for patients when cardiac arrhythmia management is warranted. Implantable components of the S-ICD System include the EMBLEM S-ICD pulse generator and the EMBLEM S-ICD subcutaneous electrode.

The programmer is a non–sterile, non–implantable, tablet computer controlled by a graphic user interface (GUI) displayed on a touchscreen. The programmer is powered by either AC line power or an internal lithium ion battery pack. The programmer uses a connected RF telemetry wand to communicate wirelessly with the S-ICD pulse generator in order to adjust programmable settings and to collect patient data. The EMBLEM S-ICD programmer is also compatible with the Cameron Health (Model 1010) SQ-Rx pulse generator. The programmer features and functions described in this manual apply to the Boston Scientific S-ICD System as well as to the Cameron Health S-ICD System.

The S-ICD System is designed for ease of use and simplicity of patient management. The S-ICD System has a number of automatic functions designed to reduce the amount of time required for implantation, initial programming and patient follow-up.

Intended Use of Programmer

The programmer is intended to communicate with the implanted pulse generator using wireless telemetry. The programmer software controls all such telemetry functions.

Indications for Use

The S-ICD System is intended to provide defibrillation therapy for the treatment of life-threatening ventricular tachyarrhythmias in patients who do not have symptomatic bradycardia, incessant ventricular tachycardia, or spontaneous, frequently recurring ventricular tachycardia that is reliably terminated with anti-tachycardia pacing.

Contraindications

Unipolar stimulation and impedance-based features are contraindicated for use with the S-ICD System.
Related Information

Before using the S-ICD system, read and follow all instructions, warnings, and precautions provided in this manual and in the manuals for the other system components, including the applicable S-ICD pulse generator, subcutaneous electrode, and electrode implant tools user’s manuals.

This guide may contain reference information for pulse generator model numbers that are not currently approved for sale in all geographies. For a complete list of model numbers approved in your geography, consult with your local sales representative. Some model numbers may contain fewer features; for those devices, disregard descriptions of the unavailable features. Descriptions found within this manual apply to all device tiers unless otherwise noted.


Programmer Warnings and Precautions

The following warnings and precautions apply specifically to the Model 3200 programmer component of the S-ICD System.

Programmer Warnings

General

- Modifications. No modification of this equipment is allowed unless approved by Boston Scientific.
- Programmer is MR Unsafe. The programmer 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 programmer be brought into the MRI scanner room, the control room, or the MRI site Zone III or IV areas.
- High temperatures. Do not subject the programmer to temperatures outside of the -10° C to 55° C (14° F to 131° F) storage range. Exposure to high temperatures may cause the programmer to overheat or ignite, and may possibly reduce its performance and service life.
- Extreme temperatures. Do not discard the programmer in a fire, incinerate, or subject it to temperatures that exceed 100° C (212° F). This could cause the programmer to explode.
- Do not immerse. Do not immerse the programmer in liquid of any kind. If the programmer does get wet, contact customer service for information about returning the programmer to

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Boston Scientific. Do not attempt to dry the programmer in an oven, microwave, or dryer because this poses a risk of overheating or explosion.

**Conditions for Operation**

- **Secure programmer.** Ensure this programmer is used by professionals trained or experienced in device implant and/or follow-up procedures. Take appropriate measures to prevent unauthorized use or tampering of the programmer.

- **Use only the supplied external power supply.** Use the programmer only with the external power supply packaged with the programmer. Using other power supplies may cause damage to the programmer.

- **Electric shock.** To avoid risk of electric shock, the programmer’s external power supply must only be connected to a grounded electrical outlet.

- **Damaged programmer or power supply.** Never use a damaged external power supply or a damaged programmer. Doing so could result in user injury, patient injury, or a lack of therapy delivery.

- **Interference with nearby equipment.** By design, the programmer emits radio frequencies in the 402-405 MHz and 2.4 GHz bands. This may interfere with nearby medical or office equipment. When using the programmer, closely monitor equipment in the vicinity to verify normal operation. It may be necessary to take mitigation measures, such as reorienting or relocating the programmer or shielding the location.

- **Interference with programmer communication.** The presence of other equipment operating in the same frequency bands used by the programmer (402-405 MHz for the pulse generator and 2.4 GHz for the printer) may interfere with communication. Interference can occur even if the other equipment complies with the International Special Committee on Radio Interference (CISPR) emission requirements. This RF interference can be reduced by increasing the distance between the interfering device and the programmer and pulse generator or printer. If communication problems persist, refer to the Troubleshooting section of this manual.

- **Use of non-approved accessories.** The use of any accessories with the programmer other than those specified by Boston Scientific in this manual may result in increased emissions or decreased immunity of the programmer and may cause decreased functionality or unintended operational behavior of the programmer. Anyone connecting such accessories to the programmer 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.
• **Programmer location.** Do not use the programmer adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, check the programmer for normal operation in that configuration.

**Programmer Precautions**

**General**

• **Wand use.** Use only the Model 3203 telemetry wand with the programmer.
• **Do not disassemble.** Do not disassemble or alter any parts of the programmer.
• **Device communication.** Use only the designated programmer and software application to communicate with the pulse generator.
• **Intended users.** The programmer is intended for use by or under the direction of healthcare professionals only.
• **Sensitive Information.** To prevent sensitive personal information from being transmitted to inappropriate devices or printers when using Bluetooth™ wireless connections, make certain to only connect with known Bluetooth™ devices.

**Storage and Handling**

• **Mishandling.** Mishandling (such as dropping or crushing) could damage the programmer. If you suspect damage to the programmer, contact your Boston Scientific representative or the customer service department for instructions and return packaging.
• **Broken or cracked screen.** The display on the programmer is made of glass or acrylic and could break if the programmer is dropped or if it receives significant impact. Do not use if screen is broken or cracked as this could cause injury.
• **Magnet handling.** Do not place a magnet on the programmer.
• **Data Storage.** The programmer and digital data storage media, such as microSD™ memory cards, used with the programmer may contain sensitive personal information. These should be handled in accordance with applicable privacy and security policies and regulations.

**Implantation**

• **Telemetry wand.** The wand is a non-sterile device. Do not sterilize the wand. The wand must be contained in a sterile barrier before use in the sterile field.

The Bluetooth™ word mark and logos are registered trademarks owned by Bluetooth SIG, Inc., and any use of such marks is under license.

microSD™ is a trademark or registered trademark of SD-3C, LLC.
• **Programmer must remain outside the sterile field.** The programmer is non-sterile and cannot be sterilized. It must remain outside the sterile field.

*Conditions for operation*

• **Disconnecting the programmer.** Mains isolation is achieved by disconnecting the external power supply power cord from the AC electrical outlet. Do not position the programmer or the external power supply in a manner that would make it difficult to disconnect that cord.

• **Programmer use.** The programmer is not waterproof or explosion-proof and cannot be sterilized. Do not use it in the presence of flammable gas mixtures containing anesthetics, oxygen, or nitrous oxide.

• **Confirm communication.** Confirm that the programmer is in communication with the intended implanted S-ICD pulse generator.

• **Electrostatic discharge.** The programmer may be affected by ESD. If ESD occurs and the programmer's functionality is affected, attempt to reset the programmer or contact Boston Scientific for instructions. Do not touch or connect the telemetry wand to the programmer unless ESD precautionary procedures are used.

*S-ICD System Warnings and Precautions*

The following warnings and precautions apply to the S-ICD System as a whole. For additional warnings and precautions that are specific to other individual components of the system, and/or to the process of implanting the system, refer to the manual of the relevant system component.

*S-ICD System Warnings*

*General*

• **Component Compatibility.** All Boston Scientific S-ICD implantable components are designed for use with the Boston Scientific or Cameron Health S-ICD System only. Connection of any S-ICD System components to a non-compatible component has not been tested and could result in failure to deliver life-saving defibrillation therapy.

• **Backup defibrillation protection.** Always have external defibrillation equipment and medical personnel skilled in CPR available during implant and follow up testing. If not terminated in a timely fashion, an induced ventricular tachyarrhythmia can result in the patient’s death.

• **Pulse generator interaction.** Using multiple pulse generators could cause pulse generator interaction, resulting in patient injury or a lack of therapy delivery. Test each system
individually and in combination to help prevent undesirable interactions. Refer to the appropriate S-ICD pulse generator manual for more information.

Post-Implant

- **Magnet Response.** Use caution when placing a magnet over the S-ICD pulse generator because it suspends arrhythmia detection and therapy response. Removing the magnet resumes arrhythmia detection and therapy response.

- **Magnet response with deep implant placement.** In patients with a deep implant placement (greater distance between the magnet and the pulse generator) magnet application may fail to elicit the magnet response. In this case the magnet cannot be used to inhibit therapy.

- **Diathermy.** Do not expose a patient with an implanted S-ICD System to diathermy. The interaction of diathermy therapy with an implanted S-ICD pulse generator or electrode can damage the pulse generator and cause patient injury.

- **Magnetic Resonance Imaging (MRI) exposure.** EMBLEM S-ICD devices are considered MR Conditional. For these devices, unless all of the MRI Conditions of Use are met, MRI scanning of the patient does not meet MR 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 manual are not MR conditional. Do not expose patients with non-MR Conditional devices to MRI scanning. Strong magnetic fields may damage the pulse generator and/or subcutaneous electrode, possibly resulting in injury to or death of the patient.

- **Protected environments.** Advise patients to seek medical guidance before entering environments that could adversely affect the operation of the active implantable medical device, including areas protected by a warning notice that prevents entry by patients who have a pulse generator.

- **Sensitivity settings and EMI.** The pulse generator may be more susceptible to low frequency electromagnetic interference at induced signals greater than 80 uV. Oversensing of noise due to this increased susceptibility could lead to inappropriate shocks and should be taken into consideration when determining the follow-up schedule for patients exposed to low frequency electromagnetic interference. The most common source of electromagnetic interference in this frequency range is the power system for some European trains which operate at 16.6 Hz. Particular attention should be given to patients with occupational exposure to these types of systems.
Clinical Considerations

- **Longevity.** Battery depletion will eventually cause the S-ICD pulse generator to stop functioning. Defibrillation and excessive numbers of charging cycles shorten the battery longevity.
- **Pediatric Use.** The S-ICD System has not been evaluated for pediatric use.
- **Available Therapies.** The S-ICD System does not provide long-term bradycardia pacing, Cardiac Resynchronization Therapy (CRT) or Anti-Tachycardia Pacing (ATP).

Implantation

- **Upper extremity injury.** During arrhythmia induction, the induction current and subsequent shock may result in forceful contraction of the pectoralis major muscle which can exert significant acute forces on the glenohumeral joint as well as on the clavicle. This, in conjunction with a tightly restrained arm, may result in injury to the clavicle, shoulder, and arm, including dislocation and fracture.
- **Avoid shock at implant.** Verify the device is in Shelf mode or Therapy Off to prevent the delivery of unwanted shocks to the patient or the person handling the device during the implant procedure.

Device Programming

- **Sensing adjustment.** Following any sensing parameter adjustment or any modification of the subcutaneous electrode, always verify appropriate sensing.
- **Patients hear tones coming from their device.** Patients should be advised to contact their physician immediately if they hear tones coming from their device.
- **Programming for supraventricular tachyarrhythmias (SVTs).** Determine if the device and programmed parameters are appropriate for patients with SVTs because SVTs can initiate unwanted device therapy.

S-ICD System Precautions

Environmental and Medical Therapy Hazards

- **Avoid electromagnetic interference (EMI).** Advise patients to avoid sources of EMI because EMI may cause the pulse generator to deliver inappropriate therapy or inhibit appropriate therapy. Moving away from the source of the EMI or turning off the source usually allows the pulse generator to return to normal operation. Examples of potential EMI sources found in hospital and medical environments are:
» Radio transmitters
» Electronic surveillance or security systems
» Medical treatments and diagnostic tests in which an electrical current is passed through the body, such as TENS, electrocautery, electrolysis/thermolysis, electrodiagnostic testing, electromyography, or nerve conduction studies
» Any externally applied device that uses an automatic lead detection alarm system (e.g., an EKG machine)

Hospital and Medical Environments

- **External defibrillation.** External defibrillation or cardioversion can damage the pulse generator or subcutaneous electrode. To help prevent damage to implanted system components, consider the following:
  » Avoid placing a pad (or paddle) directly over the pulse generator or subcutaneous electrode. Position the pads (or paddles) as far from the implanted system components as possible.
  » Set energy output of external defibrillation equipment as low as clinically acceptable.
  » Following external cardioversion or defibrillation, verify pulse generator function (“Post-Therapy Pulse Generator Follow Up” on page 13).

- **Cardiopulmonary resuscitation.** Cardiopulmonary resuscitation (CPR) may temporarily interfere with sensing and may cause delay of therapy.

- **Electrical interference.** Electrical interference or “noise” from devices such as electrocautery and monitoring equipment may interfere with establishing or maintaining telemetry for interrogating or programming the device. In the presence of such interference, move the programmer away from electrical devices, and ensure that the wand cord and cables are not crossing one another. Electrical interference or “noise” from concomitant implanted devices such as a ventricular assist device (VAD), drug pump, or insulin pump may interfere with establishing or maintaining telemetry for interrogating or programming the pulse generator. In the presence of such interference, place the wand over the pulse generator and shield both with a radiation-resistant material.

- **Ionizing radiation therapy.** It is not possible to specify a safe radiation dosage or guarantee proper pulse generator function following exposure to ionizing radiation. Multiple factors collectively determine the impact of radiation therapy on an implanted pulse generator, including proximity of the pulse generator to the radiation beam, type and energy level of the radiation beam, dose rate, total dose delivered over the life of the pulse
generator, and shielding of the pulse generator. The impact of ionizing radiation will also vary from one pulse generator to another and may range from no changes in function to a loss of therapy. Sources of ionizing radiation vary significantly in their potential impact on an implanted pulse generator. Several therapeutic radiation sources are capable of interfering with or damaging an implanted pulse generator, including those used for the treatment of cancer, such as radioactive cobalt, linear accelerators, radioactive seeds, and betatrons. Prior to a course of therapeutic radiation treatment, the patient’s radiation oncologist and cardiologist or electrophysiologist should consider all patient management options, including increased follow-up and device replacement.

Other considerations include:

» Shield the pulse generator with a radiation-resistant material, regardless of the distance between the pulse generator and the radiation beam.

» Determining the appropriate level of patient monitoring during treatment

Evaluate pulse generator operation during and following the course of radiation treatment to exercise as much device functionality as possible (“Post-Therapy Pulse Generator Follow Up” on page 13). The extent, timing, and frequency of this evaluation relative to the radiation therapy regimen are dependent upon current patient health, and therefore should be determined by the attending cardiologist or electrophysiologist.

Pulse generator diagnostics are performed automatically once per hour, so pulse generator evaluation should not be concluded until pulse generator diagnostics have been updated and reviewed (at least one hour after radiation exposure). The effects of radiation exposure on the implanted pulse generator may remain undetected until some time following exposure. For this reason, continue to monitor pulse generator function closely and use caution when programming a feature in the weeks or months following radiation therapy.

- **Electrocautery and RF Ablation.** Electrocautery and RF ablation may induce ventricular arrhythmias and/or fibrillation, and may cause inappropriate shocks and inhibition of post-shock pacing. Additionally, exercise caution when performing any other type of cardiac ablation procedure in patients with implanted devices. If electrocautery or RF ablation is medically necessary, observe the following to minimize risk to the patient and device:

  » Program the pulse generator to Therapy Off mode.

  » Have external defibrillation equipment available.

  » Avoid direct contact between the electrocautery equipment or ablation catheters and the pulse generator and subcutaneous electrode.
» Keep the path of the electrical current as far away as possible from the pulse generator and subcutaneous electrode.

» If RF ablation and/or electrocautery is performed on tissue near the device or subcutaneous electrode, verify pulse generator function (“Post-Therapy Pulse Generator Follow Up” on page 13). For electrocautery, use a bipolar electrocautery system where possible and use short, intermittent, and irregular bursts at the lowest feasible energy levels.

When the procedure is finished, return the pulse generator to Therapy On mode.

- **Lithotripsy.** Extracorporeal shock wave lithotripsy (ESWL) may cause electromagnetic interference with or damage to the pulse generator. If ESWL is medically necessary, consider the following to minimize the potential for encountering interaction:
  » Avoid focusing the lithotripsy beam near the pulse generator implant site.
  » Program the pulse generator to Therapy Off mode to prevent inappropriate shocks.

- **Ultrasound energy.** Therapeutic ultrasound (e.g., lithotripsy) energy may damage the pulse generator. If therapeutic ultrasound energy must be used, avoid focusing near the pulse generator site. Diagnostic ultrasound (e.g., echocardiography) is not known to be harmful to the pulse generator.

- **Conducted electrical current.** Any medical equipment, treatment, therapy, or diagnostic test that introduces electrical current into the patient has the potential to interfere with pulse generator function. Medical therapies, treatments, and diagnostic tests that use conducted electrical current (e.g., TENS, electrocautery, electrolysis/thermolysis, electrodiagnostic testing, electromyography, or nerve conduction studies) may interfere with or damage the pulse generator. Program the device to Therapy Off mode prior to the treatment, and monitor device performance during the treatment. After the treatment, verify pulse generator function (“Post-Therapy Pulse Generator Follow Up” on page 13).

- **Transcutaneous Electrical Nerve Stimulation (TENS).** TENS involves passing electrical current through the body, and may interfere with pulse generator function. If TENS is medically necessary, evaluate the TENS therapy settings for compatibility with the pulse generator. The following guidelines may reduce the likelihood of interaction:
  » Place the TENS electrodes as close together and as far away from the pulse generator and subcutaneous electrode as possible.
  » Use the lowest clinically-appropriate TENS energy output.
  » Consider cardiac monitoring during TENS use. Additional steps can be taken to help reduce interference during in-clinic use of TENS:
If interference is suspected during in-clinic use, turn off the TENS unit.

Do not change TENS settings until you have verified that the new settings do not interfere with pulse generator function.

If TENS is medically necessary outside the clinical setting (at-home use), provide patients with the following instructions:

» Do not change the TENS settings or electrode positions unless instructed to do so.
» End each TENS session by turning off the unit before removing the electrodes.
» If the patient receives a shock during TENS use, they should turn off the TENS unit and contact their physician. Follow these steps to use the programmer to evaluate pulse generator function during TENS use:
  1. Program the pulse generator to Therapy Off mode.
  2. Observe real-time S-ECGs at prescribed TENS output settings, noting when appropriate sensing or interference occurs.
  3. When finished, turn off the TENS unit and reprogram the pulse generator to Therapy On mode.

You should also perform a thorough follow-up evaluation of the pulse generator following TENS, to ensure that device function has not been compromised (“Post-Therapy Pulse Generator Follow Up” on page 13). For additional information, contact Boston Scientific using the information on the back cover.

• **Electronic Article Surveillance (EAS) and Security Systems.** Advise patients how to avoid impact to cardiac device function due to antitheft and security gates, tag deactivators, or tag readers that include radio frequency identification (RFID) equipment. These systems may be found at the entrances and exits of stores, at checkout counters, in public libraries, and in point-of-entry access control systems. Patients should avoid lingering near or leaning against antitheft and security gates and tag readers. In addition, patients should avoid leaning against checkout counter-mounted and handheld tag deactivation systems. Antitheft gates, security gates, and entry control systems are unlikely to affect cardiac device function when patients walk through them at a normal pace. If the patient is near an electronic antitheft, security, or entry control system and experiences symptoms, they should promptly move away from nearby equipment and inform their doctor.

• **Elevated Pressures.** The International Standards Organization (ISO) has not approved a standardized pressure test for implantable pulse generators that experience hyperbaric oxygen therapy (HBOT). Elevated pressures due to HBOT may damage the pulse
generator. Prior to starting an HBOT program, the patient’s attending cardiologist or electrophysiologist should be consulted to fully understand the potential consequences relative to the patient’s specific health condition. More frequent device follow-up may be warranted in conjunction with HBOT. Evaluate pulse generator operation following high pressure exposure (“Post-Therapy Pulse Generator Follow Up” on page 13). The extent, timing, and frequency of this evaluation relative to the high pressure exposure are dependent upon current patient health, and should be determined by the attending cardiologist or electrophysiologist. Refer to the appropriate pulse generator manual for additional information about device-specific high pressure testing results. If you have additional questions, contact Boston Scientific using the information on the back cover.

**Follow-up Testing**

- **Low shock impedance.** A reported shock impedance value of less than 25 ohms from a delivered shock could indicate a problem with the device. The delivered shock may have been compromised, and/or any future therapy from the device may be compromised. If a reported impedance value of less than 25 ohms is observed, correct functioning of the device should be verified.

- **Conversion testing.** Successful VF or VT conversion during arrhythmia conversion testing is no assurance that conversion will occur post-operatively. Be aware that changes in the patient’s condition, drug regimen, and other factors may change the DFT, which may result in nonconversion of the arrhythmia post-operatively. Verify with a conversion test that the patient’s tachyarrhythmias can be detected and terminated by the pulse generator system if the patient’s status has changed or parameters have been reprogrammed.

- **Follow-up considerations for patients leaving the country.** Pulse generator follow-up considerations should be made in advance for patients who plan to travel or relocate post-implant to a country other than the country in which their device was implanted. Regulatory approval status for devices and associated programmer software configurations varies by country; certain countries may not have approval or capability to follow specific products. Contact Boston Scientific, using the information on the back cover, for help in determining feasibility of device follow-up in the patient’s destination country.

**Explant and Disposal**

- **Handling at explant.** Before explanting, cleaning, or shipping the device, complete the following actions to prevent unwanted shocks, overwriting of important therapy history data, and audible tones:
  - Program the pulse generator to Therapy Off mode
» Disable the beeper, if available.
» Clean and disinfect the device using standard biohazard handling techniques.

Supplemental Precautionary Information

• Post-Therapy Pulse Generator Follow Up. Following any surgery or medical procedure with the potential to affect pulse generator function, you should perform a thorough follow up, which may include the following:
  » Interrogating the pulse generator with a programmer
  » Reviewing stored events, fault codes, and real-time S-ECGs prior to saving all patient data
  » Testing the subcutaneous electrode impedance
  » Verifying battery status
  » Printing any desired reports
  » Verifying the appropriate final programming prior to allowing the patient to leave the clinic
  » Ending session

S-ICD System Clinical Investigation

A summary of the S-ICD System Clinical Investigation, including observed adverse events, is provided in the S-ICD User’s Manual.

Potential Adverse Events

Potential adverse events related to implantation of the S-ICD System may include, but are not limited to, the following:

• Acceleration/induction of atrial or ventricular arrhythmia
• Adverse reaction to induction testing
• Allergic/adverse reaction to system or medication
• Bleeding
• Conductor fracture
• Cyst formation
EMBLEM™ S-ICD PROGRAMMER: GENERAL DESCRIPTION

- Death
- Delayed therapy delivery
- Discomfort or prolonged healing of incision
- Electrode deformation and/or breakage
- Electrode insulation failure
- Erosion/extrusion
- Failure to deliver therapy
- Fever
- Hematoma/seroma
- Hemothorax
- Improper electrode connection to the pulse generator
- Inability to communicate with the pulse generator
- Inability to defibrillate or pace
- Inappropriate post-shock pacing
- Inappropriate shock delivery
- Infection
- Injury to or pain in upper extremity including clavicle, shoulder, and arm
- Keloid formation
- Migration or dislodgement
- Muscle/nerve stimulation
- Nerve damage
- Pneumothorax
- Post-shock/post-pace discomfort
- Premature battery depletion
- Random component failures
- Stroke
- Subcutaneous emphysema
• Surgical revision or replacement of the system
• Syncope
• Tissue redness, irritation, numbness or necrosis

If any adverse events occur, invasive corrective action and/or S-ICD System modification or removal may be required.

Patients who receive an S-ICD System may also develop psychological disorders that include, but are not limited to, the following:
• Depression/anxiety
• Fear of device malfunction
• Fear of shocks
• Phantom shocks
Programmer Setup

Packaging

Programmer components include:

- Model 3200 Programmer with pre-loaded software
- Model 3203 Telemetry Wand
- Model 3204 External power supply and AC power cord

Visually inspect the packaging to ensure the contents are complete. Do not use if there is evidence of damage.

In case of damage return the product to Boston Scientific. For return packaging and instructions, contact Boston Scientific using the information on the back cover of this manual.

Programmer Controls and Connections

Figure 1: Controls and external connections
Charging the Programmer

The programmer is primarily intended to be operated while connected to the AC-powered external power supply, but may also be operated on battery power provided that the internal battery is adequately charged. The programmer is recharged whenever it is connected to the AC-powered external power supply. When not in use, it is recommended that the programmer remain connected to the external power supply in order to maintain an adequate battery charge.

**Note:** *Current session data may be lost if a 45 minute period of inactivity occurs during an active telemetry session and the programmer is not connected to AC power.*

Typical charge time for a fully discharged battery is 5 hours. However, more time may be required if the programmer is in use while being recharged.

The Battery Status indicator located on the upper right corner of the screen displays the status of the main battery power when the unit is in use:

- All four bars are illuminated (green) – The battery is 100% charged
- Three bars are illuminated (green) – The battery is 75% charged
- Two bars are illuminated (yellow) – The battery is 50% charged
- One bar is illuminated (red) – The battery is 25% charged

The programmer displays one of the following alert screens as battery power gets progressively lower.

- Programmer Battery Low
- Programmer Battery Critical
- Out Of Power

To charge the programmer:

1. Connect the external power supply cable to the programmer (Figure 1).
2. Plug the external power supply cord into an AC power outlet.

**Warning:** *Use the programmer only with the external power supply packaged with the programmer. Using other power supplies may cause damage to the programmer.*

**Warning:** *To avoid risk of electric shock, the programmer's external power supply must only be connected to a grounded electrical outlet.*
Using the Programmer

Turning the Programmer On

The programmer power button is located in the recess above and behind the left corner of the screen (Figure 1). Press and hold the button until the display screen is active.

Note: If the programmer cannot be turned on while it is connected to AC power via the external power supply, first unplug the external power supply cord from the programmer. Press and hold the programmer power button until the display screen is active. AC power via the external power supply can then be reconnected.

Changing the Programmer Volume Level

The volume level of programmer-generated sounds may be temporarily adjusted using the volume control (Figure 1). This level is automatically reset when the programmer is restarted.

Placing the Programmer in Suspend Mode

The programmer has a Suspend Mode which is activated automatically to conserve power. The display will be blank when this mode is in effect.

The programmer enters Suspend Mode whenever:
- The power button is momentarily pressed and released
- The programmer is not connected to the external power supply, it is not in active communication with an S-ICD pulse generator, and no user activity has occurred for 15 minutes

Momentarily pressing the power button will resume normal operation.

Turning the Programmer Off

There are two ways to turn the programmer off:

1. Press and hold the power button until the System shutdown menu appears. Select Power off from the popup and confirm by pressing OK.
2. From the programmer start-up screen, press the Power Off button and select OK at the confirmation prompt.
Using the Programmer Touch Screen

The programmer is equipped with an LCD touch screen. The screen can be adjusted to the desired viewing angle by using the kick-stand located on the back of the programmer. All interaction with the programmer is conducted using the fingers to touch the appropriate areas on the screen. Scroll on-screen lists by sliding a finger up and down the list. An on-screen keyboard is presented whenever text entry is required.

**Caution:** The display on the programmer is made of glass or acrylic and could break if the programmer is dropped or if it receives significant impact. Do not use if screen is broken or cracked as this could cause injury.

Using the Wand

The Model 3203 wand ("the wand") makes it possible for this programmer to communicate with the pulse generator.

**Caution:** Use only the Model 3203 telemetry wand with the programmer.

**Caution:** The wand is a non-sterile device. Do not sterilize the wand. The wand must be contained in a sterile barrier before use in the sterile field.

**Caution:** The programmer is non-sterile and cannot be sterilized. It must remain outside the sterile field.

To connect the wand to the programmer, slide the wand cable connector over the communication connector port located on the rear edge of the programmer (Figure 1).

To disconnect the wand, grasp the wand cable connector and gently pull it straight off the communication connector port.

**Note:** Do not pull or yank on the cable to disconnect the wand from the programmer. Such action could cause hidden damage to the cable. A damaged cable might reduce wireless communication capabilities and require a replacement wand.

Optimal telemetry depends on the wand being placed directly over the implanted pulse generator. Although it may appear that the programmer is in communication with the pulse generator at greater distances, programming should always be performed with the wand placed directly over the implanted pulse generator.

**Warning:** The presence of other equipment operating in the same frequency bands used by the programmer (402-405 MHz for the pulse generator and 2.4 GHz for the printer) may
Interference can occur even if the other equipment complies with the International Special Committee on Radio Interference (CISPR) emission requirements. This RF interference can be reduced by increasing the distance between the interfering device and the programmer and pulse generator or printer. If communication problems persist, refer to the Troubleshooting section of this manual.

When telemetry loss occurs, the display screen will turn yellow and a message will appear with the text “Communication Loss” to alert the user. Reposition the wand to establish communication. The programmer will return to the screen that was active before telemetry loss if the pulse generator is found and programming can continue.

**Note:** If communication cannot be reestablished, the session should be ended and restarted by scanning for the pulse generator.

**Navigation**

The programmer’s graphic user interface (GUI) facilitates management and control of the S-ICD System. The Navigation Bar and on-screen icons at the top of the screen allow the user to navigate programming software screens. In addition, a continuous subcutaneous electrocardiogram (S-ECG) is displayed along the bottom of the screen during Online (active) communication with the pulse generator.

**Screen Header**

When the programmer is Offline (inactive communication), the screen header displays the Battery Status Indicator.

When viewing Offline Stored Sessions, the screen header displays:

- Patient name
- Therapy On/Off
- Battery status indicator

When the programmer is Online (active communication), the screen header displays:

- Therapy On/Off
- Patient name
- Patient heart rate
- Programmer Battery and Telemetry status indicator
• Screen title
• Rescue shock icon

**Navigation Bar**

The Navigation Bar is the primary method for navigating the Online programmer screens. The bar is located along the top edge of the programmer screen and chosen screens appear with their selection icon highlighted.

Table 1 (page 22) provides a list of the programmer icons and their corresponding descriptions.

**Restarting the Programmer**

The programmer’s operating system is self-monitoring and is generally able to sense many system error conditions and automatically initiate a restart sequence in response. Follow the on-screen instructions to complete the programmer-initiated restart sequence.

The programmer may need to be manually restarted if:
• You cannot exit a screen
• The operating system stops responding

A manual restart is accomplished by pressing and holding the power button until the system shutdown menu appears on the screen. Select Restart from the popup and confirm by pressing OK.

If the programmer does not respond to a restart process, contact Boston Scientific using the information on the back cover of this manual.
Table 1: Icon descriptions

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
<th>User Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Main Menu Icon]</td>
<td>Main Menu Icon</td>
<td>Allows user to return to the main menu.</td>
</tr>
<tr>
<td>![Automatic Setup Icon]</td>
<td>Automatic Setup Icon</td>
<td>Allows user to access the Automatic Setup menu.</td>
</tr>
<tr>
<td>![Device Settings Icon]</td>
<td>Device Settings Icon</td>
<td>Allows user to access the S-ICD device settings screen.</td>
</tr>
<tr>
<td>![Device Status Icon]</td>
<td>Device Status Icon (open folder and closed folder)</td>
<td>Allows user to access the S-ICD device status screen. User can view number of shocks delivered since the last update as well as the S-ICD device battery life.</td>
</tr>
<tr>
<td>![Patient View Icon]</td>
<td>Patient View Icon</td>
<td>Allows user to access the patient chart screen. User can view information on the S-ICD device battery life.</td>
</tr>
<tr>
<td>![Captured and Stored Episodes S-ECG Icon]</td>
<td>Captured and Stored Episodes S-ECG Icon</td>
<td>Allows user to access captured S-ECG and stored episode screens.</td>
</tr>
<tr>
<td>![Induction Test Icon]</td>
<td>Induction Test Icon</td>
<td>Allows user to access induction screen.</td>
</tr>
<tr>
<td>![Battery &amp; Telemetry Meter]</td>
<td>Battery &amp; Telemetry Meter</td>
<td>Left side of the meter allows user to view the programmer's battery status. The right side of the meter allows viewing of telemetry signal strength.</td>
</tr>
<tr>
<td>![Capture S-ECG]</td>
<td>Capture S-ECG</td>
<td>Allows user to capture a live S-ECG.</td>
</tr>
<tr>
<td>![S-ECG Display Settings]</td>
<td>S-ECG Display Settings</td>
<td>Allows user to modify the zoom and sweep speed on the live S-ECG.</td>
</tr>
<tr>
<td>![Heart Rate Icon]</td>
<td>Heart Rate Icon</td>
<td>Allows user to view current heart rate.</td>
</tr>
<tr>
<td>![Rescue Shock Icon]</td>
<td>Rescue Shock Icon</td>
<td>Allows user to administer a rescue shock</td>
</tr>
<tr>
<td>![Option Selection Switch]</td>
<td>Option Selection Switch</td>
<td>Allows user to select one of two options, e.g. A or B</td>
</tr>
</tbody>
</table>
Configuring the Programmer

Configuring Programmer Settings

The programmer should be configured before communication with a pulse generator is attempted. This includes setting the date and time format, time zone, language and printer. Once these settings are configured during the initial setup process they become the default parameters and will not normally need to be changed with each session.

To configure the programmer settings:

1. Select the Programmer Settings button on the programmer start-up screen (Figure 2) to display the Programmer Settings screen (Figure 3).

![Programmer start-up screen](image)

**Figure 2: Programmer start-up screen**
2. Select the corresponding line to access each setting. The settings that can be configured include:
   - Date and time format
   - Time zone
   - Language
   - Printer

**Date and Time Format**

To set the date and time format:

1. Select Set Date and Time Format on the Programmer Settings screen (Figure 3). The Date and Time Settings screen appears.
2. Select the desired date format.
3. Select the Save button to save the changes and return to the Programmer Settings screen, or select Cancel to return to the Programmer Settings screen without saving the changes.
**Time Zone**

The time zone setting controls two S-ICD System parameters, one for the programmer (the time shown on screens and printed reports), the other for pulse generators (the electronic filter that is intended to minimize electromagnetic interference [EMI]).

Choosing the correct time zone setting for the programmer will result in the electronic filter of interrogated pulse generators being set to the appropriate regional electrical power line frequency.

Specifically, the pulse generator line frequency filter is automatically programmed to either 50 Hz or 60 Hz, based on the time zone setting of the interrogating programmer.

To set the time zone

1. Select Set Time Zone on the Programmer Settings screen. The time zone selection screen appears (Figure 4)
2. Select the time zone button for the zone in which the programmer will be used. A checkmark will appear in the selected button.
3. Select the Save button to save the changes and return to the Programmer Settings screen, or select Cancel to return to the Programmer Settings screen without saving the changes.

In the rare cases where a single time zone setting includes regional power line frequency differences, two line frequency options are available. Choose the option with the correct frequency for the region where the programmer is located.

Because a programmer will set the time zone (and electronic frequency filter) of pulse generators it interrogates to match its own time zone setting, be aware that travelling patients whose devices are interrogated in time zones or countries other than the one in which they reside may need to have their pulse generator time zone reset upon returning home.
Language Preference

To set the language preference:

1. Select Set Language on the Programmer Settings screen. The Language Settings screen appears. Scroll the list and select a language.

2. Select the Save button to save the changes or select Cancel to return to the Programmer Settings screen without saving the changes. If the language is changed the programmer will automatically restart and return to the Startup screen.

Printer Selection

The programmer communicates with the printer via Bluetooth™ wireless technology. Only Boston Scientific-approved printers can be paired and used with the programmer. To select the printer to be paired and used with the programmer:

1. Ensure the printer is on and, depending on your specific printer, that the wireless function is enabled or the wireless adapter is in the printer’s USB port.

2. Select Printer Setup on the Programmer Settings screen. The Printer Setup screen (Figure 5) will appear with a previously configured printer displayed as the default printer. If a default printer has not already been selected and configured, the screen
EMBLEM™ S-ICD PROGRAMMER: OPERATION

will be empty and the programmer will scan the area to locate wireless printers. A Scan Progress Bar will appear informing the user that the programmer is currently scanning for printers.

![Printer Setup screen](image)

**Figure 5: Printer Setup screen**

3. Select the printer of choice from among those found during the scan. If none were found, a window will appear stating that there are no printers. Select the Scan Again button or the Cancel button to return to the Programmer Settings screen.

4. Select the desired printer from the list and enter the name using the on-screen keyboard (up to 15 characters). A unique printer identifier should appear with the printer selection.

5. Select the Save button to save the changes and return to the Programmer Settings screen, or select Cancel to return to the Programmer Settings screen without saving the changes. A confirmation screen will appear when the printer setup is completed.

   **Note:** Refer to “Troubleshooting” section for information about printer problems.
**Programmer Software Version**

To view the programmer’s software version:


2. The Programmer Software Version information screen displays the current version of the programmer software. Select the Continue button to return to the Programmer Settings screen.

   **Note:** The printed reports also contain the programmer software versions.

**Bluetooth™ Data Export**

The programmer can be configured to wirelessly export patient data to desktop or notebook computers that are equipped with Bluetooth™ wireless technology. The programmer and each computer must be individually paired in order to use the wireless data export function. The procedure for pairing the programmer with a computer is different from the procedure used to pair the programmer with the printer.

   **Note:** Data transfer is supported for computers running Windows. The data transfer feature is not available for tablets or smartphones.

1. Ensure the computer to be paired has been made discoverable, since the programmer searches for nearby computers during the pairing process.

   **Note:** Detailed instructions for accomplishing this are found in the Microsoft Windows help files under the general heading of “Why can’t I connect my Bluetooth device to my computer?”

2. Once the target computer has been made discoverable, select the Export Programmer Data button on the programmer start-up screen. The Export Programmer Data Over Bluetooth screen will appear. Select the Set Up Authorized Computers button to scan for nearby computers and begin the pairing process.

3. When the scan is complete the screen will list the discovered computers (the three computers having the strongest Bluetooth™ signals) under the heading Unauthorized Computers Nearby (Figure 6). Choose the computer you wish to pair with and press the plus button next to it to complete the pairing process.
4. During the pairing process, both the programmer and the computer will present identical numeric passkeys and both machines will ask you to confirm that the two numbers are the same. The passkey is only presented while pairing and is used to verify that the correct machines are being paired.

5. Successful pairing is indicated when the listing for the subject computer appears in the Authorized Computers column instead of the Unauthorized Computers Nearby column.

6. Authorized computers may be renamed if desired. Press and hold the listing for the computer until the Rename an Authorized Computer popup appears.

---

**Figure 6:** Choosing a computer to authorize for Bluetooth™ data transfer
Programmer Modes of Operation

**Online Behavior**

The programmer’s interface varies according to whether the programmer is Online (actively communicating) or Offline (not communicating) with a selected pulse generator.

An Online session begins when the programmer establishes a telemetry link with a specific pulse generator. A yellow alert screen is displayed if the telemetry signal is lost between the programmer and the pulse generator for more than five seconds during active communication. This may occur if the wand is moved out of the telemetry communications range or if noise or interfering objects inhibit communication. Programming commands, including Rescue Shocks, will not be available until telemetry is reestablished.

Telemetry reconnection may occur automatically if the reason for the telemetry loss has been remedied, e.g. moving the wand back into telemetry range of the pulse generator or removing the source of interference or noise. Restart the session if the telemetry link does not resume within one minute.

**Note:** *When in active communication with a pulse generator, the programmer emits an audible notification to indicate that the pulse generator is preparing to deliver a shock, whether that shock is commanded or is in response to a detected arrhythmia. The notification continues until the shock is either delivered or aborted.*

**Offline Behavior**

The programmer is Offline when it is not actively communicating with a pulse generator. Programmer settings can be accessed and stored patient sessions can be viewed and/or printed during Offline sessions.

**Stored Patient Sessions**

During a patient follow-up visit, the programmer will retrieve data from the pulse generator memory. The programmer can store up to 50 patient sessions. When the 51st session occurs, the programmer will automatically replace the oldest stored session with the new data. A stored session includes the following information:

- Captured S-ECG Reports (including Induction S-ECGs)
- Episode History (including any downloaded episodes)
- Patient Data
- Programmed Device Settings
To view stored patient sessions:
1. From the programmer start-up screen, select Stored Patient Sessions.
2. Select the desired patient session.

Modes of Operation for the Pulse Generator
The pulse generator has the following modes of operation:

- Shelf
- Therapy On
- Therapy Off
- MRI Protection Mode

Shelf Mode
The Shelf mode is a low power consumption state intended for storage only. When a pulse generator in Shelf mode is interrogated by a programmer, it exits Shelf mode and defaults to Therapy Off mode. A full-energy capacitor reformation is performed and the pulse generator is prepared for set-up. Once the pulse generator is taken out of Shelf mode, it cannot be reprogrammed back into Shelf mode.

Therapy On Mode
The Therapy On mode is the primary operating mode of the pulse generator, allowing automatic detection of, and response to, ventricular tachyarrhythmias.

Therapy Off Mode
The Therapy Off mode disables automatic therapy delivery while still allowing manual control of shock delivery. Programmable parameters may be viewed and adjusted via the programmer. The subcutaneous electrogram (S-ECG) may be displayed or printed from this mode.

The pulse generator defaults to Therapy Off mode when it is taken out of Shelf mode.

Note: Manual and rescue shock therapy are available when the device is set to Therapy On or Therapy Off mode and is actively communicating with a pulse generator, but only after the initial Setup process is complete. Refer to Automatic Setup on page 39.
MRI Protection Mode

MRI Protection Mode is available in EMBLEM S-ICD devices.

MRI Protection Mode modifies certain pulse generator functions in order to mitigate risks associated with exposing the S-ICD system to the MRI environment. Choosing MRI Protection Mode will initiate a sequence of screens to assess the patient’s eligibility and readiness to undergo an MR Conditional MRI scan. Refer to the Summary Report to find out whether the device has been in MRI Protection Mode. For a complete description of MRI Protection Mode, a list of MR Conditional devices, and additional information about the ImageReady S-ICD System, refer to the MRI Technical Guide.

Prior to the patient undergoing an MRI scan, an ImageReady S-ICD System must be programmed to the MRI Protection Mode using the programmer. In MRI Protection Mode:

- Tachycardia therapy is suspended
- A Time-out feature is nominally set to 6 hours, with programmable values of 6, 9, 12, and 24 hours
- Beeper is disabled

MRI Protection Mode is terminated by manual exit or through the user-programmed automatic MRI Protection Time-out period (refer to the MRI Technical Guide for MRI Protection Mode programming instructions.) Rescue Shock will also terminate MRI Protection Mode. When MRI Protection Mode is exited, all parameters (except for the Beeper) return to the previously programmed settings.

**Note:** The Beeper can be reenabled after exiting MRI Protection Mode.

Connecting and Disconnecting from the S-ICD Pulse Generator

This section provides the information necessary for selecting, connecting to, and disconnecting from the pulse generator.

**Caution:** Use only the designated Boston Scientific S-ICD programmer and appropriate software application to communicate with and program the S-ICD pulse generator.

**Scanning for Pulse Generators**

1. Select the Scan For Devices button on the programmer start-up screen (Figure 2). The Scan Progress Bar is displayed during the scanning process, at the conclusion of which the Device List screen appears. Select the Cancel button at any time to end the scanning process.
2. When the scanning process is complete, a list of all pulse generators detected (up to 16) will be displayed on the Device List screen (Figure 7). The devices that are in Shelf mode will be displayed as “Not Implanted.” Any devices that were previously taken out of Shelf mode are displayed either as “Implanted” or with the stored patient name.

![Device List screen](image)

**Figure 7: Device List screen (scrollable list)**

3. If the desired pulse generator is not listed, select the Scan Again button to re-initiate the scanning process. Select the Cancel button to return to the programmer start-up screen.

   **Note:** Refer to the Inability to Communicate With the Pulse Generator heading within the Troubleshooting section for further assistance.

**Connecting to a Pulse Generator**

Select the desired pulse generator from the Device List screen (Figure 7) to initiate the communication session.

   **Note:** Regardless of how many pulse generators are located by a scan, the user must select a specific pulse generator from the list in order to begin active communication.
Connecting to a Pulse Generator in Shelf Mode

1. The programmer connects to the selected pulse generator after the selection is made. A window will appear indicating connection is in process.

2. The Device Identification screen appears once communication is established with the pulse generator.

   Note: The Device Identification screen is visible only while connecting to a pulse generator in Shelf mode.

3. The device model and serial numbers are automatically acquired and displayed during the initial scanning process. Select Continue to remove the device from Shelf mode and prepare for implantation, or select Cancel to return to the Device List screen.

Connecting to an Implanted Pulse Generator

If an implanted pulse generator is chosen from the Device List screen, the following connection sequence occurs:

1. The programmer connects to the selected pulse generator after the selection is made. A window will appear indicating connection is in process.

2. The Device Status screen appears once communication is established with the pulse generator (Figure 16).

Ending a Patient Session

To end an Online patient session and return the programmer to its Offline operation mode:

1. Select the Main Menu icon on the Navigation Bar. The Main Menu screen appears.

2. Select the End Session button (Figure 8).
3. Warning messages (Figure 9) will be displayed to the user if any of the following conditions exists:

- Therapy mode is programmed to Off
- Reference S-ECG has not been acquired
- Automatic or Manual Setup has not been completed
- Optimization has not been completed. This message is displayed if Setup Optimization was not performed during the Automatic Setup process
4. Select the Continue button to end the patient session and return to the programmer start-up screen, or select Cancel to remain Online and return to the Main Menu screen.

**Note:** Once the Continue button is selected, the session is stored and communication is terminated.

**Note:** A telemetry session must be terminated using the End Session process as described in steps 1 through 4 above in order for data obtained during that session to be saved. If the programmer is powered off during a session, either automatically or manually, session data will not be saved.

**Note:** In order to confirm that Therapy Mode is set to On upon disconnection, always use the End Session process and review all displayed warning messages.
Programming the Pulse Generator at Implant

This section provides the information necessary for programming the pulse generator during an implant.

**Caution:** Use only the Model 3203 telemetry wand with the programmer.

**Caution:** The wand is a non-sterile device. Do not sterilize the wand. The wand must be contained in a sterile barrier before use in the sterile field.

**Caution:** The programmer is non-sterile and cannot be sterilized. It must remain outside the sterile field.

**Caution:** Confirm that the programmer is in communication with the intended implanted S-ICD pulse generator.

**Entering Electrode Information**

The programmer maintains information on the implanted electrode. To record this information for a patient’s new or replacement electrode:

1. Select the Main Menu icon.
2. Select the Implant button.
3. Select the Automatic Setup icon in the Navigation Bar. The Automatic Setup screen appears (Figure 12).
4. Select Set Electrode ID button.
Figure 10: Select the Set Electrode ID button to enter electrode information

Note: ECG and heart rate information is not present on the Automatic Setup and Electrode ID setup screens until the electrode has been connected to the pulse generator.

5. Enter the electrode model and serial number.
6. Select the Program button to save the information. A confirmation screen will appear during communication with the device. Select Cancel to cancel information storage and return to the Automatic Setup screen.

Creating the Patient Chart

This chart contains reference information for the patient. To set up the patient chart:

1. Select the Main Menu icon on the Navigation Bar.
2. Select the Implant button.
3. Select the Patient View icon to access the Patient View screen (Figure 11).
4. The pulse generator model and serial numbers appear on the first line of the chart. The electrode model and serial numbers appear on the second line of the chart. The implant date appears on the third line of the chart. Using the on-screen keyboard, enter the following patient information:
- Patient Name: up to 25 characters
- Doctor Name: up to 25 characters
- Doctor Info: up to 25 characters
- Notes: up to 100 characters

![Patient View screen]

**Figure 11:** Patient View screen

*Note:* The Notes field will automatically wrap the text with the presence of a space between any characters within the first line.

5. Select the Save button to update the pulse generator with the patient information.

*Note:* Failure to save the new patient information will result in loss of the entered data.

**Automatic Setup**

Before the S-ICD device can be activated, it must go through an initial Automatic Setup Process at the time of the implant.
The Automatic Setup Process is initiated as follows:

1. Select the Main Menu icon.
2. Select the Implant button.
3. Select the Automatic Setup icon on the navigation bar. The Automatic Setup screen appears. Select the Automatic Setup button on this screen to advance to the next screen.
4. Select Continue if the patient's heart rate is less than 130 bpm (Figure 12). For rates greater than 130 bpm, select the Cancel button and refer to the Manual Setup section on page 64.

5. Once started the Automatic Setup will:
   - Perform the shock electrode integrity check to measure electrode impedance. Normal sub-threshold impedance range is < 400 Ohms.
   - Select the best sensing configuration. SMART Pass will be automatically configured based on the amplitude of the ECG signals in the selected vector. The sense electrode configuration appears on the printed report and can be viewed via the Manual Setup process. The status of SMART Pass (On/Off) is displayed on the SMART
Settings programmer screen and Summary Report (for more information on SMART Charge and SMART Pass, see SMART Settings on page 66).

- Select the appropriate gain selection. The selected sense gain appears on the Printed Report and can be viewed via the Manual Setup process.

![Automatic Setup Screen](image)

**Figure 13: Measuring electrode impedance**

The overall progress of the Automatic Setup process is shown in the status bar (Figure 13). When each function is complete, the arrow next to the function moves to a down position.

6. The Automatic Setup optimization process will be initiated. The programmer will display a message requesting that the patient sit up. If Automatic Setup is being performed during implant, or if the patient is unable to sit up for some other reason, this step can be omitted by selecting the Skip button. If desired, Automatic Setup can be repeated during a follow-up session to include the optimization step.

7. Select the Continue button to finish the Automatic Setup process. A confirmation screen will appear when Automatic Setup is complete.

8. Following the optional optimization process, the Acquire Reference S-ECG screen is displayed. Select the Continue button to acquire a reference S-ECG.
9. Once the Reference S-ECG acquisition process begins, a status screen appears. The process may take up to one minute, during which the patient should remain still. During this process, a template of the patient’s baseline QRS complex is stored in the pulse generator. Select Cancel at any time to end Reference S-ECG acquisition. When acquisition is complete, select the Continue button.

Programming Therapy Parameters

Once Automatic Setup has been completed, the pulse generator therapy parameters may be selected.

To set the therapy parameters:
1. Select the Main Menu icon on the Navigation Bar.
2. Select the Implant button.
3. Select the Device Settings icon on the Navigation Bar to display the Device Settings screen (Figure 14).

![Device Settings screen](image)

**Figure 14: Device Settings screen**

4. Select the desired therapy mode using the On/Off Therapy switch.
5. Select and drag the Conditional Shock Zone (yellow) and Shock Zone (red) slider bars to set the desired zone configuration.

**Note:** The S-ICD System Clinical Investigation demonstrated a significant reduction in inappropriate therapy with the activation of the Conditional Shock Zone prior to hospital discharge (see the S-ICD User’s Manual for a summary of the Clinical Investigation).

- The Shock Zone is programmable between 170 and 250 bpm in steps of 10 bpm.
- The Conditional Shock Zone is programmable between 170 and 240 bpm in steps of 10 bpm. Enhanced detection criteria are automatically enabled when the Conditional Shock Zone is programmed.
- When programming both the Shock Zone and Conditional Shock Zone, maintain at least a 10 bpm difference between the two zones. If the Conditional Shock Zone slider (yellow) is dragged over the Shock Zone slider (red), the two sliders will merge to create a single Shock Zone.

6. If post-shock pacing is desired, set the Post Shock Pacing switch to the On position. (Post-shock bradycardia pacing occurs at a non-programmable rate of 50 bpm for up to 30 seconds. Pacing is inhibited if the intrinsic rate is greater than 50 bpm.)

7. Select the Program button to apply the changes and program the pulse generator. A message confirming that the pulse generator settings were successfully programmed will appear. Select the Continue button to return to the Device Settings screen.

8. If the pulse generator does not accept the programming, a message with instructions will appear on the Device Settings screen. Press the Continue button after following the instructions.

**Warning:** The presence of other equipment operating in the same frequency bands used by the programmer (402-405 MHz for the pulse generator and 2.4 GHz for the printer) may interfere with communication. Interference can occur even if the other equipment complies with the International Special Committee on Radio Interference (CISPR) emission requirements. This RF interference can be reduced by increasing the distance between the interfering device and the programmer and pulse generator or printer. If communication problems persist, refer to the Troubleshooting section of this manual.

9. Once programming is confirmed, select the Continue button to proceed to the next operation.
Defibrillation Testing

Once the pulse generator is implanted and Therapy Mode is programmed On, defibrillation testing may be conducted. Prior to arrhythmia induction during implant procedure, the following recommendations for arm positioning are intended to reduce the potential for injury of the clavicle, arm and shoulder in the event of forceful muscle contraction:

- Avoid tight strapping of the arm to the arm board, and consider loosening arm restraints.
- Remove any wedge elevation below the torso, if used during the implant procedure, taking care to preserve the sterile field.
- Create a smaller angle of arm abduction from the torso by adducting the arm as close to the torso as feasible, taking care to preserve the sterile field. Temporarily place the hand in a neutral position while the arm is in a more adducted position, reverting to a supinated position if the arm needs to be abducted again.

Warning: During arrhythmia induction, the induction current and subsequent shock may result in forceful contraction of the pectoralis major muscle which can exert significant acute forces on the glenohumeral joint as well as on the clavicle. This, in conjunction with a tightly restrained arm, may result in injury to the clavicle, shoulder, and arm, including dislocation and fracture.

Warning: Always have external defibrillation equipment and medical personnel skilled in CPR available during implant and follow up testing. If not terminated in a timely fashion, an induced ventricular tachyarrhythmia can result in the patient’s death.

Caution: Successful VF or VT conversion during arrhythmia conversion testing is no assurance that conversion will occur post-operatively. Be aware that changes in the patient’s condition, drug regimen, and other factors may change the DFT, which may result in nonconversion of the arrhythmia post-operatively. Verify with a conversion test that the patient’s tachyarrhythmias can be detected and terminated by the pulse generator system if the patient’s status has changed or parameters have been reprogrammed.

Note: Defibrillation testing is recommended at implant, replacement, and concomitant device implants to confirm the ability of the S-ICD System to sense and convert VF.
**Note:** When the Hold to Induce button is pressed during defibrillation testing, the programmer begins capturing the episode data generated during the test. This data is available for viewing and printing (see Capturing and viewing S-ECG Strips on page 58 and Captured S-ECG Report on page 52.)

To induce VF and test the S-ICD System:

1. Select the Main Menu icon on the Navigation Bar to access the Main Menu.
2. Select the Patient Test button to setup the induction test (Figure 15).
3. Select either standard (STD) or reverse (REV) polarity.
4. Select and drag the red marker to set the desired shock energy for the first delivered shock. The shock energy may be programmed from 10 to 80 J. A 15 J safety margin is recommended for defibrillation testing.

**Figure 15:** Setting the desired first shock energy for defibrillation testing

5. Select the Continue button to display the next Induction Test screen or select the Cancel button to return to the Main Menu screen.

**Note:** Ensure that noise markers (“N”) are not present on the S-ECG prior to induction. The presence of noise markers may delay detection and therapy delivery.
6. Select and hold the Hold To Induce button for the desired duration.

The following functions occur during the test:

- The S-ICD System induces ventricular fibrillation using 200 mA alternating current (AC) at 50 Hz. Induction continues until the Hold To Induce button is released (up to a maximum of 10 seconds per attempt).

  **Note:** *If necessary, the induction can be terminated by disconnecting the wand from the programmer.*

- Arrhythmia detection and the Live S-ECG are suspended during induction. Once the Hold to Induce button is released, the programmer displays the patient’s rhythm.

- Upon detection and confirmation of an induced arrhythmia, the S-ICD System automatically delivers a shock at the programmed energy output and polarity.

  **Note:** *When in active communication with a pulse generator, the programmer emits an audible notification to indicate that the pulse generator is preparing to deliver a shock, whether that shock is commanded or is in response to a detected arrhythmia. The notification continues until the shock is either delivered or aborted.*

- If the shock fails to convert the arrhythmia, re-detection occurs and subsequent shocks are delivered at the pulse generator’s maximum energy output (80 J).

  **Note:** *Evaluate the sensing markers during the induced rhythm that follows the release of the Hold To Induce button. The S-ICD System uses a lengthened rhythm detection period. Consistent tachy “T” markers indicate that tachyarrhythmia detection is occurring, and that capacitor charging is imminent. If a high degree of amplitude variation is noted during the arrhythmia, a slight delay may be expected prior to capacitor charging or shock delivery.*

  **Note:** *The pulse generator can deliver a maximum of five shocks per episode. An 80 J rescue shock can be delivered at any time prior to therapy delivery by pressing the Rescue Shock icon.*

- The programmer will start capturing S-ECG data when the Hold to Induce button is pressed. The Captured S-ECG will cover the time period six seconds before the button is pressed and up to 102 seconds afterwards, for a maximum total of 108
seconds. The Induction S-ECGs will be viewable and printable from the Captured S-ECG screen, labeled as “Induction S-ECG”.

7. At any time prior to therapy delivery, the programmed energy may be aborted by selecting the red Abort button.

8. Select the Exit button to return to the Main Menu screen.

Performing a Follow-up

Sensing Configuration and Automatic Setup

It is not necessary to perform Automatic Setup at each follow-up. If Sensing Optimization was skipped during the original implant setup, it may be performed during a follow-up.

Sensing should be re-evaluated if Automatic Setup is performed and results in a vector change. After the setup process is complete, evaluate the streaming S-ECG during a pectoral exercise. Sensing performance during high rate exercises can also be performed. Acceptable sensing will yield “S” markers synchronous to all QRS complexes. If other markers are noted, use the Manual Setup process to evaluate other sensing configurations.

Caution: Following any sensing parameter adjustment or any modification of the subcutaneous electrode, always verify appropriate sensing.

Note: If Manual Setup was previously used to override a sensing configuration, careful consideration should be taken when selecting Automatic Setup.

If an update to the reference S-ECG is desired due to a change in the patient’s resting ECG, follow the Acquire Reference S-ECG instructions.

Viewing Pulse Generator Status

Once communication is established, the programmer displays the Device Status screen which contains information regarding the current episodes and battery status of the pulse generator.

To navigate to this screen from another location:

1. Select the Main Menu icon.
2. Select the Follow Up button.
3. Select the Device Status icon on the Navigation Bar to display the Device Status screen.
4. The Device Status screen will appear showing an overview of all pulse generator activity since the last communication session (Figure 16).

![Device Status screen](image)

**Figure 16:** Device Status screen

The Device Status overview reports:
- Date of the last follow-up session
- Total number of shocks delivered since the last follow-up session
- Total number of treated episodes since the last follow-up session
- Total number of untreated episodes since the last follow-up session
- Total number of Stored AF episodes since the last follow-up session that are available for review

*Note:* Choosing the “View” button in the Episodes row allows navigation directly to the list of stored episodes (Figure 17).

- Remaining pulse generator battery life

**Viewing Stored Episodes**

The pulse generator stores episodes, which can be viewed during a patient’s follow-up session. EMBLEM S-ICD (Model A209) and Cameron Health (Model 1010) pulse generators store up to 25 treated and 20 untreated tachycardia episodes. EMBLEM MRI S-ICD (Model A219) pulse generators
store S-ECGs for up to 20 treated and 15 untreated tachycardia episodes, as well as up to 7 AF episodes. When the maximum number of episodes is reached, the most recent episode replaces the oldest stored episode of the same type. The first treated episode is never overwritten.

**Note:** *Spontaneous episodes that occur while the pulse generator is communicating with the programmer will not be stored.*

To view stored episodes:

1. Select the Main Menu icon.
2. Select the Follow Up button.
3. Select the Captured and Stored Episodes S-ECG icon from the Navigation Bar.
4. Select the Episodes option to access the Episodes screen (Figure 17).
5. Select an episode from the list. The selected episode will be downloaded from the pulse generator and displayed.

**Note:** *In order to be available for printing, episodes must first be individually selected and viewed from the Episodes Screen.*

![Episodes screen (scrollable list)](image_url)
6. The display screen for each selected episode also displays the programmed parameters and the stored S-ECG data at the time of episode declaration.

7. Select the Continue button on the display screen for the selected episode to return to the Episodes screen.

The following details are available for each episode:

**Treated Episodes**

Up to 128 seconds of S-ECG data is stored for each Treated Episode:

- **Pre-episode S-ECG:** Up to 44 seconds
- **First shock:** Up to 24 seconds of pre-shock S-ECG and up to 12 seconds of post-shock S-ECG
- **Subsequent shocks:** 6 seconds of pre-shock and 6 seconds post-shock S-ECG

**Untreated Episodes**

An Untreated Episode is defined as any high-rate episode that spontaneously terminates during the charging process, before a shock is delivered.

Up to 128 seconds of S-ECG data is stored for each Untreated Episode:

- **Pre-episode S-ECG:** 44 seconds of pre-episode S-ECG
- **Episode S-ECG:** Up to 84 seconds of tachycardia S-ECG data

**Printing Reports from the Programmer**

**Printing Reports**

Patient reports can be printed before or after a patient session is ended. It is recommended that a final report be printed immediately following the implant procedure. There are three patient reports:

- Summary Report
- Captured S-ECG Report
- Episode Reports

To print patient reports from either an Online or Offline session:
1. Select the Main Menu icon to display the Main Menu screen.
2. Select the Print Reports button to display the Print Reports screen (Figure 18).

![Print Reports screen]

**Figure 18:** Print Reports screen

3. Select the desired report type. A checkmark will appear next to the selected report. Report types are described below.
4. Select the Print button to print the selected report.
5. Select the Cancel button to return to the previously accessed screen.

**Summary Report**

To print a summary report, select the Summary Report option on the Print Reports screen and press the Print button. The report will print for either the current active session (if the programmer is Online) or for the chosen stored session (if the programmer is Offline.)

The Summary Report includes the following information:

- Printed Report Date
- Programmer Software Version
- Pulse generator Software Version
EMBLEM™ S-ICD PROGRAMMER: OPERATION

- Patient Name
- Date of Last Follow-Up
- Date of Current Follow-Up
- Implant Date
- Pulse generator Model/Serial Number
- Electrode Model/Serial Number
- Therapy Parameters
- SMART Charge Delay
- SMART Pass status (On/Off)
- Programmed Gain Settings and Sensing Configuration
- Pulse generator Integrity Check, if applicable
- Initial Shock Polarity Configuration
- Days with measured AF
- Estimate of measured AF
- Beeper Status (if disabled)
- MRI Information
- Episode Summary: Since Last Follow-Up and Since Initial Implant
- Battery Status
- Electrode Impedance Measurement

**Captured S-ECG Report**

To print a Captured S-ECG report:

1. Select the Captured S-ECG Reports option from the Print Reports screen.
2. A scrollable list containing both Captured S-ECG and Induction S-ECG strips is displayed (Figure 19). Select the desired S-ECG(s) to be printed by placing a checkmark next to the selection(s).
3. Select Continue to return to the Print Reports screen.
4. Select the Print button to print the selected report and return to the previously accessed screen.
5. Select the Cancel button to return to the previously accessed screen without printing the report.

**Episode Reports**

To print an Episode Report:

1. Select the Episode Reports option on the Print Reports screen.
2. The Select Episode Reports screen appears showing a list of the stored episodes (Figure 20). Select the episode(s) to be printed. A checkmark appears next to the selected episode(s).

   **Note:** In order to be available for printing, episodes must have been individually selected and viewed from the Episodes Screen (Figure 17).
3. Select Continue to return to the Print Reports screen. Either 12 seconds or 44 seconds of pre-episode S-ECG data may be selected using the radio buttons below the Episode Reports row. The default value for Episode Report Onset is 12 seconds.

4. Select the Print button to print the selected report and return to the previously accessed screen.

5. Select the Cancel button to return to the previously accessed screen without printing the report.

Export Patient Data

Patient data saved on the programmer may be exported to a desktop or notebook computer using either of two means: wirelessly, over a pre-configured Bluetooth™ pairing, or with a Model 3205 microSD™ log data card. For information about Bluetooth™ pairing between the programmer and a desktop or laptop computer, see the Bluetooth™ Data Export section on page 28.

**Export using Bluetooth™ wireless technology**

Ensure that the programmer and the intended recipient computer are within 10 meters (33 feet) of each other before attempting a Bluetooth™ wireless data transfer.
1. Select the Export Programmer Data button on the programmer start-up screen. The Export Programmer Data Over Bluetooth screen will appear.

2. Select one of the three export options (Export Today’s Data, Export Last Seven Days, Export All). The “Select a receiving computer” pop-up window will appear.

   **Note:** *The Export Today’s Data and Export Last Seven Days options typically take less time to complete than the Export All option.*

3. The pop-up contains a scrollable list of all of the computers the programmer has been paired with. Select the intended receiving computer from the list to begin the transfer.

   **Note:** *Although every paired computer is listed in the scrollable box, only those within 10 meters (33 feet) of the programmer can participate in a file transfer. Once one of the three export options has been selected, the programmer will prepare the file transfer package and attempt the wireless transfer. An error message will appear if the transfer cannot be completed. Should this occur, move the programmer to within 10 meters of the intended receiving computer or choose another computer within that distance. Restart the export process by selecting one of the three export options on the Export Programmer Data Over Bluetooth screen.*

---

**Export using a microSD™ card**

Data may also be exported using a microSD™ card. For security reasons, the programmer will only export data to Model 3205 microSD™ log data cards. Using any other microSD™ card will cause an error message (invalid card) to appear.

1. Navigate to the programmer start-up screen.

2. Insert the card into the microSD™ slot according to the instructions presented in *Appendix A: Insertion and Removal of the microSD™ Card*. The instructions are also provided with Model 3205 microSD™ log data card. A Copy Data screen will appear when the microSD™ card has been properly inserted and recognized.

   **Note:** *An invalid card error message will appear if any card other than a Model 3205 microSD™ log data card is inserted. The message may also appear if the programmer does not recognize the Model 3205 card after insertion. Should this occur, remove the card and select the OK button on the error screen. Wait for the programmer start-up screen to reappear and then reinsert the card.*
3. Select the Copy Data button on this screen and the next screen.

4. A confirmation screen is presented when the copy process is complete. Selecting the OK button will return the programmer to its start-up screen.

5. Remove the microSD™ card according to the instruction sheet (Appendix A).

   **Note:** Navigating away from and returning to the start-up screen with the microSD™ card still inserted will cause the export process to begin again.

**S-ECG Features**

The programmer provides the capability to view, adjust and capture the streaming S-ECG from the pulse generator.

**S-ECG Rhythm Strip Markers**

The system provides annotations to identify specific events on the S-ECG. These markers are shown in the S-ECG Markers on Programmer Display Screens and Printed Reports table (Table 2).
Table 2: **S-ECG Markers on Programmer Display Screens and Printed Reports**

<table>
<thead>
<tr>
<th>Description</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charging *</td>
<td>C</td>
</tr>
<tr>
<td>Sensed Beat</td>
<td>S</td>
</tr>
<tr>
<td>Noisy Beat</td>
<td>N</td>
</tr>
<tr>
<td>Paced Beat</td>
<td>P</td>
</tr>
<tr>
<td>Tachy Detection</td>
<td>T</td>
</tr>
<tr>
<td>Discard Beat</td>
<td>•</td>
</tr>
<tr>
<td>Return to NSR*</td>
<td>🎯</td>
</tr>
<tr>
<td>Shock</td>
<td>⚡️</td>
</tr>
<tr>
<td>Episode data compressed or not available</td>
<td>🌈</td>
</tr>
</tbody>
</table>

* Marker present on printed report but not on programmer display screen.

**S-ECG Scale Settings**

To adjust the real-time S-ECG amplitude and display speed scale settings:

1. Select the S-ECG Display Settings icon located to the right of the Live S-ECG window. The S-ECG Settings screen is displayed.

2. Select and drag the Gain or Sweep Speed Scale bars as desired (Figure 21). The S-ECG scale will change according to the selected setting. The gain setting controls the visual gain. The programmer defaults to Full Range for pulse generators with a gain setting of 1x and to 1/2 Range for pulse generators with a gain setting of 2x. The Sweep Speed slider controls the display speed of the scrolling Live S-ECG. The nominal sweep speed setting is 25 mm/sec.
Note: Amplitude settings and display speed adjustments on scrolling real-time S-ECG and Captured S-ECGs affect the display screen settings only and have no impact on the pulse generator settings for sensing.

Figure 21: Adjusting Gain and Sweep Speed

Capturing and viewing S-ECG Strips

The programmer can display and store real-time S-ECG rhythm strips. The programmer saves a maximum of fifteen recordings generated by:

1. Manually-captured twelve-second S-ECGs using the Capture S-ECG button which include:
   • 8.5 seconds before activation of the Capture S-ECG button
   • 3.5 seconds after activation of the Capture S-ECG button
2. S-ECGs captured automatically during induction testing which include:
   • 6 seconds before the induce button is pressed
   • up to 102 seconds after the induce button is pressed
Note: The S-ICD suspends detection of sensed events for 1.6 seconds after a shock has been delivered. As a result, the S-ECG rhythm strip will not contain event markers during this 1.6 second post-shock interval.

If an additional recording is required, then the oldest previous recording is replaced with the new recording.

To manually capture a new S-ECG rhythm strip:

1. Select the Capture S-ECG button located to the left of the Live S-ECG window. The S-ECG will scroll across the display screen. Calipers appear below the Captured S-ECG rhythm strip. Each 12-second recording is date and time stamped according to the programmer’s date and time setting.

   Note: Induction S-ECGs are automatically generated during induction testing without additional user input.

2. Select and move the calipers across the S-ECG strip to measure intervals as desired.

3. Select the Continue button to return to the previously accessed screen.

It is also possible to capture S-ECGs corresponding to all three sense vectors (Primary, Secondary, and Alternate) by using the Capture All Sense Vectors button on the Utilities screen (Figure 22).

Viewing previously-captured S-ECGs

When the programmer is Online:

1. Select the Main Menu icon.

2. Select the Follow Up button.


4. Select one Captured S-ECG or Induction S-ECG from the list. The S-ECG details screen appears.

5. Select and drag the calipers to view details.

6. Select the Continue button to return to the Captured S-ECG list screen.
When the programmer is Offline:

1. Select the Stored Patient Sessions button from either the programmer start-up screen or the Main Menu.
2. Select the desired stored patient session.
3. Select one Captured S-ECG from the list. The Captured S-ECG Details screen appears.

   **Note:** Not all stored patient sessions contain captured S-ECGs. A message to that effect is presented when such patient sessions are opened. In this event select the Main Menu icon, then select the End Session button. This action returns you to the programmer start-up screen.

4. Select and drag the calipers to view details.
5. Select the Continue button to return to the Captured S-ECG list screen.

**Utilities Menu**

The programmer Utilities menu provides access to additional device features. These may include Acquire Reference S-ECG, Capture All Sense Vectors, Beeper Control, Manual Setup, SMART Settings, and AF Monitor.

To access the Utilities menu during an Online session:

1. Select the Main Menu icon to display the Main Menu screen.
2. Select the Utilities button. The Utilities screen appears (Figure 22).
Acquire Reference S-ECG

To acquire a manual Reference S-ECG:

1. From the Utilities screen (accessible from the Main Menu screen), select the Acquire Reference S-ECG button to access the Acquire Reference S-ECG screen.

2. Select Continue to acquire a Reference S-ECG. The programmer will begin acquiring the Reference S-ECG. A message will appear requesting that the patient remain still. The reference S-ECG QRS template is recorded and stored in the pulse generator.

3. Select the Continue button to complete the process and return to the Utilities screen. The Cancel button can be used at any time to end S-ECG acquisition and return to the Utilities screen.

Capture All Sense Vectors

The Capture All Sense Vectors button on the Utilities screen configures temporary programmer settings that allow you to capture S-ECGs generated from each of the three sense vectors (Primary, Secondary, and Alternate). This process takes approximately one minute. The programmer returns to its original settings configuration after all S-ECGs have been captured.
To capture the three sense vectors:

1. From the Utilities screen (accessible from the Main Menu screen), select the Capture All Sense Vectors button.

2. The Capturing 12 Second S-ECG screen will appear and display the status of the sense vector capture process.

Once captured, the three S-ECGs can be viewed by following the steps outlined in Viewing previously-captured S-ECGs on page 59.

**Beeper Control**

The pulse generator has an internal warning system (beeper) that may emit an audible tone to alert the patient to certain device conditions that require prompt consultation with the physician. These conditions may include:

- Electrode impedance out of range
- Prolonged charge times
- Failed Device Integrity Check
- Irregular battery depletion

This internal warning system is automatically activated at time of implant. Once triggered, if the beeper is enabled, tones beep for 16 seconds every nine hours until the trigger condition has been resolved. If the triggering condition reoccurs, then the tones will once again alert the patient to consult the physician.

**Caution:** Patients should be advised to contact their physician immediately whenever they hear beeping tones coming from their device.

**Note:** Access to the Reset Beeper display screen is enabled only when an alert condition occurs. If an alert condition is activated, a notification screen will appear upon connection.

**Warning:** The Beeper may no longer be usable following an MRI scan. Coming in contact with the strong magnetic field of an MRI scanner may cause a permanent loss of the Beeper volume. This cannot be recovered, even after leaving the MR scan environment and exiting MRI Protection Mode. Before an MRI procedure is performed, a physician and patient should weigh the benefit of the MR procedure against the risk of losing the Beeper. It is strongly recommended that patients are followed on LATITUDE NXT after
Reset Beeper

To reset the Beeper, select the Beeper Control button from the Utilities screen (accessible from the Main Menu) to open the Set Beeper Function screen.

Select the Reset Beeper button to suspend audible beeping tones triggered by an alert condition. If the alert condition is not corrected, the audible beeping tones will be reactivated during the next automatic S-ICD System self-check.

Disable Beeper (SQ-Rx devices)

In SQ-Rx devices, Beeper Control allows for the deactivation of beeping on alert conditions (Disable Beeper). Perform the following steps to disable the Beeper:

**Note:** The Disable Beeper function is only available once device ERI or EOL is reached.

1. From the Utilities screen, select Beeper Control to open the Set Beeper Function screen.
2. Select Disable Beeper to disable the Beeper for the device.

**Note:** This will permanently disable all beeping on alert conditions for the SQ-Rx device. However, this will not affect Beeper functionality for when a magnet is placed on the device or when a programmer connects to the device.

Enable/Disable Beeper (EMBLEM S-ICD devices)

In EMBLEM S-ICD devices, the Beeper must be tested before being enabled or disabled. Perform the following steps to test the Beeper:

**Note:** For EMBLEM S-ICD devices, the Test Beeper function is only available when beeping for an alert condition is not active.

1. From the Utilities screen, select Beeper Control.
2. Select the Test Beeper button from the Set Beeper Function screen.
3. Evaluate if the Beeper is audible using a stethoscope.
4. If the Beeper is audible, select the Yes, Enable Beeper button. If the Beeper is not audible or you wish to permanently disable beeping functionality, select the No, Disable Beeper button.

   **Note:** This will disable beeping functionality for alert conditions, for when a magnet is placed over the device, and for when a programmer connects to the device.

If the Beeper is not audible to the patient, it is strongly recommended that the patient has a follow-up schedule of every three months either on LATITUDE NXT or in-clinic to monitor device performance.

For additional information regarding the Beeper, refer to the MRI Technical Guide or contact Boston Scientific using the information on the back cover.

**Manual Setup**

Manual Setup enables the user to perform the electrode integrity test and select the electrode sensing configuration and gain setting in the pulse generator. During Manual Setup, the system will also automatically enable SMART Pass if appropriate.

1. From the Utilities screen (accessible from the Main Menu screen), select the Manual Setup button. The Measure Impedance screen appears.
2. Select the Test button to perform the electrode integrity test.
3. Select the Continue button.
4. There are three available sensing vectors that can be manually selected from the Manual Setup screen (Figure 23):
   - **Primary:** Sensing from the proximal electrode ring on the subcutaneous electrode to the surface of the active pulse generator
   - **Secondary:** Sensing from the distal sensing electrode ring on the subcutaneous electrode to the surface of the active pulse generator
   - **Alternate:** Sensing from the distal sensing electrode ring to the proximal sensing electrode ring on the subcutaneous electrode

The gain setting adjusts the sensed S-ECG signal sensitivity. It may be manually selected with the Select Gain switch on the Manual Setup screen.
To program the manually selected sense configuration:

1. Select the Program button to save the sense vector and gain settings.

2. Select the Continue button. When the continue button is selected, the device will automatically evaluate if SMART Pass should be enabled. Refer to the S-ICD User’s Manual for additional information about SMART Pass. For assistance, contact Boston Scientific using the information on the back cover.

3. The Acquire Reference S-ECG process is automatically enabled during the Manual Setup Process. Select the Continue button to acquire a reference S-ECG. A confirmation screen will appear when the captured reference S-ECG is acquired.
**SMART Settings**

The SMART Settings screen allows the user to access information and functions for the SMART Charge and SMART Pass features.

**SMART Charge**

Through the SMART Charge feature, the pulse generator charge initiation sequence adapts to the occurrence of non-sustained ventricular arrhythmia episodes by delaying capacitor charging. This conserves battery life and may prevent unnecessary shocks for non-sustained arrhythmias. Refer to the pulse generator manual for further information about the SMART Charge feature.

SMART Charge is enabled automatically when an untreated ventricular arrhythmia episode is recorded. Resetting returns the SMART Charge value to zero. To reset the SMART Charge feature:

1. From the Utilities screen (accessible from the Main Menu screen), select the SMART Settings button. The SMART Settings screen appears (Figure 24).

2. Select the Reset button to reset the SMART Charge to zero or press Cancel to return to the Utilities Menu without resetting the SMART Charge.
3. A confirmation window will appear with the message: “SMART Charge successfully reset.”

4. Press the Continue button to return to the Utilities screen.

**Disabling SMART Pass**

The SMART Pass feature is designed to reduce oversensing while still maintaining an appropriate sensing margin. The device continuously monitors the ECG signal amplitude and automatically disables SMART Pass if under-sensing is suspected.

SMART Pass can be manually disabled if under-sensing is suspected by selecting the Disable button on the SMART Settings screen.

**Note:** If SMART Pass is disabled, another automatic or manual setup must be performed to re-enable the feature.

**AF Monitor**

The AF Monitor feature is designed to assist in the diagnosis of atrial fibrillation.

The AF Monitor feature can be enabled/disabled using the On/Off switch accessed through the AF Monitor button on the Utilities screen. Select the Program button to apply the changes and program the pulse generator.

The following statistics are available on the programmer screen by selecting the AF Monitor button:

- **Days with measured AF:** Provides the number of days within the last 90 where AF was detected
- **Estimate of measured AF:** Provides the total percent of detected AF within the last 90 days

Refer to the S-ICD User’s Manual for further information about AF Monitor.

**Additional Programmer Functions**

**Rescue Shock**

The Rescue Shock icon is available in the navigation bar on the programmer display when the Setup Process is complete and a pulse generator is actively communicating with the programmer. During active communication, a maximum (80 J) rescue shock can be delivered upon programmer command.
To deliver a rescue shock:

1. Select the red Rescue Shock Icon at the top of the programmer screen. The Rescue Shock screen appears (Figure 25).

2. Select the Shock button to begin charging the pulse generator for a Rescue shock. A red background screen with the word “Charging” will appear. Selecting the Abort button will prevent delivery of a rescue shock and will return to the Device Settings screen.

3. A confirmation screen will appear with notification that the shock was delivered successfully along with the corresponding shock impedance.

   **Caution:** A reported shock impedance value of less than 25 ohms from a delivered shock could indicate a problem with the device. The delivered shock may have been compromised, and/or any future therapy from the device may be compromised. If a reported impedance value of less than 25 ohms is observed, correct functioning of the device should be verified.

If for any reason the shock could not be delivered, a red background screen will appear with a message stating “The shock could not be delivered.”

   **Note:** In the event telemetry is lost, pulse generator commands—including Rescue Shocks—will not be available until telemetry is reestablished.
**EMBLEM™ S-ICD PROGRAMMER: OPERATION**

*Manual Shock*

Manual Shock allows the user to deliver a synchronized shock during a sinus rhythm, an atrial rhythm or a ventricular rhythm. The shock energy level is user-configured in the 10 to 80 joule range and the polarity is also user-configured (Figure 26). Manual shock may also be utilized at a low energy to assess system impedance/integrity either at implant or as warranted by patient condition. A manual shock may be administered with the Therapy Mode set to On or Off.

To access Manual Shock, select the Patient Test button on the main menu. The Induction Test screen will appear. Select the Manual Shock icon in the navigation bar at the top of the screen to view the Manual Shock Test screen.

![Manual Shock Test Screen](image)

*Figure 26: Manual shock*

*S-ICD System Magnet Use*

The Boston Scientific magnet Model 6860 (the magnet) is a non-sterile accessory that may be used to temporarily inhibit the delivery of therapy from the pulse generator if necessary. The Cameron Health magnet Model 4520 may be used interchangeably with the Boston Scientific magnet for this purpose.

For detailed information about using the magnet, refer to the appropriate S-ICD User’s Manual.
Other behaviors of magnet application:

- Inhibit shock therapy delivery
- Terminate post-shock pacing therapy
- Prohibit arrhythmia induction testing
- Activate the pulse generator’s beeper with each detected QRS complex for 60 seconds if beeper is turned on and is audible

**Warning:** Use caution when placing a magnet over the S-ICD pulse generator because it suspends arrhythmia detection and therapy response. Removing the magnet resumes arrhythmia detection and therapy response.

**Warning:** In patients with a deep implant placement (greater distance between the magnet and the pulse generator) magnet application may fail to elicit the magnet response. In this case the magnet cannot be used to inhibit therapy.

**Caution:** Do not place a magnet on the programmer.

**Note:** A programmer commanded Rescue Shock can override the use of the magnet as long as the magnet was in place prior to the initiation of the programming command. If the magnet is applied after the initial command, the Rescue Shock will be terminated.

**Note:** Magnet application does not affect wireless communication between the pulse generator and the programmer.

**MAINTENANCE**

**Charging the Programmer**

When not in use, it is recommended that the programmer remain connected to the external power supply, which in turn is connected to the AC mains. This will ensure that the internal battery is adequately charged.

**Cleaning the Programmer**

Keep the programmer away from dust and dirt. Do not use harsh chemicals, cleaning solvents, or strong detergents to clean the programmer and wand.

To clean the programmer and wand as needed:

1. Turn the programmer off.
2. Gently wipe the programmer screen with a soft, clean, dry cloth.
3. Clean the programmer plastic case and the wand by wiping them with an isopropyl alcohol-moistened cloth.

4. Dry the programmer immediately to remove residue.

Service

There are no user-accessible or user-serviceable parts or components in the programmer. If any service, repair, or replacement of internal components is needed, the programmer must be returned to Boston Scientific. For instructions and return packaging, contact Boston Scientific using the information on the back cover of this manual.

When requesting service, please provide information concerning the nature of the failure and the manner in which the equipment was used when the failure occurred. The model number and serial number should also be provided.

Maintenance Check

Prior to each use, you should perform a visual inspection and verify the following:

- Mechanical and functional integrity of the programmer, cables, and accessories.
- Legibility and adherence of the programmer labels.
- That the programmer start-up screen appears a few seconds after you turn on the programmer. (The normal power-up process verifies that the programmer has passed its internal checks and is ready for use.)

Safety Measurements

National regulations may require that the user, manufacturer, or manufacturer representative periodically perform and document safety tests of the programmer. If such testing is required in your country, follow the testing interval and extent of testing as regulated in your country. If you do not know the national regulations in your country, please contact Boston Scientific using the information on the back cover of this manual. If IEC/EN 62353 is a required standard in your country, but no specific testing or interval is specified, it is recommended that you perform these safety tests using the direct method as specified in IEC/EN 62353 at an interval of every 24 months. Test values are shown in the Nominal Specifications table (Table 11).

Programmer End of Life
The programmer and accessories are designed to provide years of service under typical use. To dispose of, return, or exchange a programmer, contact Boston Scientific using the information on the back cover of this manual. Do not dispose of the programmer in the trash or at electronics recycling facilities.

TROUBLESHOOTING

This section presents potential programmer issues and the possible solutions. Of note, restarting the programmer can often resolve many of the issues listed below. The programmer can be restarted by pressing and holding the power button until the system shutdown menu appears and then choosing the “Restart” option.

Contact Boston Scientific using the information on the back cover of this manual for additional assistance.

Inability to Print

If unable to print, follow the steps below:

1. Ensure that the printer is turned On and that it contains paper and a sufficient ink supply.
2. Check printer feed for paper jam.
3. Ensure, as applicable, that the wireless function is enabled on the printer or that the Bluetooth™ wireless adapter is fully inserted into the USB slot on the printer.

No Printer Available

The No Printer Available screen will appear if a printer was not set up. Select the Try Again button or refer to the Printer Selection section for instructions.

Touch Screen Inactive while Connected to AC Power

If the touch screen does not function while the programmer is connected AC power via the external power supply, disconnect and reconnect the external power supply and restart the programmer.
Loss of Communication with Printer

When communication between the programmer and the printer fails, a Printing Error screen will appear with a message stating “Error while printing reports. Press ‘Continue’ to try printing any remaining reports, or ‘Cancel’ to cancel the current print job.”

If this occurs:

1. Select the Try Again button to reconnect to the printer.
2. Ensure, as applicable, that the wireless function is enabled on the printer or that the Bluetooth™ wireless adapter is fully inserted into the USB slot on the printer.
3. Move the programmer closer to the printer.
4. Move any devices and the associated cables that may be interfering with the RF communication.

Inability to Communicate With the Pulse Generator

If the programmer is unable to communicate with the pulse generator, follow the steps below:

1. Attempt to reposition the wand.
2. Select Scan For Devices from the programmer start-up screen or select Scan Again from the Device List screen to locate the desired device.
3. Move any equipment and associated cables that may be interfering with RF communication.
4. If available, attempt to communicate using a different S-ICD System programmer and/or wand.
5. Apply a pulse generator magnet to the pulse generator to elicit beeper tones. Remove the magnet and re-attempt communication.
EMBLEM™ S-ICD PROGRAMMER: COMPLIANCE STATEMENTS

COMPLIANCE STATEMENTS

EMI/RFI
This equipment has been tested and found to comply with the applicable limits for medical devices, IEC 60601-1-2:2007 or Active Implantable Medical Device Directive 90/385/EEC.

Although this testing shows the device to provide reasonable protection against harmful interference in a typical medical installation, there is no guarantee that interference will not occur in a particular installation. If the device does cause harmful interference the user is encouraged to try and correct the interference by the following measures:

• Reorient or relocate the device
• Increase the separation between the devices
• Connect the equipment to an outlet on a different circuit
• Contact Boston Scientific using the information on the back cover of this manual.

Essential Performance
In order for the Model 3200 Programmer to meet its intended use, it must interrogate and maintain a communications link with an S-ICD pulse generator as well as being able to appropriately detect touch screen button presses. Therefore those functions that pertain to communications with the implanted cardioverter defibrillator and detection of touch screen presses are considered essential performance.

Federal Communications Commission (FCC) Compliance

FCC ID: ESCRM320014
Contains: FCC ID ESCCRM70514

This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions:

1. This device may not cause harmful interference, and
2. this device must accept any interference received, including interference that may cause undesired operation

This transmitter is authorized by rule under the Medical Device Radiocommunication Service (in part 95 of the FCC Rules) and must not cause harmful interference to stations operating in the 400.150–406.000 MHz band in the Meteorological Aids (i.e. transmitters and receivers used to communicate weather data), the Meteorological Satellite, or the Earth Exploration Satellite Services and must accept interference that may be caused by such stations, including interference that may cause undesired operation. This transmitter shall be used only in accordance with the FCC Rules governing the Medical Device Radiocommunication Service. Analog and digital voice communications are prohibited. Although this transmitter has been approved by the Federal Communications Commission, there is no guarantee that it will not receive interference or that any particular transmission from this transmitter will be free from interference.
CAUTION: Changes or modifications not expressly approved by Boston Scientific could void the user's authority to operate the equipment.

Industry Canada (IC) Compliance

IC 4794A-CRM32004
Contains: IC 4794A-CRM7054

This device complies with Industry Canada license-exempt RSS standard(s). Operation is subject to the following two conditions:

- This device may not cause harmful interference, and
- This device must accept any interference received, including interference that may cause undesired operation.

CAUTION: Changes or modifications not expressly approved by Boston Scientific could void the user's authority to operate the equipment.

This device may not interfere with stations operating in the 400.150-406.000 MHz band in the meteorological aids, meteorological-satellite, and earth exploration-satellite services, and must accept any interference received, including interference that may cause undesired operation.
EMBLEM™ S-ICD PROGRAMMER: DECLARATIONS TABLES

Table 3: Declaration Electromagnetic Emission

The Model 3200 programmer is intended for use in the electromagnetic environment specified below. The customer or the user of the Model 3200 programmer should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions CISPR 11</td>
<td>Group 1</td>
<td>The Model 3200 programmer uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF Emissions CISPR 11</td>
<td>Class A</td>
<td>The Model 3200 programmer is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic Emissions IEC 61000-3-2</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>Voltage Fluctuations/Flicker Emissions IEC 61000-3-3</td>
<td>Complies</td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Declaration Electromagnetic Immunity Part 1

The Model 3200 programmer is intended for use in the electromagnetic environment specified below. The customer or the user of the Model 3200 should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge (ESD) IEC 61000-4-2</td>
<td>± 6 kV contact ± 8 kV air</td>
<td>± 6 kV contact ± 8 kV air</td>
<td>Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, then the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>Electrical Fast Transient/Burst IEC 61000-4-4</td>
<td>± 2 kV for power supply lines ± 1 kV for input/output lines</td>
<td>± 2 kV for power supply lines ± 1 kV for input/output lines</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>± 1 kV line(s) to line(s) ± 2 kV line(s) to earth</td>
<td>± 1 kV line(s) to line(s) ± 2 kV line(s) to earth</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>Voltage Dips, Short interruptions, and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>&lt;5% $U_i$, (&gt;95% dip in $U_i$) for 0.5 cycles 40% $U_i$, (60% dip in $U_i$) for 5 cycles 70% $U_i$, (30% dip in $U_i$) or 25 cycles &lt;5% $U_i$, (&gt;95% dip in $U_i$) for 5 sec</td>
<td>&lt;5% $U_i$, (&gt;95% dip in $U_i$) for 0.5 cycles 40% $U_i$, (60% dip in $U_i$) for 5 cycles 70% $U_i$, (30% dip in $U_i$) for 25 cycles &lt;5% $U_i$, (&gt;95% dip in $U_i$) for 5 sec</td>
<td>Mains power quality should be that of a typical commercial or hospital environment. If the user of the Model 3200 programmer requires continued operation during power mains interruptions, it is recommended that the Model 3200 programmer be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>Power Frequency (50/60 Hz) Magnetic Fields IEC 61000-4-8</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

NOTE: $U_i$ is the AC mains voltage prior to application of the test level.
Table 5: Declaration Electromagnetic Immunity Part 2

The Model 3200 programmer is intended for use in the electromagnetic environment specified below. The customer or the user of the Model 3200 programmer should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>3 Vrms</td>
<td>3 V</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Model 3200 programmer, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended Separation Distance</td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>150 kHz to 80 MHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>3 V/m</td>
<td>3 V/m</td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range. Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
<tr>
<td>IEC 61000-4-3</td>
<td>80 MHz to 2.5 GHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>where $P$ is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and $d$ is the recommended separation distance in meters (m).</td>
</tr>
</tbody>
</table>

Note 1: At 80 MHz and 800 MHz, the higher frequency range applies.
Note 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

$^{a}$ Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Model 3200 programmer is used exceeds the applicable RF compliance level above, the Model 3200 programmer should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Model 3200 programmer.

$^{b}$ Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m
Table 6: Recommended Separation Distances

Recommended separation distances between portable and mobile RF communications equipment and the Model 3200 programmer

The programmer is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the programmer can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the programmer as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 KHz to 80 MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>( d = \frac{3.5}{\sqrt{P}} )</td>
<td>( d = \frac{3.5}{E_i\sqrt{P}} )</td>
</tr>
<tr>
<td>0.01</td>
<td>0.117</td>
</tr>
<tr>
<td>0.1</td>
<td>0.369</td>
</tr>
<tr>
<td>1</td>
<td>1.17</td>
</tr>
<tr>
<td>10</td>
<td>3.69</td>
</tr>
<tr>
<td>100</td>
<td>11.7</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance \( d \) in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where \( P \) is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

Note 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Table 7: EMI/RFI Information: Programmer-to-Pulse Generator Communication

<table>
<thead>
<tr>
<th>Specification</th>
<th>Medical Implant Communications Service (MICS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency band</td>
<td>402-405 MHz</td>
</tr>
<tr>
<td>Modulation type</td>
<td>FSK</td>
</tr>
<tr>
<td>Radiated Power</td>
<td>&lt;25 μW</td>
</tr>
<tr>
<td>Bandwidth</td>
<td>&lt; 300 KHz</td>
</tr>
</tbody>
</table>

Table 8: EMI/RFI Information: Bluetooth™ Wireless Printing and Data Transfer

<table>
<thead>
<tr>
<th>Specification</th>
<th>Bluetooth™ wireless technology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency band</td>
<td>2.402-2.480 GHz</td>
</tr>
<tr>
<td>Modulation type</td>
<td>GFSK, π/4-DQPSK, 8DPSK</td>
</tr>
<tr>
<td>Radiated power</td>
<td>&lt;20 mW</td>
</tr>
<tr>
<td>Bandwidth</td>
<td>&lt; 1.5 MHz</td>
</tr>
</tbody>
</table>
**Table 9: Product Guidelines**

<table>
<thead>
<tr>
<th>Component</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DC Power</strong></td>
<td></td>
</tr>
<tr>
<td>Battery pack type</td>
<td>4000 mAh 3.7 volt lithium-ion battery pack</td>
</tr>
<tr>
<td>Charge time</td>
<td>Approximately 5 hours</td>
</tr>
<tr>
<td><strong>Power Supply</strong></td>
<td></td>
</tr>
<tr>
<td>Input</td>
<td>100 – 240 VAC, 50 – 60 Hz, 0.5A</td>
</tr>
<tr>
<td>Output</td>
<td>5.5 VDC, 3.64A</td>
</tr>
<tr>
<td></td>
<td>Power: 20 W</td>
</tr>
<tr>
<td>Manufacturer/Model</td>
<td>Elpac Power Systems MWA020005A</td>
</tr>
<tr>
<td><strong>Environment</strong></td>
<td></td>
</tr>
<tr>
<td>Temperature</td>
<td>15°C to +38°C</td>
</tr>
<tr>
<td></td>
<td>(+59°F to +100°F)</td>
</tr>
<tr>
<td></td>
<td>-10°C to +55°C</td>
</tr>
<tr>
<td></td>
<td>(+14°F to +131°F)</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>5% to 93% maximum at 40°C, non-condensing</td>
</tr>
<tr>
<td></td>
<td>5% to 93% maximum at 40°C, non-condensing</td>
</tr>
<tr>
<td>Atmospheric pressure</td>
<td>50 kPa to 106 kPa</td>
</tr>
<tr>
<td></td>
<td>(7.252 psi to 15.374 psi)</td>
</tr>
</tbody>
</table>

**Table 10: Specifications**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimensions Width x Depth x Height</td>
<td>24.0 cm x 12.7 cm x 2.6 cm</td>
</tr>
<tr>
<td></td>
<td>9.4 in x 5.0 in x 1.0 in</td>
</tr>
<tr>
<td>Weight</td>
<td>.6 kg, 1.3 lbs</td>
</tr>
<tr>
<td>Standard Screen Display</td>
<td>WVGA, 1024 x 600 pixels, 16M TFT</td>
</tr>
<tr>
<td>Table 11: Nominal Specifications (with device connected to external power supply)</td>
<td></td>
</tr>
<tr>
<td>--------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td><strong>Characteristic</strong></td>
<td><strong>Nominal</strong></td>
</tr>
<tr>
<td>Earth resistance</td>
<td>Not accessible</td>
</tr>
<tr>
<td>Earth leakage current</td>
<td>5 mA Normal Condition (NC)</td>
</tr>
<tr>
<td></td>
<td>10 mA Single Fault Condition (SFC)</td>
</tr>
<tr>
<td>Patient leakage current</td>
<td>100 μA Normal Condition (NC)</td>
</tr>
<tr>
<td></td>
<td>500 μA Single Fault Condition (SFC) (mains on applied parts)</td>
</tr>
<tr>
<td><strong>Electrical Safety Testing – IEC 62353:2008 allowed values</strong></td>
<td></td>
</tr>
<tr>
<td>Protective Earth Resistance</td>
<td>Not accessible</td>
</tr>
<tr>
<td>Equipment leakage — direct method</td>
<td>500 μA</td>
</tr>
<tr>
<td>Patient leakage current — direct method (Wand, BF)</td>
<td>&lt;= 5000 μA</td>
</tr>
<tr>
<td>Insulation resistance</td>
<td>Not accessible</td>
</tr>
<tr>
<td><strong>Safety Features</strong></td>
<td></td>
</tr>
<tr>
<td>Defibrillator protection</td>
<td>to 5000 V, 400 J</td>
</tr>
</tbody>
</table>
Table 12: Packaging and Device Symbols: Model 3200 Programmer

The following symbols may be used on the Model 3200 programmer, its accessories, and their packaging.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Specification</th>
<th>Symbol</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Follow instructions for use at <a href="http://www.bostonscientific-elabeling.com">www.bostonscientific-elabeling.com</a></td>
<td></td>
<td>Type BF Applied Part</td>
</tr>
<tr>
<td></td>
<td>Electrostatic discharge</td>
<td></td>
<td>Non-ionizing electromagnetic radiation</td>
</tr>
<tr>
<td></td>
<td>Temperature limitations</td>
<td></td>
<td>Humidity limitations</td>
</tr>
<tr>
<td></td>
<td>Atmospheric pressure limitations</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SN</td>
<td>Serial Number</td>
<td></td>
<td>Date of Manufacture</td>
</tr>
<tr>
<td></td>
<td>Reference Number</td>
<td></td>
<td>Authorized Representative in the European Community</td>
</tr>
<tr>
<td>LOT</td>
<td>Lot number</td>
<td></td>
<td>Non-sterile</td>
</tr>
<tr>
<td></td>
<td>Power plug storage</td>
<td></td>
<td>Door, open</td>
</tr>
<tr>
<td></td>
<td>Consult instructions for use</td>
<td></td>
<td>Proper insertion of microSD™ card</td>
</tr>
<tr>
<td></td>
<td>WEEE – Waste, Electrical, and Electronic Equipment (WEEE). Indicates separate</td>
<td></td>
<td>External power supply port</td>
</tr>
<tr>
<td></td>
<td>collection for electrical and electronic equipment (i.e., do not throw this</td>
<td></td>
<td>Mark for nationally recognized testing for safety</td>
</tr>
<tr>
<td></td>
<td>device in the trash).</td>
<td></td>
<td>standards</td>
</tr>
<tr>
<td>MR</td>
<td>MR Unsafe</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Limited Warranty

A limited warranty may apply to this programmer. For warranty eligibility and to obtain a copy of the limited warranty, contact Boston Scientific using the information on the back cover.
Lift the microSD slot cover and swing it away. Confirm that there is no card in the slot.

Place the microSD card into the empty slot with the lettering facing the same direction as the programmer screen.

Gently press the card into the slot until it latches into place. There will be increasing spring resistance followed by a softly audible click which indicates that the latch is activated.

The properly mounted card will be flush with the slot opening surface.

Close the microSD slot cover.

Lift the microSD slot cover and swing it away. Verify that a card is in the slot.

Press the edge of the card with your fingernail until the latch releases and you hear a click. Let the card gradually come out until you hear a second click, then remove your finger.

At this point it is safe to remove the card, although it may be difficult to grasp the card with your fingers. In this case the spring tensioner inside the slot can be used to eject the card.

Use a fingertip to slightly press the card back into the slot, then quickly slide your finger away from the slot. Note that the card may travel some distance when it is ejected.

Repeat this procedure as needed until the card is ejected.

Close the microSD slot cover.

Model 3200 S-ICD Programmer

Insert microSD  Remove microSD

APPENDIX A: INSERTION AND REMOVAL OF THE microSD™ CARD