## Ellipse™

Fortify Assura™

Tiered-therapy Cardioverter/Defibrillator

Quadra Assura™, Quadra Assura MP™

Unify Assura™

Cardiac Resynchronization Device, Tiered-therapy Cardioverter/Defibrillator



CAUTION: Federal (USA) law restricts this device to sale by or on the order of a physician.

WARNING: This product can expose you to chemicals including ethylene oxide, which is known to the State of California to cause cancer and birth defects or other reproductive harm. For more information, go to www.P65Warnings.ca.gov.

™ Indicates a trademark of the Abbott group of companies.

‡ Indicates a third-party trademark, which is property of its respective owner.

Pat. http://www.abbott.com/patents

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# **Device Description**

This manual describes the following Abbott Medical pulse generators:

Table 1. Single-chamber pulse-generator descriptions

Name	Model Number	Description	Connector Type	Delivered Energy (approx.)	MRI Status
Fortify Assura VR	CD1357-40C	Single-chamber ICD with RF telemetry, Parylene coating	DF-1/IS-1	40 J	Untested
Fortify Assura VR	CD1357-40Q	Single-chamber ICD with RF telemetry	DF4-LLHH	40 J	MR Conditional
Ellipse VR	CD1411-36C	Single-chamber ICD with RF telemetry, Parylene coating	DF-1/IS-1	36 J	Untested
Ellipse VR	CD1411-36Q	Single-chamber ICD with RF telemetry	DF4-LLHH	36 J	MR Conditional

Table 2. Dual-chamber pulse-generator descriptions

Name	Model Number	Description	Connector Type	Delivered Energy (approx.)	MRI Status
Fortify Assura DR	CD2357-40C	Dual-chamber ICD with RF telemetry, Parylene coating	DF-1/IS-1	40 J	Untested
Fortify Assura DR	CD2357-40Q	Dual-chamber ICD with RF telemetry	DF4-LLHH/IS-1	40 J	MR Conditional
Ellipse DR	CD2411-36C	Dual-chamber ICD with RF telemetry, Parylene coating	DF-1/IS-1	36 J	Untested
Ellipse DR	CD2411-36Q	Dual-chamber ICD with RF telemetry	DF4-LLHH/IS-1	36 J	MR Conditional

Table 3. CRT-D pulse generator descriptions

Name	Model Number	Description	Connector Type	Delivered Energy (approx.)	MR Status
Unify Assura	CD3357-40C	CRT-D with RF telemetry, Parylene coating	DF-1/IS-1	40 J	Untested
Unify Assura	CD3357-40Q	CRT-D with RF telemetry	DF4-LLHH/IS-1	40 J	Untested
Quadra Assura	CD3365-40C	CRT-D with RF telemetry, Parylene coating	DF-1/IS-1/ IS4-LLLL	40 J	Untested
Quadra Assura	CD3365-40Q	CRT-D with RF telemetry	DF4-LLHH/ IS4-LLLL/IS-1	40 J	MR Conditional
Quadra Assura MP	CD3369-40C	CRT-D with RF telemetry, Parylene coating	DF-1/IS-1/IS4- LLLL	40 J	Untested
Quadra Assura MP	CD3369-40Q	CRT-D with RF telemetry	DF4-LLHH/IS4- LLLL/IS1	40J	MR Conditional

The pulse generator, along with compatible, commercially available leads, constitutes the implantable portion of the ICD and CRT-D systems. The lead systems are implanted using either transvenous or transthoracic techniques. The Abbott Medical Merlin™ Patient Care System (PCS) with software model 3330 version 24.1.1 (or greater), a Merlin™ Antenna (for devices with RF communication), and a telemetry wand constitute the external portion of the ICD and CRT-D systems.

Models with the "Q" suffix are functionally equivalent in all respects to the same model without the "Q" suffix, except for the header. Models without the "Q" suffix use DF-1 lead connectors for the high-voltage leads, and models with the "Q" suffix use a single DF4-LLHH lead connector for the high-voltage leads and for the low voltage RV lead.

SJ4-LLHH is equivalent to DF4-LLHH. SJ4 and DF4 connectors comply with ISO 27186:2010(E).SJ4-LLLL is equivalent to IS4-LLLL. SJ4 and IS4 connectors comply with ISO 27186:2010(E).

## **Indications and Usage**

Abbott Medical ICDs and CRT-Ds are intended to provide ventricular antitachycardia pacing and ventricular defibrillation for automated treatment of life-threatening ventricular arrhythmias.

AF Suppression™ pacing is indicated for suppression of paroxysmal or persistent atrial fibrillation in patients with the above ICD indication and sinus node dysfunction.

In patients indicated for an ICD, CRT-Ds are also intended:

 to provide a reduction of the symptoms of moderate to severe heart failure (NYHA Functional Class III or IV) in those patients who remain symptomatic despite stable, optimal medical therapy (as defined in the

- clinical trials section included in the Merlin PCS on-screen help) and have a left ventricular ejection fraction less than or equal to 35% and a prolonged QRS duration
- to maintain synchrony of the left and right ventricles in patients who have undergone an AV nodal ablation for chronic (permanent) atrial fibrillation and have NYHA Class II or III heart failure

## **MR Conditional System**

An MR Conditional device is conditionally safe for use in the MRI environment when used in a complete MR Conditional system and according to instructions in the MRI-Ready Systems Manual for the Abbott Medical MR Conditional system.

#### **Contraindications**

Contraindications for use of the pulse generator system include ventricular tachyarrhythmias resulting from transient or correctable factors such as drug toxicity, electrolyte imbalance, or acute myocardial infarction.

## **Warnings and Precautions**

**Resuscitation Availability.** Do not perform device testing unless an external defibrillator and medical personnel skilled in cardiopulmonary resuscitation (CPR) are readily available.

**Lead system.** Do not use another manufacturer's lead system without demonstrated compatibility as undersensing cardiac activity and failure to deliver necessary therapy may result.

**Avoiding shock during handling.** Disable tachyarrhythmia therapy (Enable/ Disable Tachy Therapy) or program tachyarrhythmia therapies Off during surgical implant and explant or post-mortem procedures as well as when disconnecting leads as the device can deliver a serious shock if you touch the defibrillation terminals while the device is charged.

**Additional pacemaker implanted.** These devices provide bradycardia pacing. If another pacemaker is used, it should have a bipolar pacing reset mode and be programmed for bipolar pacing to minimize the possibility of the output pulses being detected by the device.

**Modifying the device.** This device has been tested for compliance to FCC regulations. Changes or modifications of any kind not expressly approved by Abbott Medical could void the user's authority to operate this device.

**Suboptimal radio frequency (RF) communication.** The Merlin PCS indicates the quality of the RF communication by the telemetry strength indicator LEDs on both the Merlin PCS and the Merlin Antenna. Below is a list of potential causes to suboptimal radio communication:

Table 4. Possible causes and solutions for suboptimal RF communication

Possible Causes	Solutions
The Merlin Antenna orientation/location is suboptimal.	Move or reorient the Merlin Antenna slightly. Make sure that the front of the Merlin Antenna faces the implantable device.
People or objects interfere with the communication between	Make sure that the space between the Merlin Antenna and the device is free from interfering objects/people.

Table 4. Possible causes and solutions for suboptimal RF communication

Possible Causes	Solutions
the Merlin Antenna and the device.	
The Merlin Antenna is too far away from the device.	Move the Merlin Antenna closer to the device.
Someone is holding the Merlin Antenna.	Place the Merlin Antenna on a flat surface. Do not hold the Merlin Antenna.
Other products in the vicinity are causing electromagnetic interference (EMI).	Power off or remove equipment that could cause EMI.
The Merlin Antenna cable is wound around the Merlin Antenna.	Make sure the Merlin Antenna cable is not wound around the Merlin Antenna.

## Magnetic Resonance Imaging (MRI)

 MR Conditional devices. Testing has demonstrated that the Abbott Medical MR Conditional system is conditionally safe for use in the MRI environment when used according to the instructions in the MRI-

- Ready Systems Manual. The Abbott Medical MR Conditional system includes an Abbott Medical MR Conditional pulse generator connected to one or more Abbott Medical MR Conditional leads.
- MR Untested devices. "Untested" indicates that the device has not been tested and its use in an MR environment is not determined. For more information, please consult the MRI-Ready Systems Manual.

## Sterilization, Storage and Handling

**Resterilization.** Do not resterilize and re-implant explanted pulse generators.

**Use before date.** Do not implant the device after the "use before" date because the battery may have reduced longevity.

If package is damaged. Do not use the device or accessories if the packaging is wet, punctured, opened or damaged because the integrity of the sterile packaging may be compromised. Return the device to Abbott Medical.

**Device storage.** Store the device in a clean area, away from magnets, kits containing magnets, and sources of electromagnetic interference (page 17) to avoid device damage. Store the device at room temperature of 72°F (22°C). Excursions are permitted between 59°F and 86°F (15°C and 30°C). During transportation and handling, the device can be exposed to temperature excursions between -4°F and 140°F (-20°C and 60°C) that lasts up to 24 hours. Storage outside of this range may result in device reset.

**Temperature Equilibration.** After cold storage, allow the device to reach room temperature before charging the capacitors, programming, or implanting the device because cold temperature may affect initial device function.

## Follow-up Testing

Ensure that an external defibrillator and medical personnel skilled in cardiopulmonary resuscitation (CPR) are present during post-implant device testing should the patient require external rescue.

Be aware that the changes in the patient's condition, drug regimen, and other factors may change the defibrillation threshold (DFT), which may result in nonconversion of the arrhythmia. Successful conversion of ventricular fibrillation or ventricular tachycardia during arrhythmia conversion testing is no assurance that conversion will occur post-operatively.

## **Implantation and Device Programming**

Do not position a magnet over the device as that suspends detection and treatment (unless the device has been programmed to ignore the magnet).

Replace the device when the battery reaches the elective replacement indicator (ERI).

1. If it has not already been done, prepare a pocket for the pulse generator.

WARNING: To avoid any risk of accidental shock, make sure that tachyarrhythmia therapies are off before handling the pulse generator. Do not program the pulse generator on until it is inserted in the pocket.

WARNING: For reliable data transmission, implant the pulse generator at a depth not to exceed 5 cm. For patient comfort, do not implant the pulse generator within 1.25 cm of bone unless you cannot avoid it.

2. Insert the lead pins into their receptacles, past the setscrew opening.

If necessary, use sterile lubricant on the insulated shoulder of the lead connectors.

**Properly inserted, the plug heads protrude only a few millimeters from the header.** Do not use forceps or other tools to insert the plug as these can damage its silicone insulation.

- When connecting leads to the pulse generator, make sure that you plug the correct lead into the
  correct lead receptacle. For sensing and pacing, this is important to ensure that atrial and
  ventricular signals are correctly recorded and that pacing pulses are delivered in the desired
  chamber.
- Use and fasten the appropriate lead receptacle plug in an unused lead receptacle. Refer to Spare Parts and Accessories for a list of available lead receptacle plugs.

WARNING: If you are using a single defibrillation lead with only one defibrillation coil, make sure that the lead is in the receptacle for the RV (DF-1) lead. Lubricate and insert the DF-1 plug into the receptacle for the SVC (DF-1) lead. If the lead is not in the RV receptacle, the can and the lead will have the same polarity and there will be no current flow.

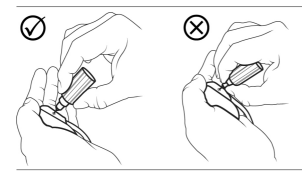
When the DF4-LLHH lead receptacle is plugged, disable tachyarrhythmia therapy.

- For IS4/DF4 leads and lead receptacles, do not use silicone oil, mineral oil, or any substance other than sterile saline, water, or heparinized saline as a lubricant.
- For IS-1 and DF-1 leads and lead receptacles the use of a lubricant is optional.
- For dual-chamber and CRT-D devices, if you are not using an atrial sense/pace lead, lubricate and insert an IS-1 receptacle plug into the receptacle for the atrial sense/pace lead.
- For CRT-D devices, if you are not using a left ventricular pacing lead, lubricate and insert an IS-1 plug into the receptacle for the LV lead.

Carefully insert the tip of the torque driver into the setscrew and turn the handle clockwise until you hear at least three clicks. Setscrews are installed in the pulse generator at the time it is shipped.

CAUTION: Exercise caution when turning the setscrew, which may be backed out of the connector if turned counterclockwise for more than two rotations.

Grip the torque wrench by the large part of the handle as shown in the figure on the left below. On applicable wrench models, do not grip the torque wrench by the smaller, narrower part of the handle as shown on the right.



Turn the wrench clockwise until it clicks. The wrench is torque-limited and when used correctly will not allow excessive tightening.

Coil any excess lead length underneath the pulse generator in the implant pocket.

Program the device parameters as specified in the Merlin™ PCS on-screen help.

The results of the DAVID Study demonstrated that, for patients with standard indications for ICD therapy, no indication for cardiac pacing and an EF < 40%, dual-chamber pacing offers no clinical advantage over backup

VVI pacing and may be associated with worsening heart failure. ¹ When programming the device to dual-chamber pacing modes, give particular attention to setting the pacing parameters (such as the A-V delay) to promote intrinsic conduction and minimize the amount of ventricular pacing.

## **Patient Registration Form**

Fill out and return the Patient Registration Form to register the patient and facilitate patient tracking.

#### **Patient Identification Card**

Fill out the Patient Identification Card as provided in the outer box with the patient's name, implant date, and your name or healthcare facility information along with the contact phone number. Record device information (model number, serial number, and UDI number), and implanted leads information (model and length), or apply the labels provided with the sterile package, in the designated section of the card. Fill out the location of the device and leads in the corresponding location fields on the Patient Identification Card. Give the completed card to the patient. This will serve as a temporary card until Abbott Medical mails a permanent card directly to the patient. To obtain a replacement card if a patient loses or damages their card, contact Abbott Medical Technical Support.

<sup>&</sup>lt;sup>1</sup> Wilkoff BL, Cook JR, Epstein AE, Greene L, Hallstrom AP, Hsia H, Kutalek SP, Sharma A. Dual-Chamber Pacing or Ventricular Backup Pacing in Patients With an Implantable Defibrillator: The Dual Chamber and VVI Implantable Defibrillator (DAVID) Trial. JAMA. December 25, 2002; Vol 288, No. 24:3115-3123.

## **Pulse Generator Explant and Disposal**

Interrogate the device and turn all therapies off before explanting, cleaning or shipping the device to prevent unwanted shocks.

Return all explanted pulse generators and leads to Abbott Medical.

Never incinerate the device because of the potential for explosion. Explant the device before cremation.

## **Environmental and Medical Therapy Hazards**

Instruct patients to avoid devices which generate a strong electric or magnetic interference (EMI). EMI could cause device malfunction or damage, resulting in non-detection or delivery of unneeded therapy. Moving away from the source or turning it off will usually allow the pulse generator to return to its normal mode of operation.

## **Hospital and Medical Environments**

**Electrosurgical cautery.** Electrosurgical cautery could induce ventricular arrhythmias and/or fibrillation, or may cause device malfunction or damage. If electrocautery is necessary, keep the current path and groundplate as far away from the pulse generator and leads as possible.

**External defibrillation.** External defibrillation may damage the pulse generator or may result in temporary and/or permanent myocardial damage at the electrode-tissue interface as well as temporarily or permanently elevated pacing capture thresholds. Minimize current flowing through the pulse generator and

lead system by following these precautions when using external defibrillation on a patient with a pulse generator:

- Position defibrillation paddles as far from the pulse generator as possible (minimum of 13 cm)
- Use the lowest clinically appropriate energy output
- Confirm pulse generator function following any external defibrillation

**High radiation sources.** Do not direct high radiation sources such as cobalt 60 or gamma radiation at the pulse generator. If a patient requires radiation therapy in the vicinity of the pulse generator, place lead shielding over the device to prevent radiation damage and confirm its function after treatment.

**Lithotripsy.** Lithotripsy may permanently damage the pulse generator. Avoid it unless the therapy site is not near the pulse generator and leads.

**Diathermy.** Avoid diathermy, even if the device is programmed off, as it may damage tissue around the implanted electrodes or may permanently damage the pulse generator.

**Ultrasound therapy.** The device should not be exposed to therapeutic levels of ultrasound energy as the device can inadvertently concentrate the ultrasound field and cause harm that might not be immediately detectable. Diagnostic ultrasound treatment is not known to affect the function of the device.

**Transcutaneous Electrical Nerve Stimulation (TENS).** TENS may interfere with device function. To reduce interference, place the TENS electrodes close to one another and as far from the device/lead system as possible. Monitor cardiac activity during TENS use.

**Radiofrequency ablation.** RF ablation in a patient with a pulse generator may cause device malfunction or damage.

Minimize RF ablation risks by:

- Programming all tachyarrhythmia therapies off
- Avoiding direct contact between the ablation catheter and the implanted lead or pulse generator
- Positioning the groundplate so that the current pathway does not pass near the pulse generator system,
   i.e., place the groundplate under the patient's buttocks or legs
- Having external defibrillation equipment available

**RF Operating Frequencies.** Nearby equipment emitting strong magnetic fields can interfere with RF communication, even if the other equipment complies with CISPR emission requirements. The operating characteristics are as follows:

MICS band: 402-405 MHz. The effective radiated power is below the limits as specified in:

• Europe: EN ETSI 301 839

USA: FCC 47 CFR Part 95; 95.601-95.673 Subpart E, 95.1201-95.1221 Subpart I

FCC ID: RIASJMRF

WARNING: 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 (that is, 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.

The following is applicable to Canada only:

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.

## **Home and Occupational Environments**

**High-voltage power transmission lines.** High-voltage power transmission lines may generate enough EMI to interfere with pulse generator operation if approached too closely.

**Communication equipment.** Communication equipment such as microwave transmitters or high-power amateur transmitters may generate enough EMI to interfere with pulse generator operation if approached too closely.

**Home appliances.** Home appliances in good working order and properly grounded do not usually produce enough EMI to interfere with pulse generator operation. There are reports of pulse generator disturbances caused by electric hand tools or electric razors used directly over the pulse generator implant site.

**Industrial equipment.** A variety of industrial equipment produce EMI of sufficient field strength and modulation characteristics to interfere with proper operation of the pulse generator. These include, but are

not limited to: arc welders; induction furnaces; very large or defective electric motors; and internal combustion engines with poorly shielded ignition systems.

# Electronic Article Surveillance, Security and Logistical Systems, and Metal Detectors

Advise patients that the Electronic Article Surveillance/Anti-theft (EAS), RFID and Metal Detector systems such as those at the point of sale and entrances/exits of stores, libraries, banks, etc., emit signals that may interact with the device. It is very unlikely that these systems will interact with their device significantly. However, to minimize the possibility of interaction, advise patients to simply walk through these areas at a normal pace and avoid lingering near or leaning on these systems. Even so, the ICD and CRT-D systems contain a metal that may set off the airport security system alarm. If the alarm does sound, the patient should present security personnel with their patient identification card. If security personnel conduct a search with a handheld wand, the patient should ask that they perform the search quickly, stressing that they should avoid holding the wand over the device for a prolonged period.

#### **Cellular Phones**

The pulse generator has been tested for compatibility with handheld wireless transmitters in accordance with the requirements of ISO 14117. This testing covered the operating frequencies (450 MHz - 3 GHz) and pulsed modulation techniques of all of the digital cellular phone technologies in worldwide use today. Based

on the results of this testing, the pulse generator should not be affected by the normal operation of cellular phones.

#### **Potential Adverse Events**

Possible adverse events (in alphabetical order) associated with the system, include, but are not limited to the following:

- Acceleration of arrhythmias (caused by device)
- Air embolism
- Allergic reaction
- Bleeding
- Cardiac tamponade
- · Chronic nerve damage
- Death
- Frosion
- Exacerbation of heart failure
- Excessive fibrotic tissue growth
- Extracardiac stimulation (phrenic nerve, diaphragm, chest wall)
- Extrusion
- Fluid accumulation
- Formation of hematomas or cysts

- Inappropriate shocks
- Infection
- Keloid formation
- Lead abrasion and discontinuity
- Lead migration/dislodgment
- Myocardial damage
- Pneumothorax
- Shunting current or insulating myocardium during defibrillation with internal or external paddles
- Potential mortality due to inability to defibrillate or pace
- Thromboemboli
- Venous occlusion
- Venous or cardiac perforation.

Patients susceptible to frequent shocks despite antiarrhythmic medical management may develop psychological intolerance to an ICD or CRT-D system that may include the following:

- Dependency
- Depression
- Fear of premature battery depletion
- Fear of shocking while conscious
- · Fear of losing shock capability
- Imagined shocking (phantom shock).

#### **Clinician Use Information**

#### WARNING:

- For devices without the Low Frequency Attenuation Filter, the default Atrial Sensitivity setting and
  the lowest possible setting of Ventricular Sensitivity, 0.2 mV, may be more susceptible to EMI,
  according to testing required by CENELEC standard EN45502-2-2. The devices comply with the
  electromagnetic compatibility requirements of CENELEC standard EN45502-2-2 at atrial and
  ventricular sensitivities of 0.3 mV and less sensitive settings.
- For devices with the Low Frequency Attenuation Filter, the default Atrial Sensitivity setting, the lowest possible setting of Ventricular Sensitivity, 0.2 mV, and the Ventricular Sensitivity setting of 0.3 mV when the Low Frequency Attenuation Filter is On, may be more susceptible to EMI, according to testing required by CENELEC standard EN45502-2-2. The devices comply with the electromagnetic compatibility requirements of CENELEC standard EN45502-2-2 at atrial sensitivities of 0.3 mV, ventricular sensitivities of 0.3 mV (Low Frequency Attenuation Filter OFF) or ventricular sensitivities of 0.4 mV (Low Frequency Attenuation Filter On), and less sensitive settings.

## **Physician Training**

Physicians should be familiar with sterile pulse generator implant procedure and with follow-up evaluation and management of patients with an ICD or CRT-D (or should refer the patient to such a physician).

## **Maintaining Device Effectiveness**

### **Device Storage**

**FOR SINGLE USE ONLY.** Do not resterilize and re-implant explanted pulse generators.

Abbott Medical has sterilized the pulse generator with ethylene oxide prior to shipment. Contact Abbott Medical if resterilization is necessary.

Do not implant the device when:

- It has been dropped on a hard surface because this could have damaged pulse generator components.
- The sterility indicator within the inner package is purple, because it might not have been sterilized.
- Its storage package has been pierced or altered, because this could have rendered it non-sterile.
- It has been stored or transported outside the environmental temperature limits.
  - An electrical reset condition may occur at temperatures below -20°C.
  - After cold storage, allow the device to reach room temperature before charging the capacitors, programming, or implanting the device because cold temperature may affect initial device function.
- Its "use before" date has expired, because this can adversely affect pulse generator longevity or device sterility.

Do not resterilize the pulse generator using an autoclave, gamma-irradiation, organic cleaning agents (e.g., alcohol, acetone, etc.), or ultrasonic cleaners.

#### **Sterilization Instructions**

Contact Abbott Medical if resterilization is necessary.

#### **Directions for Use**

Pulse generator operating characteristics should be verified at the time of implantation and recorded in the patient file. Complete the Patient Registration Form and return it to Abbott Medical as it provides necessary information for warranty purposes and patient tracking.

Copies of this user's manual can be obtained by contacting your Abbott Medical representative.

## Radiopaque Identification

Each pulse generator has an X-ray absorptive marker for non-invasive identification.

Table 5. X-ray ID codes for the device models described in this manual

Device Model	X-ray ID Model Code
CD1411-36C/36Q, CD2411-36C/36Q	SJM KF

Table 5. X-ray ID codes for the device models described in this manual

#### **Device Model**

#### X-ray ID Model Code

CD1357-40C/40Q, CD2357-40C/ SJM KC 40Q, CD3357-40C/40Q, CD3365-40C/40Q, CD3369-40C/40Q

## **Package Contents**

The pulse generator is supplied in a sterile tray for introduction into the operating field. The tray contains:

- One pulse generator (with all tachyarrhythmia therapies off) with pre-installed setscrews
- Torque driver.

The outer box contains:

Literature.

## **Technical Support**

Abbott Medical maintains 24-hour phone lines for technical questions and support:

1 818 362 6822

- 1 800 722 3774 (toll-free within North America)
- + 46 8 474 4147 (Sweden)
- + 61 2 9936 1200 (Australia)
- medical.abbott/manuals

For additional assistance, call your local Abbott Medical representative.

## **Additional Information**

For additional information on this device, see the programmer's on-screen help.

## **Symbols**

The symbols below and harmonized symbols may be found on the product or product label. For harmonized symbols, refer to the Universal Symbols Glossary at https://medical.abbott/manuals.

Symbol	Description
VVED - DDDR Dual-chamber ICDs	NBD - NBG Code; NBD - ventricular shocking, ventricular antitachycardia pacing, electrogram detection, dual-chamber bradycardia pacing; NBG - dual-chamber pacing, dual-chamber sensing, dual response, ratemodulated

Symbol	Description
VVEV - VVIR Single-chamber ICDs	NBD - NBG Code; NBD - ventricular shocking, ventricular antitachycardia pacing, electrogram detection, ventricular bradycardia pacing; NBG - ventricular pacing, ventricular sensing, inhibited response, rate-modulated
VVED - DDDRV CRT-Ds	NBD - NBG Code; NBD - ventricular shocking, ventricular antitachycardia pacing, electrogram detection, dual-chamber bradycardia pacing; NBG - dual-chamber pacing, dual-chamber sensing, dual response, ratemodulated, biventricular pacing
LLHH	Quadripolar connector (low voltage, low voltage, high voltage, high voltage)
LLLL	Quadripolar connector (low voltage, low voltage, low voltage, low voltage)
4	Shipped settings. The pulse generator is shipped with all functions off
	Australian Communications and Media Authority (ACMA) and New Zealand Radio Spectrum Management (RSM) Regulatory Compliance Mark (RCM)
	This equipment is certified for type certification pursuant of Article 38-24 of the Japan Radio Law

Symbol	Description
	Korea Certification mark for electrical devices
	This device complies with part 15 of the FCC Rules.
FC RIASJMRF	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.
IC: 7067A-SJMRF	Industry Canada certification
+	Accessories
	Product literature
<b>3</b>	Manufacturing facility

Symbol	Description
	Importer
Made in USA	Made in USA
Made in Malaysia	Made in Malaysia
UDI	Unique device identification number
MD	Medical Device
medical.abbott/manuals	Follow instructions for use on this website
三台	Patient identification card label
<b>†</b> ?	Patient identification
31	Date

Symbol	Description
W,	Healthcare center or physician
•	Physician telephone
	Lead model/length
0	Location of implant



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