

Current Accel™ VR ICD

Implantable Cardioverter Defibrillators (ICDs)
with InvisiLink™ Wireless Telemetry

MODELS CD1215-30 AND CD1215-36



SPECIFICATIONS

- AutoCapture™ Pacing System offers the maximum in threshold adaptability and patient safety with ventricular Beat-by-Beat™ capture confirmation. The AutoCapture™ Pacing System automatically delivers a 5.0 V backup safety pulse when noncapture is detected, and it may be programmed to either a bipolar or unipolar configuration.
- Exclusive DeFT Response™ technology tools provide more clinically proven, non-invasive options for managing high DFTs.
 - Programmable pulse widths allow the user to tailor the shock to the individual patient, making shocks more efficacious.¹
 - SVC shocking electrode can be quickly and non-invasively activated or deactivated with the press of a button.
 - 36 J delivered energy (model CD1215-36) provides unsurpassed energy for defibrillation.
 - Four programmable tilt options are available to accommodate variances among patients.²
 - Together, these features may help to prevent additional surgeries.
- Exclusive SenseAbility™ feature, with Decay Delay and Threshold Start, provides the flexibility to fine-tune sensing to individual patient needs and help eliminate oversensing of T waves, fractionated QRS complexes, and other extraneous signals.
- Exclusive Morphology Discrimination plus AV Rate Branch SVT discrimination feature helps reduce the risk of inappropriate ICD shocks and is intended to promote fast, accurate diagnosis and delivery of therapy. Clinical data states that this combination resulted in a sensitivity of 100% with a specificity of 85%.³
- Exclusive DC Fiber™ induction has a documented 95,5% success rate for inducing fibrillation on the first induction as compared with a 72,7% success rate for Shock-on-T.⁴
- Exercise Trend Diagnostic provides insight into the patient's disease state progression and exercise activity.
- Up to 45 Minutes of Continuous, Fully Annotated Stored Electrograms, including up to 60 seconds of pre-trigger information per electrogram.
 - Preferential EGM storage capability allows prioritisation of episode storage.
- Exclusive vibratory Patient Notifier allows even patients with hearing problems to be alerted to a low battery, lead-related complications and more.
- Automatic Daily High-Voltage (HV) Lead Integrity Test is designed to automatically test the HV lead on a daily basis to ensure therapy delivery for optimal patient safety.

- Multiple hardware and software system safeguards for added security and patient comfort.
- The capability to program multiple ATP schemes per zone has the potential to increase the success of ATP prior to requiring a shock.
- InvisiLink™ Wireless Telemetry, in conjunction with the Merlin@home™ transmitter and Merlin.net™ PCN, allows for seamless remote monitoring and follow-up. InvisiLink™ RF telemetry uses a dedicated range of frequencies designated for medical devices called the MICS (Medical Implant Communications Service) frequency band, which helps reduce the interference seen on frequencies used by common household electronics.

Indications: The devices are intended to provide ventricular antitachycardia pacing and ventricular defibrillation for automated treatment of life-threatening ventricular arrhythmias.

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:

Implantation Procedure. The physician should be familiar with all components of the system and the material in this manual before beginning the procedure. Ensure that a separate standby external defibrillator is immediately available. Implant the pulse generator no deeper than 5 cm to ensure reliable data transmission. For patient comfort, do not implant the pulse generator within 1,25 cm of bone unless you cannot avoid it.

Device Replacement. Replace the pulse generator within three months of reaching the 2.45 V indication. Replace the pulse generator immediately upon reaching 2.45 V if there is frequent high-voltage charging and/or one or more of the pacing outputs are programmed above 2.5 V.

Battery Incineration. Do not incinerate pulse generators as they contain sealed chemical power cells and capacitors that may explode. Return explanted devices to St. Jude Medical.

High-Voltage Can. Ensure that tachyarrhythmia therapy is programmed Off before handling the pulse generator to avoid any risk of accidental shock. Do not program tachyarrhythmia therapies On until the pulse generator is inserted in the pocket. For effective defibrillation, perform all defibrillation testing with the can in the pocket.

Magnetic Resonance Imaging (MRI). Avoid MRI devices because of the magnitude of the magnetic fields and the strength of the radiofrequency (RF) fields they produce.

Device Storage. Store the pulse generator at temperatures between 10° and 45°C. Do not subject it to temperatures below -20° or over 60°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.

Device Communication. Communication with the device can be affected by electrical interference and strong magnetic fields. If this is a problem, turn off nearby electrical equipment or move it away from the patient and the programmer. If the problem persists, contact St. Jude Medical.

Lead Impedance. Do not implant the pulse generator if the acute defibrillation lead impedance is less than 20 ohms or the lead impedance of chronic leads is less than 15 ohms. Damage to the device may result if high-voltage therapy is delivered into an impedance less than 15 ohms.

Suboptimal RF Communication. The Merlin™ PCS indicates the quality of the RF communication by the telemetry strength indicator LEDs on both the programmer and the Merlin Antenna.

Disconnecting Leads. Connecting or disconnecting sense/pacer leads can produce electrical artifacts that can be sensed by the pulse generator. To prevent detection of artifacts, reprogram the pulse generator to tachyarrhythmia therapy Off. Before disconnecting the leads from a pulse generator in the operating room; Before a post-mortem examination; Whenever there are no leads connected to it; When sense/pacer leads are connected but are not implanted in a patient. If a programmer is not available, use a magnet to prevent delivery of tachyarrhythmia therapy in response to detected disconnection artifacts. Place the magnet over the pulse generator before disconnecting the leads. Do not remove it until the leads are reconnected.

External Equipment for Arrhythmia Induction. If external equipment is used for arrhythmia induction through the pulse generator header and leads, apply rectified AC current through the high-voltage ports, not the sense/pacer ports, to avoid damaging the sense/pacer function; disconnect the external equipment from the pulse generator before any therapy is delivered; otherwise, damage to the device is likely to occur. Place a magnet over the device until the external equipment can be disconnected.

Adverse Events:

Implantation of the pulse generator system, like that of any other device, involves risks, some possibly life-threatening. These include but are not limited to the following: acute hemorrhage/bleeding, air emboli, arrhythmia acceleration, cardiac or venous perforation, cardiogenic shock, cyst formation, erosion, exacerbation of heart failure, extrusion, fibrotic tissue growth, fluid accumulation, hematoma formation, histotoxic reactions, infection, keloid formation, myocardial irritability, nerve damage, pneumothorax, thromboemboli, venous occlusion. Other possible adverse effects include mortality due to: component failure, device-programmer communication failure, lead abrasion, lead dislodgment or poor lead placement, lead fracture, inability to defibrillate, inhibited therapy for a ventricular tachycardia, interruption of function due to electrical or magnetic interference, shunting of energy from defibrillation paddles, system failure due to ionising radiation. Other possible adverse effects include mortality due to inappropriate delivery of therapy caused by: multiple counting of cardiac events including T-waves, P-waves, or supplemental pacemaker stimuli. Among the psychological effects of device implantation are imagined pulsing, dependency, fear of inappropriate pulsing, and fear of losing pulse capability.

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



ST. JUDE MEDICAL™

MORE CONTROL. LESS RISK.

PHYSICAL SPECIFICATIONS		
Models	CD1215-30	CD1215-36
Telemetry	RF	RF
Delivered Energy	30 J	36 J
Volume (cc)	37	41
Weight (g)	73	79
Size (mm)	75 x 50 x 13	76 x 50 x 14
Defibrillation Lead Connections	DF-1	DF-1
Sense/Pace Lead Connections	IS-1	IS-1
High Voltage Can	Electrically active titanium can	Electrically active titanium can

PARAMETER	Settings
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Sensing/Detection	
SenseAbility™ Technology	Automatic Sensitivity Control adjustment for atrial and ventricular events (Post-Sensed, Atrial) 50; 62.5; 75; 100%; (Post-Paced, Atrial) 0, 2-3, 0 mV; (Post-Sensed, Ventricular) 50; 62.5; 75; 100%; (Post-Paced, Ventricular) Auto; 0, 2-3, 0 mV (Post-Sense/Post-Pace, Atrial/Ventricular) 0-220
Threshold Start	VT-1, VT-2, VF
Decay Delay	(Post-Sense/Post-Pace, Atrial/Ventricular) 0-220
Ventricular Sense Refractory (ms)	125, 157
Detection Zones	VT-1, VT-2, VF
SVT Discriminators	Sudden Onset, Interval Stability, Morphology Discrimination (MD) with Manual or Automatic Template Update
Reconfirmation	Continuous sensing during charging

Antitachycardia Pacing Therapy	
ATP Configurations	Ramp, Burst, Scan; 1 or 2 schemes per zone
Burst Cycle Length	Adaptive, Readaptive or Fixed
Min. Burst Cycle Length (ms)	150-400 in increments of 5
Number of Bursts	1-15
Number of Stimuli	2-20
Add Stimuli per Burst	On, Off
ATP Pulse Amplitude (V)	7.5 Independent from Bradycardia and Post-Therapy Pacing
ATP Pulse Width (ms)	1, 0 or 1.5 Independently programmable from Bradycardia and Post-Therapy Pacing

High Voltage Therapy	
Waveform	Biphasic, Monophasic
RV Polarity	Cathode (-), Anode (+)
Electrode Configuration	RV to Can, RV to SVC/Can

Bradycardia Pacing	
Permanent Modes	VVI(R), Pacer Off
Temporary Modes	Off, VVI, VOO
Rate-Adaptive Sensor	On, Off, Passive
Programmable Rate Parameters	Off, Base Rate (min ⁻¹), Rest Rate (min ⁻¹), Maximum Sensor Rate (min ⁻¹), Pulse Amplitude (RV) (V), Pulse Width (ms), Hysteresis Rate (min ⁻¹), Rate Hysteresis with Search
Ventricular AutoCapture™ Pacing System	On, Off

Post-Therapy Pacing (Independently programmable from Bradycardia and ATP)	
Post-Shock Pacing Mode	Off, VVI
Post-Shock Base Rate (min ⁻¹)	30-100 in increments of 5
Post-Shock Pacing Duration (min)	Off, 0, 5; 1; 2, 5; 5, 7, 5; or 10
Post-Shock Ventricular	
Pacing Amplitude (V)	0, 25-7, 5
Post-Shock Pacing Pulse Width (ms)	0, 5; 0, 1-1, 5

Device Testing/Induction Methods	
DC Fiber™ Pulse Duration (sec)	0, 5-5, 0
Burst Fiber Cycle Length (ms)	20-100
Noninvasive Programmed Stimulation (NIPS)	2-25 stimuli with up to three extrastimuli

Patient Notifiers	
Programmable Notifiers (On, Off)	Device at ERI, Charge Time Limit Reached, Possible HV Circuit Damage, Ventricular Lead Impedance Out of Range
Device Reset	On
Entry into Backup VVI Mode	On
Vibration Duration (sec)	2, 4, 6, 8, 10, 12, 14, 16
Number of Vibrations per Notification	2
Number of Notifications	1-16
Time Between Notifications (hours)	10, 22

Electrograms and Diagnostics	
Stored Electrograms	Up to 45 minutes including up to one minute programmable pre-trigger data per VT/VF diagnosis/detection electrograms; triggers include diagnosis, therapy, PMT termination, PC shock delivery, noise reversion, magnet reversion, and morphology template verification
Therapy Summary	Diagram of therapies delivered
Episodes Summary	Directory listing of up to 60 episodes with access to more details including stored electrograms
Lifetime Diagnostics	History of bradycardia events and device initiated charging
Ventricular HV Lead Impedance Trend	Multi-Vector Trend Data
Histograms	Event Histogram, Ventricular Heart Rate Histogram
Real-Time Measurements (RTM)	Pacing lead impedances, high-voltage lead impedances, unloaded battery voltage, and signal amplitudes

- 1 Mouchawar G, Kroll M, Val-Mejias JE et al. ICD waveform optimization: a randomized prospective, pair-sampled multicenter study. PACE 2000; 23 (Part II):1992-1995.
- 2 Sweeney MO, Natale A, Volosin KJ, et al. Prospective randomized comparison of 50%/50% versus 65%/65% tilt biphasic waveform on defibrillation in humans. PACE 2001; 24:60-65.
- 3 Sperzel J, Meine M et al. A new automatic update function of the morphology template used for SVT/VT discrimination in an ICD. Europace Supplements; Vol. 3, July 2002;A 131, #1515.
- 4 Sharma AD, O'Neill PG, Fain E, et al. Shock on T versus DC for induction of ventricular fibrillation: a randomized prospective comparison. 21st Annual Scientific Session North American Society of Pacing and Electrophysiology (NASPE). Poster presentation published in meeting proceedings. Washington D.C., U.S.A. May 2000.

ATRIAL FIBRILLATION CARDIAC RHYTHM MANAGEMENT CARDIAC SURGERY CARDIOLOGY NEUROMODULATION

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Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country. Unless otherwise noted, ™ indicates that the name is a trademark of, or licensed to, St. Jude Medical or one of its subsidiaries. ST. JUDE MEDICAL, the nine-squares symbol and MORE CONTROL. LESS RISK. are trademarks and service marks of St. Jude Medical, Inc. and its related companies. ©2008 St. Jude Medical, Inc. All Rights Reserved. Item No. GMCRCM376EN