Implantable Cardioverter Defibrillator (ICD) Devices

Fortify[™] ST VR

Single-Chamber Implantable Cardioverter Defibrillator (ICD)

Product Highlights

- The ST monitoring diagnostic provides information on significant ST segment changes for improved insight in decision making
- ShockGuard[™] technology with DecisionTx[™] programming, designed to reduce inappropriate therapy and minimise the need for programming adjustments at implant
- Unique 40 J delivered energy safety shock option can provide a greater DFT safety margin
- The DF4 connector is designed to simplify implants by streamlining defibrillation connections into a single terminal pin and reducing the number of set screws
- QHR[™]* chemistry battery provides greater capacity for enhanced longevity and stable charge times
- Antitachycardia pacing (ATP) while charging and prior to charging in the VF zone further extends the programming options for terminating tachyarrhythmias without a high-voltage shock
- The low frequency attenuation filter is designed to enhance sensing performance and may reduce the possibility of oversensing T waves
- DeFT Response[™] technology offers the most noninvasive options for managing high DFTs
- The SenseAbility[™] feature provides the flexibility to fine-tune programming around T-wave oversensing without decreasing sensitivity
- Vibratory patient notifier enables patients with hearing problems to be alerted to a low battery, lead-related complications and more

Ordering Information

Contents: Cardiac pulse generator

Model Number	Dimensions (H x W x T, mm)	Weight (g)	Volume (cc)	Connector Defibrillation	Connector Sense/Pace
CD1235-40	73 x 40 x 14	76	35	DF1	IS-1
CD1235-40Q	71 x 40 x 14	75	35	DF4	DF4

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.

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, cardiagenic shock, cyst formation, erosion, exacerbation of heart failure, extrusion, fibrotic tissue growth, fluid accumulation, hematoma formation,

nistotoxic reactions, intection, keloid formation, myöcärdiai irritability, herve damäge, pneumothorax, thromboembolii, 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 Twaves, P waves, or supplemental pacemaker stimuli. Among the psychological effects of device implantation are imagined pulsing, dependency, fear of inanoronizate nulsing and itera of losing on use canability. inappropriate pulsing, and fear of losing pulse capability. Refer to the User's Manual for detailed indications, contraindications, warnings, precautions and potential



histotoxic reactions, infection, keloid formation, myocardial irritability, nerve damage, pneumothorax,



Merlin@home Transmitter Compatible

Customer Support: 46-8-474-4756

Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of Biter summary: Prior to using these devices, please review the instructions for Use for a complete listing of indications, contraindications, warnings, procautions, 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 registered and unregistered trademarks and service marks of St. Jude Medical, Inc. and its related companies. ©2011 St. Jude Medical, Inc. All rights reserved.



Fortify[™] ST VR Single-Chamber Implantable Cardioverter Defibrillator (ICD)

Product Specifications

PHYSICAL SPECIFICATIONS				
Models	CD1235-40	CD1235-40Q		
Telemetry	RF	RF		
Delivered/Stored Energy (J)	40/45	40/45		
Volume (cc)	35	35		
Weight (g)	76	75		
Size (mm)	73 x 40 x 14	71 x 40 x 14		
Defibrillation Lead Connections	DF1	DF4		
Sense/Pace Lead Connections	IS-1	DF4		
High-Voltage Can	Electrically active titanium can	Electrically active titanium can		
PARAMETERS	SETTINGS			
Sensing/Detection				
Sense <i>Ability</i> ™ Technology	nology Automatic Sensitivity Control adjustment for atrial and ventricular even			
Low Frequency Attenuation	On; Off			
Threshold Start	(Post-Sensed; Ventricular) 50; 62,5; 75; 100%;			
	(Post-Paced; Ventricular) Auto; 0,	2-3,0 mV		
Decay Delay	(Post-Sense/Post-Pace; Ventricul	ar) 0-220		
Ventricular Sense Refractory (ms)	125; 157	125; 157		
Detection Zones	VT-1; VT-2; VF			
SVT Discriminators	Sudden Onset; Interval Stability; Morphology Discrimination (MD)			
	with Manual or Automatic Templat	e Update		
Reconfirmation	Continuous sensing during charging	ng		
Antitachycardia Pacing Therapy				
ATP Configurations	Ramp; Burst; Scan; 1 or 2 scheme	s per VT zone		
ATP in VF Zone	ATP While Charging; ATP Prior to Charging; Off			
ATP Upper Rate Cutoff	150 - 300 bpm			
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 programm	1,0 or 1,5 Independently programmable from Bradycardia		
	and Post-Therapy Pacing	-		
High-Voltage Therapy				
High-Voltage Output Mode	Fixed Pulse Width; Fixed Tilt			
Waveform	Biphasic; Monophasic			
RV Polarity	Cathode (-); Anode (+)			
Electrode Configuration				
Bradycardia Pacing				
Permanent Modes	VVI(R); Pacer Off			
Temporary Modes	Off; VVI; VOO			
Rate-Adaptive Sensor	(Post-Sense/Post-Pace; Ventricul	ar) 0-220		
Programmable		min ⁻¹); Maximum Sensor Rate (min ⁻¹);		
Rate and Delay Parameters	Pulse Amplitude (RV) (V); Pulse Width (RV) (ms); Hysteresis Rate (min ⁻¹);			
	Data Ilustracia with Carach			

Rate Hysteresis with Search

On; Off

Post-Shock Pacing Mode	Off; VVI 30-100 in increments of 5 Off; 0,5; 1; 2,5; 7,5; or 10				
Post-Shock Base Rate (min-1)					
Post-Shock Pacing Duration (min)					
Device Testing/Induction Methods					
DC Fibber™ Pulse Duration (sec)	0,5-5,0				
Burst Fibber 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; High-Voltage Lead Impedance Out of Range; % V pacing; CorVue Congestion Trigger				
Device Parameter 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; 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 includir				
	stored electrograms				
Lifetime Diagnostics	History of bradycardia events and device-initiated charging				
Ventricular HV Lead Impedance Trend					
Histograms	Event Histogram; Ventricular Heart Rate Histogram;				
	Exercise and Activity Trending				
Real-Time Measurements (RTM)	Pacing lead impedances; high-voltage lead impedances; unloaded batter				
CT Monitoring	voltage; and signal amplitudes ST Histogram Data; ST Deviation Trend; ST Episode Log				
ST Monitoring CorVue™ Congestion Monitoring	ST Histogram Data; ST Deviation Trend; ST Episode Log On; Off				
CorVue Congestion Trigger	8-18 days				

*QHR is a trademark of Greatbatch, LTD.

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Ventricular AutoCapture™

Pacing System

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