Reveal LINQ™
INSERTABLE CARDIAC MONITOR
Enhanced Features and Algorithms Guide

POWERFUL CARDIAC MONITORING
SMALL. SIMPLE. CONNECTED. PRECISE.
Introduction

This Interactive PDF guide is designed to give you the basic technical details on the Reveal LINQ™ Insertable Cardiac Monitor (ICM) enhanced features and algorithms.

At the completion of this guide, you will be able to:
• Identify Reveal LINQ ICM enhanced features and algorithms
• Describe the clinical need that each feature addresses and list the benefits for each feature
• Describe the benefits of the new P-SENSE detection enhancement
The Evolution of Reveal® ICM

From a Diagnostic Device to a Patient Management Tool

1998: Reveal ICM
- 3-year battery
- Added to the Medtronic CareLink® Network
- MR-Conditional

2000: Reveal Plus ICM
- AF detection
- Cardiac Compass®

2007: Reveal DX ICM

2009: Reveal XT ICM

2011: FullView®
- Improvements for viewing and collecting data

Powered by FullView Software
Miniaturized Reveal® ICM Device

Breakthrough Technology

3-year monitoring remote management*

87% smaller and wireless transmissions

* Under the following usage scenarios:
  • Average of 1 auto-detected episode per day
  • Average of 1 patient-activated episode per month
  • Less than or equal to 6 months shelf life (between device manufacture and insertion)

Note: Under maximum shelf storage time (12 months), longevity is reduced by approximately 3 months.
Introducing…
Reveal LINQ™ ICM System

Powerful Cardiac Monitoring

**Powerfully Small**
87% smaller than Reveal® XT ICM, with 20% more data memory

**Powerfully Simple**
Simplified insertion procedure with < 1 cm incision provides the most discreet cardiac monitoring option, with improved cosmetic appearance for greater patient acceptance

**Powerfully Connected**
Only wireless insertable cardiac monitoring system that continuously collects and trends data for up to 3 years, with automatic CareAlert® Notifications

**Powerfully Precise**
Clinically actionable, easy-to-read CareLink® reports reduce the data management burden
Feature Overview
A miniaturized device that is powered by Reveal® performance

New Features
• 1.2 cc
• 59 min ECG storage
• Wireless one-way telemetry
• Titanium nitride coated electrodes to improve sensing
• Enhanced atrial arrhythmia detection
• Nominals customized by patient type
• Incision and insertion tools for a minimally invasive insertion
• Medtronic CareAlert® notifications

Leveraging Reveal XT ICM Capabilities
• 3-year longevity*
• MR-Conditional
• Daily trended diagnostics via Cardiac Compass®

* Under the following usage scenarios:
  • Average of 1 auto-detected episode per day
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Note: Under maximum shelf storage time (12 months), longevity is reduced by approximately 3 months.
Reveal LINQ™ ICM Provides More Customized Solutions

ECG Data Storage: 59 Minutes Total

Clinical Goal

Increased patient-activated ECG memory options to provide additional time, where needed, for patients to use their Patient Assistant to help establish a symptom-rhythm correlation.

Up to 30 minutes of patient-activated episodes.

4 episodes @ 7.5 min. each
3 episodes @ 10 min. each
2 episodes @ 15 min. each

27 minutes of automatically detected episodes

Episode types: Pause, Brady, Tachy

• Up to 27 episodes: 30 seconds of ECG data recorded before detection and up to 27 seconds prior to the end of the episode

Atrial episodes: AT/AF

• Two minutes of ECG data recorded before detection

Two minutes of longest AF episode stored since last interrogation in addition to the 27 minutes of automatically detected episodes.
Symptomatic Episode Duration

NOTE: Stored symptomatic events will be cleared in reprogramming Symptomatic Episode Duration.

All patient and clinical data are fictitious and for demonstration purposes only.
Clinical Data

**Value of Increased and Flexible Patient-Activated Event Memory Storage**

Clinical Data

Value of Increased and Flexible Patient-Activated Event Memory Storage

Clinical Data

Value of Increased and Flexible Patient-Activated Event Memory Storage

Clinical Data

Customized Solutions

Patient-Activated Episodes Marked with “S” on Cardiac Compass®
To help correlate symptomatic events with other clinical data.

![Cardiac Compass Report](image)
Clinical Goal

Arrhythmia detection parameters can be set up automatically when patient’s Date of Birth and clinician’s Reasons for Monitoring are entered on the programmer during device setup.

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<thead>
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Simplified Setup

Removed the following Programming options:
- ECG Recording ON/OFF
- VT Stability and Onset programming
- FVT Interval (Rate), FVT Duration (non-programmable)*

*NOTE: FVT zone fixed at 260 ms with NID 30/40

Renamed:
- Asystole to “Pause”
- VT/FVT to “Tachy”
## Detection/Termination Criteria

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<tr>
<th>Episode Type</th>
<th>Detection</th>
<th>Termination</th>
</tr>
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<tbody>
<tr>
<td>Pauses</td>
<td>No R-waves for *3 sec</td>
<td>12 R-waves</td>
</tr>
<tr>
<td>Tachy</td>
<td>*16 consecutive beats &gt; programmed rate</td>
<td>8 consecutive beats slower than the detection rate</td>
</tr>
<tr>
<td></td>
<td>FVT: non-Programmable</td>
<td></td>
</tr>
<tr>
<td></td>
<td>30/40 beats &gt; 231 bpm (260 ms)</td>
<td></td>
</tr>
<tr>
<td>Brady</td>
<td>*4 beats &lt; *30 bpm (2,000 ms)</td>
<td>4 beats faster than the detection rate</td>
</tr>
<tr>
<td>AT/AF</td>
<td>Must be &gt; recording threshold</td>
<td>Evaluate R-R intervals every 2 min</td>
</tr>
<tr>
<td></td>
<td>• Evaluate R-R intervals every 2 min</td>
<td></td>
</tr>
<tr>
<td>Symptom</td>
<td>Patient Activation (button press) using their Patient Assistant</td>
<td>1 min post-activation</td>
</tr>
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* Programmable parameters

Click on each one of these three options for more details.
Pause Detection

Clinical Goal

Reveal LINQ™ ICM’s ability to continuously monitor if the patient’s heart rhythm stops and no ventricular events are sensed for a programmable period of time.

All patient and clinical data are fictitious and for demonstration purposes only.
Pause Detection

Detected Pause Due to Diminishing R-Waves: Identification and Rejection

Decreasing and small R-waves may lead to Pause detection.

Low Signal Evidence (LSE) count algorithm will help prevent false positive Pause detection due to decreasing R-waves.

All patient and clinical data are fictitious and for demonstration purposes only.
Detected Pause due to Diminishing R-Waves
Identification and Rejection Details

Reveal LINQ™ ICM’s ability to distinguish between diminishing R-waves and a true asystolic pause.

Algorithm identifies diminishing R-waves before detection:

- “Low Signal Evidence” counter is incremented by sensed R-waves prior to the pause which are < 2X the minimum sensing threshold, and decremented by signals above it
- Pause detection is rejected if the Low Signal Evidence > 0 on the beat before the long pause
- Pause reject-episode marker: \( \frac{A}{D} \)

\[2 \times \text{Sensing floor} = 70 \, \mu\text{V}\]
\[\text{Sensing floor} = 35 \, \mu\text{V}\]

Note: This algorithm is only active if sensing is programmed to 25, 35, or 50 \( \mu\text{V} \)
Tachy Detection

FVT Zone
Non-programmable
Rate: > 231 bpm (260 ms)
Duration: 30/40 beats

All patient and clinical data are fictitious and for demonstration purposes only.
Detection/Termination
Ventricular Episode Storage

Detection: 30 sec
Termination: 27 sec

ECG Suspension: 27 sec

Pauses:
- 30 sec
- 27 sec

At/AF: IMPROVED ARRHYTHMIA DETECTION

TACHY: PAUSE

Medtronic
Reveal LINQ™
INSERTABLE CARDIAC MONITOR

21
Tachy Detection – Noise Rejection Algorithm

Clinical Goal

Reveal LINQ™ ICM’s ability to recognize and ignore noise that may trigger Tachy detection.

• 150 ms blanking only scheme

All patient and clinical data are fictitious and for demonstration purposes only.
Tachy Detection – Noise Rejection Algorithm

Noise Rejection Algorithm

• At the FVT detection point, if at least one R-R is < 220 ms in the last 12 beats then Reveal LINQ™ ICM counts the number of signal deflections in the prior 0.78 seconds (> 20 signal inflections clears the FVT counters)
• Adds episode marker for FVT Rejection (  \[ \text{F-D} \] )

> 20 signal inflections in 0.78 seconds clears FVT and VT counters

NID met AND at least one RR of the last 12 is < 220 ms. Look back and count signal inflections
AF Detection

Clinical Goal

Specificity of AF detection through discrimination of true AF from other irregular rhythms.

Two minute Lorenz Plots of RR intervals. Lorenz plots are a way to graphically represent correlation structures in an RR interval time series.
Overall Goal for AF Detection Enhancements

• Preserve AF burden accuracy
  – Preserve Sensitivity to AF detection

• Reduce episode review burden
  – Detection is optimized for accurate detection, fast and simple follow-up
  – Nominal programming based on patient type
  – Enhanced episode storage scheme
P-SENSE Detection Enhancement

The P-SENSE detection enhancement is programmed through the Ectopy Rejection.

The P-SENSE detection enhancement is programmed through the Ectopy Rejection.
Algorithm Enhancement

P-SENSE

NOTE: This is the current operation of the AF detection algorithm in Reveal® XT with FullView®.
Algorithm Enhancement

P-SENSE

Reveal® ICM ECG → P-Wave Feature Extraction Algorithm → P-Wave Evidence Algorithm → Ectopy Rejection Programmed to Nominal

Nominal

RR Intervals → AF Detection Algorithm → Ectopy Rejection

AT/AF

PAUSE

TACHY

IMPROVED ARRHYTHMIA DETECTION
Algorithm Enhancement

P-SENSE

- RR Intervals
- AF Detection Algorithm
- RR Interval Based Ectopy Rejection Algorithm
- P-Wave Feature Extraction Algorithm
- P-Wave Evidence Algorithm
- Ectopy Rejection Programmed to Aggressive

- Reveal® ICM ECG
- Aggressive
- Ectopy Rejection

Improvement Arrhythmia Detection

PAUSE

TACHY

AT/AF
Algorithm Enhancement

P-SENSE

Reveal® ICM ECG → P-Wave Feature Extraction Algorithm → P-Wave Evidence Algorithm

RR Intervals → AF Detection Algorithm

Ectopy Rejection Programmed to Nominal

Nominal → Ectopy Rejection

Aggressive → Ectopy Rejection Programmed to Off

RR Interval Based Ectopy Rejection Algorithm

Reveal ICM ECG → P-Wave Feature Extraction Algorithm → P-Wave Evidence Algorithm

Ectopy Rejection Programmed to Aggressive
P-SENSE Details

- P-wave feature extraction algorithm includes:
  - Averaging ECG for a set of 4 beats meeting rate and regularity criteria (dependent of ectopy rejection setting)
  - Identifying p-wave, atrial flutter waves, and noise from morphologic features of average ECG
  - Identifying p-wave evidence if there is one p-wave and absence of atrial flutter waves or noise

- P-wave evidence algorithm includes:
  - Accumulation of P-wave evidence over two-minute detection period
  - Reduction of AF evidence, computed from Lorenz plot based algorithm, by P-wave evidence prior to detection

Evidence of "1P between 2R"

P-wave Averaging

RR > 780 ms
RR Interval Based Ectopy Rejection Algorithm

- “Ectopy Rejector” better discriminates true AF episodes from episodes of bigeminy and trigeminy.
- Runs of bigeminy or trigeminy (see ECG below) commonly translate into a density of points in segments 5, 9, 8, and 11 of the Lorenz plot.
- AF will not be detected if the evidence of ectopy is great enough at the end of each two-minute detection window.
AF Detection
Enhanced Episode Storage Scheme

**New:** Longest AF Only storage option (default for syncope patients)

Only AF episodes that meet the minimum duration setting will create an AF log entry.

**Atrial Episodes**
Optimizing AF Detection

Purpose: Reduce burden of episode review

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P-SENSE Performance

First 150 patients from XPECT study; 5,937 hours of Holter monitoring; 52 patients with 752 hours of AF in 303 episodes

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<th>FullView® Nominal</th>
<th>AF Monitoring</th>
<th>AF Diagnosis</th>
<th>Non-AF</th>
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<tr>
<td>Duration Sensitivity</td>
<td>98.0%</td>
<td>97.9%</td>
<td>97.6%</td>
<td>95.9%</td>
</tr>
<tr>
<td>(AF detected)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration Specificity</td>
<td>97.4%</td>
<td>98.8%</td>
<td>99.0%</td>
<td>99.5%</td>
</tr>
<tr>
<td>(non-AF not over-detected)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration PPV</td>
<td>84.4%</td>
<td>92.1%</td>
<td>93.5%</td>
<td>96.3%</td>
</tr>
<tr>
<td>(true AF detected)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Episode Sensitivity</td>
<td>91.8%</td>
<td>91.1%</td>
<td>90.8%</td>
<td>88.1%</td>
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<tr>
<td>(AF episodes detected)</td>
<td></td>
<td></td>
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<tr>
<td>Episode PPV</td>
<td>66.6%</td>
<td>71.6%</td>
<td>73.2%</td>
<td>84.3%</td>
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<tr>
<td>(detected episodes with true AF)</td>
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Improved Arrhythmia Detection

P-SENSE Performance Comparative Analysis

Reveal® XT
AF false positive episode detected based on R-R variability only (FullView® AF Detection algorithm)

Reveal LINQ™ ICM P-SENSE
Enhanced rhythm discrimination in the presence of R-R irregularity

All patient and clinical data are fictitious and for demonstration purposes only.
Programming AT/AF Detection Parameters

AF Only Detection
To program AT/AF detection parameters:
1. Select Parameters
2. Select AT/AF Detection

AT/AF Detection type in AF Only, AF Detection in Balanced Sensitivity, and Ectopy Rejection in Nominal setting will work best for the majority of patients. Medtronic recommends these settings for optimal atrial fibrillation burden detection.* If false positives are noted in AF Only mode (for example: irregular sinus rhythm or sinus with frequent PACs), consider reprogramming detection to a less sensitive value, and Ectopy Rejection to “Aggressive”. If it is desired to increase sensitivity to detecting atrial fibrillation, consider reprogramming detection to a more sensitive value. Only if it is suspected or known that the patient has atrial tachycardia or atrial flutter does Medtronic recommend programming AT/AF Detection type to ATAF for a short duration of time, and after diagnosis of AT reprogram back to AF Only mode.

* Burden is defined as the cumulative time in AT/AF. Time in AT/AF (Quick Look™ screen, Cardiac Compass®, and AT/AF Summary) will report total time of AF episodes when programmed to AF Only. When programmed to AT/AF time in AT/AF is reported as the combined total time of AT and AF episodes.
To discriminate very regular AT rhythms from very regular sinus rhythm, selectable lower rate cutoffs can be added to the “Detect Very Regular AT Rhythms” algorithm.

Consider the following if very regular sinus rhythms are being detected as AT:
- Intrinsic intervals > 900 ms program “On-Rates ≥ 67 bpm” (see above ECG strip and intervals example)
- Intrinsic intervals < 900 ms program “On-Rates ≥ 100 bpm”
- Intrinsic intervals < 600 ms program “Off”

All patient and clinical data are fictitious and for demonstration purposes only.
AF False Positive Detection

Due to ectopy rhythms (i.e., sinus arrhythmia, PACs, PVCs, bigeminy, trigeminy)

If false positive detections due to ectopy rhythms: Consider one, two, or all three options below:

1. If Ectopy Rejection is currently Off – consider programming to “Nominal”; If Ectopy Rejection is currently Nominal – consider programming to “Aggressive”

2. Program AT/AF ECG Recording Threshold to a longer ECG.

3. Consider programming AF Detection to “Less” or “Least” Sensitive.

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Zoom Capability

- Zoom function to analyze stored ECGs at appropriate scale and amplitude
- Calibration pulse to quickly visualize R-wave signal amplitude/quality during stored ECG analysis
- P-Wave Zoom Capability

Note: ECG Gain ranges from ± 0.05mV to ± 2.0mV (with ± 0.5mV the default)
Flexible Printing Options

Clinical Goal

Flexibility in how much of a stored episode is printed from the 2090 Programmer (to alleviate too much, or too little, data being printed).

- Option allows you to print the “Displayed ECG” plus the prior 0, 30, 60 or 120 seconds for stored episodes

Note: This printing function is available for internal strip-chart-recorder, external printer and print to PDF file function.
Medtronic MyCareLink Patient Monitor and the Medtronic CareLink® Network are indicated for use in the transfer of patient data from Medtronic implantable cardiac devices. These products are not a substitute for appropriate medical attention in the event of an emergency. Data availability and alert notifications are subject to Internet connectivity and access, and service availability. The MyCareLink Patient Monitor must be on and in range of the device. Alert notifications are not intended to be used as the sole basis for making decisions about patient medical care.

**Brief Statement**
See the device manual for detailed information regarding the implant procedure, indications, contraindications, warnings, precautions, and potential adverse events.