YOUR APP-BASED DEVICE MANAGEMENT SOLUTION

Reveal LINQ™ Mobile Manager
Supporting the Reveal LINQ™ and LINQ II™ Insertable Cardiac Monitors
The Reveal LINQ Mobile Manager is an innovative, app-based device management system for Reveal LINQ and LINQ II insertable cardiac monitors (ICMs). It enables procedure simplicity and programmer portability for device activation, programming, and device check activities.

EXCLUSIVE APP-BASED TECHNOLOGY

- Guided Workflow Animations
- Mobile Convenience for Programming

SECURE
Dedicated protection of patient health data:
- Data encryption and built-in security
- Authentication and password security
- Remotely disable data flow if tablet is lost or stolen

INTUITIVE
An easy-to-use device management system:
- Connectivity status feedback
- Device package scanning functionality
- System feedback loop for pre-enrollment success

INTEGRATED
A single workflow solution for:
- Device programming
- CareLink™ network pre-enrollment
- Patient education
- Device check with episode adjudication

All patient and clinical data are fictitious and for demonstration purposes only.
VISUALS FOR PATIENT MANAGEMENT

THE REVEAL LINQ MOBILE MANAGER SYSTEM

- Medtronic ICM
- Patient Connector
- Mobile Manager Application
- CareLink Network

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Indications, Safety, and Warnings

If you are located in the United States, please refer to the brief statement below to review applicable indications, safety, and warning information. See the device manual for detailed information regarding the implant procedure, indications, contraindications, warnings, precautions, and potential complications/adverse events. For further information, please call Medtronic at 763-514-4000 and/or consult the Medtronic website at medtronic.com.

If you are located outside the United States, see the device manual for detailed information regarding instructions for use, the implant procedure, indications, contraindications, warnings, precautions, and potential adverse events. If using an MRI SureScan™ device, see the MRI SureScan technical manual before performing an MRI. For further information, contact your local Medtronic representative and/or consult the Medtronic website at medtronic.com.

Consult instructions for use at this website. Manuals can be viewed using a current version of any major internet browser. For best results, use Adobe Acrobat Reader® with the browser.

Important Reminder: This information is intended only for users in markets where Medtronic products and therapies are approved or available for use as indicated within the respective product manuals. Content on specific Medtronic products and therapies is not intended for users in markets that do not have authorization for use.

www.medtronic.com/manuals

Brief Statement

Reveal LINQ™ and LINQ II™ Insertable Cardiac Monitors with Reveal LINQ™ Mobile Manager System

Indications

The Reveal LINQ and LINQ II™ Insertable Cardiac Monitors (ICMs) are insertable automatically-activated and patient-activated monitoring systems that record subcutaneous ECG and are indicated in the following cases:

- patients with clinical syndromes or situations at increased risk of cardiac arrhythmias
- patients who experience transient symptoms such as dizziness, palpitation, syncope, and chest pain that may suggest a cardiac arrhythmia

The device has not been tested specifically for pediatric use.

Reveal LINQ Mobile Manager System

The Reveal LINQ Mobile Manager app is intended for programming and interrogating the Reveal LINQ and LINQ II™ Insertable Cardiac Monitors. The patient connector is intended to be used with Medtronic apps to interrogate, analyze, and/or program implantable Medtronic devices. The patient connector uses Bluetooth® technology to transmit data to the Reveal LINQ Mobile Manager app for further processing. The patient connector is intended for use by healthcare professionals in a clinical or hospital environment.

Contraindications

There are no known contraindications for the insertion of the Reveal LINQ or LINQ II ICM or for the Reveal LINQ Mobile Manager system. However, the patient’s particular medical condition may dictate whether or not a subcutaneous, chronically implanted device can be tolerated.

Warnings and Precautions

Reveal LINQ and LINQ II™ Insertable Cardiac Monitors

Patients with the Reveal LINQ and LINQ II ICMs should avoid sources of diathermy, high sources of radiation, electrocautery, external defibrillation, lithotripsy, therapeutic ultrasound, and radiofrequency ablation to avoid electrical reset of the device, and/or inappropriate sensing as described in the Medical Procedure and EMI Warnings, Precautions, and Guidance manual. MRI scans should be performed only in a specified MR environment under specified conditions as described in the Reveal LINQ or LINQ II MRI Technical Manual.

Reveal LINQ Mobile Manager System

Before inserting the device, verify that the patient connector and tablet are fully charged. The patient connector and tablet may run out of power during the insertion procedure if they are not fully charged. You will not be able to program or interrogate the patient’s device until the patient connector and the tablet have power. Only use the patient connector to communicate with the intended implanted device.

Use of wireless devices — The patient connector incorporates radio-frequency (RF) communications components which may affect other devices and equipment in the medical environment.

Radio-frequency (RF) interference — Portable and mobile RF communications equipment can interfere with the operation of the patient connector. There is no guarantee that it will not receive interference or that any particular transmission from this system will be free from interference.

Security — Maintain adequate physical security of the patient connector to prevent unauthorized use that could lead to harm to patients.

Environmental precautions — To ensure safe and effective operation, use the device with care to avoid damage to the patient connector from environmental factors that may impair its function.

Potential Adverse Events or Potential Complications

Potential adverse events of the Reveal LINQ and LINQ II ICMs include, but are not limited to, device rejection phenomena (including local tissue reaction), device migration, infection, and erosion through the skin. See the device manuals for detailed information regarding the implant procedure, indications/intended use, contraindications, warnings, precautions, and potential complications/adverse events. For further information, please call Medtronic Technical Services at (800) 328-2518 and/or consult Medtronic’s website at www.medtronic.com.

Caution: Federal law (USA) restricts these devices to sale by or on the order of a physician.