



UNLOCK THE POTENTIAL

Real World HeartLogic™ Workflows

Clinic: Cardiovascular Medicine Advanced Heart Failure **Hospital Affiliation**: University of Kansas Medical Center **Location**: Kansas City, KS & surrounding metropolitan area

148 HeartLogic Patients¹

- 9% of HL patients in Alert (active alert-based management)
- o 91% of HL patients out of Alert (standard remote monitoring schedule)
- 88% of HL patients regularly remotely transmitting physiologic data



About Leslie Steinkamp & Dr. Andrew Sauer

Leslie Steinkamp is the Heart Failure Remote Monitoring Clinical Program Coordinator at The University of Kansas Health System. With over 10 years of working with heart failure patients, she has helped develop the Heart Failure Remote Monitoring Clinical Program workflows for the health system.

Dr. Sauer is a cardiologist who specializes in advanced heart failure therapies, including heart transplantation and mechanical circulatory support devices. He is also an Associate Professor and Director of the Center for Advanced Heart Failure and Transplantation at the University of Kansas Medical Center.

HeartLogic Program Overview

"We believe every patient in Kansas and the Kansas City surrounding region should have access to the very best heart failure therapies available anywhere in the world. This is explicitly stated in our program mission. We believe best guideline-based heart failure medical therapies can be increased by leveraging technology that supplements the telemedicine revolution for heart failure therapies. By using such technology, we believe this increases the access the community can have to make our team more readily available. We believe this allows us to get upstream with the longitudinal care continuum, notably intervening for patients before they become acutely or critically ill using the HeartLogic technology. Having more access to our team using telemedicine supplemented by remote monitoring data allows us to reduce disparities in care often exploited by issues related to transportation or copays at the time of clinic visit which often reduces total visits/encounters for more vulnerable populations, including patients living in rural/remote regions and minority or socio-economically challenged groups. Finally, we believe using HeartLogic remote monitoring technology enhances our ability to participate in shared care with referring cardiologists, including our partners in electrophysiology, cardiothoracic surgery, and general cardiology throughout the community."

- Dr. Andrew Sauer

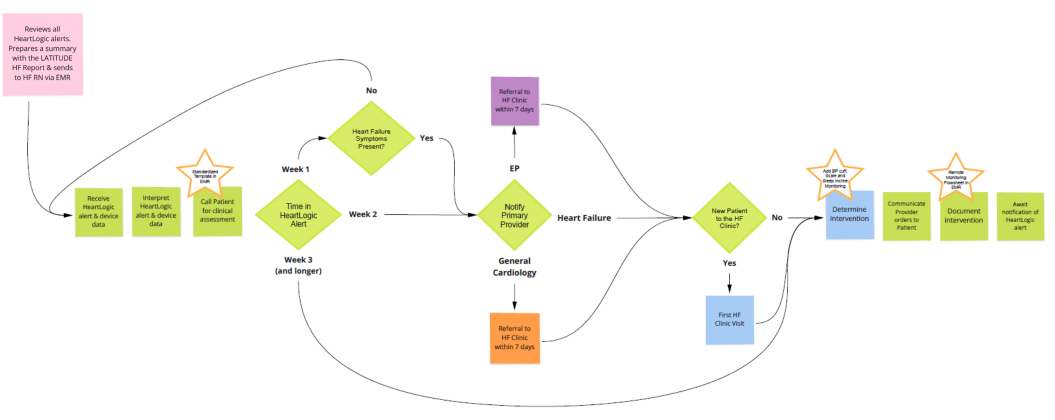
Leslie Steinkamp's Top 3 Pieces of Advice for a Successful HeartLogic Workflow

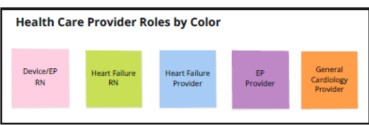
- 1. Co-management is KEY. HF + EP + GC shared care.
- 2. Build Consistency. Consistent documentation. Consistent patient interaction. Consistent clinician communication.
- 3. Demonstrate HeartLogic's capabilities & build a belief network: transitioning from re-active to pro-active intervention.





University of Kansas Health System's HeartLogic Workflow





Workflows, Processes, Proposed Intervention, & Results from HeartLogic Workflows are not necessarily predictive of results in other situations and may vary.

Workflow depicted above documented - November 2020



EDUCARE

UNLOCK THE POTENTIAL



The device/EP RN receives HeartLogic alerts from the LATITUDE NXT Patient Management system. They review the Heartlogic index & data, record the findings in their EMR, and send the heart failure diagnostic information to a designated Heart Failure RN within the Cardiovascular Medicine Advanced Heart Failure clinic.

Heart Failure RN (HF RM Program Coordinator)

The HF RN is specifically designated to receive all HeartLogic alerts & sensor data from the device clinic. They review every HeartLogic alert with a protocol based on the time a patient has been in alert. The HF RN is the primary contact for patients in assessing symptoms and compliance to their plan of care. Per KU's protocol, the HF RN generates an auto-referral for patients with a HeartLogic alert to be evaluated by the advanced Heart Failure clinic.

Heart Failure Provider

The Advanced Heart Failure Provider establishes care to introduce the patient to their team. During the visit, they assess the HeartLogic data and may add additional remote diagnostic tools (eg: Blood Pressure Cuff, Weight, Sleep Incline Sensor) to aid in titrating medication changes remotely. The Heart Failure team monitors and determines the intervention for HeartLogic patients while in alert status.

EP/General Cardiology (Primary) Provider

The EP and General Cardiology teams have a co-management partnership for their HeartLogic patients. Upon one of their patient's receiving a HeartLogic alert, the provider can choose to manage the alert or follow the established automatic referral protocol to the Advanced Heart Failure team.

Objective Data to Guide Therapy

The Cardiovascular Medicine Advanced Heart Failure team continuously seeks to optimize patient outcomes with GDMT (Guideline Directed Medical Therapy). The multiple sensors and trended index of HeartLogic help provide additional physiologic insights for the team to optimize medications to match the needs of each patient.

The 3A Process for University of Kansas Health System

ALERT

HeartLogic alerts are compiled by the KU device clinic and triaged to a designated HF RN within the Advanced Heart Failure program

ASSESSMENT

HeartLogic data in conjunction with a remote patient interview for clinical symptoms is conducted by the HF RN. The assessment is documented via standard EMR templates.

ACTION

The action is determined by the Provider with recommendations from the HF RN's clinical assessment. The HF RN communicates the recommended intervention to the patient. The HF and primary care teams co-manage patients going forward.



CRT-D Systems -RESONATE™HF, RESONATE™, RESONATE™X4, VIGILANT™, VIGILANT™X4, MOMENTUM™, MOMENTUM™ X4

INDICATIONS AND USAGE

These Boston Scientific Cardiac Resynchronization Theracy Delibrillators (CRT-Ds) are indicated for patients with heart failure who receive stable cotimal pharmacologic therapy (OPT) for heart failure and who meet any one of the following classifications: Moderate to severe heart failure (NYHA Class III-IV) with EF < 35% and QRS duration > 120 ms; or left bundle branch block (LBBB) with QRS duration > 130 ms. EF < 30%, and mild (NYHA Class II) ischemic or nonischemic heart failure or asymptomatic (NYHA Class I) ischemic heart failure

CONTRAINDICATIONS

Read this manual thoroughly before implantation to avoid damage to the pulse generator and/or lead. For single patient use only. Do not reuse, reprocess, or resterilize. Always have external delibrillation equipment available during implant and electrophysiologic testing. Ensure that an external delibrillator and medical personnel skilled in CPR are present during post-implant device testing should the patient require external The standard of the standard o inhibition. Physicians should use medical discretion when implantable medical device in patients who present with slow VT. Advise patients to seek medical guidance before entering environments that could adversely affect the operation of the active implantable medical device, unless all of the patient of colenital adverse events applicable when the Conditions of Use are met or not met, refer to the MRI Technical Guide. Do not subject a patient with an implanted pulse generator and/or lead to diathermy. If desired, ensure that Patient Triopered Monitor (PTM) is enabled prior to sending the pagient home by confirming the magnet response is programmed to Store EGM. Once the PTM feature has been frigored and the magnet response set to Inhibit therapy the patient should not reapply the magnet.

PRECAUTIONS

The Specific information on precautions, refer to the following sections of the product labeling: clinical considerations, sterilization and storage, implantation, device programming, environmental and medical therapy hazards, hospital and medical environments, follow up testing, explant and disposal, supplemental precautionary information. Advise patients to avoid sources of EMI because EMI may cause the pulse generator to deliver inappropriate therapy or inhibit appropriate therapy.

POTENTIAL ADVERSE EVENTS

Extracardiac stimulation (muscle/nerve strains) and pulse generator and/or lead implant experience, the following alphabetical list includes the possible adverse evets associated with the included devices: Air embolism; Allergic reaction; Bleeding; Bradycardia; Cardiac tamponade; Chronic nerve damage; Component failure; Conductor coil fracture, Death; Electrolyte imbalance/dehydration; Elevated thresholds; Erosion; Excessive fibrotic tissue growth; Extracardiac stimulation (muscle/nerve stimulation); Failure to convert an induced arrhythmia; Fluid accumulation; Foreign body rejection phenomena; Formation of hematomas or seromas; Heart block; Inability to delibrillate or pace; Inappropriate therapy (e.g., shocks and antitachycardia pain; Incomplete dependent pain; Incomplete pa The factorial role, related in the factorial role of the factorial

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only, Prior to use, please see the complete "Directions for Use" for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator's Instruction

ICD Systems - RESONATE™ HF. RESONATE™ EL. PERCIVA™ HF. PERCIVA™, VIGILANT™ EL. MOMENTUM™ EL ICD

INDICATIONS AND USAGE

Boston Scientific implantable cardioverter defibrillators (ICDs) are intended to provide ventricular antitachycardia pacing (ATP) and ventricular defibrillation for automated treatment of life-threatening ventricular arrhythmias.

CONTRAINDICATIONS

Use of these Boston Scientific pulse generators are contraindicated for the following; patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication (MI), electrocution, drowning; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication, electrocyte imbalance, hypoxia, sepsis; or patients whose ventricular tachyarrhythmias have a transient cause, such as: digitalis intoxication in the contraction of the cont

Read this manual thoroughly before implantation to avoid damage to the pulse generator and/or lead. For single patient use only, Do not reuse, reprocess, or resterlize. Always have external defibrillation equipment available during implant and electrophysiologic testing. Ensure that an external defibrillator and medical personnel skilled in CPR are present during post-implant device testing should the patient require external resource. Dr not use this pulse generator with another pulse generator. Program the pul Connector Tool is not present on the lead. Do not directly contact the lead terminal, other than the terminal with any surgical instruments or electrical connections, such as PSA (alligator) clips, EGC connections, beneats, and clarges, beneats, and clarges, beneats, encoded the prevents of electrical terminal, other than the terminal pin, even when the lead cap is in place. Do not use atrial tracking modes in patients with chronic refract tarchyarrhythmias. Tracking of atrial arrhythmias could result in ventricular tarchyarrhythmias. Advise patients to seek medical guidance before entering environments that could adversely affect the operation of the active implantable medical device, including areas protected by a warning notice that prevents entry by patients who have a pulse generator. RESONATE HF, RESONATE, PERCIVA, VIGILANT and MOMENTUM devices are considered Mis Conditional. For these devices, unless all of the MiRI canning of the patient does not meet Mir Conditional requirements for the implanted system, and significant harm to or depose patients with non-Mir Conditional devices to MiRI scanning. For potential adverse events applicable when the Conditions of Use are met or not meet, refer to the MiRI Technical Guide. Do not subject a patient with an implanted pulse generator and/or lead to diathermy. If desired, ensure that Patient Triggered Monitor is enabled prior to sending the patient home. Once the Patient Triggered Monitor feature has been triggered by the magnet and an EGM has been stored, or after 60 days have elapsed from the day that Store EGM was enabled, the patient should not apply the magnet.

For specific information on precautions, refer to the following sections of the product labeling: clinical considerations, sterilization and storage, implantation, device programming, environments, follow up testing, explant and disposal, supplemental precautionary information.

Based on the literature and on pulse generator and/or lead implant experience, the following alphabetical list includes the possible adverse evets associated with the included devices: Air embolism; Allergic reaction; Bleeding; Bradycardia; Cardiac tamponade; Chronic nerve damage; Component failure; Conductor coil fracture; Death; Elevated thresholds; Erosion; Excessive fibrotic tissue growth; Extracardiac stimulation (muscle/nerveners). based on the literature of components for a literature of the components and in the literature of the components of the (PMT); Pericardial rub, effusion; Pneumothorax; Pulse generator migration; Shunting current during defibrillation with internal or external paddles; Syncope; Tachyarrhythmias and early, recurrent atrial fibrillation; Thrombosis/thromboemboli; Valve damage; Vasovagal response; Venous occlusion; Venous trauma (e.g., perforation, dissection, erosion); Worsening heart failure For a list of potential adverse events associated with MRI scanning, refer to the MRI Technical Guide

Patients may develop psychological intolerance to a pulse generator system and may experience the following: Dependency; Depression; Fear of premature battery depletion; Fear of a device malfunction

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete "Directions for Use" for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator's Instructions

LATITUDE™ NXT Patient Management System

The LATITUDE" NXT Patient Management System is intended for use to remotely communicate with a compatible pulse generator from Boston Scientific CRM and transfer data to a central database. The LATITUDE NXT System provides patient data that can be used as part of the clinical evaluation of the patient

The LATITUDE NXT Patient Management System is contraindicated for use with any implanted device, other than a compatible Boston Scientific implanted devices are compatible with the LATITUDE NXT System. For contraindications for use related to the implanted device, refer to the System Guide for the Boston Scientific implanted device being interrogated

Alerts may appear on the LATITUDE NXT website on a daily basis. Primary notification of alert conditions is through the View Patient List page on the LATITUDE NXT website. The clinician needs to log onto the LATITUDE NXT website in order to receive alerts. Although secondary notification through email and SMS and the secondary notification feature does not eliminate or reduce the messages is available, these reminders are dependent on external systems and may be delayed or not occur. The secondary notification feature does not eliminate or reduce the messages is available, these reminders are dependent on external systems and may be delayed or not occur. The secondary notification feature does not eliminate or reduce the messages is available, these reminders are dependent on external systems and may be delayed or not occur. The secondary notification feature does not eliminate or reduce the messages is available, these reminders are dependent on external systems and may be delayed or not occur. The secondary notification feature does not eliminate or reduce the messages is available, the secondary notification feature does not eliminate or reduce the messages is available, the secondary notification feature does not eliminate or reduce the messages is available.

delayed or not occur at all under various conditions, which include but are not limited to the following: System limitations; the Communicator is not able to connect to the LATITUDE NXT server through the configured phone system; the implanted device and the Communicator among establish and complete a telemetry session; the Communicator is damaged or malfunctions; the patient is not complete a telemetry session; the Communicator is not able to connect to the LATITUDE NXT server through the configured phone system; the implanted device and the Communicator is not using the LATITUDE NXT System as described in the patient manual; if subscribed to the LATITUDE Cellular Data Plan, missing two or more payments discontinues the subscription; the clinic user can identify any patients that are not being monitored as described above by using the Not Monitored filter on the View Patient Lis

ADVERSE EFFECTS:

SYSTEM LIMITATIONS:

The LATITUDE NXT System does not provide continuous real-time monitoring, As a remote monitoring, as a remote monitoring, as a remote monitoring, as a remote monitoring based on clinician. These factors that can hinder, delay, or prevent acquisition and delivery of implanted device, sensor, and patient information as intended by the clinician. These factors include: implanted device clock; patient environment; cellular data service; telephone system; communicator memory capacity; clinic environment; schedule/configuration changes; or data processing

Communicator is unable to interrogate the implanted device or if the Communicator is unable to contact the LATITUDE NXT server to upload data, up to 14 days may elabse before the LATITUDE NXT server detects these conditions and informs the clinic user that monitoring is not occurring. If both of these conditions occur at the same time, this notification could take up to 28 days, implanted device data and alert notification may be

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only Prior to use please see the complete "Directions for Use" for more information on Indications. Contraindications. Warnings. Precautions. Adverse Events, and Operator's Instructions.

Workflows, Processes, Proposed Intervention, & Results from HeartLogic Workflows are not necessarily predictive of results in other situations and may vary. All trademarks are the property of their respective owners

Sources

- Boston Scientific Internal Data. November, 2020.
- https://www.kansashealthsystem.com/care/specialties/heart-care/about-us

