

ANDROID® Setup Guide

MyLATITUDE[™]

PATIENT APP

GETTING STARTED

Below is a list of everything you need to set up the MyLATITUDE Patient App:

- ☐ Your smartphone or tablet.
- ☐ Your Google Play ID and password.
- ☐ Your implanted device model and serial number.

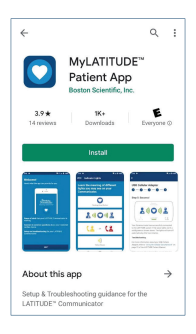
This information can be found on your patient ID card.

The MyLATITUDE™ Patient App is compatible with Android version 4.4 and up.

The requirements for your mobile device or operating system (OS) version will change over time. You may need to update or replace your mobile device and OS to use the app.

STEP 1

From your mobile device, navigate to mylatitudeapp.com. Scroll down and click on the Google Play Store badge to download the application from the Google Play Store.



STEP 2

You will be directed to the App Store.

- Tap "INSTALL" to download the app. The MyLATITUDE Patient App is free to download and use.
- If your mobile device is not compatible, the "INSTALL" button will be disabled.



STEP 3

Once downloaded, the MyLATITUDE Patient App icon will appear on your mobile device screen. Look for this icon and tap it to open the app.

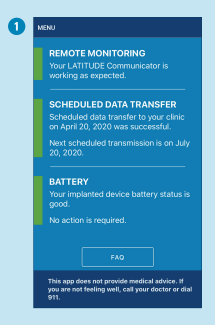
STEP 4

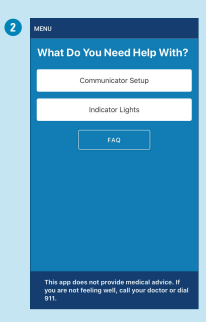
Follow the steps to setup the app:

- Review the Terms of Use and Data Privacy Policy.
- Enter your date of birth, implanted device model number, and implanted device serial number.
- Accept notifications if you would like to be notified if your LATITUDE™ Communicator becomes disconnected.

STEP 5

After completing step 4, you will see one of the below screens.





- 1 If you see this screen or a screen similar to this, you entered your information correctly. No further action is needed to complete the setup of the MyLATITUDE Patient App.
- If you see this screen, you may have entered your information incorrectly. Tap 'MENU', then 'My Information' to review the information you entered. Ensure all of the information you entered is correct by verifying your device information against your Medical Device ID Card. Tap 'MENU', then 'Home' to return to the home screen. If all of your information is correct but you are still seeing this screen, call Boston Scientific Patient Services at 1.866.484.3268.

LATITUDE™ NXT Patient Management System

INTENDED USE: The LATITUDETM 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.

CONTRAINDICATIONS: The LATITUDE NXT Patient Management System is contraindicated for use with any implanted device other than a compatible Boston Scientific implanted device. Not all 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.

PRECAUTIONS: 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 text 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 need to check the website. Implanted device data and alerts are typically available for review on the LATITUDE NXT website within 15 minutes of a successful interrogation. However, data uploads may take significantly longer (up to 14 days). If the 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 elapse 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 delayed or not occur.

at all under various conditions, which include but are not limited to the following: System limitations; the Communicator is unplugged; the Communicator is not able to connect to the LATITUDE NXT server through the configured phone system; the implanted device and the Communicator cannot establish and complete a telemetry session; the Communicator is damaged or malfunctions; the patient is not compliant with prescribed use or 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 List.

ADVERSE EFFECTS: None known

SYSTEM LIMITATIONS: The LATITUDE NXT System does not provide continuous real-time monitoring. As a remote monitoring system, the LATITUDE NXT System provides periodic patient monitoring based on clinician configured settings. There are many internal and external 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. 92436260 (Rev. A)

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.



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Medical Professionals: 1.800.CARDIAC (227.3422) Patients and Families: 1.866.484.3268

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