

TOP 5 THINGS TO KNOW

ABOUT THE NEW FDA PMA REGULATION



What is PMA?

PMA, or premarket approval, is the FDA's regulatory review process, designed to ensure the safety and effectiveness of medical devices. The FDA recently announced that accessories and service for any defibrillator that is not approved under PMA will no longer be available after February 3, 2022.



Which ZOLL defibrillators are not FDA-approved?

The ZOLL M Series® and E Series® monitor/defibrillators are not FDAapproved under PMA. In 2012 and 2015, respectively, ZOLL discontinued these devices, and therefore no PMA application was



Which ZOLL defibrillators are FDA-approved?

- AED Plus®, AED Pro®, ZOLL AED 3® and ZOLL AED 3 BLS
- Powerheart® G3 and Powerheart® G5
- R Series[®]
- X Series[®]





4 How does PMA affect me?

All defibrillators need to be FDA-approved by the above date. Take inventory of your AED and professional defibrillator fleet and check them against the FDA-approved devices list: You can find this list at www.fda.gov/medical-devices/cardiovasculardevices/automated-external-defibrillators-aeds.



Where can I find more information about transitioning to FDA-approved devices?

ZOLL is offering several upgrade programs in order to meet all of our customer needs.

All these devices are FDA approved.



















Contact your SSI Representative for all your Zoll Needs!

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