



Urgent Field Safety Notice

Defibrillators

Return for engineering evaluation

April 2024

Medtronic Reference: FA1416: [EU Manufacturer Single Registration Number \(SRN\): US-MF-000019977](#)

Dear [Account Manager/Healthcare Professional](#),

Medtronic is retrieving the device(s) listed below from your inventory. Your Medtronic representative will collect the device(s) and [arrange for/provide](#) a replacement device(s).

Product Name	Manufacturer's Catalog Number	GTIN	Lot Number(s)
ICD-VR DVMD3D4 PRIMO MRI	DVMD3D4	00763000611965	CWL620432S

Medtronic's internal review processes identified that this device(s) may have undergone a specific manufacturing sequence that requires additional engineering evaluation. No other products in your inventory are being retrieved for this evaluation.

No immediate risk to patients has been identified. In the unlikely event that a patient has received one of the identified device(s), there are no recommended changes to the standard follow-up care protocol.

[Please acknowledge receipt of this letter by signing below.](#) No other action is required.

If you have questions regarding this communication, please contact your Medtronic representative. Thank you for your attention to this matter,

Egidijus Petrauskas Dr.

Medtronic Sr. Field development specialist