

HeartStart FR3. Defibrillator (2018-10-02)

Report a Concern
(<http://www.healthycanadians.gc.ca/report-signalez/index-eng.php>)

Starting date:	October 2, 2018
Posting date:	November 2, 2018
Type of communication:	Medical Device Recall
Subcategory:	Medical Device
Hazard classification:	Type II
Source of recall:	Health Canada
Issue:	Medical Devices
Audience:	General Public, Healthcare Professionals, Hospitals
Identification number:	RA-68188

- Reason
- Affected products

Affected products

HeartStart FR3. Defibrillator

Reason

The recalled AEDs may not meet their IPx5 water ingress performance specification.

Affected products

HeartStart FR3. Defibrillator

Lot or serial number

begin with C16J, C16K, C17A & C17B

Model or catalog number

861388

Companies

Manufacturer

Philips Medical Systems
 22100 Bothell Everett Highway
 Bothell
 98021
 UNITED STATES

Date modified: 2018-11-02