

[Month DD, YYYY]

URGENT MEDICAL DEVICE FIELD SAFETY NOTICE

Reference Number: OT 1210922

Datascope Cardiosave Hybrid Intra-Aortic Balloon Pumps (IABP), CS 100 Intra-Aortic Ballon Pumps (IABP), and CS300 Intra-Aortic Ballon Pumps (IABP) with Maquet-Branded D900 Ultrasonic Doppler and Maquet-Branded D900 Ultrasonic Dopplers sold separately.

Product Name	CARDIOSAVE Hybrid, CS100, and CS300 – Maquet- Branded D900 Ultrasonic Doppler			
Product Code	0154-01-0001, Maquet-Branded D900 Ultrasonic Doppler			
UDI-DI	N/A – Doppler is a Cardiosave Hybrid, CS100, and CS300 Accessory			
Manufacturing Dates for All:	0154-01-0001, Maquet-Branded D900 Ultrasonic Doppler	Manufactured since 12- Dec-2012		
Distribution Dates for All:	0154-01-0001, Maquet-Branded D900 Ultrasonic Doppler	Distributed since12-Dec- 2012		

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Dear Risk Manager,

Datascope Corp., a subsidiary of Getinge, is initiating a voluntary Medical Device Correction for the Maquet-Branded D900 Ultrasonic Doppler supplied with Cardiosave Hybrid Intra-Aortic Balloon Pump (IABP), CS100 IABP, and CS300 IABP devices, and can be purchased separately. Maquet-Branded D900 Ultrasonic Dopplers distributed within affected countries are provided with Instructions for Use (IFU) that are not available in local language translation. This voluntary Correction only affects the countries where IFUs are required in a language other than English. Updated IFUs will be made available in the manner described below when translation is complete.

Identification of the issue:

During an internal review of Datascope's Cardiosave registrations and labeling, Datascope/Getinge identified that the IFU supplied with Maquet-Branded D900 Ultrasonic Dopplers to customers in affected countries is not available in the local language required by regulation in affected countries.

The Ultrasonic Doppler IFU is currently supplied in English only.

Risk To Health:

The IFU not being available in the local language does not pose a risk to health, as it does not impact the user's ability to properly use the device.

Actions to be taken by the user:

Our records indicate that you may have one or more Cardiosave IABPs, CS100 IABPs, and/or CS300 IABPs in your facility that is installed with the Maquet-Branded D900 Ultrasonic Doppler, and/or a separate Maquet-Branded D900 Ultrasonic Doppler.

NO DEVICES NEED TO BE RETURNED.

- Complete and sign the attached Response Form (Page X) to acknowledge that you
 have received and understand this notification. Return the completed form to
 Datascope/Getinge by e-mailing a scanned copy or by faxing the form to your local
 Datascope/Getinge Representative or office.
- Datascope/Getinge expects the necessary IFU translations will be completed by September 30, 2025. At which point we will provide a copy to your facility in your local language.



 The IFU that is currently in your possession can be used until an updated IFU in your local language is provided. If you have any questions or need clarification regarding any IFU content, please contact your local Datascope/ Getinge Sales/Service Unit (SSU) representative to obtain assistance.

SSU add local contact information to this letter>

Please forward this information to all current and potential Cardiosave Hybrid Intra-Aortic Balloon Pump (IABP), CS100 IABP, CS300 IABP, and Maquet-Branded D900 Ultrasonic Doppler users within your hospital/facility.

If you are a distributor who has shipped any affected products to customers, please forward this letter to their attention for appropriate action.

Actions to be taken by Datascope/Getinge:

Datascope/Getinge is initiating this Medical Device Correction to notify Cardiosave Hybrid IABP, CS100 IABP, CS300 IABP, and Maquet-Branded D900 Ultrasonic Doppler Users of this Ultrasonic Doppler IFU translation issue.

Datascope/Getinge is currently working to translate the IFUs to the following local languages required by affected country requirements.

Albanian	Bulgarian	Chinese	Croatian	Czech	Danish
Dutch	Estonian	Finnish	French	German	Greek
Hungarian	Italian	Japanese	Korean	Latvian	Lithuanian
Macedonian	Norwegian	Polish	Portuguese	Romanian	Russian
Serbian	Slovak	Slovenian	Spanish	Swedish	Turkish
Ukrainian					

Once the translation is completed, Datascope/Getinge will provide the IFU to your facility in your local language.



We apologize for any inconvenience this Medical Device Correction may cause	We	apologize	for any	/ incon	venience	this	Medical	Device	Correction	mav	/ cause
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Sincerely,

Ojas Zatakia

Sr. Director, Quality Assurance

Getinge/Datascope Cardiac Assist



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0154-01-0001, Maquet-Branded D900 Ultrasonic Doppler distributed since 12-Dec-2012

ADD ACCOUNT# [FACILITY NAME STREET ADDRESS

CITY, STATE, ZIP CODE]

New Facility Contact Name:

I acknowledge that I have read and understand this Medical Device Correction Letter for the affected Datascope Cardiosave Hybrid Intra-Aortic Balloon Pumps, CS100 Intra-Aortic Balloon Pumps, and/or CS300 Intra-Aortic Balloon Pumps with Maquet-Branded D900 Ultrasonic Doppler and/or a separate Maquet-Branded D900 Ultrasonic Doppler at this facility. I confirm that all users of the above-mentioned products at this facility have been notified accordingly.

Please provide the required information and signature below.

Facility Representative Information:					
Signature:			Date:		
Name:			Phone:		
E-Mail Address:					
Title:			Department:		
Hospital Name:					
Address, City a	nd State <u>:</u>				
We have scrapped	d/discarde	d our affe	cted product:		
Circle one	YES	NO			
We have sold/moved our affected product to another facility:					
Circle one YES NO					
If you answered YES above: please provide new facility information below.					
New Facility Name:					
New Facility Address:					

New Facility Phone #:



Return the completed form by FAX to INSERT LOCAL SSU FAX NUMBER or by EMAIL to INSERT LOCAL SSU EMAIL ADDRESS