29th July, 2020

URGENT MEDICAL DEVICE RECALL – REMOVAL
Datascope/Getinge IAB  Potential Endotoxin Contamination

<table>
<thead>
<tr>
<th>Linear 7.5Fr 25cc IAB</th>
<th>Sensation 7Fr 34cc IAB</th>
<th>MEGA 7.5Fr 30cc IAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear 7.5Fr 40cc IAB</td>
<td>Sensation 7Fr 40cc IAB</td>
<td>MEGA 7.5Fr 40cc IAB</td>
</tr>
<tr>
<td>Linear 7.5Fr 34cc IAB</td>
<td>Sensation Plus 7.5 Fr 40cc IAB</td>
<td>MEGA 8Fr 50cc IAB</td>
</tr>
</tbody>
</table>

KIT Part Numbers and LOT numbers: The IAB Kit lots that were shipped to you and may potentially contain the serial numbers above are listed in the consignee list.

Affected IAB Serial Numbers: Affected serial numbers are listed in the consignee list.

IAB Manufacturing Dates: [February 3, 2017] through [February 21, 2020]

KIT Distribution Dates: [February 9, 2017] through [May 21, 2020]

Risk Manager
Customer name
Street address
City, State, Zip Code

PLEASE FORWARD THIS INFORMATION TO ALL CURRENT AND POTENTIAL DATASCOPE/GETINGE IAB USERS WITHIN YOUR HOSPITAL / FACILITY. IF YOU HAVE FURTHER DISTRIBUTED ANY OF THE AFFECTED PRODUCTS, FORWARD THIS INFORMATION TO THE RECIPIENT.

Dear Risk Manager,

Datascope/Getinge is initiating a voluntary Recall-Removal involving certain Intra-Aortic Balloon Catheters (IABs) that may not meet the requirement for endotoxin per AAMI ST72. Datascope/Getinge performs functional testing on a small number of units from every lot prior to sterilization and these functionally-tested units may pose an elevated risk of endotoxin contamination compared to normal production IABs. These functionally-tested units can be identified by serial number and represent less than 1% of total IABs distributed in this timeframe.

Identification of the issue
The issue is not detectable by the end user. Datascope/Getinge has determined that the affected IABs may not meet the requirement for endotoxin per industry Standard AAMI ST72. All affected IABs can be identified using the Product Labeling and serial number on the Y-fitting of the IAB, quarantined and returned to Datascope/Getinge.

Risk to Health
Physicians would not be able to detect elevated levels of endotoxin before a device is used on a patient. Moreover, since patients in need of this therapy are at higher risk for a systemic inflammatory response including fever, the sole presence of such signs and symptoms usually would not allow
identification of a pyrogenic device as root cause. Several drugs administered during and after procedures or during intensive care stay may even limit the extent of activation of endotoxin-associated cascades. To date, Getinge/Maquet has not received any complaints or adverse events regarding this issue.

**Actions to be taken:**
Our records indicate that you received potentially affected IAB Serial Numbers identified in Table 1. Please complete the steps below:

- Monitor patients for pyrogens reaction/humoral immune response/coagulation and complement cascades/inflammation. Monitor and treat any signs of inflammation according to your facility’s protocols and clinical judgment.

- Please examine your inventory immediately, remove and quarantine any unexpired affected IAB following the steps below:
  - Identify any unexpired IAB Kits referencing the *part and lot numbers* listed in Table 1. The Kit part and lot numbers can be found on the Outer Shelf Carton as circled in RED in the *example* below. REF denotes part number and LOT denotes lot/batch number.
  - Using steps below, Identify any IAB Y-fittings with *serial numbers* listed in Table 1.
    - Remove tape on **one** side of the Shelf Carton.
    - Hinge the box open, keeping the other side taped.
    - Do not remove any items from the carton. Instead, Lift the Accessories up and locate the Y-Fitting through the clear Mylar overwrap.
    - Read the serial number off the IAB Y-Fitting. (See below for example. IAB SN will appear is the area circled in RED.)
• If you find an IAB with a serial number listed in Table 1, immediately place the entire Kit in quarantine for return and replacement or credit.

• If the affected serial number is not found, close the carton and re-seal. Cut out and affix one of the enclosed adhesive labels to designate the kit has been checked for affected recall Serial Numbers and may be placed back in stock. 40 peel-off labels have been provided. Color copies may be made and affixed with tape if more than 40 are needed.

• If you have unexpired affected product to return, please contact Getinge Customer Service at 888- 9GETUSA (888-943–8872) (option 2) between the hours of 9 AM and 6 PM Eastern Standard Time to request a return authorization number (RMA) and shipping instructions. Pack the product to be returned with the appropriate return documents and, using the shipping instructions provided, arrange for pickup with the designated delivery service provider.

• Whether you have affected product or not, please complete and return the form (including all pages) to acknowledge this recall by e-mailing a scanned copy to palash.saxena@getinge.com and Mubashir.javed@getinge.com

• This voluntary recall only affects specific IAB serial numbers manufactured between February 3, 2017 and February 21, 2020. No other products are affected by this voluntary recall.

Adverse reactions or quality problems experienced with the use of this product may be reported to the FDA’s MedWatch Adverse Event Reporting program either online, by regular mail or by fax using the following:

• Online: www.fda.gov/medwatch/report.htm
• Regular mail: Download form www.fda.gov/MedWatch/getforms.htm or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form
• Fax: 1-800-FDA-0178

We sincerely apologize for any inconvenience this recall may cause. If you have any questions, please contact your Maquet/Getinge representative or call the Maquet/Getinge Customer Support at 888- 9GETUSA (888-943–8872) (option 2) between the hours of 9 AM and 6 PM Eastern Standard Time.

This recall is being made with the knowledge of the U.S. Food and Drug Administration.

Sincerely,

Maryanna Krivak

Regulatory Affairs and Quality Compliance Field Actions
USA Shared Services
URGENT MEDICAL DEVICE RECALL – REMOVAL
Datascope/Getinge IAB Endotoxin Issue Recall
Customer Response Form

Linear 7.5Fr 25cc IAB   Sensation 7Fr 34cc IAB   MEGA 7.5Fr 30cc IAB
Linear 7.5Fr 40cc IAB   Sensation 7Fr 40cc IAB   MEGA 7.5Fr 40cc IAB
Linear 7.5Fr 34cc IAB   Sensation Plus 7.5 Fr 40cc IAB   MEGA 8Fr 50cc IAB
Sensation Plus 8Fr 50cc IAB

ACCT#
UST NAME
STREET
CITY, ST 12455

If you have any un-expired affected SERIAL NUMBERS for return and replacement or credit, please complete the table below by circling any serial numbers if found. If you have affected product to return, please contact Getinge Customer Service at 888-9GETUSA (888-943–8872) (option 2) between the hours of 9 AM and 6 PM Eastern Standard Time to request a return authorization number (RMA) and shipping instructions.

Table 1: Sales History of IAB Kits and Lots:

<table>
<thead>
<tr>
<th>Distribution Date</th>
<th>Finished Goods Kit Part Number</th>
<th>Finished Goods Batch</th>
<th>Serial Number</th>
</tr>
</thead>
</table>

ACKNOWLEDGMENT:
By signing below, I acknowledge that I have read and understand this Medical Device Recall Notice for the recalled Maquet/Getinge Intra-Aortic Balloon Catheter Endotoxin recall. I confirm that all IAB users at this facility have been notified accordingly. RMA Number (if applicable): ________________________________

Confirm the Facility Name you are responding for

<table>
<thead>
<tr>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Phone</td>
</tr>
<tr>
<td>Title</td>
<td>Department</td>
</tr>
</tbody>
</table>

Return this form by email to palash.saxena@getinge.com and Mubashir.javed@getinge.com