24 March 2015

**URGENT MEDICAL DEVICE RECALL (Removal)**
**IMMEDIATE ACTION REQUIRED**
LAAx, Inc. TIGERPAW® System II

Product Distributed from 1 April 2013 – 26 February 2015 – ALL LOT NUMBERS

<table>
<thead>
<tr>
<th>Products/Model Numbers:</th>
<th>Model Numbers</th>
<th>Lot Numbers</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7 Connector</td>
<td>9 Connector</td>
</tr>
<tr>
<td>TIGERPAW System II – 7 &amp; 9 connector configurations</td>
<td>C-TP-1507</td>
<td>C-TP-1509</td>
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**PLEASE FORWARD THIS INFORMATION TO YOUR HOSPITAL STAFF WHO MAY USE THE LAAx Inc. TIGERPAW SYSTEM II IN YOUR FACILITY.**

Dear Risk Manager,

This notification is to inform you of an urgent voluntary medical device recall involving the removal of all lot numbers of the LAAx, Inc. TIGERPAW System II, 7 and 9 connector configurations. Maquet Medical Systems, as the distributor of the LAAx, Inc. TIGERPAW System II, is notifying all customers of the recall.

**Reason for Recall:**
The TIGERPAW System II is indicated for the occlusion of the left atrial appendage (LAA), under direct visualization, in conjunction with other open cardiac surgical procedures. The TIGERPAW System II consists of a compliant LAA occlusion fastener and delivery tool.

The Maquet post-marketing vigilance program has identified issues associated with the TIGERPAW System II. There has been an increase in the frequency of field reports involving the following issues which represent a concern:

- Incomplete closure of the TIGERPAW System II fastener that may result in tissue tears and/or bleeding.
- Possible tear on the left atrial wall during the use of the device.

**Potential Patient Impact:**
The left atrial appendage is a very delicate structure and because of its anatomical location, any surgical intervention to correct bleeding could potentially cause serious consequences and, in extreme circumstances, it could be life-threatening.

**There has been no concern identified with the long term safety of implanted Fasteners.**
Action Required:
A review of our records indicates that you have received one or more of the TIGERPAW System II products affected by this recall. Consistent with this voluntary recall we are requiring the return of all TIGERPAW System II products.

Please examine your inventory immediately to determine if you have any TIGERPAW System II products. If so, please remove the TIGERPAW System II products, quarantine them and place in a secure location.

Your Maquet Representative will contact you shortly to assist with the return of any product. You will be issued a credit by Maquet after the returned product is received.

**Because you have received one or more of the TIGERPAW System II products affected by this recall you must acknowledge receipt of this notification by completing and returning the attached Response Form.**

**Note:** If you have acted as a distributor for the TIGERPAW System II product, please immediately contact those accounts, advise them of the recall situation, and have them return all stock to you. Then complete the attached Response Form and return to Maquet.

The voluntary recall has been communicated to the Health Authority in your country.

We appreciate your understanding and thank you for your continued support. We apologize for any inconvenience this may have caused.

If you have any additional questions, please contact your local Maquet office.

Thank you for your cooperation and immediate assistance.

Sincerely,

Karen LeFevere
Director of Regulatory Affairs and Field Action Compliance
Maquet Medical Systems
45 Barbour Pond Drive
Wayne, New Jersey 07470

Attachment: Medical Device Recall Response Form (page 3)
MEDICAL DEVICE RECALL (Removal) RESPONSE FORM
Product Handling Instructions
EMAIL TO: Tigerpaw2015@maquet.com

Product Description: LAAx, Inc. TIGERPAW® System II

Please complete the information below and return this form, acknowledging your receipt and understanding of this communication, by e-mailing a scanned copy to Tigerpaw2015@maquet.com.

If you do not have any TIGERPAW System II product then please check here [ ] sign below and return the form.

If you currently have any TIGERPAW System II product please list all lot numbers and quantity that will be returned, in the table below. If additional space is needed please copy this form as necessary.

<table>
<thead>
<tr>
<th>Model Number</th>
<th>Lot Number</th>
<th>Quantity Returning</th>
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<tr>
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If you have any product, please contact Maquet Customer Service for an RMA number and shipping instructions to return the product. Maquet Customer Service can be reached via email at INTL.cscustserv@maquet.com or via telephone at +1-408-635-0700.

Please enter your RMA number: ________________

By signing this form, I acknowledge that I have reviewed and understand this Medical Device Recall (Removal) Notification and have notified all relevant users in our facility and confirm the product status.

Signature: ____________________________ Date: ____________________________

Print Name: ____________________________

Title and Department: ____________________________

Hospital Name: ____________________________

Hospital Address: ____________________________

Country: ____________________________

Telephone Number: ____________________________