Urgent Field Safety Notice

MiniCap Extended Life PD Transfer Sets
FA-2021-058
Device Correction

January 2022

Dear Sir/Madam,

Problem Description
Baxter Healthcare Corporation is issuing a Device Correction for the MiniCap Extended Life PD transfer sets listed below. The following products may cause damage (for example, leaking or cracking), if they come into direct contact with the transfer set:

- Cleaning products such as hand sanitizer, or those containing, but not limited to, hydrogen peroxide, bleach, alcohol or antiseptic agents
- Solvents intended to remove adhesive residue, such as those containing acetone, toluene, xylene, or cyclohexanone

Baxter will be updating the Instructions for Use (IFU) to include a warning against the use of these cleaning products and solvents.

Affected Product

<table>
<thead>
<tr>
<th>Product Code</th>
<th>Description</th>
<th>Lot #</th>
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<tbody>
<tr>
<td>5C4482</td>
<td>Transfer Set (MiniCap Extended Life PD Transfer Set (6”) with Twist Clamp)</td>
<td>All lots within expiry</td>
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<tr>
<td>R5C4482</td>
<td>MINICAP EXTEND LIFE PD TRANSFER SET</td>
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<tr>
<td>R5C4483</td>
<td>MINICAP EXTD LIFE TRANSFER SET</td>
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<td>R5C4484</td>
<td>MINICAP EXTD LIFE TRANSFER SET</td>
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<td>R5C4482E</td>
<td>MINICAP EXTEND LIFE PD TRANSFER SET</td>
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Hazard Involved
A damaged or leaking transfer set could result in microbial contamination of the sterile fluid path. This may predispose patients to peritonitis. Baxter has received 13 reports of peritonitis possibly related to this issue. Additional hazards that may result include delay in therapy and exposure to bodily fluids.

Actions to be taken by Customers
1. If you are using one of the above-mentioned cleaning products or solvents, please discontinue use of this product or solvent immediately. Please ensure all home patients are aware of proper cleaning practices. Clinicians who are not using any of the above-mentioned cleaning products or solvents may continue to use Baxter transfer sets.
2. If you have patients who have used the cleaning products or solvents listed above and have identified damage to their transfer set, please replace their transfer set and contact Baxter.

3. Baxter will be updating the Instructions for Use (IFU) for all Luer transfer sets to instruct patients not to allow cleaning products or solvents to come into contact with the transfer set.

4. If you purchased this product directly from Baxter, complete the enclosed Baxter Customer Reply Form and return it to Baxter by either faxing it or scanning and e-mailing it or sending it by post to, even if you don’t have any inventory. Returning the customer reply form promptly will confirm your receipt of this notification and prevent you from receiving repeat notices.

5. If you purchased this product from a distributor, please note that the Baxter customer reply form is not applicable. If a reply form is provided by your distributor or wholesaler, please return it to the supplier according to their instructions.

6. If you distribute this product to other facilities or departments within your institution, please forward a copy of this communication to them.

7. If you are a dealer, wholesaler, distributor/reseller, or original equipment manufacturer (OEM) that distributed any affected product to other facilities, please notify your customers of this Device Correction in accordance with your customary procedures.

Further information and support

For general questions regarding this communication, contact Baxter.

The local Ministry of Health (MOH) has been notified of this action.

We apologize for any inconvenience this may cause you and your staff.

Sincerely,

Baxter Healthcare Corporation