Central Drugs Standard Control Organization

Directorate General of Health Services
Ministry of Health and Family Welfare
Government of India

(Medical Devices and Diagnostics Division)

Email: ddcimd-cdsco@nic.in

Food & Drugs Administration Bhawan, Kotla Road, New Delhi.

Date: 21st July 2022

F.No.29/Misc/03/2022-DC (192)

MEDICAL DEVICE ALERT

DEVICE

Adsorba Hemoperfusion Cartridge 300 C (Product code – 101223, Lot no. 1-419, 1-430, 1-439, 1-440, 1-441) and Adsorba 300 C (Product code – 115264, Lot no. 1-507, 1-508, 1-509, 1-512, 1-513, 1-514)

BACKGROUND

Hemoperfusion should be considered if: 1. After taking life threatening amounts of adsorbable drugs, deep coma and one of the following symptoms are observed: hypoventilation, hypotonia, hypothermia, worsening of the clinical state despite conservative medical management; 2. the patient has taken drugs of which the amount, composition and kind are unknown and the patient is deeply comatose. The Adsorba has proven a high degree of efficacy for the following drugs: barbiturates, organophosphates, bromocarbamide, paracetamol, ethchlorvynol, meprobamate, phenacetin, methaqualone, salicylate. The use of hemoperfusion as a supplementary treatment does not mean that other conventional methods of treatment should be omitted; measures such as gastric lavage, establishment of free airway and assisted respiration, controlled electrolyte and water balance, and forced diuresis should be administered whenever indicated. Futher more it might be necessary to monitor carefully the blood levels of vital substances or drugs which also could be adsorbed during the hemoperfusion treatment. Access to the blood stream for hemoperfusion treatment can be obtained by normal hemodialysis methods.

Baxter Healthcare Corporation is issuing an Urgent Medical Device Recall for the the Adsorba Hemoperfusion Cartridge 300 C and Adsorba 300 C due to the presence of particulate matter within the cartridge.

Reason for Recall

 There was recall due to the potential presence of particulate matter within the cartridge.

Health Hazard

If particulate matter is not detected before use, the particles may reach the vascular system of the patient with potential serious adverse health consequences.

Who May Be Affected

- Any person with who have used an affected Adsorba Hemoperfusion Cartridge 300 C and Adsorba 300 C
- Health care providers who treat people using the affected Adsorba Hemoperfusion Cartridge 300 C and Adsorba 300 C

Note: CDSCO have not received any complaints from the market on this issue.

FURTHER DETAILS & CONTACTS

M/s Baxter India Pvt. Ltd, Gurgaon, Haryana had issued an Urgent Medical Device Recall which is attached herewith this alert.

M/s Baxter India Pvt. Ltd.

5th Floor, Tower A Building 9, DLF phase III. DLF cyber city, Gurgaon 122002 Haryana, India India product complaints@baxter.com Tel +91-124-4500200 Fax +91-124-4500200

Urgent Medical Device Recall

June 23, 2022

Dear Director of Materials Management:

Problem Description

Baxter Healthcare Corporation is issuing an Urgent Medical Device Recall for the Adsorba Hemoperfusion Cartridge 300 C and ADSORBA 300C APAC listed below due to the potential presence of particulate matter within the cartridge. The affected lot numbers are listed in the enclosed Attachment A.

Affected Product

Product Code	Product Description	Lot Numbers	UDI	Expiration
	Adsorba	- Tullibels		Dates
101223	Hemoperfusion Cartridge 300 C	See	07332414015473	0.
115264	ADSORBA 300C APAC	Attachment A	07332414118174	See Attachment A

Hazard Involved

If particulate matter is not detected before use, the particles may reach the vascular system of the patient with potential serious adverse health consequences. There have been no complaints or patient injury associated with this issue.

Actions to be taken by Customers

- Locate and remove all affected product lots from your facility. The product code and lot number can be found on the individual product and shipping carton.
- 2. Contact your Baxter sales representative for sales return and credit.
- 3. If you purchased this product directly from Baxter, complete the enclosed Baxter customer reply form mention the impacted product code, lot and associated lot quantity along with your contact details Name of facility location etc. and return it to Baxter by e-mailing it to India product complaints@baxter.com or sales representative email, even if you do not have any inventory returning the customer reply form promptly will confirm your receipt of this notification and prevent you (customer reply form), you will receive a phone call from Baxter sales or clinical representative on behalf of Baxter to confirm your receipt of this notification.
- 4. If you purchased this product from a distributor, contact your distributor for return and credit. Please note that the Baxter customer reply form is not applicable in this situation. If a reply form is provided by your distributor or wholesaler, please return it to the supplier according to their instructions.
- If you distribute this product to other facilities or departments within your institution, please forward a copy of this communication to them.

Baxter

6. If you are a dealer, wholesaler, distributor/reseller, or original equipment manufacturer (OEM) that distributed any affected product to other facilities, please conduct a consumer-level recall of the affected product that you distributed to customers and check the associated box on the reply form.

Further information and support For general questions regarding this communication, contact Baxter Sales Representatives.

The India Ministry of Health (MOH) will be notified of this action. Any adverse events or quality problems experienced with the use of these products may be reported using one of the following options:

Contact your Baxter sales representative.

Emailing to Baxter at: India product complaints@baxter.com.

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

Sincerely,

Ramadoss

Digitally signed by

Ramadoss

Sathyanaraya Sathyanarayanan

Date: 2022.06.24

09:55:58 +08'00'

Sathyanarayanan Ramadoss

Dir, QA-EA Mfg & Prod Surv-AP APAC QA

Baxter India Pvt Ltd.

Enclosure: Baxter Customer Reply Form

Attachment A: Affected Lot Numbers

Attachment A: Affected Lot Numbers

Adsorba Products

Product Code			
	Lot Number	Expiration Date	
101223	1-419	29-Feb-24	
101223	1-430 1-439		
101223		30-Apr-24	
101223		30-Apr-24	
101223	1-440	31-May-24	
	1-441	31-May-24	
115264	1-507	30-Sep-24	
115264	1-508	30-Sep-24	
115264	1-509	30-Sep-24	
115264	1-512		
115264	1-513	31-Oct-24	
115264		31-Oct-24	
_20207	1-514	31-Oct-24	