

March 19, 2019

## Urgent: Hospira, Inc. Initiates a Voluntary Recall of Select Lots of 8.4% Sodium Bicarbonate Injection, USP

Dear ASD Healthcare Customer:

Hospira, Inc., a Pfizer company, (“Hospira”) is initiating a voluntary recall of select lots of 8.4% Sodium Bicarbonate Injection, USP. The affected lot information is detailed below.

As a precautionary measure, this voluntary recall is being conducted due to the potential presence of particulate matter, confirmed as glass. The administration of a glass particulate, if present in an intravenous drug, may result in local irritation or swelling in response to foreign material. More serious potential outcomes would include blockage and clotting in blood vessels, which may be life-threatening if a critical organ is affected.

The recall of lots referenced below are being conducted to the **Hospital/Institution level**.

**Hospira provided the following affected lot information for 8.4% Sodium Bicarbonate Injection, USP:**

| NDC          | Lot Number | Exp. Date | Strength    | Configuration / Count   |
|--------------|------------|-----------|-------------|-------------------------|
| 0409-6625-02 | 79-238-EV  | 1JUL2019  | 50mEq/50 mL | Case Pack 4 x 25, 50 mL |
| 0409-6625-02 | 79-240-EV  | 1JUL2019  | 50mEq/50 mL | Case Pack 4 x 25, 50 mL |
| 0409-6625-02 | 80-088-EV  | 1AUG2019  | 50mEq/50 mL | Case Pack 4 x 25, 50 mL |

There have been no reports of serious injury or infection to date. **To reduce any risk, we ask for your cooperation in taking the following action:**

- **Review your Hospira Sodium Bicarbonate Injection, USP inventory** immediately to identify and isolate affected inventory in order to prevent future use.
- **Contact Stericycle at 1-800-805-3093** to obtain a BRC to initiate the return process.

- All returns are requested to be completed within six months of this notice date.

**For the full recall notice, please view the attached document on the following pages.**

If you have any questions or concerns, please contact ASD Healthcare's customer service at **800.746.6273**.



## URGENT: DRUG RECALL

March 19, 2019

### 8.4% Sodium Bicarbonate Injection, USP 50 mEq/50mL (1 mEq/mL)

| NDC          | Lot Number | Expiration Date | Strength     | Configuration/Count    |
|--------------|------------|-----------------|--------------|------------------------|
| 0409-6625-02 | 79-238-EV  | 1JUL2019        | 50 mEq/50 mL | Case Pack 4 x 25, 50mL |
| 0409-6625-02 | 79-240-EV  | 1JUL2019        | 50 mEq/50 mL | Case Pack 4 x 25, 50mL |
| 0409-6625-02 | 80-088-EV  | 1AUG2019        | 50 mEq/50 mL | Case Pack 4 x 25, 50mL |

Dear Customer:

Hospira, Inc., a Pfizer company, ("Hospira") is voluntarily recalling the above referenced lots of 8.4% Sodium Bicarbonate Injection, USP due to the potential presence of particulate matter, confirmed as glass. The administration of a glass particulate, if present in an intravenous drug, may result in local irritation or swelling in response to the foreign material. More serious potential outcomes would include blockage and clotting in blood vessels, which may be life-threatening if a critical organ is affected. To date, Hospira has not received reports of any adverse events associated with this issue for these lots. The risk is reduced by the possibility of detection, as the label contains a clear statement directing the healthcare professional to visually inspect the product for particulate matter and discoloration prior to administration.

**FEDERAL REGULATION 21 CFR 7.49 (d) RESPONSIBILITY OF RECIPIENT STATES: "CONSIGNEES THAT RECEIVE A RECALL COMMUNICATION SHOULD IMMEDIATELY CARRY OUT THE INSTRUCTIONS SET FORTH BY THE RECALLING FIRM ..."HOSPIRA RECOMMENDS THAT YOU RESPOND TO THIS RECALL EVEN IF YOU DO NOT HAVE THE RECALLED PRODUCT. TO RESPOND, COMPLETE THE REQUESTED INFORMATION ON THE ATTACHED BUSINESS REPLY CARD (BRC) AND RETURN AS DIRECTED, WITHIN FIVE (5) BUSINESS DAYS.**

If you have any questions about responding to this letter, please contact Stericycle Inc. at 1-800-805-3093 (Mon.-Fri. 8am-5pm ET).

The recall of the above-referenced lots are being conducted to the **Hospital/Institution level**.

Our records indicate you may have received shipment of the affected product between **August 2017 and September 2017**. Please check your stock immediately against the table above. If you have any of the affected product in your inventory, please stop distribution immediately and promptly return it to Stericycle using the

Hospira, Inc., a Pfizer company  
275 North Field Drive  
Lake Forest, IL 60045  
(224) 212-2000  
[www.pfizerinjectables.com](http://www.pfizerinjectables.com)



label provided with this letter. **All returns are requested to be completed within six months of this notice date.** To ensure proper and timely credit, follow the instructions on the return label for returning the product.

If you have further distributed the recalled product, please notify any accounts or additional locations which may have received the recalled product from you. Please conduct a sub-recall to those accounts and communicate this recall information immediately. Please request they immediately cease distribution of the affected product and promptly contact Stericycle at 1-800-805-3093 to obtain a BRC to initiate the return process.

Please contact Pfizer Customer Service at 1-844-646-4398 (Mon.-Fri. 8am-7pm ET) or your Pfizer representative regarding product availability and for questions regarding this market action.

This recall is being conducted with the knowledge of the U.S. Food and Drug Administration. We appreciate your immediate attention and cooperation and sincerely regret any inconvenience this action may cause you. If you have any questions regarding this recall you may contact Pfizer using the below information.

| Contact                    | Contact Information                                              | Areas of Support                               |
|----------------------------|------------------------------------------------------------------|------------------------------------------------|
| Pfizer Medical Information | 1-800-438-1985, option 3<br>(8am – 7pm ET Monday through Friday) | Medical Inquiries                              |
| Pfizer Safety              | 1-800-438-1985, option 1<br>(24 hours a day 7 days per week)     | To report adverse events or product complaints |

Sincerely,

Navin Katyal  
General Manager, Pfizer Injectables  
Pfizer Essential Health

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275 North Field Drive  
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