FDA Recall – Clearlink Basic Solution Set with Duovent

Purpose of this communication:

- We are writing to inform you that Baxter Healthcare Corporation is recalling Clearlink Basic Solution Set with Duovent after increased customer reports of leaks.

What do I need to know?

- As the majority of the Clearlink Basic Solution Sets with Duovent are used for the delivery of hazardous drugs (chemotherapy), leakage could expose healthcare personnel, patients, and others to potentially hazardous drugs that may be toxic and/or are irritants. These leaks may also allow air into the set or breach the sterile fluid pathway, thereby increasing the risk of air embolism and contaminated infusions, respectively. Patients may suffer delayed or interrupted therapy or may not receive the necessary amount of their medication. These issues could lead to serious injury or death.

What do I need to do?

- Review the following recall notice: [Baxter Healthcare Corporation Recalls Clearlink Basic Solution Set with Duovent for Risk of Leaks That May Expose Providers and Patients to Hazardous / Toxic Substances: FDA](#)
- Notify impacted patients and facilitate the repair, replacement and/or resolution of the recall, according to the guidelines issued by the manufacturer in the FDA notification.
- Confirm if you have any CareCentrix patients affected by this recall and notify the CareCentrix Quality Department by faxing information to: 919-792-6718 Attn: Quality Department.

Thank you in advance for your cooperation and continued partnership.