

DUPLEX® Container

Directions for Use

"PEEL"

PEEL THE FOIL STRIP FROM THE DRUG CHAMBER



"SNAP"

SQUEEZE THE BAG TO OPEN THE FIRST SEAL BETWEEN THE DILUENT AND DRUG POWDER



"SHAKE"

SHAKE THOROUGHLY TO MIX



"SNAP"

SQUEEZE THE BAG TO OPEN THE SECOND SEAL, RELEASING SOLUTION TO THE SET PORT



"GO"

REMOVE THE FOIL TAB FROM THE SET PORT, SPIKE BAG AND START ADMINISTRATION



Label and Inspect:

- Apply patient-specific label on foil side of container. Do not cover any portion of foil strip with patient label.
- Unfold the DUPLEX Container. Use only if container and seals are intact.
- Visually inspect diluent chamber for particulate matter. To inspect the drug powder for foreign matter or discoloration, peel foil strip from drug chamber. Protect from light after removal of foil strip.

NOTE: Product does not require refrigeration prior to activation.

Reconstitute:

- Unfold the DUPLEX Container and point the set port in a downward direction.
- Starting at the hanger tab end, fold the DUPLEX Container just below the diluent line, trapping all air above the fold. To activate, squeeze the folded diluent chamber until the seal between the diluent and powder opens, releasing diluent into the drug powder chamber.
- Shake the diluent-powder mixture until the drug powder is completely dissolved.
- Visually inspect the reconstituted solution for particulate matter.

Administer:

- Point the set port in a downward direction.
- Starting at the hanger tab end, fold the DUPLEX Container just below the solution line, trapping all air above the fold. Squeeze the folded DUPLEX Container until the seal between the solution and set port opens, releasing solution to set port.
- Using aseptic technique, remove the foil tab cover from the set port and attach sterile administration set. Refer to Directions for Use accompanying the administration set.

Precautions:

- Use only if prepared solution is clear and free from particulate matter.
- Do not use in series connection.
- Do not introduce additives into the DUPLEX Container.
- Do not freeze.
- Refer to product package insert for complete Directions for Use and prescribing information.

DUPLEX Containers are not made with DEHP, PVC or natural rubber latex.

Product Description	NDC No.	REF No.
500 mg Meropenem for Injection USP and Sodium Chloride Injection USP	00264-3183-11	3183-11
1 g Meropenem for Injection USP and Sodium Chloride Injection USP	00264-3185-11	3185-11
1 g Cefazolin for Injection USP and Dextrose Injection USP	00264-3103-11	3103-11
2 g Cefazolin for Injection USP and Dextrose Injection USP	00264-3105-11	3105-11
3 g Cefazolin for Injection USP and Dextrose Injection USP	00264-3107-11	3107-11
1 g Cefoxitin for Injection and Dextrose Injection	00264-3123-11	3123-11
2 g Cefoxitin for Injection and Dextrose Injection	00264-3125-11	3125-11
1 g CefTRIaxONE for Injection and Dextrose Injection	00264-3153-11	3153-11
2 g CefTRIaxONE for Injection and Dextrose Injection	00264-3155-11	3155-11
1 g Cefepime for Injection USP and Dextrose Injection USP	00264-3193-11	3193-11
2 g Cefepime for Injection USP and Dextrose Injection USP	00264-3195-11	3195-11
2.25 g Piperacillin and Tazobactam for Injection USP and 50 mL of 0.45% Sodium Chloride Injection USP	0264-3446-11	3446-11
3.375 g Piperacillin and Tazobactam for Injection USP and 50 mL of 0.3% Sodium Chloride Injection USP	0264-3448-11	3448-11
4.5 g Piperacillin and Tazobactam for Injection USP and 100 mL of 0.45% Sodium Chloride Injection USP	0264-3450-22	3450-22