

INSTRUCTIONS FOR USE

Measured Volume Fluid Administration Set (POLYPEDIA)

Materials Used

- Polypropylene, Acrylonitrile Butadiene Styrene, PVC, PE, Stainless Steel, Polycarbonate, Glass Fibre.

Indications

- Infusion of IV fluids, administration of other drugs and medicines.
- Maintaining hydration in patients who are unable to take sufficient volume of oral fluids.

Contraindications

- Blood Transfusion.
- Administration of highly viscous fluids.
- Not to be used in patients with known hypersensitivity to any of the materials used.

Directions for Use

- Close the upper clamp and regulating clamp (lower clamp).
- Remove anti kink device before using the product from bottom.
- Remove the spike cover and insert the spike into the I.V. fluid container to its full length.
- Open the upper clamp and airvent to allow about 25 ml solution into the graduated burette. Close the upper clamp.
- Squeeze the drip chamber and adjust the fluid level till 1/3rd filled.
- Remove the cap of the connector of IV set and slightly open lower clamp to remove air from the tubing. Close the regulating clamp.
- Open the upper clamp to allow solution flow into graduated chamber until the desired volume is obtained. Do not fill the chamber beyond its maximum capacity. Close the upper clamp and connect the set with IV device.
- Open the lower clamp. Adjust drop rate and control infusion with the regulating clamp. Allow the infusion to continue.
- The valve will shut off the flow when solution level in graduated chamber is zero.
- When more solution is required, close the regulating clamp and open the upper clamp to fill graduated chamber until desired level. Close the upper clamp and squeeze the drip chamber a little to release the float valve.

Cautions

- Do not use if protective cap is loose or missing.
- Do not use if package is opened or damaged.
- Use injection site and Y site only for injecting the medicines using aseptic techniques.
- Close the airvent during periods of interrupted infusion therapy.
- For gravity feed only.
- The product should be used only by qualified healthcare professionals.
- Do not re-sterilize. Discard the set after single use.
- Reuse and cleaning of product may alter their structural and mechanical properties. It may lead to infection or other illness/injury.
- The product should be replaced and disposed of as per facility approved protocol or CDC guidelines.
- Store in between 5°C to 35°C, avoid excessive heat, protect from direct sunlight and moisture.
- Poly Medicure Limited will not be responsible for any direct incidental or consequential damages resulting from reuse of product.



Cautions



See Instructions for use



Product reference/Art. No.



For single use only



Do not Resterilize



Do not use if packaging or product has been damaged or contaminated



Batch Number



Date of Manufacturing



Use by / Expiry date



Sterilised by Ethylene Oxide



Single Sterile Barrier System



European Authorised Representative



Manufactured by



Non Pyrogenic



Storage Condition



Keep Away from Sunlight



Keep Dry



Medical Devices

Not made with natural rubber latex
Not made with DEHP plasticizers

 **POLYMED**

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