

Polyflush syringe DS
0.9% sodium chloride (0.9% NaCl)
Double sterile (DS) = Sterile Fluid + Sterile Syringe
INSTRUCTIONS FOR USE

GB

MATERIALS USED

Polymerized SBR, Sodium Chloride

INDICATION AND CLINICAL USE

Prefilled Syringe with Saline solution is intended to be used only for flushing of indwelling vascular access devices (VAD's) for maintaining patency of VAD's.

CONTRAINDICATION

None Known

ADVERSE REACTION

No known adverse reaction known when the product is indicated.

DIRECTIONS FOR USE

Use aseptic technique throughout the procedure.

1. Open the pack and remove syringe.

2. Wash hands and prepare aseptic barrier and

disinfection prior to use according to Warnings and Precautions.

Check that syringe tip cap is in place. Inspect the clarity of

solution visually.

3. Remove the stopper cap to release the stopper seal (Fig.1b).

4. Unscrew tip cap from the syringe ensuring that there is no touch contamination of the syringe luer connection (Fig. 2).

5. Push syringe plunger into the air. Do not use device if no touch contamination has occurred.

6. Connect Polyflush Syringe to vascular access device,

taking care that there is no touch contamination of the connection (Fig. 4).

7. Push syringe plunger to flush the required volume of saline

following institution's policy (Fig. 5). In case of extreme plunger

resistance, it is recommended that excess force is not exerted.

After use, flush the syringe including any unused solution in

accordance with institution's biohazardous waste policy or CDC

guidelines (Fig. 6).

ACTION

0.9% Sodium Chloride solution is a sterile, aqueous solution having approximately the same osmotic pressure and composition as extracellular fluids. It is non-irritating to tissues. It is used to flush vascular access devices in order to maintain catheter patency and to decontaminate the device.

WARNINGS AND PRECAUTIONS

1. Read carefully instructions for use before using the product.

2. The product should be used only by qualified healthcare

3. Not to be used for injections as this is intended for flushing only.

4. Check the expiration date of product before use. Do not use if

5. Do not use if tip/plunger is damaged, as it indicates syringe

leakage or if sterile

6. Do not use if packaging or product has been damaged or

7. Visually inspect the contents of each prefilled syringe for

presence of saline, clear content, particulate matter,

discoloration, discolouration and leakage prior to use. Do not use if

8. Do not allow air to be trapped in fluid path.

9. Do not re-sterilize. Discard the set after single use.

10. For single use in a single patient only. Reuse of single use

device may result in cross-contamination.

Contamination and/or limited functionality of the device may

lead to injury, illness or inflammation.

PolyMedic Limited will not be responsible for any direct incidental

or consequential damage resulting from reuse of product.

STORAGE AND STABILITY

Prefilled Syringes with 0.9% sodium chloride solution should be stored

between 5°C and 25°C, protect from direct sunlight and freezing.



	Cautions		See Instructions for use		For single use only
	Do not use		Batch Number		Date of Manufacturing
	Do not sterilize		STERILE R		European Authorised Representative
	Use by / Expiry		STERILE R		European Authorised Representative
	Non Pyrogenic		Storage Condition		Keep Away from Sunlight
	Storage Condition		Keep Dry		Single sterile barrier system

Спринцовка Polyflush DS
0.9% натриев хлорид (0.9% NaCl)

Двойна стерилна (DS) = Стерилна течуна + стерилна спринцовка

ИНСТРУКЦИИ ЗА УПОТРЕБА

CE 0123

ИЗПОЛЗУВАНИ МАТЕРИАЛАМ

Полимеризиран SBR, Sodium Chloride

ИНДИКАЦИИ А КЛИНИЧКА УПОТРЕБА

Prefilled Syringe with Saline solution is intended to be used only for

flushing of indwelling vascular access devices (VAD's) for maintaining

patency of VAD's.

CONTRAINDICATION

None Known

ADVERSE REACTION

No known adverse reaction known when the product is indicated.

DIRECTIONS FOR USE

Use aseptic technique throughout the procedure.

1. Open the pack and remove syringe.

2. Wash hands and prepare aseptic barrier and

disinfection prior to use according to Warnings and Precautions.

Check that syringe tip cap is in place. Inspect the clarity of

solution visually.

3. Remove the stopper cap to release the stopper seal (Fig.1b).

4. Unscrew tip cap from the syringe ensuring that there is no touch

contamination of the syringe luer connection (Fig. 2).

5. Push syringe plunger into the air. Do not use device if no

touch contamination has occurred.

6. Connect Polyflush Syringe to vascular access device,

taking care that there is no touch contamination of the connection (Fig. 4).

7. Push syringe plunger to flush the required volume of saline

following institution's policy (Fig. 5). In case of extreme plunger

resistance, it is recommended that excess force is not exerted.

After use, flush the syringe including any unused solution in

accordance with institution's biohazardous waste policy or CDC

guidelines (Fig. 6).

ACTION

0.9% Sodium Chloride solution is a sterile, aqueous solution having

approximately the same osmotic pressure and composition as

extracellular fluids. It is non-irritating to tissues. It is used to flush

vascular access devices in order to maintain catheter patency and to

decontaminate the device.

WARNINGS AND PRECAUTIONS

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2. The product should be used only by qualified healthcare

3. Not to be used for injections as this is intended for flushing only.

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or consequential damage resulting from reuse of product.

STORAGE AND STABILITY

Prefilled Syringes with 0.9% sodium chloride solution should be stored

between 5°C and 25°C, protect from direct sunlight and freezing.

Proplachovaci stříkačka Polyflush DS
0.9% chlorid sodný (0.9% NaCl)

Dvojité sterilita (DS) = sterilní kapalina + sterilní stříkačka

INSTRUKCE NA UPOŘEPU

CE 0123

KÓD VÝROBECKY

Polypropylen SBR, Chlorid sodný

INDIKACE A KLINICKÁ UPOŘEPA

Proplachovací stříkačka sestavená z fyziologického roztoku, který je určen

k udržení patency indwelling vascular access devices (VAD's) pro

přemístění a udržení patency.

CONTRAINDIKACE

Ne je známé.

ADVERSE REACTION

Ne je známé.

DIRECTIONS FOR USE

Užívání je bezpečné po celém období užití.

1. Otevřete balení a vyměňte stříkačku.

2. Před použitím, vymaďte výrobu a očistěte s rukavicemi a mydlem.

3. Provedete dlaněmi na výrobce a mydlem.

4. Check, že je správně umístěna výroba.

5. Použijte výrobu.

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