

**DIANEAL LOW CALCIUM WITH DEXTROSE- sodium chloride, sodium lactate, calcium chloride, magnesium chloride and dextrose injection, solution
DIANEAL PD-2 WITH DEXTROSE- sodium chloride, sodium lactate, calcium chloride, magnesium chloride and dextrose injection, solution**

Baxter Healthcare Corporation

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

DIANEAL with Dextrose

Health Care Provider Letter

IMPORTANT PRESCRIBING INFORMATION

DATE: October 21, 2024

Subject: Temporary importation of DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose (2000 mL) and DIANEAL Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose (2000 mL) from Guangzhou, China for use in Automated Peritoneal Dialysis to address drug shortages

Dear Healthcare Professional,

Due to the current critical shortage of DIANEAL PD-2 and DIANEAL Low Calcium (2.5 mEq/L) Peritoneal Dialysis Solution in the United States (US) market, Baxter Healthcare Corporation (Baxter) is coordinating with the U.S. Food and Drug Administration (FDA) to temporarily import DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose and Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose (2000 mL) from Baxter's manufacturing facility in Guangzhou, China. FDA has not approved this product manufactured by Baxter's Guangzhou, China facility.

You may be provided with additional letters for other DIANEAL imported peritoneal dialysis solutions you receive. Please read each letter in its entirety because each letter may contain different, product-specific information.

Baxter has initiated temporary importation of DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose and Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose (2000 mL) for use in Ambulatory Peritoneal Dialysis (APD) therapy as described in the table below. This product is manufactured by Baxter's manufacturing facility in Guangzhou, China and is marketed in Hong Kong. At this time, importation or distribution of DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose and Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose (2000 mL) in the United States by any entity other than Baxter or its authorized distributor(s) is considered a violation of the Federal Food, Drug, and Cosmetic Act and is subject to enforcement by the FDA.

Effective immediately, and during this temporary period, Baxter will offer the following imported products from Baxter's facility in Guangzhou, China:

China Imported Product Name and Description	APD Solution Volume	Product Code	Bags per Carton	NDC Code
DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose	2000 mL	6AB5177E	6 bags	NDC 0941-0698-01 (bag) NDC 0941-0698-06 (carton)
DIANEAL Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose	2000 mL	6AB9747E	6 bags	NDC 0941-0696-01 (bag) NDC 0941-0696-06 (carton)

It is important to note the following:

- DIANEAL PD-2 with 2.5% Dextrose and DIANEAL Low Calcium (2.5mEq/L) PD Solution with 4.25% Dextrose imported from China will only be available in 2000 mL volume for APD, so there will need to be adaptation to the PD prescription for some patients.
- DIANEAL Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose contains 2.5 mEq/L of calcium compared to DIANEAL PD-2 Peritoneal Dialysis Solution with 4.25% Dextrose which contains 3.5 mEq/L of calcium. Patients receiving China-imported DIANEAL Low Calcium (2.5mEq/L) PD Solution solution with 4.25% dextrose should have their serum calcium levels monitored for the development of hypocalcemia.
- There are no other clinically relevant differences in the DIANEAL drug composition between the U.S.-manufactured and China-manufactured product (see Table 1, below). As such, clinical practice for usage, administration, and dosage for the China imported product is the same as with DIANEAL PD-2 and DIANEAL Low Calcium (2.5 mEq/L) Peritoneal Dialysis Solution manufactured in the US. Please refer to the FDA-approved Dianeal Peritoneal Dialysis Solution Prescribing Information for reference.
- The Luer-lock connector on the China imported product functions the same as and is fully compatible with peritoneal dialysis sets marketed in the United States. However, the U.S. product has color-coded pull rings covering the luer to identify the dextrose concentration, while the China-imported product has a blue protective tip protector which is the same for all dextrose concentrations. The frangible is green in the imported product but blue in U.S. product. Users of the China imported product should check the product label to ensure that they are using the correct dextrose concentration. See Table 1 for more details of product differences.
- The China imported product may include barcodes on the shipping carton; however, the **barcodes may not register accurately in the U.S. scanning systems**. There are no barcodes on the solution containers of the China imported product. Alternative procedures should be followed to assure that the correct drug product is being used in all systems and processes and administered to individual patients. For example, institutions should consider manually inputting the product into their systems and confirm that barcode systems do not provide incorrect information when the product is scanned. Please note that the imported solutions bags do not have barcodes.

Before prescribing, healthcare providers should be aware of some key differences in the container packaging and labeling between the China imported products and the FDA-approved products which are stated in the product comparison tables at the end of this letter as follows:

- Table 1: Key differences of DIANEAL Peritoneal Dialysis Solution for APD therapy
- Table 2: Label images of DIANEAL 2000 mL APD product presentations

Reporting Adverse Events

To report adverse events associated with these imported products, please call Baxter at 1-866-888-2472, or fax: 1-800-759-1801. Adverse events or quality problems experienced with the use of these imported products may also be reported to the FDA's MedWatch Adverse Event Reporting program either online, by regular mail or by fax:

- Complete and submit the report **Online**: www.fda.gov/medwatch/report.htm
- **Regular mail or Fax**: Download form www.fda.gov/MedWatch/getforms.htm or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form, or submit by fax to 1-800-FDA-0178 (1-800-332-0178)

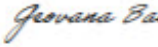
To report product quality issues, please report to:
[Baxter - Product Feedback Portal](https://productfeedback.baxter.com/) (<https://productfeedback.baxter.com/>).

**Please refer to the FDA approved full prescribing information for Dianeal Peritoneal Dialysis Solution at
[DailyMed \(nih.gov\)](#)**

If you have any questions about the information contained in this letter or the use of imported DIANEAL PD-2 with 2.5% Dextrose and DIANEAL Low Calcium (2.5mEq/L) PD Solution with 4.25% Dextrose (2000 mL), please contact Baxter's Medical Information Service at 1-888-736-2543.

To place an order, please contact Baxter's Center for Home Care Services by calling 1-800-284-4060.

Sincerely,


Electronically signed by:
Geovana Basso
Reason: Ok.
Date: Oct 21, 2024 14:22
CDT

Geovana Basso, M.D.
Director of Americas Medical Affairs
Baxter Healthcare Corporation
One Baxter Parkway
Deerfield, Illinois 60015

Baxter and Dianeal are registered trademarks of Baxter International Inc.

Attachments:

Product Comparison Tables 1 and 2

Table 1. Key differences of DIANEAL Peritoneal Dialysis Solutions for APD therapy

	Imported Product (Guangzhou, China) DIANEAL Low Calcium 4.25% Dextrose	Imported Product (Guangzhou, China) DIANEAL PD-2 2.5% Dextrose	U.S. FDA Approved Product DIANEAL Low Calcium	U.S. FDA Approved Product DIANEAL PD2
Product name	DIANEAL Low Calcium (2.5 mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose	DIANEAL PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose	DIANEAL Low Calcium (2.5mEq/L) Peritoneal Dialysis Solution with 2.5% Dextrose	DIANEAL PD2 Peritoneal Dialysis Solution with 2.5% Dextrose
Labeled Fill Volume	2000 mL	2000 mL	2000 mL and 3000 mL solution fill volumes available	1000, 2000 mL and 3000 mL solution fill volumes available
Container Type	Ambu-Flex (PVC) container with luer-lock and blue tip connector cap	Ambu-Flex (PVC) container with luer-lock and blue tip connector cap	AMBU-FLEX II (PVC) container with luer-lock connector and colored pull ring caps	AMBU-FLEX II (PVC) container with luer-lock connector and colored pull ring caps
Bags per Carton	6 bags	6 bags	2000 mL: 6 bags 3000 mL: 4 bags	1000 mL: 12 bags 2000 mL: 6 bags 3000 mL: 4 bags
Indications	DIANEAL Low Calcium peritoneal dialysis solutions is indicated for use in chronic renal failure patients being maintained on peritoneal dialysis	Peritoneal dialysis is indicated for patients in acute or chronic renal failure when nondialytic medical therapy is judged to be inadequate (Vaamonde and Perez 1977). It may also be indicated in the treatment of certain fluid and electrolyte disturbances, and for patients intoxicated with certain poisons and drugs (Knepshield et al. 1977). However, for many substances other methods of detoxification have been reported to be more effective than peritoneal dialysis (Vaamonde and Perez; Chang 1977)	DIANEAL peritoneal dialysis solutions are indicated for patients in acute or chronic renal failure.	DIANEAL peritoneal dialysis solutions are indicated for patients in acute or chronic renal failure.
Active Ingredient - Dextrose (Glucose)	Dextrose Hydrous, USP 4.25g / 100mL	Dextrose Hydrous, USP 2.5g / 100mL	1.5% Dextrose: Dextrose Hydrous, USP 1.5g / 100mL 2.5% Dextrose: Dextrose Hydrous, USP 2.5g / 100mL 4.25% Dextrose: Dextrose Hydrous, USP 4.25g / 100mL	1.5% Dextrose: Dextrose Hydrous, USP 1.5g / 100mL 2.5% Dextrose: Dextrose Hydrous, USP 2.5g / 100mL 4.25% Dextrose: Dextrose Hydrous, USP 4.25g / 100mL
Active Ingredients - Electrolytes	Sodium Chloride, USP 538 mg / 100mL Sodium Lactate 448 mg / 100 mL Calcium Chloride, USP 18.3 mg / 100 mL Magnesium Chloride, USP 5.08 mg / 100 mL	Sodium Chloride, USP 538 mg / 100mL Sodium Lactate 448 mg / 100 mL Calcium Chloride, USP 25.7 mg / 100 mL Magnesium Chloride, USP 5.08 mg / 100 mL	Sodium Chloride, USP 538 mg / 100mL Sodium Lactate 448 mg / 100 mL Calcium Chloride, USP 18.3 mg / 100 mL Magnesium Chloride, USP 5.08 mg / 100 mL	Sodium Chloride, USP 538 mg / 100mL Sodium Lactate 448 mg / 100 mL Calcium Chloride, USP 25.7 mg / 100 mL Magnesium Chloride, USP 5.08 mg / 100 mL

Imported Product (Guangzhou, China) DIANEAL Low Calcium 4.25% Dextrose		Imported Product (Guangzhou, China) DIANEAL PD-2 2.5% Dextrose	U.S. FDA Approved Product DIANEAL Low Calcium	U.S. FDA Approved Product DIANEAL PD2
Electrolyte Content per Liter	Sodium 132 mEq/L Calcium 2.5 mEq/L Magnesium 0.5 mEq/L Chloride 95 mEq/L Lactate 40 mEq/L	Sodium 132 mEq/L Calcium 3.5 mEq/L Magnesium 0.5 mEq/L Chloride 98 mEq/L Lactate 40 mEq/L	Sodium 132 mEq/L Calcium 2.5 mEq/L Magnesium 0.5 mEq/L Chloride 95 mEq/L Lactate 40 mEq/L	Sodium 132 mEq/L Calcium 3.5 mEq/L Magnesium 0.5 mEq/L Chloride 96 mEq/L Lactate 40 mEq/L
pH	5.2 (4.5 to 6.5)	5.2 (4.5 to 6.5)	5.2 (4.0 to 6.5)	5.2 (4.0 to 6.5)
Additional Information	4.25% Dextrose: Osmolarity 483 mOsmol/L (Calc)	2.5% Dextrose: Osmolarity 396 mOsmol/L (Calc)	2.5% Dextrose: Osmolarity 395 mOsmol/L (Calc) 4.25% Dextrose Osmolarity 483 mOsmol/L (Calc)	2.5% Dextrose: Osmolarity 396 mOsmol/L (Calc) 4.25% Dextrose: Osmolarity 485 mOsmol/L (Calc)
Storage Conditions	Store at room temperature (25°C/77°F): Brief exposure up to 40°C (104°F) does not adversely affect the product.	Store at room temperature (25°C/77°F): Brief exposure up to 40°C (104°F) does not adversely affect the product.	Store at room temperature (25°C/77°F) Brief exposure up to 40°C does not adversely affect the product	Store at room temperature (25°C/77°F) Brief exposure up to 40°C does not adversely affect the product
Expiration Dating	24 months	24 months	24 months	1000 mL: 18 months 2000 mL: 24 months 3000 mL: 24 months
Container Closure System				

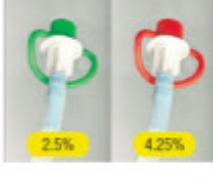
Imported Product (Guangzhou, China) DIANEAL Low Calcium 4.25% Dextrose		Imported Product (Guangzhou, China) DIANEAL PD-2 2.5% Dextrose	U.S. FDA Approved Product DIANEAL Low Calcium	U.S. FDA Approved Product DIANEAL PD2
Container Closure				
Container Closure Differences:	<ul style="list-style-type: none"> One green frangible at luer connector Blue protective tip protector 	<ul style="list-style-type: none"> One green frangible at luer connector Blue protective tip protector 	<ul style="list-style-type: none"> One blue frangible at luer connector Pull ring cap color-coded to solution dextrose concentration: <ul style="list-style-type: none"> Green = 2.5% dextrose Red = 4.25% dextrose 	<ul style="list-style-type: none"> One blue frangible at luer connector Pull ring cap color-coded to solution dextrose concentration: <ul style="list-style-type: none"> Green = 2.5% dextrose Red = 4.25% dextrose

Table 2. Label images of DIANEAL 2000 mL APD product presentations

Comparative container labels are presented below for DIANEAL Peritoneal Dialysis Solution in the 2000 mL fill volume. Labels for other US approved solution fill volumes differ only by product code / NDC / Fill Volume / Barcode. There are no differences in composition or other safety-related information.

Imported Product (Guangzhou, China) 2000mL DIANEAL PD Solution	US FDA Approved Product DIANEAL Low Calcium PD Solution	US FDA Approved Product DIANEAL PD-2
DIANEAL Low Calcium with 4.25% Dextrose PD Solution  2000ml <small>(APPROX 80mL EXCESS) 3000mL NOMINAL SIZE CONTAINER</small> Baxter Dianeal® Low Calcium(2.5mEq/L) Peritoneal Dialysis Solution With 4.25% Dextrose <small>EACH 100mL CONTAINS: 4.25 g DEXTROSE HYDROLYSIS USP 53.8 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L: SODIUM -132 CALCIUM -2.5 MAGNESIUM -0.5 CHLORIDE -132 LACTATE -40 OSMOLALITY -483 mOsm/L (CALC) STERILE NON-PYROGENIC</small> <div style="border: 1px solid black; padding: 2px;">POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</div> <small>WARNING: EXTENSIVE USE OF THIS SOLUTION DURING ONE PERITONEAL DIALYSIS PROCEDURE CAN RESULT IN SIGNIFICANT REMOVAL OF WATER FROM THE PATIENT. FOR INTRAPERITONEAL ADMINISTRATION ONLY. USE AS DIRECTED BY PHYSICIAN.</small> <small>CAUTIONS: SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY. DISCARD IF LEAKS ARE FOUND. DO NOT USE UNLESS SOLUTION IS CLEAR. DISCARD UNUSED PORTION. STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (UNDER 25°C) UNTIL READY TO USE. AVOID EXCESSIVE HEAT. SEE INSERT.</small> <small>請儲存於攝氏25度以下</small> Ambu-Flex™ III CONTAINER PL-146® PLASTIC <small>MANUFACTURED BY BAFTER HEALTHCARE (GUANGZHOU) CO LTD GUANGZHOU CHINA (AN AFFILIATE OF BAXTER WORLD TRADE INC USA) HK-42532 DIRECTIONS TO BE USED AS DIRECTED BY THE PHYSICIAN 用法 請依照醫生指示使用 Prescription Drug 處方藥物 Manufacturer Address: Jiaoyuan Road, Dongji Industrial District, GETDD, Guangzhou, P.R. China</small>	L5B9747  2000 mL <small>(APPROX 80mL EXCESS) 3000 mL NOMINAL SIZE CONTAINER</small> Baxter Dianeal Low Calcium (2.5 mEq/L) Peritoneal Dialysis Solution with 4.25% Dextrose <small>EACH 100mL CONTAINS: 4.25 g DEXTROSE HYDROLYSIS USP 53.8 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L: SODIUM -132 CALCIUM -2.5 MAGNESIUM -0.5 CHLORIDE -132 LACTATE -40 OSMOLALITY -483 mOsm/L (CALC) STERILE NONPYROGENIC</small> <div style="border: 1px solid black; padding: 2px;">POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</div> <small>SEE PACKAGE INSERT FOR DOSAGE INFORMATION USE AS DIRECTED BY PHYSICIAN FOR INTRAPERITONEAL ADMINISTRATION ONLY WARNING: EXTENSIVE USE OF THIS SOLUTION DURING ONE PERITONEAL DIALYSIS PROCEDURE CAN RESULT IN SIGNIFICANT REMOVAL OF WATER FROM THE PATIENT.</small> <small>CAUTIONS: SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY. DISCARD IF LEAKS ARE FOUND. DO NOT USE UNLESS SOLUTION IS CLEAR. DISCARD UNUSED PORTION. Rx ONLY STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (20°C/77°F) UNTIL READY TO USE. AVOID EXCESSIVE HEAT. SEE INSERT.</small> Ambu-Flex II CONTAINER PL 146 PLASTIC <small>BAFTER DANEAL AMBU-FLEX II AND PL 146 ARE TRADEMARKS OF BAXTER INTERNATIONAL, INC BAXTER HEALTHCARE CORPORATION DESPFIELD, IL 60011 USA MADE IN USA</small>	L5B5187  2000 mL <small>(APPROX 80mL EXCESS) 3000 mL NOMINAL SIZE CONTAINER</small> Baxter Dianeal PD-2 Peritoneal Dialysis Solution with 4.25% Dextrose <small>EACH 100 mL CONTAINS: 4.25 g DEXTROSE HYDROLYSIS USP 53.8 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L: SODIUM -132 CALCIUM -2.5 MAGNESIUM -0.5 CHLORIDE -132 LACTATE -40 OSMOLALITY -483 mOsm/L (CALC) STERILE NONPYROGENIC</small> <div style="border: 1px solid black; padding: 2px;">POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</div> <small>SEE PACKAGE INSERT FOR DOSAGE INFORMATION USE AS DIRECTED BY PHYSICIAN FOR INTRAPERITONEAL ADMINISTRATION ONLY WARNING: EXTENSIVE USE OF THIS SOLUTION DURING ONE PERITONEAL DIALYSIS PROCEDURE CAN RESULT IN SIGNIFICANT REMOVAL OF WATER FROM THE PATIENT.</small> <small>CAUTIONS: SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY. DISCARD IF LEAKS ARE FOUND. DO NOT USE UNLESS SOLUTION IS CLEAR. DISCARD UNUSED PORTION. Rx ONLY STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (20°C/77°F) UNTIL READY TO USE. AVOID EXCESSIVE HEAT. SEE INSERT.</small> Ambu-Flex II CONTAINER PL 146 PLASTIC <small>BAXTER DANEAL AMBU-FLEX II AND PL 146 ARE TRADEMARKS OF BAXTER INTERNATIONAL, INC BAXTER HEALTHCARE CORPORATION DESPFIELD, IL 60011 USA MADE IN USA</small>

Imported Product (Guangzhou, China) 2000mL DIANEAL PD Solution	US FDA Approved Product DIANEAL Low Calcium PD Solution	US FDA Approved Product DIANEAL PD-2
<p>DIANEAL PD-2 with 2.5% Dextrose PD Solution</p> <p>6AB5177E 2000ml (APPROX 80ml EXCESS) 3000ml NOMINAL SIZE CONTAINER</p> <p>Baxter</p> <p>Dianeal® PD-2 Peritoneal Dialysis Solution With 2.5% Dextrose</p> <p>EACH 100ml CONTAINS 2.5 g DEXTROSE HYDROUS USP 538 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 20.7 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L SODIUM -132 CALCIUM -3.5 MAGNESIUM -0.5 CHLORIDE 96 LACTATE 40 OSMOLALITY 366 mOsmol/L(CALC) STERILE NON PYROGENIC</p> <p>POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</p> <p>READ PACKAGE INSERT FOR FULL INFORMATION FOR INTRAPERITONEAL ADMINISTRATION ONLY</p> <p>CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND DO NOT USE UNLESS SOLUTION IS CLEAR DISCARD UNUSED PORTION</p> <p>STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (UNDER 25°C) UNTIL READY TO USE AVOID EXCESSIVE HEAT SEE INSERT</p> <p>請儲存於攝氏25度以下</p> <p>Ambu-Flex™ III CONTAINER PL-146® PLASTIC</p> <p>MANUFACTURED BY BAFTER HEALTHCARE (GUANGZHOU) CO LTD GUANGZHOU CHINA (AN AFFILIATE OF BAXTER WORLD TRADE INC USA) HK-42533</p> <p>DIRECTIONS TO BE USED AS DIRECTED BY THE PHYSICIAN</p> <p>用法 請依照醫生指示使用</p> <p>Prescription Drug 處方藥物</p> <p>Manufacturer Address: Jiaoyan Road, Dongji Industrial District, GETDD,Guangzhou, P.R. China</p>  <p>01100000000000000000000000000000</p> <p>L5B9727 ① ② 2000 ml (APPROX 80 ml EXCESS) 3000 ml, NOMINAL SIZE CONTAINER</p> <p>Baxter</p> <p>Dianeal Low Calcium (2.5 mEq/L) Peritoneal Dialysis Solution with 2.5% Dextrose</p> <p>EACH 100 mL CONTAINS 2.5 g DEXTROSE HYDROUS USP 538 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L SODIUM -132 CALCIUM -3.5 MAGNESIUM -0.5 CHLORIDE -96 LACTATE -48 OSMOLALITY -366 mOsmol/L(CALC) STERILE NONPYROGENIC</p> <p>POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</p> <p>SEE PACKAGE INSERT FOR DOSAGE INFORMATION</p> <p>USE AS DIRECTED BY PHYSICIAN</p> <p>FOR INTRAPERITONEAL ADMINISTRATION ONLY</p> <p>CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND DO NOT USE UNLESS SOLUTION IS CLEAR DISCARD UNUSED PORTION Rx ONLY</p> <p>STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (25°C/77°F) UNTIL READY TO USE AVOID EXCESSIVE HEAT SEE INSERT</p> <p>Ambu-Flex II CONTAINER PL-146 PLASTIC</p> <p>BAFTER DIANEAL AMBU-FLEX II AND PL-146 ARE TRADEMARKS OF BAXTER INTERNATIONAL INC</p> <p>BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA</p> <p>Low Calcium 2.5% Dextrose</p> <p>L5B5177 ① ② 2000 ml (APPROX 80 ml EXCESS) 3000 ml, NOMINAL SIZE CONTAINER</p> <p>Baxter</p> <p>Dianeal PD-2 Peritoneal Dialysis Solution with 2.5% Dextrose</p> <p>EACH 100 mL CONTAINS 2.5 g DEXTROSE HYDROUS USP 538 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 20.7 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH 5.2 (4.5 TO 6.5) mEq/L SODIUM -132 CALCIUM -3.5 MAGNESIUM -0.5 CHLORIDE -96 LACTATE -48 OSMOLALITY -366 mOsmol/L(CALC) STERILE NONPYROGENIC</p> <p>POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN</p> <p>SEE PACKAGE INSERT FOR DOSAGE INFORMATION</p> <p>USE AS DIRECTED BY PHYSICIAN</p> <p>FOR INTRAPERITONEAL ADMINISTRATION ONLY</p> <p>CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND DO NOT USE UNLESS SOLUTION IS CLEAR DISCARD UNUSED PORTION Rx ONLY</p> <p>STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (25°C/77°F) UNTIL READY TO USE AVOID EXCESSIVE HEAT SEE INSERT</p> <p>Ambu-Flex II CONTAINER PL-146 PLASTIC</p> <p>BAFTER DIANEAL AMBU-FLEX II AND PL-146 ARE TRADEMARKS OF BAXTER INTERNATIONAL INC</p> <p>BAXTER HEALTHCARE CORPORATION DEERFIELD IL 60015 USA MADE IN USA</p> <p>PD-2 2.5% Dextrose</p>		

PACKAGE/LABEL PRINCIPAL DISPLAY PANEL

6AB5177E

2000ml

(APPROX 80ml EXCESS)
3000ml NOMINAL SIZE CONTAINER

Baxter

**Dianeal® PD-2
Peritoneal Dialysis Solution
With 2.5% Dextrose**

EACH 100ml CONTAINS 2.5 g DEXTROSE HYDROUS USP 538 mg
SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 25.7 mg CALCIUM
CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH5.2 (4.5 TO 6.5)
mEq/L SODIUM 132 CALCIUM 3.5 MAGNESIUM 0.5 CHLORIDE 96
LACTATE 40 **OSMOLARITY** 396 mOsmol/L(CALC)
STERILE NON PYROGENIC

POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE
DIRECTION OF A PHYSICIAN

READ PACKAGE INSERT FOR FULL INFORMATION
FOR INTRAPERITONEAL ADMINISTRATION ONLY

CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH
MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND
DO NOT USE UNLESS SOLUTION IS CLEAR
DISCARD UNUSED PORTION

STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM
TEMPERATURE (UNDER 25°C) UNTIL READY TO USE
AVOID EXCESSIVE HEAT SEE INSERT

請儲存於攝氏25度以下

Ambu-Flex™ III CONTAINER PL-146® PLASTIC

MANUFACTURED BY
BAXTER HEALTHCARE (GUANGZHOU) CO LTD
GUANGZHOU CHINA
(AN AFFILIATE OF BAXTER WORLD TRADE INC USA)

HK-42533
DIRECTIONS TO BE USED AS DIRECTED BY THE PHYSICIAN

用法 請依照醫生指示使用

Prescription Drug 處方藥物

Manufacturer Address:

Jiaoyuan Road, Dongji Industrial District,
GETDD,Guangzhou, P.R. China

6AB5177E 2000ml
(APPROX 80ml EXCESS)
3000ml NOMINAL SIZE CONTAINER

BaxterLogo

**Dianeal® PD-2
Peritoneal Dialysis Solution
With 2.5% Dextrose**

EACH 100ml CONTAINS 2.5 g DEXTROSE HYDROUS USP 538 mg
SODIUM CHLORIDE USP 448 mg SODIUM LACTATE 25.7 mg CALCIUM
CHLORIDE USP 5.08 mg MAGNESIUM CHLORIDE USP pH5.2 (4.5 to 6.5)
mEq/L SODIUM 132 CALCIUM 3.5 MAGNESIUM 0.5 CHLORIDE 96
LACTATE 40 **OSMOLARITY** 396 mOsmol/L(CALC)
STERILE NON PYROGENIC

POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE
DIRECTION OF A PHYSICIAN

READ PACKAGE INSERT FOR FULL INFORMATION
FOR INTRAPERITONEAL ADMINISTRATION ONLY
CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH
MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND
DO NOT USE UNLESS SOLUTION IS CLEAR
DISCARD UNUSED PORTION

STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM
TEMPERATURE (UNDER 25°C) UNTIL READY TO USE
AVOID EXCESSIVE HEAT SEE INSERT

Ambu-Flex™ III CONTAINER PL-146 ® PLASTIC

MANUFACTURED BY
BAXTER HEALTHCARE (GUANGZHOU) CO LTD
GUANGZHOU CHINA
(AN AFFILIATE OF BAXTER WORLD TRADE INC USA)

HK-42533

DIRECTIONS TO BE USED AS DIRECTED BY THE PHYSICIAN

Prescription Drug

Manufacturer Address:

Jiaoyuan Road, Dongji Industrial District,
GETDD, Guangzhou, P.R. China

2.5 PD-2 WITH 2.5% DEXTROSE 2000mlX6
2.5 LOT G00000000 EXP JAN 00

6AB5177E
S/N 0000

2.5 PD-2 WITH 2.5% DEXTROSE 2000mlX6

LOT G00000000 EXP JAN 00

6AB5177E

S/N 0000

6AB9747E

2000ml

(APPROX 80ml EXCESS)
3000ml NOMINAL SIZE CONTAINER

Baxter

**Dianeal® Low Calcium(2.5mEq/L)
Peritoneal Dialysis Solution
With 4.25% Dextrose**

EACH 100ml CONTAINS 4.25 g DEXTROSE HYDROUS USP
538 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE
18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM
CHLORIDE USP pH 5.2 (4.5 TO 6.5)
mEq/L SODIUM 132 CALCIUM 2.5 MAGNESIUM 0.5
CHLORIDE 95 LACTATE 40
OSMOLARITY 483 mOsmol/L(CALC)
STERILE NON PYROGENIC

POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE
DIRECTION OF A PHYSICIAN

WARNING EXTENSIVE USE OF THIS SOLUTION DURING
ONE PERITONEAL DIALYSIS PROCEDURE CAN RESULT IN
SIGNIFICANT REMOVAL OF WATER FROM THE PATIENT
READ PACKAGE INSERT FOR FULL INFORMATION
FOR INTRAPERITONEAL ADMINISTRATION ONLY

CAUTIONS SQUEEZE AND INSPECT INNER BAG WHICH
MAINTAINS PRODUCT STERILITY DISCARD IF
LEAKS ARE FOUND

DO NOT USE UNLESS SOLUTION IS CLEAR
DISCARD UNUSED PORTION
STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM
TEMPERATURE (UNDER 25°C) UNTIL READY TO USE
AVOID EXCESSIVE HEAT SEE INSERT

請儲存於攝氏25度以下

Ambu-Flex™ III CONTAINER PL-146®PLASTIC

MANUFACTURED BY
BAXTER HEALTHCARE (GUANGZHOU) CO LTD
GUANGZHOU CHINA
(AN AFFILIATE OF BAXTER WORLD TRADE INC USA)

HK-42532

DIRECTIONS TO BE USED AS DIRECTED BY THE PHYSICIAN

用法 請依照醫生指示使用

Prescription Drug 處方藥物

Manufacturer Address:

Jiaoyuan Road, Dongji Industrial District,
GETDD,Guangzhou, P.R. China

6AB9747E 2000ml
(APPROX 80ml EXCESS)
3000ml NOMINAL SIZE CONTAINER

BaxterLogo

**Dianeal® Low Calcium(2.5mEq/L)
Peritoneal Dialysis Solution
With 4.25% Dextrose**

EACH 100ml CONTAINS 4.25 g DEXTROSE HYDROUS USP
538 mg SODIUM CHLORIDE USP 448 mg SODIUM LACTATE
18.3 mg CALCIUM CHLORIDE USP 5.08 mg MAGNESIUM
CHLORIDE USP pH 5.2 (4.5 to 6.5)

Low Calcium 4.25% Dextrose

mEq/LSODIUM 132 CALCIUM 2.5 MAGNESIUM 0.5

CHLORIDE 95 LACTATE 40

OSMOLARITY483 mOsmol/L(CALC)

STERILE NON PYROGENIC

POTASSIUM CHLORIDE TO BE ADDED ONLY UNDER THE DIRECTION OF A PHYSICIAN

WARNINGEXTENSIVE USE OF THIS SOLUTION DURING ONE PERITONEAL DIALYSIS PROCEDURE CAN RESULT IN SIGNIFICANT REMOVAL OF WATER FROM THE PATIENT
READ PACKAGE INSERT FOR FULL INFORMATION FOR INTRAPERITONEAL ADMINISTRATION ONLY

CAUTIONSSQUEEZE AND INSPECT INNER BAG WHICH MAINTAINS PRODUCT STERILITY DISCARD IF LEAKS ARE FOUND
DO NOT USE UNLESS SOLUTION IS CLEAR
DISCARD UNUSED PORTION
STORE UNIT IN MOISTURE BARRIER OVERWRAP AT ROOM TEMPERATURE (UNDER 25°C) UNTIL READY TO USE
AVOID EXCESSIVE HEAT SEE INSERT

Ambu-Flex™ III CONTAINER PL-146 ®PLASTIC

MANUFACTURED BY

BAXTER HEALTHCARE (GUANGZHOU) CO LTD

GUANGZHOU CHINA

(AN AFFILIATE OF BAXTER WORLD TRADE INC USA)

HK-42532

DIRECTIONSTO BE USED AS DIRECTED BY THE PHYSICIAN

Prescription Drug

Manufacturer Address:

Jiaoyuan Road, Dongji Industrial District,

GETDD, Guangzhou, P.R. China

4.25 LOW CALCIUM WITH 4.25% DEXTROSE 2000mlX6
4.25 LOT G00000000 EXP JAN 00 6AB9747E S/N 0000

4.25 LOW CALCIUM WITH 4.25% DEXTROSE 2000mlX6

LOT G00000000 EXP JAN 00

6AB9747E

S/N 0000

DIANEAL LOW CALCIUM WITH DEXTROSE

sodium chloride, sodium lactate, calcium chloride, magnesium chloride and dextrose injection, solution

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0941-0696
Route of Administration	INTRAPERITONEAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DEXTROSE MONOHYDRATE (UNII: LX22YL083G) (ANHYDROUS DEXTROSE - UNII:5SL0G7R0OK)	DEXTROSE MONOHYDRATE	4.25 g in 100 mL
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37, CHLORIDE ION - UNII:Q32ZN48698)	SODIUM CHLORIDE	538 mg in 100 mL
SODIUM LACTATE (UNII: TU7HW0W0QT) (SODIUM CATION - UNII:LYR4M0NH37, LACTIC ACID, UNSPECIFIED FORM - UNII:33X04XA5AT)	SODIUM LACTATE	448 mg in 100 mL
CALCIUM CHLORIDE (UNII: M4I0D6VV5M) (CALCIUM CATION - UNII:2M83C4R6ZB, CHLORIDE ION - UNII:Q32ZN48698)	CALCIUM CHLORIDE	18.3 mg in 100 mL
MAGNESIUM CHLORIDE (UNII: 02F3473H9O) (MAGNESIUM CATION - UNII:T6V3LHY838, CHLORIDE ION - UNII:Q32ZN48698)	MAGNESIUM CHLORIDE	5.08 mg in 100 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0941-0696-06	6 in 1 CARTON	10/30/2024	
1	NDC:0941-0696-01	2000 mL in 1 BAG; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
Unapproved drug for use in drug shortage		10/30/2024	

DIANEAL PD-2 WITH DEXTROSE

sodium chloride, sodium lactate, calcium chloride, magnesium chloride and dextrose injection, solution

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0941-0698
Route of Administration	INTRAPERITONEAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SODIUM LACTATE (UNII: TU7HW0W0QT) (SODIUM CATION - UNII:LYR4M0NH37, LACTIC ACID, UNSPECIFIED FORM - UNII:33X04XA5AT)	SODIUM LACTATE	448 mg in 100 mL
CALCIUM CHLORIDE (UNII: M4I0D6VV5M) (CALCIUM CATION - UNII:2M83C4R6ZB, CHLORIDE ION - UNII:Q32ZN48698)	CALCIUM CHLORIDE	25.7 mg in 100 mL
MAGNESIUM CHLORIDE (UNII: 02F3473H9O) (MAGNESIUM CATION - UNII:T6V3LHY838, CHLORIDE ION - UNII:Q32ZN48698)	MAGNESIUM CHLORIDE	5.08 mg in 100 mL
DEXTROSE MONOHYDRATE (UNII: LX22YL083G) (ANHYDROUS DEXTROSE - UNII:5SL0G7R0OK)	DEXTROSE MONOHYDRATE	2.5 g in 100 mL
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37, CHLORIDE ION - UNII:Q32ZN48698)	SODIUM CHLORIDE	538 mg in 100 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0941-0698-06	6 in 1 CARTON	10/30/2024	
1	NDC:0941-0698-01	2000 mL in 1 BAG; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
Unapproved drug for use in drug shortage		10/30/2024	

Labeler - Baxter Healthcare Corporation (005083209)

Establishment

Name	Address	ID/FEI	Business Operations
Baxter Healthcare (Guangzhou) Co., Ltd		421040114	analysis(0941-0696, 0941-0698) , label(0941-0696, 0941-0698) , manufacture(0941-0696, 0941-0698) , pack(0941-0696, 0941-0698) , sterilize(0941-0696, 0941-0698)