

### **IMPORTANT PRESCRIBING INFORMATION**

DATE: October 22, 2024

# Subject: Temporary importation of EXTRANEAL (icodextrin) Peritoneal Dialysis Solution from Ireland for use in Continuous Ambulatory Peritoneal Dialysis to address drug shortages

Dear Healthcare Professional,

Due to the current critical shortage of EXTRANEAL (icodextrin) 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 EXTRANEAL from Baxter's manufacturing facility in Castlebar, Ireland. FDA has not approved this product manufactured by Baxter's Castlebar, Ireland facility.

You may be provided with additional letters for other 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 EXTRANEAL (icodextrin) Peritoneal Dialysis Solution for use in Continuous Ambulatory Peritoneal Dialysis (CAPD) therapy as described in the table below. This product is manufactured by Baxter's manufacturing facility in Castlebar, Ireland and is marketed in the United Kingdom and other countries within the European Union (EU). At this time, importation or distribution of EXTRANEAL (icodextrin) peritoneal dialysis solution 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 Castlebar, Ireland:

Product Name and Description	CAPD Fill Volume	Product Code	Bags per Carton	NDC Code
EXTRANEAL (icodextrin) Solution for Peritoneal Dialysis	2000 mL	FPB5268C	5 bags	0941-0717-01 (bag) 0941-0717-05 (carton)
EXTRANEAL (icodextrin) Solution for Peritoneal Dialysis	2500 mL	FPB5270	4 bags	0941-0681-01 (bag) 0941-0681-04 (carton)

#### It is important to note the following:

 There are no clinically relevant differences in the EXTRANEAL drug composition between the European-manufactured and U.S.-manufactured CAPD product (see Table 1). As such, clinical practice for usage, administration, and dosage for Extraneal with 7.5% icodextrin (manufactured in EU) products is the same as with the Extraneal with 7.5% icodextrin (manufactured in U.S.). Please refer to the FDA-approved EXTRANEAL (icodextrin) Peritoneal Dialysis Solution Prescribing Information for reference.



- Calcium and Magnesium electrolyte concentrations are identical in EXTRANEAL manufactured in the EU and U.S. but appear different as they are expressed in mmol/L (EU) and in mEq/L (U.S.).
- The Luer-lock connector on the Ireland imported product functions the same and is fully compatible with peritoneal dialysis sets marketed in the United States. Ireland imported product has a single green frangible near the Y connector which is broken at the same time during flush phase as the corresponding blue frangible in U.S.-manufactured product. The Ireland imported product has solution in the infusion line whereas U.S. manufactured product has a dry line. See Table 1 for more details of product differences.
- EXTRANEAL (icodextrin) solution for peritoneal dialysis imported carton labeling includes barcodes; however, the barcodes may not register accurately in the U.S. scanning systems. There are no barcodes on the solution containers of the Ireland 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.

Before prescribing, healthcare providers should be aware of some key differences in the container packaging and labeling between the EXTRANEAL (icodextrin) Solution for Peritoneal Dialysis products (manufactured in EU) and EXTRANEAL (icodextrin) Peritoneal Dialysis Solution (manufactured in U.S.).

Key differences are highlighted in the following Product Comparison Tables:

- Table 1: Key differences of EXTRANEAL for CAPD therapy
- Table 2: Label images of EXTRANEAL for 2000 mL and 2500 mL CAPD product presentations

#### **Reporting Adverse Events**

To report adverse events associated with the imported product, please call Baxter at 1-866-888-2472, or fax: 1-800-759-1801. Adverse events or quality problems experienced with the use of the imported product 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:** <u>www.fda.gov/medwatch/report.htm</u>
- **Regular mail or Fax:** Download form <a href="http://www.fda.gov/MedWatch/getforms.htm">http://www.fda.gov/MedWatch/getforms.htm</a> 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: <u>Baxter - Product Feedback Portal (https://productfeedback.baxter.com/)</u>.

# Please refer to the FDA approved prescribing information for EXTRANEAL (icodextrin) Peritoneal Dialysis Solution at <u>DailyMed (nih.gov)</u>.

If you have any questions about the information contained in this letter or the use of imported EXTRANEAL (icodextrin) Solution for Peritoneal Dialysis, 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 Geovana Basso Reason: Ok. Date: Oct 22, 2024 08:52 CDT

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

Baxter, EXTRANEAL and ULTRABAG (or UltraBag) are registered trademarks of Baxter International Inc.

Attachments:

Product Comparison Tables 1 and 2



	Imported Product from Castlebar, Ireland	US FDA Approved Product					
Product name	EXTRANEAL (icodextrin) Solution for Peritoneal Dialysis	EXTRANEAL (icodextrin) Peritoneal Dialysis Solution					
Labeled Fill Volume	2000 mL 2500 mL	2000 mL 2500 mL					
Container Type	TwinBag (PVC)	UltraBag Container (PVC)					
Bags per carton	2000 mL: 5 bags 2500 mL: 4 bags	2000 mL: 6 bags 2500 mL: 5 bags					
Indications	Extraneal is recommended as a once daily replacement for a single glucose exchange as part of a continuous ambulatory peritoneal dialysis (CAPD) or automated peritoneal dialysis (APD) regimen for the treatment of chronic renal failure, particularly for patients who have lost ultrafiltration on glucose solutions, because it can extend time on CAPD therapy in such patients.	EXTRANEAL (icodextrin) is indicated for a single daily exchange for the long (8- to 16- hour) dwell during continuous ambulatory peritoneal dialysis (CAPD) or automated peritoneal dialysis (APD) for the management of kidney failure in patients requiring long- term kidney replacement therapy. EXTRANEAL is also indicated to improve (compared to 4.25% dextrose) long-dwell ultrafiltration and clearance of creatinine and urea nitrogen in patients with high average or greater transport characteristics, as defined using the peritoneal equilibration test (PET)					
Active Ingredients	75 g/L Icodextrin (7.5 g/100 mL) 5.4 g/L Sodium Chloride (540 mg/100 mL) 4.5 g/L Sodium Lactate (450 mg/100 mL) 0.257 g/L Calcium Chloride (25.7 mg/100 mL) 0.051 g/L Magnesium Chloride (5.1 mg/100 mL)	7.5 g/100 mL lcodextrin 535 mg/100 mL Sodium Chloride, USP* 448 mg/100mL Sodium Lactate* 25.7 mg/100mL Calcium Chloride, USP* 5.08 mg/100 mL Magnesium Chloride, USP* * considered excipients in US drug registration					
Electrolyte Content per Liter	Sodium 133 mmol/L (equivalent to 133 mEq/L) Calcium 1.75 mmol/L (equivalent to 3.5 mEq/L) Magnesium 0.25 mmol/L (equivalent to 0.5 mEq/L) Chloride 96 mmol/L (equivalent to 96 mEq/L) Lactate 40 mmol/L (equivalent to 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					
рН	pH 5.0 – 6.0 HCl / NaOH may have been used to adjust pH	pH 5.0 – 6.0 HCl / NaOH may have been used to adjust pH					
Additional Information	Osmolarity 284 mOsm/L	Osmolarity (Calc) 282 – 286 mOsmol/L					
Storage Conditions	Do not store below 4°C	Store at 20–25°C (68–77°F). Excursions permitted to 15–30°C (59–86°F) [See USP Controlled Room Temperature]. Protect from freezing.					
Expiration Dating	24 months	18 months					
Container Closure System							
Container Closure Differences	Closure broken in Flush phase green near solution bag						



### Table 2. Comparison of EXTRANEAL (icodextrin) PD Solution Container Labels

A comparison of labels is provided below. Note, both Baxter and Vantive-branded labels are presented, which represent all imported lot labeling scenarios during this transition.

Imported Product from Castlebar, Ireland			US FDA Approved Product			
B5268		EXP 2000 ml	5 <b>B4984</b> NDC 0941-0679-52	Θ		2000 mL 30 mL EXCESS)
Baxter			Baxter			
EXTRANEAL Solution for Peritoneal Dialysi	is					
Formula 1000 ml		mmol/l	Extraneal (i			
Icodextrin       75 g         Sodium Chloride       5,4 g         Sodium (S)-lactate       4,5 g         Calcium Chloride 2H <sub>2</sub> 0       0,257 g         Magnesium Chloride 6H <sub>2</sub> 0       0,051 g         Water for Injections       Hydrochloric acid or sodium hydroxide (for pH adjustment)         For intraperitoneal use       Not for intravenous use         Use as directed by a physician       Read the package leaflet before use         Keep out of the sight and reach of children       Nonpyrogenic         Do not use unless solution is clear and container undamaged       For single use only         Once removed from the overpouch use immediately       Discard unused solution         UK:       IE and MT:         Baxter Healthcare Ltd       Kobaltweg 49, 3542C	Ca <sup>++</sup> Mg <sup>++</sup> Ci <sup>-</sup> C <sub>3</sub> H <sub>5</sub> O <sub>3</sub> <sup>-</sup> Osmolarity pH	133 1,75 0,25 96 40 284 m0sm/l 5,0 - 6,0	Peritoneal I EACH 100 mL CONTAIN SODIUM CHLORIDE US 25.7 mg CALCIUM CHLO CHLORIDE USP WAT mEq/L SODIUM 132 CHLORIDE 96 LACT) PH 5.0 - 6.0 pH MAY 1 HYDROCHLORIC ACID EXTRANEAL SOLUTION ANTIMICROBIAL AGEN OSMOLARITY (CALC) 20 STERILE NONPYRO POTASSIUM CHLOI UNDER THE DIREC SEE PACKAGE INSER USE AS DIRECTED BY FOR INTRAPERITOR	Dialysis S S S S S S S S S S S S S S	CIUITION TRIN 535 mg JM LACTATE mg MAGNESIUM N USP MAGNESIUM 0.5 STED WITH DXIDE CTERIOSTATIC OF ED ONLY ICIAN IFORMATION RATION ONLY T INNER BAG	7.5% icodextri
PL 00116/0266     PA 2299/017/001       MA1277/01101       POM       Manufactured by Baxter Healthcare S A IRL-Castlebar CB-35-04-608			DO NOT USE UNLESS DISCARD UNUSED PO Rx ONLY STORE IN MOISTURE I CARTON UNTIL READ STORE AT 20-25°C (66 PERMITTED TO 15-30° CONTROLLED ROOM FROM FREEZING UltraBag CONTAIN BAXTER EXTRANEAL TRADEMARKS OF BA BAXTER HEALTHCAF DEERFIELD IL 60015 US MADE IN USA US PAT NOS 4761237 6077836	ARRIER OVERPO Y TO USE 3-77°F) EXCURS C (59-86°F) [SE TEMPERATURE] NER F L ULTRABAG AND XTER INTERNATION RE CORPORATION SA 4573980 488670	DUCH IN SIONS E USP PROTECT PL 146 PLASTIC D PL 146 ARE DNAL INC N	PD-2



Imported Product from ( 15268 Vantive					pproved Product	
			2000 ml			
XTRANEAL Solution for Peritone	eal Dialysi	S				
ormula 1000 ml			mmol/I			
odextrin	75 g	Na+	133			
odium Chloride	5,4 g	Ca++	1,75			
odium (S)-lactate	4,5 g		0,25			
alcium Chloride 2H <sub>2</sub> 0	0,257 g		96			
agnesium Chloride 6H20	0,051 g		40			
ater for Injections /drochloric acid or sodium hydroxide (for pH adjustment)		Osmolarity pH	284 m0sm/l 5,0 – 6,0			
or intraperitoneal use ot for intravenous use sea sa directed by a physician ead the package leaflet before use sep out of the sight and reach of children onpyrogenic o not store below 4°C o not use unless solution is clear and container undamaged or single use only nee removed from the overpouch use immediately iscard unused solution antive Limited favertree Technology Park Wavertree Boulevard verpool, L7 9PE nited Kingdom _ 58711/0005						
OM			CB-35-05-250			
	RSION					

## Baxter

Imported Product from Castlebar, Ireland			US FDA Approved Product					
B5270 Baxter EXTRANEAL Solution for Peritone Formula 1000 ml		EXP 2500 ml mmol/I 133	0	5B4986 NDC 0941-0679-53 <b>Baxter</b> Extraneal (ic Peritoneal D EACH 100 mL CONTAINS	© codextrin lialysis S	© (APPROX ) olution	<b>2500</b> mL 90 mL EXCESS)	0
Sodium Chloride Sodium (S)-lactate Calcium Chloride 2H <sub>2</sub> O Magnesium Chloride 6H <sub>2</sub> O Water for Injections Hydrochloric acid or sodium hydroxide (for pH adjustment) For intraperitoneal use Not for intravenous use Use as directed by a physician Read the package leaflet before use Keep out of the sight and reach of children Nonpyrogenic Do not sure below 4°C Do not use unless solution is clear and container undamaged For single use only Once removed from the overpouch use immediately Discard unused solution UK: Baxter Healthcare Ltd Caxton Way,Thefford, Norfolk, IP24 3SE, United Kingdom IE and MT: Baxter Holding B.V., Kobaltweg 49, 3542CE, Utrecht, Netherlands PL 00116/0266 PA 2299/017/001 MA 1277/01101 [PDM] Manufactured by Baxter Healthcare S A IRL-Castlebar CB-35-04-609	5,4 g 4,5 g 0,257 g Cr 0,051 g C <sub>3</sub> H <sub>5</sub> O <sub>3</sub> <sup>-</sup> 0smolarity pH	1,75 0,25 96 40 284 m0sm/l 5,0 - 6,0	0	SODIUM CHLORIDE USS 25.7 mg CALCUM CHLO CHLORIDE USP WATE mEq/L SODIUM 132 CHLORIDE DE LACTA PH 50-6.0 PH MAY H VH7DRCHLORIC ACID O EXTRANEAL SOLUTION ANTIMICROBIAL AGENT GOMOLARITY (CALC) 28 STERILE NONPYROC POTASSIUM CHLORI THE DIRECTION OF / SEE PACKAGE INSERT USE AS DIRECTED BY FOR INTRAPERITON CAUTIONS SOUE THAT MAINTAINS PROC LACAS AS DIRECTED BY FOR INTRAPERITON CAUTIONS SOUE THAT MAINTAINS PROC DISCARD UNUSED POR RX ONLY STORE IN MOISTURE B UNITIL READY TO USE OF TEMPERATURE) PRC UNITIL BEADY TO ZO 25°C (68- 00°F) TEMPERATURE) PRC UNITIL BEADY TO ZO 25°C (68- TO 15-30°C (68-80°F) TEMPERATURE) PRC UNITIL BEADY TO ZO 25°C (68- DISCARD UNUSED POR BAXTER EXTRANEAL TRADEMARKS OF BA BAXTER EXTRANEAL TRADEMARKS OF DA BAXTER EXTRANEAL TRADEMARKS OF DA BAXTER EXTRANEAL	Atá mg son) Atá mg son) R FOR INJECTIO CALCIUM 3.5 TE 40 AVE BEEN ADJUE TE 40 AVE BEEN ADJUE R SODIUM HYDR CONTAINS NO B/ S 2286 mOsmol/L ENIC 2286 mOsmol/L ENIC 2286 mOsmol/L ENIC DE TO BE ADD A PHYSICIAN EAL ADMINIST EZE AND INSPEC EXELUTION IS CL RITION ARRRIER OVERP T7°F) EXCUR: TOTON FR ER ULTRABAG /A XTER INTERNA E CORPORATIO A	UM LACTATE <sup>®</sup> mg MAGNESIUM N USP MAGNESIUM 0.5 STED WITH OXIDE ED ONLY UNDER ED ONLY UNDER NFORMATION INFORMATION	PD-2 7.5% icodextrin	0



Imported Product from	Castlebar, Ire	eland			US	FDA Approved	Product
Lot			EXP				
B5270			2500 ml				
Vantive							
EXTRANEAL Solution for Peritone	al Dialysis						
Formula 1000 ml			mmol/l	]			
Icodextrin	75 g	Na+	133				
Sodium Chloride	5,4 g	Ca++	1,75				
Sodium (S)-lactate	4,5 g	Mg <sup>++</sup>	0,25				
Calcium Chloride 2H20	0,257 g	CI-	96				
Magnesium Chloride 6H20	0,051 g	C3H503	40				
Water for Injections		Osmolarity	284 m0sm/l				
Hydrochloric acid or sodium hydroxide (for pH adjustment)		pH	5,0-6,0				
For intraperitoneal use Not for intravenous use Use as directed by a physician Read the package leaflet before use Keep out of the sight and reach of children Nonpyrogenic Do not store below 4°C Do not use unless solution is clear and container undamaged For single use only Once removed from the overpouch use immediately Discard unused solution Vantive Limited Wavertree Technology Park 2 Wavertree Boulevard Liverpool, L7 9FE United Kingdom PL 58711/0005							
РОМ							
			CB-35-05-251	]			
VANTIVE V	ERSION						