HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use PRISMASOL and PHOXILLUM safely and effectively. See full prescribing information for PRISMASOL and PHOXILLUM.

PRISMASOL renal replacement solution
PRISMASOL Initial U.S. Approval: 2006

PHOXILLUM renal replacement solution
PHOXILLUM Initial U.S. Approval: 2015

-------------------RECENT MAJOR CHANGES-------------------
Contraindications (4) 07/2016
Warnings and Precautions (5.1, 5.2) 06/2017

-------------------INDICATIONS AND USAGE-------------------
PRISMASOL and PHOXILLUM solutions are indicated:
• As a replacement solution in Continuous Renal Replacement Therapy (CRRT) and in case of drug poisoning when CRRT is used to remove dialyzable substances (1)

-------------------DOSAGE AND ADMINISTRATION-------------------
• Therapy must be individualized based on the patient’s clinical condition, fluid, electrolyte, acid-base and glucose balance (2.2)
• Solution must be mixed prior to use (2.2)
• Use only with extracorporeal dialysis equipment appropriate for CRRT (2.3)

-------------------DOSAGE FORMS AND STRENGTHS-------------------
PRISMASOL and PHOXILLUM are available in multiple combinations of ingredients and in multiple variations of strengths. See full Prescribing Information for detailed descriptions of each formulation. (2, 3, 11)

-------------------CONTRAINDICATIONS-------------------
• Known hypersensitivities to PRISMASOL and PHOXILLUM solutions (4)

-------------------WARNINGS AND PRECAUTIONS-------------------
• Monitor hemodynamic status and fluid inputs and outputs, potassium, phosphorus, other electrolytes and acid-base balance. Abnormalities may be corrected by the use of appropriate formulations and dosage of PRISMASOL and PHOXILLUM solutions (5.1)
• Treatment may affect glucose levels. Monitor blood glucose levels.
• Antidiabetic therapy adjustment or other corrective measures may be required during treatment (5.2)

To report SUSPECTED ADVERSE REACTIONS, contact Baxter Healthcare Corporation at 1-866-888-2472 or FDA 1-800-FDA-1088 or www.fda.gov/medwatch
Revised: 06/2017

FULL PRESCRIBING INFORMATION: CONTENTS*
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2 DOSAGE AND ADMINISTRATION
  2.1 Administration Instructions
  2.2 Dosing Considerations
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3 DOSAGE FORMS AND STRENGTHS
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*Sections or subsections omitted from the full prescribing information are not listed.
FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

PRISMASOL and PHOXILLUM solutions are indicated in pediatric and adult patients for use as a replacement solution in Continuous Renal Replacement Therapy (CRRT) to replace plasma volume removed by ultrafiltration and to correct electrolyte and acid-base imbalances. They may also be used in case of drug poisoning when CRRT is used to remove dialyzable substances.

2 DOSAGE AND ADMINISTRATION

2.1 Administration Instructions

Visually inspect PRISMASOL and PHOXILLUM for particulate matter and discoloration prior to administration.

Administration should only be under the direction of a physician competent in intensive care treatment including CRRT. Use only with extracorporeal dialysis equipment appropriate for CRRT.

The prepared solution is for single patient use only.

Aseptic technique should be used throughout administration to the patient.

Discard any unused solution.

2.2 Dosing Considerations

PRISMASOL replacement solutions contain 4 different combinations of active ingredients (8 different products with varying ingredient amounts). PHOXILLUM replacement solutions contain 2 different combinations of active ingredients (2 different products with varying ingredient amounts). PRISMASOL and PHOXILLUM are supplied in a two-compartment bag that must be mixed immediately prior to use [see Dosage and Administration (2.3)]:

• Small compartment A (250 mL) containing an electrolyte solution, and
• Large compartment B (4750 mL) containing the buffer solution.

See Table 1 for the concentrations of the active ingredients (after mixing) in these 10 different replacement solutions (total volume is 5 Liters).
Table 1: Concentrations of Active Ingredients in the 8 PRISMASOL and 2 PHOXILLUM Replacement Solutions after Mixing

<table>
<thead>
<tr>
<th>PRISMASOL Replacement Solutions</th>
<th>Ca$^{2+}$ mEq/L</th>
<th>HCO$_3^-$ mEq/L</th>
<th>K$^+$ mEq/L</th>
<th>Mg$^{2+}$ mEq/L</th>
<th>Na$^+$ mEq/L</th>
<th>HPO$_4^{2-}$ mmol/L</th>
<th>Cl$^-$ mEq/L</th>
<th>Lactate mEq/L</th>
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<table>
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<th>Mg$^{2+}$ mEq/L</th>
<th>Na$^+$ mEq/L</th>
<th>HPO$_4^{2-}$ mmol/L</th>
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Ca$^{2+}$ = calcium, HCO$_3^-$ = bicarbonate, K$^+$ = potassium, Mg$^{2+}$ = magnesium, Na$^+$ = sodium, HPO$_4^{2-}$ = phosphate, Cl$^-$ = chloride; osmolarity is estimated.

The mode of therapy, solute formulation, flow rates, and length of PRISMASOL and PHOXILLUM replacement therapy in CRRT should be established by a physician based on the patient’s clinical condition, blood concentration of phosphate and other electrolytes, acid-base and glucose balance. Administer either PRISMASOL or PHOXILLUM into the extracorporeal circuit:

- Before (pre-dilution) the hemofilter or hemodiafilter,
- After (post-dilution) the hemofilter or hemodiafilter, or
- Before and after the hemofilter or hemodiafilter.

2.3 Preparing the Solution
Use only if the overwrap is not damaged, all seals are intact, frangible pin or peel seal is not broken, and the solution is clear.

The solution may be warmed to 37°C/98.6°F prior to removing the overwrap to enhance patient comfort. However, only dry heat should be used. Solutions should not be heated in water or in a microwave oven. After heating, verify that the solution remains clear and contains no particulate matter.

The solutions are supplied in two different two-compartment bags, one made of polyvinyl chloride with a frangible pin separating compartment A and B (see Figure 1) and one made of polyolefin with a peel seal separating compartment A and B (see Figure 6).

Follow the instructions below when connecting the solution bags for correct use of the access ports.
Instructions for preparing solutions supplied in a two-compartment, polyvinyl chloride (PVC) bag with a red frangible pin:

**Figure 1**

Step 1  Immediately before use, remove the overwrap from the bag and mix the solutions in the two different compartments. After removing the overwrap, inspect the container for leakage by pressing firmly on the bag. Discard the bag if any leakage is detected since sterility cannot be assured. As soon as the overwrap is removed, the reconstitution of compartments A and B should be done and the mixed solution should be used immediately.

After removal of the overwrap, the solution is stable for 24 hours including the duration of the treatment. Open the seal by breaking the red frangible pin between the two compartments of the bag. The frangible pin will remain in the bag. (See Figure 2 beside)

**Figure 2**

**Step 2**  Make sure all the fluid from the small compartment A is transferred into the large compartment B. (See Figure 3 beside)

**Figure 3**

**Step 3**  Rinse the small compartment A twice by pressing the mixed solution back into the small compartment A and then back into the large compartment B. (See Figure 3 beside)
**Figure 4**

**Step 4** When the small compartment A is empty, shake the large compartment B so that the contents mix completely. (See Figure 4 beside)
The solution is now ready to use and the bag can be hung on the equipment.

**Figure 5a**

**Step 5** The replacement line may be connected to the bag through either the luer connector or the injection connector (spike connector).

**Step 5a** The luer connector is a needle-less and swabbable connector. Remove the cap with a twist and pull motion, and connect the male luer lock on the replacement line to the female luer receptor on the bag. (See Figure 5a beside) Ensure that the connection is fully seated and tighten. The connector is now open. Verify that the fluid is flowing freely during use.

When the replacement line is disconnected from the luer connector, the connector will close and the flow of the solution will stop.

**Figure 5b**

**Step 5b** If the injection connector (spike connector) is used, first remove the snap-off cap. Then introduce the replacement line spike through the rubber septum of the bag connector. (See Figure 5b beside) Ensure that the spike is fully inserted and verify that the fluid is flowing freely during use.

**Instructions for preparing solutions supplied in a two-compartment, polyolefin bag with a peel seal:**

**Figure 6**
### Step 1
Immediately before use, remove the overwrap from the bag and mix the solutions in the two different compartments. After removing the overwrap, inspect the container for leakage by pressing firmly on the bag. Discard the bag if any leakage is detected since sterility cannot be assured. As soon as the overwrap is removed, the reconstitution of compartments A and B should be done and the mixed solution should be used immediately. After removal of the overwrap, the solution is stable for 24 hours including the duration of the treatment. Hold the small compartment with both hands and squeeze it until an opening is created in the peel seal. (See Figure 7 beside)

### Step 2
Squeeze with both hands on the large compartment until the peel seal between the two compartments is entirely open. Shake gently to mix. (See Figure 8 beside)

The solution is now ready to use and the bag can be hung on the equipment.

### Step 3
The replacement line may be connected to the bag through either of the luer connector or the injection connector (spike connector).

#### Step 3a
The luer connector is a needle-less and swabbable connector. Remove the cap with a twist and pull motion, and connect the male luer lock on the replacement line to the female luer receptor on the bag. (See Figure 9a beside)

Ensure that the connection is fully seated and tighten. The connector is now open. Verify that the fluid is flowing freely during use.

When the replacement line is disconnected from the luer connector, the connector will close and the flow of the solution will stop.

#### Step 3b
If the injection connector (spike connector) is used, first remove the snap-off cap. Then introduce the replacement line spike through the rubber septum of the bag connector. (See Figure 9b beside)

Ensure that the spike is fully inserted and verify that the fluid is flowing freely during use.

### 2.4 Adding Drugs to the Solutions
After mixing, additional drugs may be added to the bag via injection connector (spike connector) in large compartment B. In general, drugs other than phosphate should be administered through a different access line.

When introducing drugs, use aseptic techniques and mix thoroughly. Do not use if there is a color change and/or the appearance of precipitates, insoluble complexes or crystals after addition of medication.

**Phosphate:** Up to 1.2 mmol/L of phosphate can be added to the bag as potassium phosphate or sodium phosphate. The total potassium concentration of PRISMASOL solution should not exceed 4 mEq/L. Use sodium phosphate to add phosphate if the total potassium concentration in PRISMASOL solution is 4...
mEq/L.

**PHOXILLUM Solutions:**
*Phosphate:* Phosphate up to 0.2 mmol/L may be added to the solution. Use sodium phosphate if adding phosphate to bag. The total phosphate concentration should not exceed 1.2 mmol/L.

3 **DOSE FORMS AND STRENGTHS**
See Table 1 for the concentrations of the active ingredients (after mixing) in these 10 different replacement solutions [see Dosage and Administration (2.2)].

4 **CONTRAINDICATIONS**
PHOXILLUM and PRISMASOL replacement solutions are contraindicated in patients with known hypersensitivities to these products.

5 **WARNINGS AND PRECAUTIONS**
5.1 **Electrolyte and Volume Abnormalities**
PHOXILLUM and PRISMASOL solutions can affect electrolytes and volume and may result in hyperkalemia or hyperphosphatemia. Monitor hemodynamic status and fluid inputs and outputs, potassium, phosphorous, calcium, other electrolytes and acid-base balance throughout the procedure. Abnormalities may be corrected by changing the formulation of replacement solution and/or dialysate, supplementation, or adjusting flow rates appropriately [see Dosage and Administration (2)].

PHOXILLUM replacement solutions contain hydrogen phosphate, a weak acid that may increase the risk of metabolic acidosis.

5.2 **Blood Glucose Abnormalities**
The use of PRISMASOL and PHOXILLUM replacement solutions can affect blood glucose levels resulting in hypo- or hyper-glycemia depending upon the dextrose content of the replacement solution. Monitor blood glucose levels regularly. Patients may require initiation of or modification of antidiabetic therapy or othercorrective measures during treatment.

6 **ADVERSE REACTIONS**
The following adverse reactions have been identified during postapproval use with these or other similar products and therefore may occur with use of PHOXILLUM or PRISMASOL. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

- Metabolic acidosis
- Hypotension
- Acid-base disorders
- Electrolyte imbalance including calcium ionized increased (reported in PRISMASOL solutions containing calcium), hyperphosphatemia, and hypophosphatemia
- Fluid imbalance

7 **DRUG INTERACTIONS**
As with the use of other replacement solutions, blood concentrations of dialyzable drugs may be reduced by CRRT due to their removal by the hemofilter or hemodiafilter. The blood concentrations of certain drugs may need to be monitored and appropriate therapy implemented to correct for removal during treatment.

7.1 **Citrate**
When used as an anticoagulant, citrate contributes to the overall buffer load and can reduce plasma calcium levels. Select the PRISMASOL/PHOXILLUM formulation(s) accordingly.
8 \hspace{1em} \textbf{USE IN SPECIFIC POPULATIONS}

8.1 \hspace{1em} \textbf{Pregnancy}

Risk Summary

PRISMASOL and PHOXILLUM are pharmacologically inactive solutions. While there are no adequate and well controlled studies in pregnant women, appropriate administration of PRISMASOL and PHOXILLUM solutions with monitoring of fluid, electrolyte, acid-base and glucose balance, is not expected to cause fetal harm. Animal reproduction studies have not been conducted with PRISMASOL and PHOXILLUM solutions.

The estimated background risk of major birth defects and miscarriage for the indicated population are unknown. All pregnancies have a background risk of birth defect, loss or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively.

Clinical Considerations

Maintenance of normal acid-base balance is important for fetal well-being.

8.2 \hspace{1em} \textbf{Lactation}

Risk Summary

The components of PRISMASOL and PHOXILLUM solutions are excreted in human milk. Appropriate administration of PRISMASOL and PHOXILLUM solutions with monitoring of fluid, electrolyte, acid-base and glucose balance, is not expected to harm a nursing infant.

8.4 \hspace{1em} \textbf{Pediatric Use}

Safety and effectiveness have been established based on published clinical data of CRRT replacement solutions with compositions similar to PRISMASOL and PHOXILLUM used in adults and two hemofiltration studies in pediatric patients, including a study of newborns to 17 years old.

8.5 \hspace{1em} \textbf{Geriatric Use}

The experience with PRISMASOL and PHOXILLUM solutions in geriatric patients has not identified novel concerns.

11 \hspace{1em} \textbf{DESCRIPTION}

PRISMASOL and PHOXILLUM solutions are clear, sterile, free of bacterial endotoxins and contain no bacteriostatic or antimicrobial agents. These solutions are used in Continuous Renal Replacement Therapies (CRRT) as a replacement solution in hemofiltration and hemodiafiltration. Depending on the product (see Table 2), the two compartments contain:

- Calcium chloride, USP, is chemically designated calcium chloride dihydrate (CaCl$_2$ • 2H$_2$O).
- Magnesium chloride, USP, is chemically designated magnesium chloride hexahydrate (MgCl$_2$ • 6H$_2$O).
- Sodium chloride, USP, is chemically designated NaCl.
- Potassium chloride, USP, is chemically designated KCl.
- Sodium bicarbonate, USP, is chemically designated NaHCO$_3$.
- Dextrose, USP, is chemically designated D-Glucose anhydrous ($C$_6$H$_12$O$_6$) or D-Glucose monohydrate ($C$_6$H$_12$O$_6$ • H$_2$O).
- Lactic acid, USP, is chemically designated CH$_3$CH(OH)COOH.
- Dibasic sodium phosphate, USP, is chemically designated as disodium hydrogen phosphate, dihydrate (Na$_2$HPO$_4$ • 2H$_2$O).
### Table 2 – Compartment Composition (Before Mixing)

<table>
<thead>
<tr>
<th></th>
<th>Compartment A (g/L)</th>
<th>Compartment B (g/L)</th>
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<tbody>
<tr>
<td></td>
<td>Calcium Chloride ($\cdot 2H_2O$)</td>
<td>Magnesium Chloride ($\cdot 6H_2O$)</td>
</tr>
<tr>
<td><strong>PRISMASOL SOLUTIONS</strong></td>
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<tr>
<td>BGK 0/2.5</td>
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<td>B22K 4/0</td>
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</tr>
</tbody>
</table>

The pH of the final solution is in the range of 7.0 to 8.5.

### 12 CLINICAL PHARMACOLOGY

#### 12.1 Mechanism of Action

PRISMASOL and PHOXILLUM solutions are pharmacologically inactive. The electrolyte concentrations in the solutions are chosen to restore plasma levels to clinically desired concentrations or maintain plasma levels at the desired concentrations.

PRISMASOL and PHOXILLUM solutions are used as replacement solution to replace water and electrolytes removed during hemofiltration and hemodiafiltration. Bicarbonate (or precursor lactate) in the solution is used as an alkalinizing buffer to restore acid-base balance to a clinically desirable level.

#### 12.3 Pharmacokinetics

The distribution of electrolytes, bicarbonate, and dextrose is determined by the patient’s clinical condition, metabolic status, and residual renal function.

The elimination and replacement of water, electrolytes and buffer depend on the patient’s electrolyte and acid-base balance, metabolic status, residual renal function and ongoing physiologic losses through intestinal, respiratory and cutaneous routes.

### 16 HOW SUPPLIED/STORAGE AND HANDLING

PRISMASOL and PHOXILLUM solutions are supplied in a two-compartment bag made of either polyvinyl chloride (PVC) or polyolefin. The 5000 mL bag is composed of a small compartment (250 mL) and a large compartment (4750 mL). The two compartments are separated by a red frangible pin in the PVC bag and by a peel seal in the polyolefin bag.

The bag is overwrapped with a transparent overwrap. See Table 2 for the concentrations of the active ingredients in each compartment for each product [see Description (11)].
### PRISMASOL Solutions

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<thead>
<tr>
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Polyolefin

### PHOXILLUM Solutions

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<tr>
<td>PHOXILLUM B22K4/0</td>
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</tr>
</tbody>
</table>

Not all formulations may be marketed.

**Storage conditions**

Store at 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F). [See USP Controlled Room Temperature]

Do not freeze or expose to excessive heat. Do not use if precipitate has formed or if container seals have been damaged.

Manufactured for:
Baxter Healthcare Corporation
One Baxter Parkway
Deerfield, Illinois 60015

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