

DEXTROSE- dextrose injection, solution
IBA Inc.

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](#).

DEXTROSE 50 %

INDICATIONS:

As a treatment of uncomplicated primary ketosis in cattle.

DOSAGE and ADMINISTRATION:

The usual dose is 50 mL per 100 lbs. of body weight. May be repeated in 8 to 10 hours or on successive days. For intravenous use only.

FOR VETERINARY USE ONLY

Store at controlled temperature between 15°C and 30°C (59°F -86°F).

TAKE TIME OBSERVE LABEL DIRECTIONS

CONTAINS:

Dextrose • H₂O, 50% w/v
Water For Injection q.s.

KEEP OUT OF REACH OF CHILDREN

NOTE:

If there is no noticeable improvement in the condition being treated after 3 of 4 days of treatment, consult your veterinarian. Aseptic precautions should be observed such as using sterile needle and syringe. Disinfect the site of injection.

CAUTION:

Solution should be warmed to body temperature and administered slowly. This product contains no preservative. Entire contents should be used upon entering. Discard any unused portion.

IBA STOCK #494302

09NT6

Rev. 05-09

18-801-50

RMS 92-554

STERILE

NET CONTENTS: 500 mL

MANUFACTURED FOR:

IBA INC.

ANIMAL HEALTH DIVISION

27 PROVIDENCE RD.

MILLBURY, MA 01527

LOT NO.:

EXP. DATE:

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OBSERVE LABEL DIRECTIONS

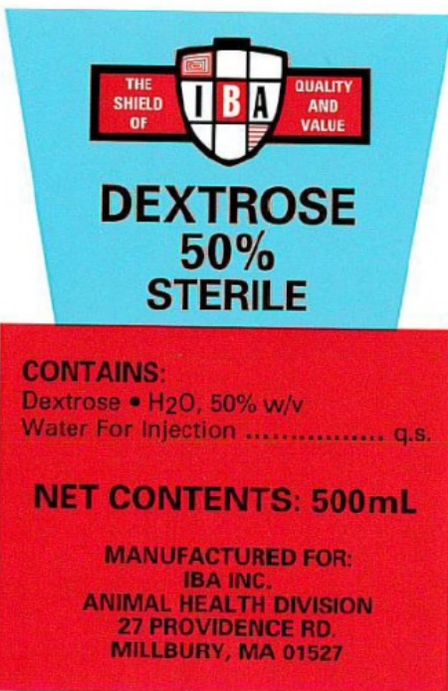
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DEXTROSE

dextrose injection, solution

Product Information

Product Type	OTC ANIMAL DRUG	Item Code (Source)	NDC:29983-2403
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DEXTROSE MONOHYDRATE (UNII: LX22YL083G) (ANHYDROUS DEXTROSE - UNII:5SL0G7R0OK)	DEXTROSE MONOHYDRATE	250 g in 500 mL

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:29983-2403-1	500 mL in 1 BOTTLE, PLASTIC		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		01/16/2018	

Labeler - IBA Inc. (019494160)

Registrant - IBA Inc. (019494160)

Establishment

Name	Address	ID/FEI	Business Operations
Nova-Tech, Inc.		196078976	manufacture, api manufacture

Revised: 1/2018

IBA Inc.