

DOCUSATE SODIUM- docusate sodium capsule, liquid filled
NuCare Pharmaceuticals, Inc.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

GC 401

Active ingredient (in each softgel)

Docusate Sodium 100 mg

Purpose

Stool Softener Laxative

Uses

- relieves occasional constipation (irregularity)
- generally produces bowel movement in 12 to 72 hours

Warnings

Ask a doctor before use if you

- have stomach pain, nausea or vomiting
- have a sudden change in bowel habits that persists over a period of 2 weeks
- are presently taking mineral oil

Stop use and ask a doctor if

- you need to use a laxative longer than 1 week
- you have rectal bleeding or fail to have a bowel movement. These could be signs of a serious condition.

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

Directions

- adults and children 12 years and older: take 1-2 softgel daily until first bowel movement; 1 softgel daily thereafter, or as directed by doctor
- children under 12: consult a doctor
- do not exceed recommended dose

Other information

- **each softgel contains:** sodium 5 mg. very low sodium
- store at 15°C-25°C (59° F-77° F)

- keep tightly closed
- product from USA or Canada
- **Tamper Evident:** Do not use if imprinted seal under cap is missing or broken.

Inactive ingredients

FD&C red #40, gelatin, glycerin, edible ink, PEG, propylene glycol, sorbitol sorbitan solution, water. Also contains D&C yellow #10 or FD&C yellow #6 (sunset yellow).

Package Label


NuCare Pharmaceuticals, Inc.

Take _____ **times a day.**

every _____ **hours**

Patent Instructions:

Orange, CA 92867

Brooklyn, NY 11204

Gerl-Care Pharmaceuticals Corp.

Distributed by:

3 6807152803

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Rev 01/01/19

NDC: 68071-5280-3

Docusate Sodium 100mg

#30 Softgels

Each softgel contains: Docusate Sodium 100mg..... Stool Softener Laxative FD & C Yellow #6 (sunset yellow).
 Warnings: Ask a doctor before use if you have stomach pain, nausea or vomiting, have a sudden change in bowel habits that persists over a period of 2 weeks, are presently taking mineral oil. Stop use and ask a doctor if, you need to use a laxative longer than 1 week, you have rectal bleeding or fail to have a bowel movement. These could be signs of a serious condition. If pregnant or breast-feeding, ask a health professional before use. Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away. Oval Clear-Red Softgel Printed: "40"

Product #: P0083030

Docusate Sodium 100mg
 Lot: 000000 NDC: 68071-5280-03
 MFR NDC: 57896-407-10 Exp.: 00-00
 Serial# 00000000002

Docusate Sodium 100mg
 Lot: 000000 NDC: 68071-5280-03
 MFR NDC: 57896-407-10 Exp.: 00-00
 Serial# 00000000002

 GTIN 00368071528039
 Serial# 00000000002
 Exp. Date 00-00
 LOT#: 000000

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

WARNING: KEEP OUT OF REACH OF CHILDREN

STORE AT CONTROLLED TEMPERATURE 59-77°F.

DOCUSATE SODIUM

docusate sodium capsule, liquid filled

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68071-5280(NDC:57896-407)
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DOCUSATE SODIUM (UNII: F05Q2T2JA0) (DOCUSATE - UNII:M7P27195AG)	DOCUSATE SODIUM	100 mg

Inactive Ingredients

Ingredient Name	Strength
FD&C RED NO. 40 (UNII: WZB9127XOA)	
GELATIN (UNII: 2G86QN327L)	
GLYCERIN (UNII: PDC6A3C0OX)	
POLYETHYLENE GLYCOL 400 (UNII: B697894SGQ)	

PROPYLENE GLYCOL (UNII: 6DC9Q167V3)				
SORBITOL (UNII: 506T60A25R)				
WATER (UNII: 059QF0KO0R)				
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)				
FD&C YELLOW NO. 6 (UNII: H77VEI93A8)				
SORBITAN (UNII: 6O92ICV9RU)				
Product Characteristics				
Color	red	Score	no score	
Shape	OVAL	Size	12mm	
Flavor		Imprint Code	401	
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68071-5280-3	30 in 1 BOTTLE; Type 0: Not a Combination Product	06/11/2020	
Marketing Information				
Marketing Category	Application Number or Monograph Citation		Marketing Start Date	Marketing End Date
OTC monograph not final	part334		01/01/2012	

Labeler - NuCare Pharmaceuticals, Inc. (010632300)

Establishment

Name	Address	ID/FEI	Business Operations
NuCare Pharmaceuticals, Inc.		010632300	repack(68071-5280)

Revised: 6/2020

NuCare Pharmaceuticals, Inc.