

**HEB GENTLE LAXATIVE- bisacodyl suppository**  
**H E B**

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**DRUG FACTS**

**Active ingredient**

Bisacodyl USP, 10 mg

**Purpose**

Stimulant Laxative

**Uses**

- For relief of occasional constipation and irregularity
- -This product generally produces bowel movement in 15 minutes to 1 hour

**Warnings**

For rectal use only.

- stomach pain, nausea or vomiting
- noticed a sudden change in bowel habits that persists over a period of two weeks

**When using this product**

May cause abdominal discomfort, faintness, rectal burning, and mild cramps

**Stop use and ask a doctor if**

- if you have rectal bleeding or fail to have bowel movement after using a laxative. This may indicate a serious condition
- you need to use a laxative for more than 1 week

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

**Keep out of reach of children**

**If swallowed, get medical help or contact a Poison Control Center right away.**

**Directions**

Adults and children 12 years of age and older  
years Children under 6

Children 6 to under 12

One suppository once daily

1/2 suppository once

daily

Ask doctor.

-Detach one suppository from the strip and remove from foil - Carefully insert one suppository well into the rectum

-Do not use more than once per day

**Other Information**

- do not store above 30°C (86°F)

**Inactive Ingredients**

hydrogenated vegetable oil



## HEB GENTLE LAXATIVE

bisacodyl suppository

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:37808-877
<b>Route of Administration</b>	RECTAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BISACODYL</b> (UNII: 10X0709Y6I) (DEACETYLBISACODYL - UNII:R09078E41Y)	BISACODYL	10 mg in 2000 mg

### Inactive Ingredients

Ingredient Name	Strength
<b>FAT, HARD</b> (UNII: 8334LX7S21)	1990 mg in 2000 mg

### Product Characteristics

<b>Color</b>	white	<b>Score</b>	
<b>Shape</b>	BULLET	<b>Size</b>	34mm
<b>Flavor</b>		<b>Imprint Code</b>	
<b>Contains</b>			

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:37808-877-08	2 in 1 CARTON	06/05/2019	
1		40 mg in 1 BLISTER PACK; Type 0: Not a Combination Product		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M007	06/05/2019	

**Labeler** - H E B (007924756)

**Registrant** - Reese Pharmaceutical Co (004172052)

### Establishment

Name	Address	ID/FEI	Business Operations
DSC Laboratories, Inc.		097807374	manufacture(37808-877)

Revised: 12/2024

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