LAXATIVE MAXIMUM STRENGTH- sennosides tablet, sugar coated L.N.K. International, Inc.

Sound Body 44-348

Active ingredient (in each tablet)

Sennosides USP, 25 mg

Purpose

Stimulant laxative

Uses

- relieves occasional constipation (irregularity)
- generally produces bowel movement in 6 to 12 hours

Warnings

Do not use

laxative products when abdominal pain, nausea, or vomiting are present unless directed by a doctor.

Ask a doctor before use if you have

noticed a sudden change in bowel habits that persists over a period of 2 weeks.

Ask a doctor or pharmacist before use if you are

taking any other drug. Laxatives may affect how other drugs work. Take this product 2 or more hours before or after other drugs.

When using this product,

do not use for a period longer than 1 week.

Stop use and ask a doctor if

rectal bleeding or failure to have a bowel movement occur after use of a laxative. These may 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 (1-800-222-

Directions

- swallow tablet(s) with a glass of water
- swallow tablet(s) whole; do not crush, break, or chew

adults and children 12 years and over	2 tablets once or twice daily
children 6 to under 12 years	1 tablet once or twice daily
children under 6 years	ask a doctor

Other information

- each tablet contains: calcium 40 mg, magnesium 5 mg
- TAMPER EVIDENT: DO NOT USE IF OUTER PACKAGE IS OPENED OR BLISTER IS TORN OR BROKEN
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- see end flap for expiration date and lot number

Inactive ingredients

acacia, calcium carbonate, carnauba wax, corn starch, dibasic calcium phosphate dihydrate, FD&C blue #1 aluminum lake, iron oxide black, magnesium stearate, methylparaben, microcrystalline cellulose, polyethylene glycol, polyvinyl alcohol, povidone, propylene glycol, propylparaben, shellac glaze, silicon dioxide, sodium benzoate, sodium lauryl sulfate, sucrose, talc, titanium dioxide

Questions or comments?

1-800-426-9391

Principal Display Panel

SOUND**BODY**™

*Compare to the active ingredient in ex•lax® Maximum Strength

NDC 50844-948-08

MAXIMUM STRENGTH

Laxative

Sennosides USP, 25 mg Stimulant Laxative Gentle, Dependable Constipation Relief

24 TABLETS

Actual Size

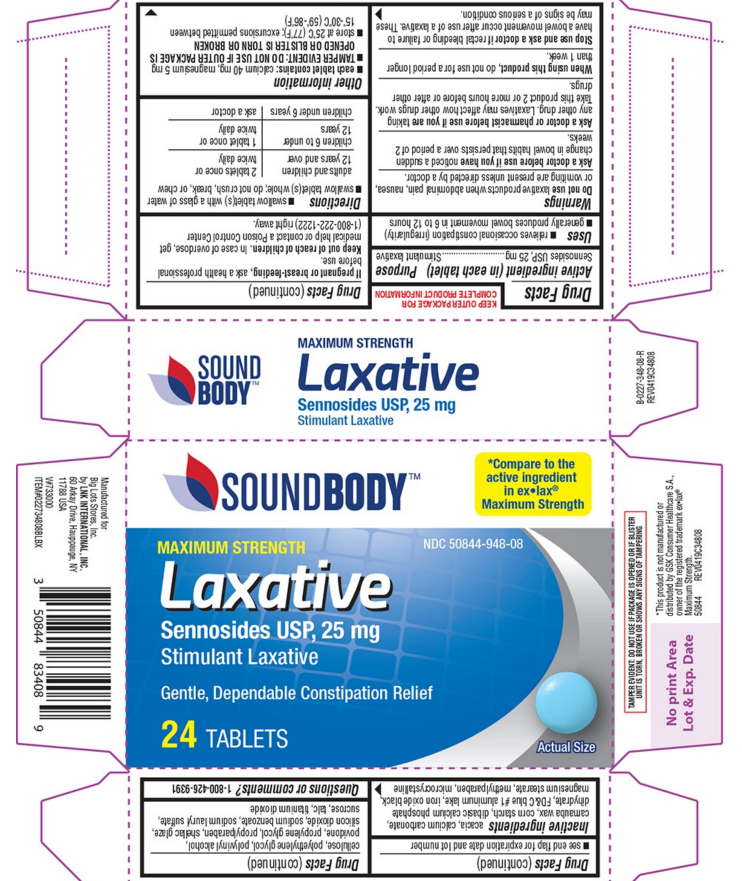
TAMPER EVIDENT: DO NOT USE IF PACKAGE IS OPENED OR IF BLISTER UNIT IS TORN, BROKEN OR SHOWS ANY SIGNS OF TAMPERING

*This product is not manufactured or distributed by GSK Consumer Healthcare S.A., owner of the registered trademark ex•lax® Maximum Strength. 50844 REV0419C34808

Manufactured for Big Lots Stores, Inc. by **LNK INTERNATIONAL, INC.** 60 Arkay Drive, Hauppauge, NY 11788 USA

LAXATIVE MAXIMUM STRENGTH





Product Inform	nation						
Product Type		HUMAN OTC	DRUG	Item Code	(Source)	NDC:50)844-948
Route of Adminis	tration	ORAL					
Noute of Adminis	tration						
Active Ingredie	nt/Activ	e Moiety					
	Inar	edient Name	3		Basis of Str	enath	Strength
SENNOSIDES (UNII: 3	-)IJX)	SENNOSIDES	5	25 mg
Inactive Ingred	lients						
		Ingredi	ent Name				Strength
ACACIA (UNII: 5C540)							
CALCIUM CARBONA	TE (UNII: H	0G9379FGK)					
CARNAUBA WAX (UN							
STARCH, CORN (UNI							
DIBASIC CALCIUM P				7GEP)			
FD&C BLUE NO. 1 A			QA3S2JM)				
METHYLPARABEN (U MICROCRYSTALLINI			206111)				
POLYETHYLENE GL				\)			
POLYVINYL ALCOHO			-	-)			
POVIDONE, UNSPEC			-				
PROPYLENE GLYCO							
PROPYLPARABEN (U	-						
SHELLAC (UNII: 46N1		,					
SILICON DIOXIDE (U		XBU4)					
SODIUM BENZOATE	(UNII: OJ24	5FE5EU)					
SODIUM LAURYL SU	LFATE (UN	III: 368GB5141J)					
SUCROSE (UNII: C15	1H8M554)						
TALC (UNII: 7SEV7J4R	1U)						
TITANIUM DIOXIDE	(UNII: 15FIX	9V2JP)					
		_					
Product Charac			c				
Color		lue	Score			no score	
Shape Flavor	R	OUND	Size	Co do		10mm 44;348	
Contains			Imprint (Lode		44,540	
Packaging							

1 NDC:50844- 948-08	2 in 1 CARTON	01/02/2003				
1	12 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			
		01/02/2002				
OTC Monograph Dr	ıg 505G(a)(3)	01/02/2003				

Labeler - L.N.K. Inte	rnational, Inc. (038154464)
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Establishment					
Name	Address	ID/FE	1		Business Operations
NK International, Inc.		038154464	4	manufacture(50844-948) , pack(50844-948)
Establishment					
Name	Address	ID/FE	1		Business Operations
LNK International, Inc.		83286783	7	manufacture(50844-948) , pack(50844-948)
Establishment					
Name	Ad	dress		ID/FEI	Business Operations
LNK International, Inc.			8328	67894	manufacture(50844-948)
Establishment					
Name	Ad	dress		ID/FEI	Business Operations

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		117025878	manufacture(50844-948)

967626305

pack(50844-948)

Revised: 10/2023

LNK International, Inc.

L.N.K. International, Inc.