

GERI-KOT- sennosides tablet
Geri-Care Pharmaceutical Corp

GC 451C 417

Active ingredient (in each tablet)

Sennosides 8.6 mg

Purpose

Laxative

Uses

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

Warnings

Do not use for more than one week unless directed by a doctor.

Ask a doctor before use if you have

- stomach pain, nausea or vomiting
- noticed a change in bowel habits that lasts over two weeks

Stop use and ask a doctor if you have rectal bleeding

or fail to have a bowel movement after use of a laxative.

These may indicate 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

- do not exceed 8 tablets in 24 hours

Age	Starting Dose	Maximum Dose
adults and children 12 years of age and older	2 tablets once a day preferably at bedtime; increase as needed, or as directed by a doctor	4 tablets in the morning and 4 tablets at bedtime
children under 12 years	ask a doctor	

- **each tablet contains:** calcium 20 mg
- **Tamper Evident:** Do not use if imprinted seal under cap is missing or broken
- store at 20°C-25°C (68°F-77°F)

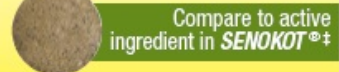
croscarmellose sodium, dicalcium phosphate, hypromellose, magnesium stearate, maltodextrin, microcrystalline cellulose, mineral oil, PEG, talc.



NCDC 57896-417-01

GERI-KOT

STANDARDIZED SENNA CONCENTRATE
NATURAL VEGETABLE
LAXATIVE



100 Tablets 8.6 mg each

Compare to active ingredient in *SENOKOT*®†

Drug Facts

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Other information

• each tablet contains: calcium 20 mg • **Tamper Evident** Do not use if imprinted seal under cap is missing or broken • store at 20°C-25°C (68°F-77°F)

Inactive ingredients:

croscarmellose sodium, dicalcium phosphate, hypromellose, magnesium stearate, maltodextrin, microcrystalline cellulose, mineral oil, PEG, talc.

Questions or comments? Call 1-800-540-3765

† This product is not manufactured or distributed by the owner of the registered trademark SENOKOT®. Dist. by: Geri-Care Pharmaceuticals Corp. 1295 Towbin Avenue, Lakewood, NJ 08701



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REV 11/23C

sennosides tablet

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:57896-417
Route of Administration	ORAL		

Ingredient Name	Basis of Strength	Strength
SENNOSIDES (UNII: 3FYP5M0IJX) (SENNOSIDES - UNII:3FYP5M0IJX)	SENNOSIDES	8.6 mg

Ingredient Name	Strength
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	

DIBASIC CALCIUM PHOSPHATE DIHYDRATE (UNII: O7TSZ97GEP)				
MALTODEXTRIN (UNII: 7CVR7L4A2D)				
MAGNESIUM STEARATE (UNII: 70097M6I30)				
TALC (UNII: 7SEV7J4R1U)				
MINERAL OIL (UNII: T5L8T28FGP)				
HYPROMELLOSES (UNII: 3NXW29V3WO)				
Product Characteristics				
Color	brown	Score	no score	
Shape	ROUND	Size	9mm	
Flavor		Imprint Code	W2	
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:57896-417-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	01/01/2024	
2	NDC:57896-417-10	1000 in 1 BOTTLE; Type 0: Not a Combination Product	01/01/2024	
3	NDC:57896-417-20	200 in 1 BOTTLE; Type 0: Not a Combination Product	01/01/2024	
Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug		505G(a)(3)	01/01/2024	

Labeler - Geri-Care Pharmaceutical Corp (611196254)

Registrant - Geri-Care Pharmaceutical Corp (611196254)

Revised: 5/2024

Geri-Care Pharmaceutical Corp