

STOMINT - atractylodes lancea root tablet
Evercarepharm Co., Ltd

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.

Drug Facts

active ingredient: atractylodis rhizoma

inactive ingredient: hovenia dulcis thunberg, crataegi fructus, puerariae radix, glycyrrhizae resina, agastachis herba, scutellariae radix, cinnamon bark

for the treatment of reflux esophagitis, chronic gastritis, acute gastritis, abdominal distension, indigestion, hangovers

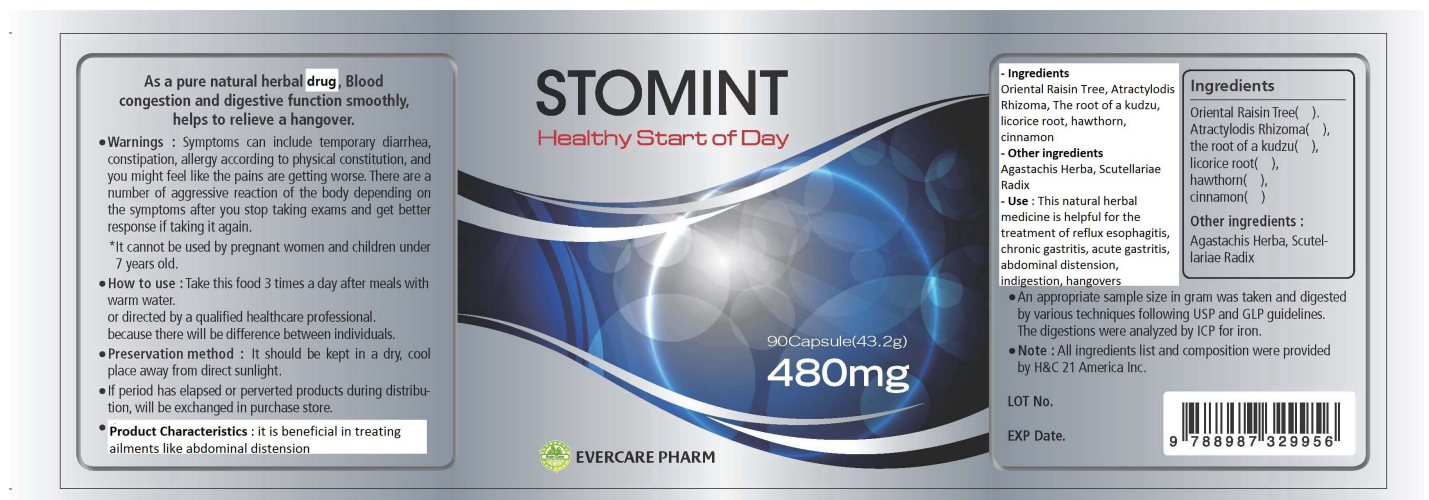
keep out of reach of the children

take 3 times a day after meals with warm water

keep in a dry, cool place

keep away from direct sunlight

do not take if you are pregnant



STOMINT

atractylodes lancea root tablet

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:42469-2001
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
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ATRACTYLODES LANCEA ROOT (UNII: CAZ6282J2O) (ATRACTYLODES LANCEA ROOT - UNII:CAZ6282J2O)			ATRACTYLODES LANCEA ROOT	70 mg
Inactive Ingredients				
Ingredient Name				Strength
HOVENIA DULCIS FRUIT (UNII: 18F22L82RE)				
CRATAEGUS MONOGYNA FRUIT (UNII: KJ2JZT1TS)				
PUERARIA MONTANA VAR. LOBATA ROOT (UNII: PET93F4I3C)				
GLYCYRRHIZA GLABRA (UNII: 2788Z9758H)				
POGOSTEMON CABLIN TOP (UNII: 2I2A73IYL7)				
SCUTELLARIA BAICALENSIS ROOT (UNII: 7J95K7ID2S)				
CHINESE CINNAMON (UNII: WS4CQ062KM)				
Product Characteristics				
Color	white (white)	Score	no score	
Shape	OVAL (tablet)	Size	20mm	
Flavor		Imprint Code	3;hp;x	
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:42469-2001-1	90 in 1 BOTTLE, WITH APPLICATOR		
Marketing Information				
Marketing Category	Application Number or Monograph Citation		Marketing Start Date	Marketing End Date
unapproved drug other			06/06/2012	

Labeler - Evercarepharm Co., Ltd (557812798)

Registrant - Evercarepharm Co., Ltd (557812798)

Establishment			
Name	Address	ID/FEI	Business Operations
Evercarepharm Co., Ltd		557812798	manufacture