MENSTRUAL RELIEF- acetaminophen, caffeine, pyrilamine maleate tablet, film coated

Geiss, Destin & Dunn Inc.

GoodSense 44-390

Active ingredients (in each caplet)

Acetaminophen 500 mg Caffeine 60 mg Pyrilamine maleate 15 mg

Purpose

Pain reliever Diuretic Antihistamine

Uses

for the temporary relief of these symptoms associated with menstrual periods:

- bloating
- headache
- water-weight gain
- cramps
- backache
- fatigue
- muscle aches

Warnings

Liver warning: This product contains acetaminophen. Severe liver damage may occur if you take

- more than 4,000 mg of acetaminophen in 24 hours
- with other drugs containing acetaminophen
- 3 or more alcoholic drinks every day while using this product

Allergy alert: Acetaminophen may cause severe skin reactions. Symptoms may include:

- skin reddening
- blisters
- rash

If a skin reaction occurs, stop use and seek medical help right away.

Do not use

- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.
- if you have ever had an allergic reaction to this product or any of its ingredients

Ask a doctor before use if you have

- difficulty in urination due to enlargement of the prostate gland
- liver disease
- glaucoma
- a breathing problem such as emphysema or chronic bronchitis

Ask a doctor or pharmacist before use if you are

- taking the blood thinning drug warfarin
- taking sedatives or tranquilizers

When using this product

- drowsiness may occur
- avoid alcoholic beverages
- alcohol, sedatives, and tranquilizers may increase drowsiness
- excitability may occur, especially in children
- use caution when driving a motor vehicle or operating machinery
- limit the use of caffeine-containing medications, foods, or beverages while taking this product because too much caffeine may cause nervousness, irritability, sleeplessness, and, occasionally, rapid heart rate. The recommended dose of this product contains about as much caffeine as a cup of coffee.

Stop use and ask a doctor if

- new symptoms occur
- redness or swelling is present
- pain gets worse or lasts more than 10 days
- fever gets worse or lasts more than 3 days

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. Prompt medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Directions

- do not take more than directed
- adults and children 12 years and over:
 - take 2 caplets with water
 - repeat every 6 hours, as needed

- do not exceed 6 caplets per day
- children under 12 years: ask a doctor

Other information

- see end flap for expiration date and lot number
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)

Inactive ingredients

corn starch, croscarmellose sodium, crospovidone, hypromellose, magnesium stearate, microcrystalline cellulose, polydextrose, polyethylene glycol, povidone, silicon dioxide, stearic acid, titanium dioxide, triacetin

Questions or comments?

1-800-426-9391

Principal Display Panel

GOODSENSE®

NDC 50804-990-08

Menstrual Relief Caplets

Acetaminophen, Caffeine,

Pyrilamine maleate

• Pain Reliever • Diuretic • Antihistamine

Multi-Symptom Relief of:

- Cramps Bloating Fatigue
- Backache
 Headache

*Compare to the active ingredients of Midol® Complete

24 Caplets

actual size

TAMPER EVIDENT: DO NOT USE IF IMPRINTED SAFETY SEAL UNDER CAP IS BROKEN OR MISSING

*This product is not manufactured or distributed by Bayer HealthCare LLC, owner of the registered trademark Midol® Complete. 50844 REV0623B39008

Distributed by: Perrigo Direct, Inc., Peachtree City, GA 30269 www.PerrigoDirect.com (1-800-426-9391) GoodSense® is a registered trademark of L. Perrigo Company.



Good Sense 44-390

Product Info	rmation							
Product Type		HUMAN OTC	DRUG	Item Co	ode (Sou	irce)	NDC:50	804-990
Route of Admin	istration	ORAL						
Active Ingred	ient/Active	e Moiety						
	Ing	redient Nan	ne			Basis of	Strengt	n Strengt
ACETAMINOPHEN	(UNII: 362091	TL9D) (ACETAMI	NOPHEN - UNI	II:36209IT	L9D)	ACETAMINC	PHEN	500 mg
CAFFEINE (UNII: 3	G6A5W338E) (CAFFEINE - UNII:	3G6A5W338E))		CAFFEINE		60 mg
PYRILAMINE MAL	E ATE (UNII: R3	5D29L3ZA) (PYF	RILAMINE - UN	II:HPE3170	O9TL)	PYRILAMINE	MALEATE	15 mg
Inactive Ingre	edients							
		Ingredie	ent Name					Strength
STARCH, CORN (L	INII: 08232NY3	3SJ)						
CROSCARMELLOS	SE SODIUM (L	JNII: M28OL1HH4	48)					
CROSPOVIDONE,	UNSPECIFIEI	D (UNII: 2578301	E561)					
HYPROMELLOSE,	UNSPECIFIE	D (UNII: 3NXW29	V3WO)					
MAGNESIUM STE	ARATE (UNII: 7	70097M6I30)						
MICROCRYSTALL	NE CELLULO	SE (UNII: OP1R3	2D61U)					
POLYDEXTROSE	UNII: VH2XOU	L2IE)						
POLYETHYLENE O	SLYCOL, UNS	PECIFIED (UNII:	3WJQ0SDW1	4)				
POVIDONE, UNSP	ECIFIED (UNII	: FZ989GH94E)						
SILICON DIOXIDE	-							
STEARIC ACID (UN	III: 4ELV7Z65A	P)						
TITANIUM DIOXID		9V2JP)						
TRIACETIN (UNII:)	(HX3C3X673)							
Product Char	acteristics	5						
Color	٧	white	Score				no score	
Shape	(OVAL	Size				17mm	
Flavor			Imprint Co	de			44;390	
Contains			-					
Packaging								
# Item Code	F	Package Des	cription			ting Star Date	t Mark	eting End Date
1 NDC:50804- 990-08	1 in 1 CARTO				03/24/20	20		
1	24 in 1 BOTT Combination	LE, PLASTIC; Tyj Product	pe 0: Not a					
Marketing	Informa	tion						

Category	Citation	Date	Date
OTC Monograph Drug	M013	03/24/2020	

labolar at a		(0760500			
Labeler - Geiss, Des	itin & Dunn Inc.	. (0760598	36)		
Establishment					
Name	Ad	dress	ID/FEI	Business Operations	
NK International, Inc.			038154464	pack(50804-990)	
Establishment					
Name	Address	ID/FE	I	Business Operations	
LNK International, Inc.		83286783	7 manufactur	re(50804-990) , pack(50804-990)	
Establishment					
Name	Ad	dress	ID/FEI	Business Operations	
NK International, Inc.			832867894	manufacture(50804-990)	
Establishment					
Name	Ad	dress	ID/FEI	Business Operations	
LNK International, Inc.			967626305	pack(50804-990)	
Establishment					
Name	Ad	dress	ID/FEI	Business Operations	
LNK International, Inc.			117025878	manufacture(50804-990)	

Revised: 2/2024

Geiss, Destin & Dunn Inc.