PAIN RELIEF EXTRA STRENGTH- acetaminophen tablet, coated L.N.K. International, Inc.

Sound Body 44-519

Active ingredient (in each gelcap)

Acetaminophen 500 mg

Purpose

Pain reliever/fever reducer

Uses

- temporarily relieves minor aches and pains due to:
 - headache
 - the common cold
 - toothache
 - backache
 - muscular aches
 - minor pain of arthritis
 - premenstrual and menstrual cramps
- temporarily reduces fever

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

- if you are allergic to acetaminophen or any of the inactive ingredients in this product
- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.

Ask a doctor before use if you have

liver disease.

Ask a doctor or pharmacist before use if you are

taking the blood thinning drug warfarin.

Stop use and ask a doctor if

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

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 gelcaps every 6 hours while symptoms last
 - do not take more than 6 gelcaps in 24 hours, unless directed by a doctor
 - do not take for more than 10 days unless directed by a doctor
- children under 12 years: ask a doctor

Other information

- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- avoid high humidity
- use by expiration date on package

Inactive ingredients

croscarmellose sodium, D&C red #33, FD&C blue #1, FD&C red #40, gelatin, hydroxypropyl cellulose, hypromellose, iron oxide black, iron oxide red, iron oxide yellow, polyethylene glycol, povidone, pregelatinized starch, propylene glycol, shellac glaze, stearic acid, titanium dioxide

Questions or comments?

1-800-426-9391

Principal Display Panel

SOUND**BODY**™

*Compare to the active ingredient in Extra Strength Tylenol® Rapid Release Gels

NDC 50844-951-20

EXTRA STRENGTH
Pain Relief

Acetaminophen 500 mg

Pain Reliever/Fever Reducer

Contains No Aspirin

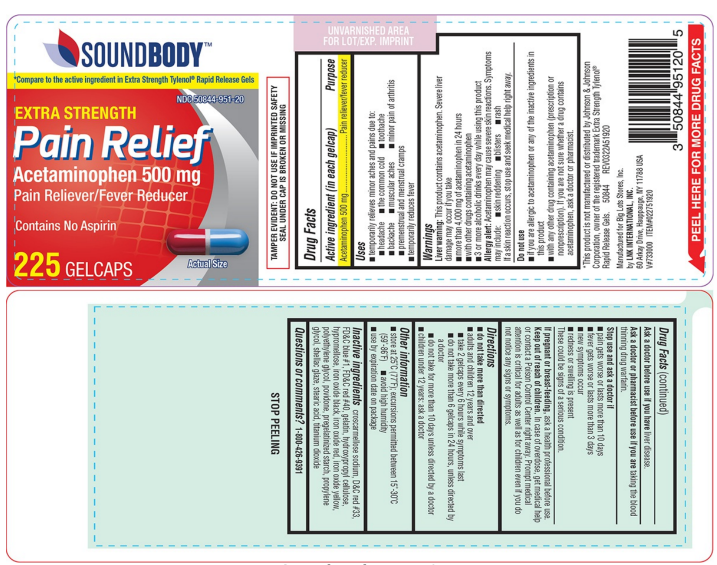
225 GELCAPS

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 Johnson & Johnson Corporation, owner of the registered trademark Extra Strength Tylenol® Rapid Release Gels. 50844 REV0322A51920

Manufactured for Big Lots Stores, Inc. by **LNK INTERNATIONAL, INC.** 60 Arkay Drive, Hauppauge, NY 11788 USA V#733000 ITEM#022751920



Sound Body 44-519

PAIN RELIEF EXTRA STRENGTH

acetaminophen tablet, coated

Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:50844-951

Route of Administration ORAL

Active Ingredient/Active Moiety

ACETAMINOPHEN (UNII: 36209ITL9D) (ACETAMINOPHEN - UNII:36209ITL9D) ACETAMINOPHEN 500 mg

Inactive Ingredients		
Ingredient Name	Strength	
CROSCARMELLOSE SODIUM (UNII: M280L1HH48)		
D&C RED NO. 33 (UNII: 9DBA0SBB0L)		

FD&C BLUE NO. 1 (UNII: H3R47K3TBD)

FD&C RED NO. 40 (UNII: WZ B9127XOA)

GELATIN, UNSPECIFIED (UNII: 2G86QN327L)

HYDROXYPROPYL CELLULOSE, UNSPECIFIED (UNII: 9XZ 8H6N6OH)

HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)

FERROSOFERRIC OXIDE (UNII: XM0M87F357)

FERRIC OXIDE RED (UNII: 1K09F3G675)

FERRIC OXIDE YELLOW (UNII: EX43802MRT)

POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)

POVIDONE, UNSPECIFIED (UNII: FZ989GH94E)

STARCH, CORN (UNII: 08232NY3SJ)

PROPYLENE GLYCOL (UNII: 6DC9Q167V3)

SHELLAC (UNII: 46N107B71O)

STEARIC ACID (UNII: 4ELV7Z65AP)

TITANIUM DIOXIDE (UNII: 15FIX9V2JP)

Product Characteristics			
Color	red, blue	Score	no score
Shape	OVAL	Size	19mm
Flavor		Imprint Code	L;5
Contains			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:50844- 951-15	1 in 1 CARTON	05/10/2004		
1		50 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product			
2	NDC:50844- 951-12	100 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	05/10/2004		
3	NDC:50844- 951-20	225 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	05/10/2004		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M013	05/10/2004	

Labeler - L.N.K. International, Inc. (038154464)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		038154464	manufacture(50844-951) , pack(50844-951)

Establishment				
Name	Address	ID/FEI	Business Operations	
LNK International, Inc.		832867837	manufacture(50844-951)	

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867894	manufacture(50844-951)

Establishment				
Name	Address	ID/FEI	Business Operations	
LNK International, Inc.		868734088	manufacture(50844-951)	

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		967626305	pack(50844-951)

Revised: 6/2023 L.N.K. International, Inc.