PAIN RELIEVER EXTRA STRENGTH- acetaminophen tablet Chain Drug Consortium

Premier Value 44-148

Active ingredient (in each tablet)

Acetaminophen 500 mg

Purpose

Pain reliever/fever reducer

Uses

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

- 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 are allergic to acetaminophen or any of the inactive ingredients in this product

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 accidental 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 tablets every 6 hours while symptoms last
 - do not take more than 6 tablets 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)
- see end flap for expiration date and lot number

Inactive ingredients

corn starch, povidone, sodium starch glycolate, stearic acid

Questions or comments?

1-800-426-9391

Principal Display Panel

Premier Value®

*COMPARE TO THE ACTIVE INGREDIENT IN EXTRA STRENGTH TYLENOL®

Extra Strength Pain Reliever Acetaminophen 500 mg

PAIN RELIEVER/FEVER REDUCER

100 Tablets

Non aspirin

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®. 50844 ORG062114812

Distributed By: Pharmacy Value Alliance, LLC 407 East Lancaster Avenue, Wayne, PA 19087

If for any reason you are not satisfied with this product, please return it to the store where purchased for a full refund.



Premier Value 44-148

PAIN RELIEVER EXTRA STRENGTH acetaminophen tablet					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68016-691		
Route of Administration	ORAL				

	tive Ingred	lient/Act	ive Moiety	/					
Ingredient Name						Basis of Strength		Strength	
ACETAMINOPHEN (UNII: 36209ITL9D) (ACETAMINOPHEN - UNII: 36209IT				TL9D) ACETAMINOPHEN		500 mg			
In	active Ingre	edients							
			Ingr	redient	Name			S	Strength
ST	ARCH, CORN (L	JNII: 082321	NY3SJ)						
РС	VIDONE, UNSP	PECIFIED (U	INII: FZ989GH	94E)					
so	DIUM STARCH	GLYCOLAT	E TYPE A PO	DTATO (L	JNII: 5856J3G2A2)				
ST	EARIC ACID (UN	NII: 4ELV7Z6	55AP)						
Pı	oduct Char	acteristi	ics						
Color			white		Score			2 pieces	
Shape			ROUND		Size			12mm	
Flavor				I	Imprint Code 44			44;148	
Co	ntains								
	ntains ackaging								
			Package	Descri	ption	Mark	eting Start Date		eting End Date
Pa #	ackaging	1 in 1 CART	•	Descri	ption	Mark	Date		
Pa #	ackaging Item Code NDC:68016-		FON		ption mbination Product		Date		
P a # 1	ackaging Item Code NDC:68016-		TON TTLE; Type 0:				Date 993		
Pa # 1	Ackaging Item Code NDC:68016- 691-60 NDC:68016-	60 in 1 BO	TON TTLE; Type 0: TON DTTLE, PLASTI	Not a Co	mbination Product	01/21/1	Date 993		
Pa # 1 1 2	Ackaging Item Code NDC:68016- 691-60 NDC:68016-	60 in 1 BO 1 in 1 CART 100 in 1 BC	TON TTLE; Type 0: TON DTTLE, PLASTI	Not a Co	mbination Product	01/21/1	Date 993		
Pa # 1 2 2	Ackaging Item Code NDC:68016- 691-60 NDC:68016-	60 in 1 BO 1 in 1 CART 100 in 1 BC Combinatio	FON TTLE; Type 0: FON DTTLE, PLASTI on Product	Not a Co	mbination Product	01/21/1	Date 993		
Pa # 1 2 2	Ackaging Item Code NDC:68016- 691-60 NDC:68016- 691-01	60 in 1 BO 1 in 1 CART 100 in 1 BC Combination	TON TTLE; Type 0: TON DTTLE, PLASTION Product	Not a Co IC; Type (mbination Product	01/21/1 01/21/1	Date 993	Marke	

Labeler - Chain Drug Consortium (101668460)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		038154464	pack(68016-691)
Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867837	manufacture(68016-691)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		868734088	manufacture(68016-691)
Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		967626305	pack(68016-691)
Revised: 4/2024			Chain Drug Consortiun