TYLENOL EXTRA STRENGTH- acetaminophen tablet, film coated Johnson & Johnson Consumer Inc.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

TYLENOL Extra Strength

Drug Facts

Active ingredient (in each caplet)

Acetaminophen 500 mg

Purpose

Pain reliever/fever reducer

Uses

- temporarily relieves minor aches and pains due to:
 - the common cold
 - headache
 - 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.

Overdose warning

In case of overdose, get medical help or contact a Poison Control Center right away. (1-800-222-1222) Quick 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 (see overdose warning)

adults and children 12 years and over	 take 2 caplets every 6 hours while symptoms last do not take more than 6 caplets in 24 hours, unless directed by a doctor do not use for more than 10 days unless directed by a doctor
children under	

12 years

Other information

- store between 20-25°C (68-77°F)
- do not use if carton is opened. Do not use if foil inner seal imprinted with

"TYLENOL" is broken or missing

Inactive ingredients

carnauba wax ¹, corn starch ¹, FD&C red no. 40 aluminum lake, hypromellose, magnesium stearate, modified starch ¹, polyethylene glycol ¹, powdered cellulose, pregelatinized starch, propylene glycol, shellac, sodium starch glycolate, titanium dioxide

1 contains one or more of these ingredients

Questions or comments?

call 1-877-895-3665 (toll-free) or 215-273-8755 (collect)

PRINCIPAL DISPLAY PANEL

NDC 50580-449-96

TYLENOL[®] FOR ADULTS

Acetaminophen Pain Reliever Fever Reducer

Extra Strength Actual Size

50 Caplets 500 mg each



TYLENOL EXTRA STRENGTH

acetaminophen tablet, film coated

Product Information

Pr	oduct Type		HUMAN	OTC DRUG	ltem Code	(Source)	NDC:505	80-449
Ro	ute of Admir	nistratio	n ORAL					
Ac	tive Ingred	lient/Ac	tive Moiet	у				
			Ingredient	Name		Basis of St	rength	Strength
AC	ETAMINOPHEN	I (UNII: 36	209ITL9D) (AC	ETAMINOPHEN - UN	II:362O9ITL9D) ACETAMINOPHE	N	500 mg
In	active Ingr	odionte						
		eulents		redient Name			C	trength
C۵	RNAUBA WAX		•				3	trength
	ARCH, CORN (U							
	&C RED NO. 4							
AL		E (UNII: LM	112606933)					
HY	PROMELLOSE,	UNSPEC	IFIED (UNII: 3M	IXW29V3WO)				
MA	GNESIUM STE	ARATE (U	NII: 70097M6I3	0)				
PO	LYETHYLENE	GLYCOL,	UNSPECIFIED	(UNII: 3WJQ0SDW1	Α)			
PO	WDERED CELI	ULOSE (l	JNII: SMD1X3X	D9M)				
	PROPYLENE GLYCOL (UNII: 6DC9Q167V3)							
	ELLAC (UNII: 46							
				JNII: H8AV0SQX4D)				
тп		DE (UNII: 1	SFIX9V2JP)					
Pr	oduct Char	acteris	tics					
			Score	core no score				
Shape		OVAL	Size		19mm			
Flavor				Imprint Code		TYLENOL;500		
Со	ntains							
Pa	ckaging							
#	Item Code		Packag	e Description		Marketing Start Date		eting End Date
1	NDC:50580- 449-00	1 in 1 CA	RTON			08/19/1984		
1	449-00	125 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination			Combination			
2	NDC:50580-	Product 1 in 1 CARTON		08/19/1984				
2	449-05	24 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination						
3	NDC:50580-	Product 2 in 1 POUCH; Type 0: Not a Combination Product			08/19/1984			
3		2 IN 1 PO	och, Type o. i	Not a Combination				
3 4	449-08 NDC:50580-	1 in 1 CA		Not a Combination		08/19/1984		
	449-08	1 in 1 CA	RTON	Not a Combination		08/19/1984		

5		2 in 1 POUCH; Type 0: Not a Combination Product	
6	NDC:50580- 449-11	50 in 1 TRAY	08/19/1984
6		2 in 1 POUCH; Type 0: Not a Combination Product	
7	NDC:50580- 449-13	3 in 1 CARTON	08/19/1984
7		2 in 1 POUCH; Type 0: Not a Combination Product	
8	NDC:50580- 449-14	2 in 1 POUCH; Type 0: Not a Combination Product	08/19/1984
9	NDC:50580- 449-15	10 in 1 VIAL, PLASTIC; Type 0: Not a Combination Product	08/19/1984
10	NDC:50580- 449-23	1 in 1 CARTON	08/19/1984
10		150 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	
11	NDC:50580- 449-31	1 in 1 CARTON	08/19/1984
11		36 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	
12	NDC:50580- 449-34	325 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	08/19/1984
13	NDC:50580- 449-35	1 in 1 CARTON	08/19/1984
13		24 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	
14	NDC:50580- 449-36	1 in 1 CARTON	08/19/1984
14		100 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	
15	NDC:50580- 449-61	1 in 1 CARTON	08/19/1984
15		225 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	
16	NDC:50580- 449-62	325 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	08/19/1984
17	NDC:50580- 449-84	2 in 1 POUCH; Type 0: Not a Combination Product	08/19/1984
18	NDC:50580- 449-85	50 in 1 TRAY	08/19/1984
18		2 in 1 POUCH; Type 0: Not a Combination Product	
19	NDC:50580- 449-86	50 in 1 TRAY	08/19/1984
19		2 in 1 POUCH; Type 0: Not a Combination Product	
20	NDC:50580- 449-87	3 in 1 CARTON	08/19/1984
20		2 in 1 POUCH; Type 0: Not a Combination Product	
21	NDC:50580- 449-12	12 in 1 PACKAGE	10/21/2014
21		10 in 1 VIAL, PLASTIC; Type 0: Not a Combination Product	
22	NDC:50580- 449-96	1 in 1 CARTON	06/25/2018
22		50 in 1 BOTTLE; Type 0: Not a Combination Product	
23	NDC:50580- 449-97	249 in 1 PACKAGE, COMBINATION; Type 0: Not a Combination Product	01/28/2019
24	NDC:50580- 449-98	110 in 1 PACKAGE, COMBINATION; Type 0: Not a Combination Product	01/28/2019

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC monograph not final	part343	08/19/1984				

Labeler - Johnson & Johnson Consumer Inc. (878046358)

Revised: 3/2023

Johnson & Johnson Consumer Inc.