PLUS PHARMA PAIN RELIEVER, FEVER REDUCER- acetaminophen tablet Unit Dose Services

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.

Drug Facts

Active ingredient (in each tablet)

Acetaminophen 325 mg

Purposes

Pain reliever/fever reducer

Uses

for the temporary relief of minor aches and pains due to:

- Headache
- Muscular aches
- Backache
- Minor pain of arthritis
- The common cold
- Toothache
- Premenstrual and menstrual cramps

Temporarily reduces fever.

Warnings

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

- adult takes more than 12 tablets in 24 hours, which is the maximum daily amount
- child takes more than 5 doses in 24 hours
- taken with other drugs containing acetaminophen
- adult has 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 non prescription). 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 the user has

liver disease.

Ask a doctor or pharmacist before use if the user is

taking the blood thinning drug warfarin.

Stop use and ask a doctor if

- Pain gets worse or lasts more than 10 days in adults and children
- Pain gets worse or lasts more than 5 days in children under 12 years
- 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:

Taking more than the recommended dose (overdose) may cause liver damage. 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

AGE	DOSE
Adults and Children 12 years and over	 Take 2 tablets every 4 to 6 hours while symptoms last Do not take more than 12 tablets in 24 hours
Children 6 - 11 years	 Take 1 tablet every 4 to 6 hours while symptoms last Do not take more than 5 tablets in 24 hours
Children under 6 voors	Do not use adult Regular Strength products in children under 6 years of age; this will provide more than the recommended dose (overdose) of acetaminophen and may cause liver damage.

Other information

- Do not use if imprinted Safety Seal under cap is broken or missing
- Store at room temperature

Inactive ingredients

Povidone, Pregelatinized Starch, Sodium Starch Glycolate, Stearic Acid.

Questions?

If you have any questions or comments, or to report an adverse event, please contact (800) 795-9775.

ACETAMINOPHEN 325MG TAB 24EA

NDC: 50436-0703-1 ACETAMINOPHEN 325 MG / 24 TAB

 WARNING: KEEP OUT OF
 LOT: XXXXX
 EXP: XXXXX
 NDC: 50436-0703-1

 ROOM TEMPERATURE. SEE PACKAGE
 Pkg by: Unit Dose Services, LLC
 DRUG: ACETAMINOPHEN 325 MG / 24 TAB

 INSERT FOR DOSAGE INFORMATION.
 Danis, FL 33004
 LOT: XXXX
 EXP: XXXXX



 Active ingredient:
 DIST. BY: PLUS PHARMA,
 LOT: XXXX
 EXP: 3

 (in each tablet)
 COMMACK, NY 11725
 NDC: 50436-0703-1

 Acetaminophen 325 mg.
 MFG NDC: 51645-0703-10
 DRUG: ACETAMINOPHEN 325 MG
 MFG LOT: XXXXX

NDC: 50436-0703-1 DRUG: ACETAMINOPHEN 325 MG / 24 TAB LOT: XXXXX EXP: XXXXX

NDC: 50436-0703-1 DRUG: ACETAMINOPHEN 325 MG / 24 TAB LOT: XXXXX EXP: XXXXX

325 MG / 24 TAB LOT: XXXXX EXP: XXXXX

PLUS PHARMA PAIN RELIEVER, FEVER REDUCER

acetaminophen tablet

Product Inf	ormation						
Product Type	2	HUMAN OTC DRUG	Item Code (Source)	NDC:50436-0703	NDC:50436-0703(NDC:51645-703)		
Route of Adm	inistration	ORAL					
Active Ingre	edient/Active Moi	ety					
Ingredient Name					ength	Strength	
ACETAMINO PHEN (UNII: 36209 ITL9D) (ACETAMINO PHEN - UNII: 36209 ITL9D)					EN	325 mg	
Inactive Ing	redients	Ingredient Na	me			Strength	
DO VIDO NES (I	JNII: FZ989GH94E)	Ingredient Na	me			Strength	
STARCH, COR	N (UNII: O8232NY3SJ)						
	N (UNII: O8232NY3SJ) CH GLYCOLATE TYP	PE A CORN (UNII: AG9B6	65PV6B)				
SODIUM STAR		PE A CORN (UNII: AG9B6	65PV6B)				
SODIUM STAR	CH GLYCOLATE TYP	P E A CORN (UNII: AG9B6	55PV6B)				
SO DIUM STAR STEARIC ACID	CH GLYCOLATE TYP (UNII: 4ELV7Z65AP)	PE A CORN (UNII: AG9B	55PV6B)				
SODIUM STAR STEARIC ACID Product Cha	CH GLYCOLATE TYP (UNII: 4ELV7Z65AP)	PE A CORN (UNII: AG9B6	55PV6B) Scor	e	no s	5C 0 FE	
SODIUM STAR	CH GLYCOLATE TYP (UNII: 4ELV7Z65AP) aracteristics	, , , , , , , , , , , , , , , , , , ,		-	no s 10 m		

Contains			
Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:50436-0703-1	24 in 1 BOTTLE; Type 0: Not a Combination Product	07/04/2016	
Marketing Info	ormation		
Marketing Categor	y Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not fin	al part343	03/27/2006	

Labeler - Unit Dose Services (831995316)

Registrant - Unit Dose Services (831995316)

Establishment

Name	Address	ID/FEI	Business Operations
Unit Dose Services		831995316	REPACK(50436-0703), RELABEL(50436-0703)

Revised: 7/2016

Unit Dose Services