

**ARTHRITIS PAIN RELIEVER- acetaminophen tablet, film coated, extended release**  
**Dispensing Solutions, Inc.**

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**Drug Facts**

**ACTIVE INGREDIENT (IN EACH CAPLET)**

Acetaminophen USP, 650 mg

**PURPOSE**

Pain reliever/fever reducer

**USES**

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

**WARNINGS**

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

- more than 6 caplets in 24 hours, which is the maximum daily amount
- with other drugs containing acetaminophen
- 3 or more alcoholic drinks every day while using this product

**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	take 2 caplets every 8 hours with water swallow whole - do not crush, chew, split or dissolve do not take more than 6 caplets in 24 hours do not use for more than 10 days unless directed by a doctor
under 18 years of age	ask a doctor

**OTHER INFORMATION**

- store at 20 - 25° C (68 - 77° F). Avoid excessive heat 40° C (104° F).
- see end panel for batch number and expiration date (*Applicable only for Bottle Carton*)
- **TAMPER EVIDENT: DO NOT USE IF IMPRINTED SEAL IS BROKEN OR MISSING FROM BOTTLE.**
- **THIS PACKAGE FOR HOUSEHOLDS WITHOUT YOUNG CHILDREN. (*for non CRC packages*)**

**INACTIVE INGREDIENTS**

croscarmellose sodium, hypromellose, magnesium stearate, microcrystalline cellulose, povidone, pregelatinized starch, propylene glycol, sodium lauryl sulfate, stearic acid, titanium dioxide

**QUESTIONS?**

call **1-800-406-7984**

**Contains No Aspirin**

**Keep the carton. It contains important information.**

Distributed by:

Ohm Laboratories Inc.

1385 Livingston Avenue

North Brunswick, NJ 08902

**PRINCIPAL DISPLAY PANEL**

**NDC 66336-0233-XX**

**EASY TO OPEN BOTTLE**

**Use only as directed.**

## See New Warnings Information

Lasts up to 8 hours

Arthritis Pain Relief

Acetaminophen Extended-Release Tablets, USP 650 mg

Pain Reliever/Fever Reducer

For the Temporary Relief of Minor Arthritis Pain

DO NOT USE WITH OTHER MEDICINES CONTAINING ACETAMINOPHEN

100 CAPLETS\*

650 mg EACH

(\*capsule-shaped tablets)

†Compare to the active ingredient of Tylenol® Arthritis Pain

†This product is not manufactured or distributed by McNeil Consumer Healthcare, Inc., The owner of the registered trademark Tylenol® is the Tylenol Company.

<p>BULK SOURCE DATA</p> <p>DIST. BY: OHM LABORATORIES INC. NORTH BRUNSWICK, NJ 08902</p> <p>PRODUCT ID: WHITE CAPSULE-SHAPED TABLET DEBOSSSED cor 116</p> <p>BULK SOURCE NDC: 51660-0333-50 MFR. LOT: XXXXXX PEDIGREE #: 19700293</p> <p>DISPENSE IN THIS TIGHT/LIGHT RESISTANT CONTAINER</p>  <p>Rev. Date: 03/11</p>	 <p><b>ACETAMINOPHEN ER 650 mg</b> <b>XX CAPLETS</b> <b>NDC 66336-0233-XX</b> <b>PRODUCT # 216-XX</b></p> <p>ACTIVE INGREDIENT (IN EACH CAPLET): ACETAMINOPHEN USP . . . . 650 mg (PURPOSES: PAIN RELIEVER/FEVER REDUCER)</p> <p>"LIVER WARNING: THIS PRODUCT CONTAINS ACETAMINOPHEN. SEVERE LIVER DAMAGE MAY OCCUR IF YOU TAKE MORE THAN 6 CAPLETS IN 24 HOURS, WHICH IS THE MAXIMUM DAILY AMOUNT; 3 OR MORE ALCOHOLIC DRINKS EVERY DAY WHILE USING THIS PRODUCT." CONTAINS NO ASPIRIN.</p> <p>COMPARE TO THE ACTIVE INGREDIENT OF TYLENOL ARTHRITIS PAIN</p> <p>LOT# SAMPLE EXP: 00-00 Rx # 23938312</p>	<p>WARNING: KEEP OUT OF CHILDREN'S REACH STORE AT 68°- 77° F. SEE USP.</p> <p>216-XX NDC 66336-0233- XX ACETAMINOPHEN ER 650 mg XX CAPLETS LOT # SAMPLE EXP: 00-00 MN 51660-0333-50 RX# 23938312</p> <p>216-XX NDC 66336-0233- XX ACETAMINOPHEN ER 650 mg XX CAPLETS LOT # SAMPLE EXP: 00-00 MN 51660-0333-50 RX# 23938312</p> <p>216-XX NDC 66336-0233- XX ACETAMINOPHEN ER 650 mg XX CAPLETS LOT # SAMPLE EXP: 00-00 MN 51660-0333-50 RX# 23938312</p>  <p>Packaged Exclusively By: <b>DISPENSING SOLUTIONS</b>  Santa Ana, CA 92704</p>
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## ARTHRITIS PAIN RELIEVER

acetaminophen tablet, film coated, extended release

### Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:66336-233(NDC:51660-333)
Route of Administration	ORAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ACETAMINOPHEN (UNII: 362O9ITL9D) (ACETAMINOPHEN - UNII:362O9ITL9D)	ACETAMINOPHEN	650 mg

### Inactive Ingredients

Ingredient Name	Strength
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
POVIDONE (UNII: FZ989GH94E)	
STARCH, PREGELATINIZED CORN (UNII: O8232NY3SJ)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
SODIUM LAURYL SULFATE (UNII: 368GB5141J)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

### Product Characteristics

Color	white	Score	no score
Shape	OVAL (Capsule Shaped)	Size	19mm
Flavor		Imprint Code	cor116
Contains			

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:66336-233-00	100 in 1 BOTTLE		
2	NDC:66336-233-30	30 in 1 BOTTLE		

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA076200	04/30/2002	

**Labeler** - Dispensing Solutions, Inc. (066070785)

**Registrant** - PSS World Medical, Inc. (101822682)

### Establishment

Name	Address	ID/FEI	Business Operations
Dispensing Solutions, Inc.		066070785	relabel(66336-233) , repack(66336-233)

Revised: 7/2013

Dispensing Solutions, Inc.