



Standard Pharmaceutical Product Information (Rx Product Only)

Introduction Type: Final Version Date: 1/22/2020

Application: ANDA

Company Name: TAGI Pharma, Inc. Application: ANDA

Application Number: NDANDA18BLA (drug); PMA15701X (med device); A208249

DUNS: 063322660

Proprietary Name (if applicable) and Established Name: Azithromycin Tablets, USP

Selling Unit NDC: 51224-122-09 Individual Unit NDC:

UDI CVX Code: MVX Code:

Description: Azithromycin Tablet, USP 500mg 9 Count Blister

Active Ingredient(s): Azithromycin Dihydrate

URL for Additional Product Information: www.tagipharma.com

Address: 722 Progressive Lane

City: South Beloit

Key Contact: Pierce Dewey

Phone Number: (815)824-7885

Product Therapeutic Classification: Antibiotic

Address 2: Room 205

IL: Zip: 61040

Email: codewy@tagipharma.com

Fax: (815)824-4828

ADDITIONAL PRODUCT INFORMATION

Is the Product... a legend device? No Yes

reverse numbered? No Yes

co-licensed? No Yes

Is the Product... Direct And Drop-Ship Unit of Use No Yes

Is the Product... No Yes

If Unit Dose, is item bar coded to unit dose for hospital scanning? No Yes

If Unit Dose NDC, indicate NDC here:

Country of Origin: China

Is this product covered under the Trade Agreements Act (TAA)? No Yes

FOR GENERIC DRUG PRODUCTS

I. Orange Book Rating: AB Authorized Generic *If Authorized Generic, other section fields are not applicable

II. Generic Equivalent to What Brand?: ZITHROMAX

DRUG SUPPLY CHAIN SECURITY ACT (DSCSA) INFORMATION

Does supplier meet DSCSA definition of manufacturer? Yes No

Is product exempt from DSCSA? No Yes

If yes, select exemption: No No No

Other exemption - Write in: No No No

Is product repackaged? No No No

Has FDA granted waiver/exemption for product? No No No

Serialized? No Yes

If not, when? No No No

Items aggregated? No No No

GTIN PRODUCT INFORMATION

Item	Level	Unit	Quantity	GTIN-14
<input checked="" type="checkbox"/>	Box/ Carton/Bundle/Inner Pack	Linear	1	00351224122003
<input checked="" type="checkbox"/>	Case	Linear	100	50351224122008
<input type="checkbox"/>	Pallet	Linear		
<input type="checkbox"/>	Linear	Linear		
<input type="checkbox"/>	Linear	Linear		
<input type="checkbox"/>	Linear	Linear		
<input type="checkbox"/>	Linear	Linear		

PRODUCT DESCRIPTION INFORMATION

Size: 9

Strength: 500mg

Dosage Form: Tablets

Product Shape: oval

Product Color: red

Product Imprint: OE:500

SPECIAL HANDLING AND STORAGE REQUIREMENTS*

a. Temperature - Indicate the USP temperature range for this product. Controlled Room - between 20 and 25 C (68° - 77° F) Temperature Range

Other Temperature Range Requirement (write in)

Is this product to be shipped to customers on ice? No Yes

Is this product to be shipped to customers on dry ice? No Yes

b. Contact for temperature excursion questions:

Name:

Number:

Group E-mail:

Special returns requirements for this product? No Yes

c. Special regulations for product in any states? No Yes

Protect product (unit of sale) from light? No Yes

d. Store product (unit of sale) upright? No Yes

e. Shelf life:

Initial shelf life at launch (if different):

ORDER INFORMATION

Unit of Sale: Bottle Box/ Carton Ampule Glass Tube Vial Liquid Sgl Vial Liquid Multi Vial Powder Sgl Vial Powder Multi Other: Write in Blister

What is the NDC selling unit? (Write-in, e.g. 1 Box of 10 Vials)

Minimum order quantity?

If Yes, how many of which package type? Each Inner/ Carton/ Pack Case

PHARMACY ORDER / BILL UNIT

Rec. sell unit to customer?

Rx billing unit to pharmacy:

(Write-in, e.g. 1 Vial)

ITEM AND PACKING INFORMATION

Item:	Weight Lbs.	Depth	Height	Width	Volume (Cube)	# Pieces:
Box/ Carton/Bundle/ Inner Pack:	0.22	4.25	3.07	2.09	27.289275	1
Case:	22	22.05	16.14	9.84	3501.92808	100
Pallet:						0
UPC:	Case:	Carton:				

COST INFORMATION

Regular Cost

Invoice Cost (NAC) (\$)

Federal Excise Tax Per Unit of Sale

As of date: 1/22/2020

WHOLESALE USE ONLY:

Vendor #:

Whsl. Code #:

Pineline Code:

Signature:

Pierce Dewey

Attach copy of SAFETY DATA SHEET (SDS) or non hazard letter. PACKAGE INSERT, LABEL AND PHOTO OF PRODUCT PACKAGING and BARCODE.

See new p. 3 for Designated Drop Ship Only.

*Please provide any additional information on page 2.



Standard Pharmaceutical Product Information (Page 2)

For Designated Drop Ship Only Products, Please Use Page 3

MATERIAL HAZARD CLASSIFICATION and TRANSPORTATION

Is this product (check all that apply):

a. Cytotoxic? No

b. CA Prop. 65 Carcinogen or Reproductive Toxicant?
 Is the product a CA Prop 65 carcinogen? No
 Is the product a CA Prop 65 reproductive toxicant? No
 Does the product label bear a CA Prop 65 warning? No

c. Contact Hazard? No

d. Does this product require special clean-up instructions?
 (if yes, attach SDS with special instructions.) No

e. Does the product contain DEHP? No

Is this product regulated for shipment by DOT or IATA?
 (if yes, answer a-e below and provide SDS) No

a. UN/Identification Number _____

b. Proper Shipping Name _____

c. DOT Hazard Class _____

d. Packing Group _____

e. Inhalation Hazard? No

Is the product restricted for air shipment? If so, indicate restriction:

Passenger

Cargo

Passenger & Cargo

Is this a reposable quantity? No

RQ Threshold: _____

Is this a marine pollutant? No

Is this product shipped utilizing an authorized DOT exception or Special Permit?
 No (if yes, identify method below)

Limited Quantity

Consumer Commodity, ORM-D

Small Quantity (49 CFR 173.4)

Special Permit: DOT-SP

Special Provision (listed in Column 7 of 49 CFR 172.101); SP# _____

Is there a REMS on this product?
 If Yes, is it managed with a pharmacy registry?
 Website URL: _____

Comments / Details: (For example, IRT/edge program?) _____

REMS: No Yes

REMS Program Manager Name: _____ Phone: _____

Supplier Manages REMS registry exclusively: _____

Wholesale distributor support: _____

Provider Name: _____

Site Enrollment Number assigned by Supplier: _____

DEA #: _____

PCPDP #: _____

NPI #: _____

Comments _____

Registry: _____

Registry Program Contact Name: _____ Phone: _____

Comments _____

RETURN INSTRUCTIONS

Contact tel. # if product received damaged: (855)225-8244

Is product returnable for credit? Yes No

URL/Link to returns policy: _____

CLASS OF TRADE RESTRICTION:

No restriction: Select YES if sold to retail pharmacy, hospitals, clinics and physician offices Yes

Restricted to retail pharmacy only: No

Restricted to hospital, clinics, and physician offices only: No

Restricted from US territories? (explain in comments) No

Comments: _____

REMS or REGISTRY RESTRICTIONS

Is there a REMS on this product? No Yes

Website URL: _____

Comments / Details: (For example, IRT/edge program?) _____

ADD'L STORAGE INFORMATION

Is the Product... No Yes

Controlled Substance? No Yes

Controlled by State(s)? No Yes

ARCOS Reportable? No Yes

Schedule No. (inc. N for non-narcotic) _____

Controlled Substance Code _____

Listed Chemical (List I or II) _____

If yes, indicate which: _____

Is it a scheduled listed chemical product?: No Yes

Hazardous Waste Identification

EPA Hazardous Waste Code: _____

SDS Hazard Classification

Organic Corrosive

Inorganic Oxidizer

Steroid/Androgen Contact Hazard

Aerosol Class: Identify NFPA Storage Level: _____

Is the product a NIOSH hazardous drug? No Yes

If yes, indicate which: _____

MISCELLANEOUS NOTES and/or Image of Product Barcode:

Release DATE _____



Standard Pharmaceutical Product Information (Page 3)

FOR DESIGNATED DROP SHIP PRODUCT ONLY - if not a designated drop ship, do not complete.

Order Method for Designated Drop Ship Product

Purchase orders may be accepted by:

a. EDI Yes No

b. Autofax Yes No

c. Fax Yes No

d. Phone only Yes No

e. Supplier Web Site only Yes No

Minimum Order Quantity: case

Supplier's Customer Service Number:

Contracted 3PL company / contact #:

Standard Order Receipt and Processing

Purchase order daily receipt cut off time by supplier: Central

Cut off time: Central

Shipping lead time of PO: Hours Days

Ships same day for next day receipt: No

Ships for second day receipt: No

Ships regular ground for 3-10 days receipt: Yes

Expedited Freight Charges or Other Designated Drop Ship Fees:

Expedited freight fees billed with each order: Yes No

Drop Ship service fee billed with each order: Yes No

Drop Ship miscellaneous fees billed: Yes No

Comments:

Overnight and Priority Overnight PO Processing

Overnight receipt available: Yes No

PO Receipt cut off time: Central

Days of week overnight is available:

<input checked="" type="checkbox"/>	Monday
<input checked="" type="checkbox"/>	Tuesday
<input checked="" type="checkbox"/>	Wednesday
<input checked="" type="checkbox"/>	Thursday
<input type="checkbox"/>	Friday

Priority Overnight receipt available: Yes No

PO Receipt Cut off time:

Class of Trade Restriction:

No restriction: Select YES if sold to retail pharmacy, hospitals, clinics and physician offices Yes No

Restricted to retail pharmacy only: Yes No

Restricted to hospital, clinics, and physician offices only: Yes No

Restricted from US territories? (explain in comments) Yes No

Comments:

Priority Overnight receipt available:

PO Receipt Cut off time:

Saturday Overnight receipt available: Yes No

PO Receipt Cut off time:

Order receipt method: Yes No

Phone #:

Fax #:

EDI:

Overnight Fees apply: Yes No

Other fees apply:

Other Data Information Required to Process PO:

Patient Procedure Date:

Physician Name:

Physician/Clinic Phone #:

Physician State License #:

Physician/Clinic DEA #:

Physician/Clinic Specialty:

Return Instructions

Contact # if product is received damaged:

Is product returnable for credit: Yes No

URL/Link to returns policy:

Special regulations or returns requirements for this product in certain states? Yes No

If so, which states? Other requirements? Comments?

Georgia, Mississippi, and North Carolina. Requests can be made via fax: (815)624-0608 or email

Miscellaneous Notes:

ADDITIONAL INFORMATION

Is product order for scheduled patient procedure?

Is product order for restocking purposes?

SAFETY DATA SHEET

1. IDENTIFICATION OF THE SUBSTANCE/PREPARATION AND THE COMPANY/UNDERTAKING

Material Name: Azithromycin Tablets

Trade Name: Azithromycin

Chemical Family: Mixture

Intended Use: Pharmaceutical product used as antibiotic agent

Manufacturer Information

Company name: CSPC Ouyi Pharmaceutical Co., Ltd.

Address No.276 Zhongshan West Road Shijiazhuang 050051, China

Telephone + 86-311-87896575

Website www.ouyipharma.com

Emergency phone number +1-877-436-7220

2. HAZARDS IDENTIFICATION

Appearance:

Red, modified oval-shaped, film-coated tablets, debossed with "OE" on one side and "250", "500" or "600" on the other side.

Statement of Hazard: Non-hazardous in accordance with international standards for workplace safety.

Short Term: Dust may cause irritation if tablets are crushed or broken. Individuals sensitive to this chemical or other materials in its chemical class may develop allergic reactions.

Known Clinical Effects: May cause effects similar to those seen in clinical use including transient diarrhea, nausea and abdominal pain.

EU Indication of danger: Not classified

Australian Hazard Classification

(NOHSC):

Non-Hazardous Substance. Non-Dangerous Goods.

Note: This document has been prepared in accordance with standards for workplace safety, which require the inclusion of all known hazards of the product or its ingredients regardless of the potential risk. The precautionary statements and warnings included may not apply in all cases.

Your needs may vary depending upon the potential for exposure in your workplace.

3. COMPOSITION/INFORMATION ON INGREDIENTS

Hazardous

Ingredient CAS Number EU EINECS/ELINCS List Classification %

Azithromycin dihydrate 117772-70-0 Not listed Not Listed 56

Starch, pregelatinized 9005-25-8 232-679-6 Not Listed *

Sodium lauryl sulfate 151-21-3 205-788-1 Not Listed *

Magnesium stearate 557-04-0 209-150-3 Not Listed *

Ingredient CAS Number EU EINECS/ELINCS List Classification %

Calcium phosphate dibasic, anhydrous 7757-93-9 231-826-1 Not Listed *

Croscarmellose sodium 74811-65-7 Not listed Not Listed *

Additional Information: * Proprietary

Ingredient(s) indicated as hazardous have been assessed under standards for workplace safety.

4. FIRST AID MEASURES

Eye Contact: Flush with water while holding eyelids open for at least 15 minutes. Seek medical attention immediately.

Skin Contact: Remove contaminated clothing. Flush area with large amounts of water. Use soap. Seek medical attention.

Ingestion: Never give anything by mouth to an unconscious person. Wash out mouth with water. Do not induce vomiting unless directed by medical personnel. Seek medical attention immediately.

Inhalation: Remove to fresh air and keep patient at rest. Seek medical attention immediately.

Symptoms and Effects of Exposure: For information on potential signs and symptoms of exposure, See Section 2 - Hazards Identification and/or Section 11 - Toxicological Information.

5. FIRE FIGHTING MEASURES

Extinguishing Media: Use carbon dioxide, dry chemical, or water spray.

Hazardous Combustion Products: Emits toxic fumes of carbon monoxide, carbon dioxide, and nitrogen oxides.

Fire Fighting Procedures: Wear approved positive pressure, self-contained breathing apparatus and full protective turnout gear. Use caution in approaching fire.

Fire / Explosion Hazards: Not determined

6. ACCIDENTAL RELEASE MEASURES

Health and Safety Precautions: Personnel involved in clean-up should wear appropriate personal protective equipment (see Section 8). Minimize exposure.

Measures for Cleaning / Collecting: Contain the source of spill if it is safe to do so. Collect spilled material by a method that controls dust generation. A damp cloth or a filtered vacuum should be used to clean spills of dry solids. Clean spill area thoroughly.

Measures for Environmental

Protections:

Place waste in an appropriately labeled, sealed container for disposal. Care should be taken to avoid environmental release.

Additional Consideration for Large

Spills:

Non-essential personnel should be evacuated from affected area. Report emergency situations immediately. Clean up operations should only be undertaken by trained personnel.

7. HANDLING AND STORAGE

General Handling: Minimize dust generation and accumulation. If tablets or capsules are crushed and/or broken, avoid breathing dust and avoid contact with eyes, skin, and clothing. When handling, use appropriate personal protective equipment (see Section 8). Wash hands and any exposed skin after removal of PPE. Releases to the environment should be avoided. Review and implement appropriate technical and procedural waste water and waste disposal measures to prevent occupational exposure or environmental releases. Potential points of process emissions of this material to the atmosphere should be controlled with dust collectors, HEPA filtration systems or other equivalent controls.

Storage Conditions: Store as directed by product packaging.

8. EXPOSURE CONTROLS / PERSONAL PROTECTION

Refer to available public information for specific member state Occupational Exposure Limits.

Analytical Method: Analytical method available for Azithromycin. Contact Pfizer Inc for further information.

Engineering Controls: Engineering controls should be used as the primary means to control exposures. General room ventilation is adequate unless the process generates dust, mist or fumes. Keep airborne contamination levels below the exposure limits listed above in this section.

Environmental Exposure Controls: Refer to specific Member State legislation for requirements under Community environmental legislation.

10 mg/m³ TWA

Spain OEL - TWA Listed

Calcium phosphate dibasic, anhydrous

Latvia OEL - TWA Listed

Australia TWA 10 mg/m³

Sodium lauryl sulfate

Azithromycin dihydrate

Pfizer OEL TWA-8 Hr: 0.3 mg/m³

Belgium OEL - TWA

Magnesium stearate

Listed

ACGIH Threshold Limit Value (TWA) 10 mg/m³ TWA

Australia TWA 10 mg/m³

Bulgaria OEL - TWA Listed

Belgium OEL - TWA Listed

Ireland OEL - TWAs Listed

Pfizer OEL TWA-8 Hr:

Lithuania OEL - TWA Listed

Czech Republic OEL - TWA Listed

Portugal OEL - TWA Listed

500µg/m³

Spain OEL - TWA Listed

Greece OEL - TWA

Sweden OEL - TWAs Listed

Listed

Ireland OEL - TWAs Listed

Starch, pregelatinized

Revision date: 08-Sep-2009

OSHA - Final PELs - TWAs: 15 mg/m³ total

5 mg/m³

8. EXPOSURE CONTROLS / PERSONAL PROTECTION

Personal Protective Equipment: Refer to applicable national standards and regulations in the selection and use of personal

protective equipment (PPE).

Hands: Impervious gloves are recommended if skin contact with drug product is possible and for bulk processing operations.

Eyes: Wear safety glasses or goggles if eye contact is possible.

Skin: Impervious protective clothing is recommended if skin contact with drug product is possible and for bulk processing operations.

Respiratory protection: If the applicable Occupational Exposure Limit (OEL) is exceeded, wear an appropriate respirator with a protection factor sufficient to control exposures to below the OEL.

9. PHYSICAL AND CHEMICAL PROPERTIES

Polymerization: Will not occur

10. STABILITY AND REACTIVITY

Stability: Stable under normal conditions of use.

Conditions to Avoid: Fine particles (such as dust and mists) may fuel fires/explosions.

Incompatible Materials: As a precautionary measure, keep away from strong oxidizers

11. TOXICOLOGICAL INFORMATION

General Information: The information included in this section describes the potential hazards of the individual ingredients.

Acute Toxicity: (Species, Route, End Point, Dose)

Acute Toxicity Comments: A greater than symbol (>) indicates that the toxicity endpoint being tested was not achievable at the highest dose used in the test.

Irritation / Sensitization: (Study Type, Species, Severity)

Rat Oral LD50 1288 mg/kg

Color: White to off-white

Azithromycin dihydrate

Mouse (F) Oral LD50 4000 mg/kg

Molecular Formula: Mixture

Mouse (M) Oral LD50 3000 mg/kg

Rat Oral LD50 > 2000 mg/kg

Molecular Weight: Mixture

Magnesium stearate

Sodium lauryl sulfate

Rat Oral LD50 > 2000 mg/kg

Eye Irritation Rabbit Severe

Rat Inhalation LC50 > 2000 mg/m³

Physical State: Film-coated tablets

Sodium lauryl sulfate

11. TOXICOLOGICAL INFORMATION

Repeated Dose Toxicity: (Duration, Species, Route, Dose, End Point, Target Organ)

Reproduction & Developmental Toxicity: (Study Type, Species, Route, Dose, End Point, Effect(s))

Genetic Toxicity: (Study Type, Cell Type/Organism, Result)

Carcinogen Status: None of the components of this formulation are listed as a carcinogen by IARC, NTP or OSHA.

12. ECOLOGICAL INFORMATION

Environmental Overview: In the environment, the active ingredient in this formulation is expected to mainly reside in the aquatic environment and slowly degrade.

Mobility, Persistence and

Degradability:

Azithromycin half life < 28 days (Aerobic Biodegradation - Water)

Bioaccumulation and Toxicity: The active ingredient in this formulation has low potential to bioaccumulate and long-term adverse effects to aquatic organisms are not expected. See aquatic toxicity data, below.

Aquatic Toxicity: (Species, Method, End Point, Duration, Result)

Aquatic Toxicity Comments: A greater than (>) symbol indicates that acute ecotoxicity was not observed at the maximum solubility. Since the substance is insoluble in aqueous solutions above this concentration, an acute ecotoxicity value (i.e. LC/EC50) is not achievable.

Bacterial Inhibition: (Species, Method, End Point, Duration, Result)

In Vitro Cytogenetics Human Lymphocytes Negative

1 Month(s) Rat Intravenous 5 mg/kg/day NOEL Liver

1 Month(s) Dog Intravenous 5 mg/kg/day NOEL Liver

Antigenicity- Passive cutaneous anaphylaxis Rabbit Negative

Azithromycin dihydrate

Antigenicity- Passive cutaneous anaphylaxis Mouse Negative

Reproductive & Fertility Rat Oral 10 mg/kg/day NOEL Fertility

Skin Irritation Rabbit Severe

Prenatal & Postnatal Development Mouse Oral 40 mg/kg/day NOEL Not Teratogenic

Azithromycin dihydrate

Prenatal & Postnatal Development Rat Oral 40 mg/kg/day NOEL Not Teratogenic

Daphnia Magna OECD EC50 48 Hours 120 mg/L

Azithromycin dihydrate

Hyallela azteca OECD LC50 96 Hours > 120 mg/L

Azithromycin dihydrate

Rainbow Trout OECD LC50 96 Hours > 84 mg/L

Azithromycin dihydrate

Green Algae OECD EC50 72 Hours 0.0037 mg/L

6 Month(s) Rat Oral 10 mg/kg/day LOEL Liver

Bacterial Mutagenicity (Ames) *Salmonella* Negative

In Vivo Cytogenetics Mouse Lymphoma Negative

6 Month(s) Dog Oral 10 mg/kg/day LOEL Liver

In Vitro Cytogenetics Mouse Negative

Antigenicity- Active anaphylaxis Guinea Pig Negative

12. ECOLOGICAL INFORMATION

13. DISPOSAL CONSIDERATIONS

Disposal Procedures: Dispose of waste in accordance with all applicable laws and regulations. Member State specific and Community specific provisions must be considered. Considering the relevant known environmental and human health hazards of the material, review and implement appropriate technical and procedural waste water and waste disposal measures to prevent occupational exposure and environmental release. It is recommended that waste minimization be practiced. The best available technology should be utilized to prevent environmental releases. This may include destructive techniques for waste and wastewater.

14. TRANSPORT INFORMATION

Not regulated for transport under USDOT, EUADR, IATA, or IMDG regulations.

15. REGULATORY INFORMATION

EU Indication of danger: Not classified

OSHA Label:

Non-hazardous in accordance with international standards for workplace safety.

Canada - WHMIS: Classifications

WHMIS hazard class:

None required

This product has been classified in accordance with the hazard criteria of the CPR and the MSDS contains all of the information

required by the CPR.

Starch, pregelatinized

Inventory - United States TSCA - Sect. 8(b) Listed

Trichoderma viride (Fungus) OECD MIC > 1000 mg/L

Australia (AICS): Listed

Material Name: Azithromycin Tablets

REACH - Annex IV - Exemptions from the obligations of Register:

Present

Clostridium perfringens (Bacterium) OECD MIC 2.0 mg/L

EU EINECS/ELINCS List 232-679-6

Calcium phosphate dibasic, anhydrous

Bacillus subtilis (Bacterium) OECD MIC 2.0 mg/L

Azithromycin dihydrate

Aspergillus niger (Fungus) OECD MIC > 1000 mg/L

15. REGULATORY INFORMATION

16. OTHER INFORMATION

Data Sources: Publicly available toxicity information. Safety data sheets for individual ingredients.

Issue Date: 09/2017

Prepared by: CSPC Ouyi Pharmaceutical Co., Ltd.

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