FDA-2021QZSXRYMI355G6ZICIKKH

Republic of the Philippines Department of Health



Civic Drive, Filinvest Corporate City, Alabang, Muntinlupa City



CERTIFICATE OF PRODUCT REGISTRATION

Pursuant to the provisions of Republic Act (R.A.) No. 3720 as amended, known as the Foods, Drugs, Devices and Cosmetics Act, and consistent with R.A. No. 6675, known as the Generics Act of 1988, and R.A. No. 9711, otherwise known as the Food and Drug Administration Act of 2009, the product described hereunder has been found to conform with the requirements and standards for marketing authorization of pharmaceutical products per existing regulations in force as of date hereof.

Registration Number : DR-XY39072

Generic Name

Brand Name

Dosage Strength & Form

Pharmacologic Category Classification

Approved Shelf-life

Storage Condition

Packaging

Manufacturer

Importer

Distributor

Glutathione

Tad-600 : 600 mg Lyophilized Powder for Injection (I.M./I.V.)

Antidote

: Prescription (Rx) Drug

: 36 months

: Store at temperatures not exceeding 25°C. Protect

from light.

: One (1) box contains: Type III clear and colorless glass vial x 10's + 4 mL (net content) Type I clear

and colorless glass ampoule x 10's (as diluent) Biomedica Foscama Industria Chimico-Farmaceutica

S.P.A.

Via Morolense, 87 03013 Ferentino, Italy : 2 World Traders Subic, Inc.

Building, 202, Lot C-3 Gateway Hub Unit Commercial Area, Subic Bay Gateway Park, Phase I,

Subic Bay Freeport Zone, Olongapo, Zambales

: 2 World Traders Subic, Inc.

Unit 202, Lot Gateway Hub Building, Commercial Area, Subic Bay Gateway Park, Phase I,

Subic Bay Freeport Zone, Olongapo, Zambales

The marketing authorization shall be valid until 16 November 2024 subject to the conditions listed on the reverse side. No change in the formulation, labelling and commercial presentation of this product shall be made at any time during the effectivity of this registration without prior written approval of this Office.

This marketing authorization is subject to suspension, cancellation or recall should any violation of R.A. No. 3720, R.A. No. 6675 and R.A. No. 9711 and/or regulations issued thereunder involving the product be committed.

Witness My Hand and Seal of this Office, this 07 September 2021.

By Authority of the Director General Per FDA Order No. 2016-005

JESUSA JOYCE N/CIRUNAY, RPh Director IV

Center for Drug Regulation and Research

REG. STATUS AMOUNT OR NUMBER

BAR CODE DOC TRACK

Initial [from Monitored Release] Php 16,680.00 1179349 18-Sep-2019







Registration Number

DR-XY39072

SPECIAL CONDITION:

Provided that nothing in the registration of the product herein granted shall be interpreted or construed as an endorsement or representation by FDA, that Registrant has the right or privilege to the use of the name or brand so registered; Registrant hereby agrees and affirms to indemnify and/or hold FDA free and harmless against any and all third-party claims on infringement of patent, trademark or intellectual property right arising from the registration of the product.

	A	This is subject to batch notification.
	В	This is subject to lot release certification.
	C	This is subject to compliance with the requirements under FDA Circular No. 2013-004 for Monitored Release (MR) drug products.
X	D	Subject to post-marketing surveillance of the marketing authorization holder's strict compliance to the Generic Labeling Requirements following the applicable provisions of A.O No. 2016-0008 for drug products for human use and A.O. No. 105 s. 1991 for veterinary drug products.
7 6	E	Submit a satisfactory Bioequivalence Study Report or Biowaiver (whichever is applicable) within the validity of this CPR in accordance with FDA Circular No. 2016-019.
	F	Dangerous Drug - To be prescribed by a PDEA S-2 licensed practitioner in a DOH (yellow) prescription form. It is a habit-forming drug.
	G	Dangerous Drug - To be prescribed by a PDEA S-2 licensed practitioner in a personalized ordinary prescription form. It is a habit-forming drug.
Call Trans	Н	Patient Information Leaflet - Appropriate information for the consumers shall be written in Filipino and/or local dialects, as appropriate.
	I	Submit a Certificate of Good Manufacturing Practice (GMP) Compliance of Foreign Drug Manufacturer(s) within the validity of this CPR in accordance with A. O. No. 2013-0022 and FDA Circular No. 2014-016.
	J	Review of the submitted Bioequivalence Study Report or Biowaiver, whichever is applicable, shall be completed by the FDA within the validity of this CPR; correspondingly, this CPR shall be revoked if product interchangeability has not been established.
X	K	Subject to satisfactory compliance to the post-approval commitments detailed in the letter accompanying this CPR.

REMARKS: