



Republic of the Philippines  
Department of Health  
**FOOD AND DRUG ADMINISTRATION**



**FDA ADVISORY**  
No. 2017-120-A

19 DEC 2022

**TO: ALL HEALTHCARE PROFESSIONALS AND THE GENERAL PUBLIC**

**SUBJECT: Lifting the Advisory on the Registered Food Product "RELUMINS Advance Nutrition Reduced L-Glutathione" under FDA Advisory No. 2017-120 with Subject "Public Health Warning Against the Purchase and Consumption of the Unregistered Food Products and Food Supplements"**

The Food and Drug Administration (FDA) informs the public that the food product **RELUMINS ADVANCE NUTRITION GLUTA 1000 BRAND Reduced L-Glutathione Complex With ALA & Rosehips Dietary Supplement Capsule** is registered by the Market Authorization Holder (MAH) **RSG FLAWLESS BEAUTY INTERNATIONAL INC.**, in accordance to existing FDA rules and regulations.



Figure 1. The Approved label of RELUMINS ADVANCE NUTRITION GLUTA 1000 BRAND Reduced L-Glutathione Complex With ALA & Rosehips Dietary Supplement with registration number **FR-4000007710175** issued on 22 April 2021 valid until **17 April 2025**.





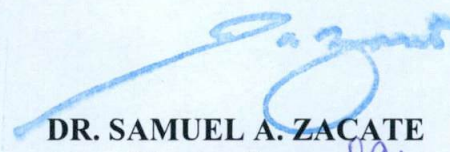

Accordingly, the list released in FDA Advisory No. 2017-120 is hereby updated to remove the aforementioned food supplement.

The public is advised to always check if a food product or food supplement is registered with the FDA by using the **FDA Verification Portal feature** accessible at <https://verification.fda.gov.ph>. You may also look for the FDA Registration number on the product label, if available or simply type the name of the product.

To report any sale or distribution of unregistered food products, kindly email us through [ereport@fda.gov.ph](mailto:ereport@fda.gov.ph).

The public health warning imposed on the remaining products listed in FDA Advisory No. 2017-120 shall remain to be upheld and shall not be affected by the issuance of this advisory. Furthermore, the issuance of this advisory shall not in any manner preclude this Office from issuing subsequent orders it may deem necessary and appropriate, should there be findings of any violation of the company to the existing laws, rules, and regulations.

Dissemination of this advisory to all concerned is hereby requested.

  
**DR. SAMUEL A. ZACATE**  
Director General 



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