



FDA ADVISORY
No. **2021-2120**

11 8 AUG 2021

TO: GENERAL CONSUMING PUBLIC

SUBJECT: Public Health Warning Against the Purchase and Use of Cosmetic Product KEVLO CLARIFYING LOTION Containing Banned Ingredient/s

The Food and Drug Administration (FDA) warns the public from purchasing and using the adulterated cosmetic product **KEVLO CLARIFYING LOTION** which tested positive for the presence of **TRETINOIN**. Such ingredient is not allowed to be part of a cosmetic product as per Annex II Part 1 of the ASEAN Cosmetic Directive. As per Administrative Order No. 13 s. 1999, products containing Tretinoin (Retinoic Acid) are classified as home remedy, over-the-counter, or prescription drug. The table below indicates the particulars of the adulterated cosmetic product/s:

PRODUCT DETAILS	PRODUCT IMAGE/S
Name of Manufacturer/ Local Company Responsible for Placing the Product in the Market: Kevlo Skinceuticals, Inc.	
Address: Landing, Catarman, Liloan, Cebu	
Country of Manufacture: Not Indicated	
Lot/Batch Number: CL-B-067 Mfg. Date: Not Indicated Exp. Date: Not Indicated	

The aforementioned adulterated product is found to be non-compliant with the existing standards, and, thus pose potential hazards to the consuming public. The use of adulterated cosmetic products may result to adverse reactions, including but not limited to, skin irritation, itchiness, anaphylactic shock and organ failure.



In light of the foregoing, the public is advised not to purchase the aforementioned adulterated cosmetic product. Always check if a product is notified with the FDA by using the FDA Verification Portal feature accessible at <https://verification.fda.gov.ph> which may be used by typing in the name of the product before the purchase and/or using the cosmetic products.

All concerned establishments are warned not to distribute adulterated cosmetic products.

All FDA Regional Field Offices and Regulatory Enforcement Units, in coordination with law enforcement agencies and Local Government Units, are requested to ensure that violative products are not sold or made available in the market or areas of their jurisdiction.

To report any sale, distribution, complaint and/or adverse event on the use of the violative cosmetic product, the online reporting facility, eReport can be accessed at ereport@fda.gov.ph, or call us at the Center for Cosmetics and Household/Urban Hazardous Substances Regulation and Research (CCHUHSRR) hotline **(02) 8857-1900 loc. 8113 or 8107**.

Dissemination of this advisory to all concerned is hereby requested.


ROLANDO ENRIQUE D. DOMINGO, MD
Director General

DTN 20210816104505