

1769021	-3566
1/0/021	-3300

CERTIFICATE OF COMPLIANCE

PRODUCT DESCRIPTION : EXCIPIAL [®] - LIPOCREME 50g						
BATCH NUMBER : 1769021						
MANUFACTURING SITE : LABORATOIRES GALDERMA, ZI MONTDESIR, ALBY SUR CHERAN, 74540, FRANCE						
AUTHORIZATION NUMBER: M 20/018 - February 12, 2020						
IMPORTING COUNTRY (DESTINATION) : FRANCE						
FORMULA NUMBER : 0769		MANUFACTURING DATE (MM/DD/YYYY) :	10/02/2021			
PRODUCT CODE: 023566		SHELF LIFE : 36 MONTHS				
		EXPIRY DATE (MM/YYYY) : 09/2024				
REFERENCE SPECIFICATION : MT.09.DOC.1433.R	00.1	ANALYSIS TYPE : Full				
ANALYSIS NUMBER : B-20211007-00042						
RELEASED UNITS QUANTITY : 13419 Units						
MEAN FILL (>= Nominal value of label claim AC.02.SOP.5005)	: 50.6 g					
TESTS	METHODS	SPECIFICATIONS	RESULTS			
Macroscopic appearance	Visual examination					
Form		Semi solid emulsion	Complies			
Color Other Properties		White Smooth, supple	Complies Complies			
Odor	Smell Examination	Characteristic (perfumed)	Complies			
Structure	Visual examination (Microscope)	No crystal visible, shiny structure <= 50μm visible	corresponds			
Total aerobic microbial count (TAMC)	Ph. Eur 2.6.12/USP 1111,61/JP G4,4.05*	NMT 100 CFU/g	< 100 CFU/g			
Total combined yeast/mould count (TYMC)	Ph. Eur 2.6.12/USP 1111,61/JP G4,4.05*		< 10 CFU/g			
Specific microorganisms	Ph. Eur 2.6.13/USP 1111,62/JP G4,4.05*					
Pseudomonas aeruginosa		Absence per gram	Absence per gram			
Staphylococcus aureus	urrant adition	Absence per gram	Absence per gram			

*All pharmacopoeia methods are carried out according to the current edition.

GALDERMA

EST. 1981



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NOTES : N/A						

Certification :

I hereby certified that above information is authentic and accurate.

This batch has been manufactured (including packaging and quality control), at the above mentioned site in full compliance with the GMP requirements of the local regulatory authorities and with the specification in the marketing authorization of the importing country.

The batch processing, packaging and analysis records were reviewed and found to be in compliance with the GMP. All major processes (manufacturing and analytical) have been validated. All deviation or changes have been properly investigated, evaluated and approved by authorized personal.

This document has been electronically signed by a qualified person :

WEIBEL Marie Date : October 18, 2021 GALDERMA