

	CERTIFICAT DE CONFORMITE	DATE
		03/06/19
		VERSION N°1

POUR LE CLIENT

NUV

NOVOMA SARL
 BATIMENT ZEPHYR AVENUE BERNARD
 31 400 TOULOUSE

Je soussigné M. WACRENIER, Président de Laboratoire PHYTOCOSMA SAS certifie que le produit cité ci-après est conforme aux spécifications établies.

Nous vous rappelons qu'il vous appartient de vérifier les conditions de distribution et d'utilisation de ces produits conformément à la législation en vigueur.

Code interne	NUVVIT03
Désignation interne	VITAMINE C LIPOSOMALE QUALI C 90 GVT0T

Code client	
Désignation client	VITAMINE C LIPOSOMALE

Numéro de lot	D16893	Numéro de BL	20882
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Date de fabrication	19/10/2023	DDM	10/2025
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Conditions de conservation	A conserver à l'abri de l'oxygène et de la lumière à une température comprise entre 15 et 25°C dans son emballage d'origine
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Le produit contient de(s) Allergène(s)	Non
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Liste des allergène(s) dans le produit

Non applicable

Le produit contient de(s) Additif(s)	Non
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Liste de(s) additif(s) dans le produit

Non applicable

Le produit est BIO	Non
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(*Produit issu de l'agriculture biologique FR-BIO-01 et process conforme à la fabrication de produits biologiques)

Le produit est sans OGM	Oui	Le produit est Ionisé	Non
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Le produit est sans Gluten	Non
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Conforme Végétarien	Oui	Conforme Végétalien	Oui
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Conforme Halal	Oui	Conforme Casher	Oui
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CONDITIONNEMENT PRODUIT

Type de produit	Gélules HPMC	Couleur	Transparente
PV interne gélules	24854	Taille	Taille 0
Lot Fournisseur gélules	1-000536	Dosage	515mg

Type de conditionnement	Piluliers	Quantité par colis	61*50
Quantité conditionnée	3065	Cartons incomplets	1*15

Autres (dont fond de bol)

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COMPOSITION PRODUIT

INGREDIENT(S)	DOSAGE	PV interne	N°LOT FOURNISSEUR
Vitamine C liposomale Quali C	475mg	24213+24885	22062022-00000-25000+26102022-00000-25000
Amidon de riz	40mg	24518	2320302850

24/10/23

Service qualité

PHYTICOSMA Laboratoires
 23 Route du Burgaud
 82600 AUCAMVILLE
 Tél. : 05 63 02 71 07
 phytocosma@orange.fr
 SAS au capital de 13 720,41 euros
 RCS MONTAUBAN 432 694 354

P 2603



Natural
VITAMINS

CERTIFICATE OF ANALYSIS

Product and Batch Informations

SUNFLOWER LIPOSOMIAL VITAMIN C

REF : VITCLIPO_1002

DT V05 - 21/12/2021

Batch	22062022-00000-25000	Origin (natural/synthetic)	Natural
N° CAS	/	Country of manufacturing	Europe
MF date	22/06/2022	Expiration date	21/06/2024

ANALYSIS ITEM	SPECIFICATION	RESULT	TEST METHOD
Active Ingredients/Substance to control			
Assay	NLT 67,5% Vitamin C	Complies	Volumetric determination
Physical/Chemical Control			
Appearance	White to creamy powder	Complies	Visual inspection
Sieve analysis	NLT 90,0% through 300 microns	Complies	Circular Vibrating Screener
Loss on drying	NMT 5%	Complies	105°C - 3 hours
Contaminant Control*			
Heavy metals	NMT 10ppm	Complies	Atomic Absorption
Lead (Pb)	NMT 3ppm	Complies	Atomic Absorption
Arsenic (As)	NMT 1ppm	Complies	Atomic Absorption
Cadmium(Cd)	NMT 1ppm	Complies	Atomic Absorption
Mercury (Hg)	NMT 0,1ppm	Complies	Atomic Absorption
Microbiological Control			
Total aerobic microbial	NMT 20 000 cfu/g	Complies	As per Eur.Ph
Tot. yeast and mould	NMT 200 cfu/g	Complies	As per Eur.Ph
Enterobacteriaceae*	NMT 100 cfu/g	Complies	As per Eur.Ph
Salmonella*	Negative/25g	Complies	As per Eur.Ph
E.Coli*	Negative/g	Complies	As per Eur.Ph
Statements			
Allergens	Allergen free		
GMO	No OGM		
Irradiation	No irradiation		
BSE/TSE	BSE/TSE Free		
Nanoparticles	Nanoparticles Free		
Vegans/Vegetarians	Suitable for vegans/vegetarians		
Packing and Storage			
Packing	Suitable for food industry		
Storage	Store in dry places and keep away from strong direct light and heat.		

*According to a control plan



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01/07/2022
ALLAIRE Laurène
Quality Department

Certificate of Analysis

General Information

Product	Remy O DR6	Production Date	12/02/2023	(dd/mm/yyyy)
Batch	2320302850	Best before	11/02/2027	(dd/mm/yyyy)
Issued by	Quality Control Management	Date CoA Issued	08/03/2023	(dd/mm/yyyy)

Results of analyses

Parameter	Result	Unit	Method ⁽¹⁾	LSL	USL
Physical and Chemical Parameters					
Moisture	7	g/100g	ISO 712	≤	14
Protein content (N*6,25) on DM	3,6	g/100g d.m.	ISO 1871 ⁽¹⁾	≤	6,0
Ash content on DM	0,2	g/100g d.m.	ISO 3593	≤	1,0
Rheological Parameters					
Starting gel point, pH as is, 6%	83	°C	Brabender	≥	60
End viscosity, pH as is, 6%	584	BU	Brabender	≥	500
Microbiological Parameters					
Salmonella (/375g)	Negative	/375g	ISO 6579		Negative
Total mesophilic bacteria (aerobic)	100	cfu/g	ISO 4833	≤	100.000
Yeasts and Moulds	<10	cfu/g		≤	1.000
Enterobacteriaceae	<10	cfu/g		≤	100

⁽¹⁾ or (acknowledged and) validated equivalent

Remarks

We herewith confirm that the product complies with the corresponding guarantees listed in its Product Sheet .

Rice starch issued from organic farming, Certisys BE-BIO-01 , Certified Organic NOP by bio.inspecta AG

CERTIFICATE OF ANALYSIS EMPTY HARD CAPSULES OF VEGETABLE ORIGIN

CUSTOMER: GOCAPS GMBH			
LOT No.: K2305001315	PRODUCT CODE: K00003	SIZE: 0	
PURCHASE ORDER NUMBER: PO2000392	CHARGE No.: 1-000536	ART No.: 56-000107	
CAPSULE COLOR / CODE: CAP - NATURAL 1-0K / BODY - NATURAL 1-0K			
PRINT: N/A	TEXT: N/A	INK COLOR: N/A	

THIS IS TO CERTIFY THAT: The hard capsules of vegetable origin (K-CAPS) manufactured by C.I. FARMACAPSULAS S.A.S. are made from cellulose ethers, which are polymers derived from vegetable sources. Our capsules are certified as Kosher and Halal, and meet all requirements of current European Pharmacopoeia (EP) and United States Pharmacopoeia (USP). Hypromellose used in the manufacturing of capsules meet specifications as described in the current United States Pharmacopoeia. Cellulose ethers are considered as Generally Recognized As Safe (GRAS) by the FDA.

(% Ingredients to % Cellulose)

Cap	%	Body	%
<p><i>Colorant and ingredients used in capsules are officially approved for use as dye in Foods, Drugs and Cosmetics and/or Drugs and Cosmetics, in the country of destination.</i></p> <p><i>Some changes in color are due to natural colorants or can occur/are within the specification. The above specifications apply to all capsules having the same size and code numbers, unless otherwise stipulated.</i></p>			

Date of Manufacture: 2023-05

Expiration Date: 2028-05

CRITERIA	METHOD / REFERENCE	SPECIFICATIONS	RESULTS
PHYSICAL			
Average Capsule Weight	DCC-MI-P003/ USP<2091>	103.00-115.00 mg	108.0
Loss on drying	DCC-MA-P027	4.00-8.00 %	4.6
Disintegration	DCC-MA-P063/ USP<701>	N.M.T. 15 min	PASSES
Residue on Ignition *	USP	N.M.T. 1.5% Transparent Capsules	PASSES
		N.M.T. 6.0% Colored Capsules	
Identification of HPMC	DCC-MA-P073 / USP	Meets USP Requirements	PASSES
ANALYTICAL			
Arsenic *	EXTERNAL	N.M.T. 0.8 ppm	PASSES
Chromium *	EXTERNAL	N.M.T. 2 ppm	PASSES
Cadmium *	EXTERNAL	N.M.T. 0.5 ppm	PASSES
Lead *	EXTERNAL	N.M.T. 0.5 ppm	PASSES
Mercury *	EXTERNAL	N.M.T. 0.1 ppm	PASSES
Cobalt *	EXTERNAL	N.M.T. 5.0 ppm	PASSES
Vanadium *	EXTERNAL	N.M.T. 10.0 ppm	PASSES
Nickel *	EXTERNAL	N.M.T. 20.0 ppm	PASSES
Total Aerobic Microbial Count	DCC-MA-P031 / USP <61>	N.M.T. 1000 cfu/g	10
Total Yeasts and Molds Count	DCC-MA-P040 / USP <61>	N.M.T. 100 cfu/g	<10
Total Coliforms	DCC-MA-P036 / USP <62>	Absence / 1 g	Absence
Salmonella	DCC-MA-P039 / USP <62>	Absence / 10 g	Absence
Escherichia Coli	DCC-MA-P036 / USP <62>	Absence / 1g	Absence
Staphylococcus aureus	DCC-MA-P037 / USP <62>	Absence / 1g	Absence
Pseudomonas aeruginosa	DCC-MA-P033 / USP <62>	Absence / 1g	Absence

*Reduced Frequency Testing

Storage Conditions: Temperature: 15°C - 30°C / Relative Humidity: 35 % - 70 % RH N.M.T. = No More Than

NOTE: The VISUAL QUALITY is superior to the established figures in the sampling plans of the ANSI/ASQ Z1.4-2013 "Procedure of sampling to inspect for attributes", using simple sampling with level III of General Inspection and Acceptable Level of Quality (AQL) of 0.010 for Critical defects, 0.040 for Major defects and 0.250 for Minor defects. They also fulfill the specifications established in the Technical Information Manual in force.


Quality Assurance

Date: 2023/06/08

Code: DCC-032G (Valid since November 1st, 2021)
Edition 7



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