

qsp 100 %

5.0000 %

3660214

PO24000003

# CERTIFICATE OF ANALYSIS

Page: 1 of 2

The capsules are produced under carefully controlled conditions. Controls are performed continuously throughout the process and guarantee that capsules conform to the highest quality standards. The capsules described below conform to the specifications as defined in the current edition of the Capsugel "Technical Reference File".

Empty DRcaps® Capsules PRODUCT DESCRIPTION

Euromar Ltd Customer:

Customer Reference: DRCAPS SIZE 0 TRANSPARENT Product Name:

Size 0, Coni-Snap, Standard Product Size: 001389.85 Product Code:

Lot Number:

02-Feb-2024 Manufacturing Date: 01-Feb-2029 Expiration Date:

CAP BODY

V43.700 Code: V43.700 Code: NATURAL TR. V700 Name:

NATURAL TR. V700 Name: Non-Print Print Type:

Cap Composition **Body Composition** Hypromellose qsp 100 % Hypromellose Gellan gum 5.0000 % Gellan gum

Due to the nature of raw materials, their sourcing, and technology improvements, the colorant composition data indicated are target values and actual values may vary to insure the consistency of lot color. Capsugel supports the expiry date if precautions for warehousing and transportation are observed (recommended: 15°C - 25°C and 35% - 65% relative humidity).

observed (recommended: 15°C - Ingredient / Reference Hypromellose Gellan gum	E Nr E464 E418	C.I. Nr	Function Structure Gelling agent	(EU)	ilatory References 231/2012, EP, JP, USP/NF, CHP 231/2012, JSFA, 21 CFR
ANALYTICAL DATA		Test Method	Units	Specifications	Re

ANALYTICAL DATA Characteristics Identification of Hypromellose Average weight Loss on drying Sulphated ash Iubricant content Total Aerobic Microbial Count Escherichia coli Salmonella Staphylococcus aureus Total Yeasts/Moulds Count Pseudomonas aeruginosa	Test Method TRF 002C TRF 100A TRF 101A TRF 200A TRF 202B TRF 500A TRF 550A TRF 550A TRF 550A TRF 540A	Units  mg % % % cfu/g	Specifications Positive 87 to 99 Not more than 9.0% Less than 6 Less than 0.5% Less than 1000 Absence in 1 gram Absence in 1 gram Less than 100 Absence in 1 gram Less than 100 Absence in 1 gram	pass * 93.3 4.6 pass * 0.04 * <20 pass * pass * pass * pass * pass *
Customer specific tests Identification of gellan gum Disintegration time	TRF 008A TRF 310A	min/sec	Positive Not less than 30 minutes	pass * pass *

<sup>\*</sup> Reduced frequency testing

#### Elemental Impurities / Heavy Metals

With reference to ICH Q3D and other applicable standards controlling levels of elemental impurities in drug products and food supplements, Capsugel empty capsule products are meeting below levels of applicable elements. Monitoring testing is in place under validated methods, as described in the current edition of Capsugel's applicable Technical Reference File. A documented risk assessment based on the ICH Q3D principles is available on www.mycapsugel.com.

all process	Unit	Acceptance Level
Element	ppm	Not more than I
Arsenic	ppm	Not more than 1
Lend	ppm	Not more than 0.5
Cadmium	ppm	Not more than 0.1
Mercury	.,	Not more than 5
Cobalt	ppm	Not more than 10
Vanadium	ppm	Not more than 20
Nickel	ppm	



## CERTIFICATE OF ANALYSIS

Page: 2 of 2

Customer Name: Euromar Ltd

Lot Nr: 3660214

## Residual Solvent Statement

In accordance with ICH Q3C residual solvent guideline, Class 3 solvents may be used according to good manufacturing practices such that their cumulative value does not exceed 5000ppm or 0.5%, under option 1 as defined in ICH Q3C, USP<467>, and EP General Text 5.4.

#### Physical Characteristics

Defect levels are in conformance with the Coni-Snap® specification for Visual attributes, as defined in the table below.

Defect Group	Class I	Class II	Class III
	Visual	Visual	Visual
Sigma Level	4.9	4.7	4.2
PPM	<290	<670	<3600

Appearance - Clean empty capsules, meeting the specified requirements of color and size. Odor - Free of disagreeable odor.

### Manufacturing Processes:

No Addition of Prescryatives No Ethylene Oxide Treatment No Irradiation Treatment