

# Vegan DDS Tranexamic

---

TECHNICAL DOSSIER

Name: Vegan DDS Tranexamic

Ref.: Vegan DDS Tranexamic\_01

Ver.: 01 (13/04/2021)



INdermal

## INDEX

PRODUCT IDENTIFICATION _____	2
INCI _____	2
COMPOSITION _____	2
ORIGIN _____	3
SPECIFICATIONS _____	3
COUNTRY OF MANUFACTURE _____	3
EXPIRATION AND STORAGE _____	3
RECOMMENDED DOSE _____	4
DISPERSABILITY _____	4
HOW TO USE _____	4
INCOMPATIBILITIES _____	4
ISO 16128-1:2016 and 16128-2:2017 _____	4
TESTS ON ANIMALS _____	5
HEAVY METALS _____	5
REACH _____	5
INGREDIENTS STATEMENTS* _____	6
REGULATIONS _____	7
CHARACTERIZATION _____	7
PRODUCTION PROCESS FLOW CHART _____	8

## PRODUCT IDENTIFICATION

**Tradename:** Vegan DDS TRANEXAMIC

**Reference:** Vegan DDS TRANEXAMIC\_01

**Description:** Tranexamic acid (2.5 %) and Niacinamide (0.5%) encapsulated in vegan deep release nano-vesicles (DDS - Deep Delivery System) to add in cosmetic, cosmeceutical or dermo pharmaceutical formulations.

## INCI

AQUA, MANNITOL, PHOSPHATIDYLCHOLINE, GLYCERIN, TRANEXAMIC ACID, CETYL ALCOHOL, DECYL GLUCOSIDE, POTASSIUM SORBATE, SODIUM BENZOATE, NIACINAMIDE, XANTHAN GUM, SODIUM CHLORIDE.

## COMPOSITION

Ingredient	CAS	% (w/w)
AQUA	7732-18-5	Q.S. 100
MANNITOL	69-65-8	4.50 – 6.50
PHOSPHATIDYLCHOLINE	8002-43-5	3.20 – 5.20
GLYCERIN	56-81-5	2.90 – 3.90
TRANEXAMIC ACID	1197-18-8	2.00 – 3.00
CETYL ALCOHOL	36653-82-4	0.60 – 0.80
DECYL GLUCOSIDE	54549-25-6	0.50 – 0.70
POTASSIUM SORBATE	24634-61-5	0.50 – 0.70
SODIUM BENZOATE	532-32-1	0.50 – 0.70
NIACINAMIDE	98-92-0	0.40 – 0.60
XANTHAN GUM	11138-66-2	0.10 – 0.30
SODIUM CHLORIDE	7647-14-5	0.05 – 0.15

## ORIGIN

Ingredient	Origin	Country of origin
AQUA	NATURAL	SPAIN
MANNITOL	VEGETABLE	WORLDWIDE
PHOSPHATIDYLCHOLINE	VEGETABLE	WORLDWIDE
GLYCERIN	VEGETABLE	SOUTH EAST ASIA and EUROPE
TRANEXAMIC ACID	SYNTHETIC	CHINA
CETYL ALCOHOL	VEGETABLE	WORLDWIDE
DECYL GLUCOSIDE	VEGETABLE	FRANCE
POTASSIUM SORBATE	SYNTHETIC	GERMANY
SODIUM BENZOATE	SYNTHETIC	NETHERLANDS
NIACINAMIDE	SYNTHETIC	CHINA
XANTHAN GUM	NATURAL	EUROPE
SODIUM CHLORIDE	SYNTHETIC	NETHERLANDS or FRANCE

## SPECIFICATIONS

**Appearance:** LIQUID  
**Color:** WHITE  
**Odor:** CHARACTERISTIC  
**pH:** 5.5 – 6.5

## COUNTRY OF MANUFACTURE

Spain

## EXPIRATION AND STORAGE

12 months if the product is kept in the original sealed container.

The product has been packed in a protective nitrogen atmosphere. Once opened, it is recommended to use all the product at once or repackage the excess using nitrogen.

Do not freeze.

## RECOMMENDED DOSE

1 % - 10 % According to the frequency of application of the final product and the intensity of the effect that you want to achieve.

Examples: Daily cream: 3% – 5 %    Serum: 4% – 8%    Mask: 3% - 7%    Ampoules: 5% - 10%

Professional use: 7% - 10%

## DISPERSABILITY

Dispersible product in aqueous media. (See incompatibilities)

## HOW TO USE

Shake gently before using.

Add to bulk during the final phase of the production process, ensuring that the temperature does not exceed 40°C to avoid degradation of the encapsulated molecules. If you need to add it to higher temperatures, please consult our technical service

Maximum homogenization: 20.000 rpm

Formulation pH: 3 – 11

## INCOMPATIBILITIES

Ethanol concentrations higher than 15% may damage liposomes (contact our technical service for advice)

Detergents may break liposomes.

Do not add to oil. In emulsions W/O or O/W add it in the aqueous solution.

## ISO 16128-1:2016 and 16128-2:2017

In accordance with the guidelines on definitions of natural and organic cosmetic ingredients (ISO 16128-1: 2016 and ISO 16 128-2: 2017), we certify according to the information provided by our suppliers and our production process, the following information

<b>Naturally origin content:</b>	97.19 %
<b>Natural content:</b>	93.87 %

## TESTS ON ANIMALS

This product has not been tested on animals by or on behalf of INdermal - Nanovex Biotechnologies SL, in accordance with European Regulation (EC) No. 1223/2009

## HEAVY METALS

Max. 0.5 ppm

## REACH

Ingredient	REACH
AQUA	Exempt (Ocurring in nature)
MANNITOL	Exempt (Annex IV Article 2(7)(a))
PHOSPHATIDYLCHOLINE	Exempt (Annex IV Article 2(7)(a))
GLYCERIN	Exempt (Annex V Article 2(7)(b))
TRANEXAMIC ACID	Exempt (<1 Tonne/year)
CETYL ALCOHOL	01-2119485905-24-XXXX
DECYL GLUCOSIDE	01-2119489418-23-XXXX
POTASSIUM SORBATE	01-2119950315-41-XXXX
SODIUM BENZOATE	01-2119460683-35-XXXX
NIACINAMIDE	01-2119968268-22-XXXX
XANTHAN GUM	Exempt (Article 2 (7a and 7b))
SODIUM CHLORIDE	01-2119485491-33-XXXX

## INGREDIENTS STATEMENTS\*

<b>Cosmetic allergens</b>	This product does not contain any of the substances introduced in Annex III of the European Cosmetic Regulation by the Seventh Amendment (EC) No. 2003/15 based on SCCNFP opinion.
<b>GMO</b>	According to our knowledge, this product does not contain any ingredient considered a genetically modified organism.
<b>CMR</b>	According to our knowledge, this product does not contain any ingredient classified as carcinogenic, mutagenic or reprotoxic according to Regulation (EC) No. 1272/2008.
<b>SVHC</b>	According to our knowledge, this product does not contain any substance considered extremely worrying.
<b>COV</b>	According to our knowledge, this product does not contain any compound considered volatile organic
<b>Phthalatos</b>	This product does not contain phthalates as an ingredient
<b>Nanomaterials</b>	This product cannot be considered a nanomaterial as defined in Regulation (EC) No. 1223/2009
<b>Formaldehyde</b>	This product does not contain formaldehydes or derivatives
<b>Palm Oil</b>	This product contains palm oil (RSPO Certified)
<b>BSE / TSE</b>	According to our knowledge, this product does not contain any compound related to Bovine Spongiform Encephalopathy or Transmissible Spongiform Encephalopathies
<b>Dioxins</b>	This product is not expected to contain Dioxins
<b>Pesticides</b>	This product is not expected to contain Pesticides
<b>Vegetarian / Vegan</b>	This product is suitable for vegetarians and vegans.
<b>Halal</b>	This product does not have Halal certification
<b>COSMOS certified</b>	This product does not have COSMOS certification

\* Information based on our manufacturing process and information provided by our suppliers

## REGULATIONS

Geographical Area	Regulation
<b>European Union</b>	This product is in accordance with Regulation (EC) No. 1223/2009
<b>USA</b>	None of the compounds of this product are prohibited or restricted by FDA regulations.
<b>Canada</b>	None of the compounds of this product are included in the Cosmetic Ingredient Hotlist of Canada (2019) of Ingredients that are Prohibited or Restricted for Use un Cosmetic Products.
<b>China</b>	All INCI names are included in the last version of IECIC list (2015)

## CHARACTERIZATION

Product characterization is based on the particle analysis using nanotechnology characterization techniques such as; Dynamic Light Scattering, Nanoparticle Track Analysis or Phase analysis Light Scattering. The main properties analyzed are; Average Size, Size Distribution, Zeta-Potential and Particle concentration.

### **Average size: Average Size = 150 – 300 nm**

Average size of nano-vesicles can be determined using Dynamic Light Scattering (DLS) and Nanoparticle Track Analysis (NTA) techniques. Both are based on the Brownian motion of the particle and the light scattering from it. They apply Stokes-Einstein equation to relate diffusion to size (Hydrodynamic diameter). Additionally, DLS offers an ensemble measurement, whereas NTA delivers a particle-by-particle measurement.

### **Size distribution: Polydispersity Index (PDI) < 0.5**

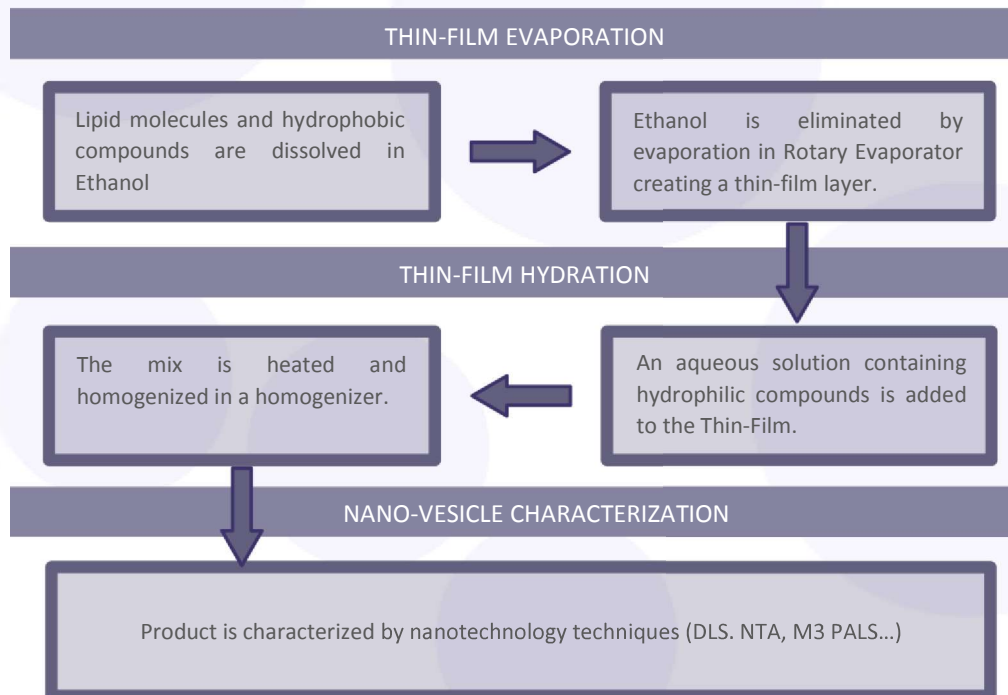
Size distribution was determined using the DLS technique. The distribution is represented by the polydispersity index PDI, whose values are in the range between 0 and 1. Values near 0 represent a monodispersed sample and values near 1 represent polydisperse sample. It can be considered that values below 0.5 have good distribution values

### **Liposomes concentration (particles/ml): $\times 10^{14}$ nanovesicles/ml**

Nanovesicle concentration can be determined by Nanoparticle Tracking Analysis (NTA) as it is a nanoparticle visualization technique that provides size, count and concentration measurements.



## PRODUCTION PROCESS FLOW CHART



**Responsibility:** The information contained in this document is, to the best of our knowledge, accurate. However, we reject any responsibility for the application and in processed materials that contain our product. Only the producer of the final product must assume full responsibility in accordance with current regulations. The content of this document is subject to change without notice unless otherwise agreed in writing. Consult with [info@nanovexbiotech.com](mailto:info@nanovexbiotech.com) to get the latest version of this document.