

Danilon equidos 1,5g granulátum A.U  
Gysz:161918 Lej:20/02/29  
Kisz:60x10g Me.:doboz  
DR. SZÉKELY KFT. részére

2151 Fót  
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Tel:27/537-100  
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**ESTEVE**

LABORATORIOS Dr. ESTEVE, S.A.U  
Pharmaceutical Plant

GMP Certificate number: NCF/1609/001/CAT

CERTIFICATE OF MANUFACTURE AND BATCH RELEASE

**Manufacturing site:** Laboratorios Dr. ESTEVE S.A.U  
San Martí, s/n - Polígono Industrial  
08107 - Martorelles (Barcelona), SPAIN  
**Name of Product , Strength, Dosage form / Pack Size (type) :**  
Danilon Equidos (Suxibuzone) 1.5 g sachets  
/ 60 x 10 g sac.  
**Packaging Batch Number:** 161918  
**Date of manufacture** (according to EMEA/CVMP453/01): 02-2016  
**Expiry Date:** 02-2020

**Results of analysis:**  
See attached Certificate of Analysis (CoA) number: 040000067904,8900000093829

**Additional information:**

- Destination (Country): Hungary

**Certification:**

I hereby certify that the above information is authentic and accurate. This batch of product has been manufactured, packaged and quality controlled at the above mentioned site in full compliance with the EU GMP requirements and with the specifications of the Marketing Authorisation, including API origin and quality. The batch processing, packaging and analysis records were reviewed and found to be in compliance with the agreed specifications.

(X) No significant deviations during manufacturing / packaging are reported.

( ) The following significant deviations were reported during manufacturing / packaging and properly investigated.

**Signature:**

  
**Cristina Colet, Deputy Technical Director**

(Qualified Person according Art. 49, Directive 2001 / 83 / EC)

Date: 13 SEP 2016

END OF REPORT - Certificate of Manufacture & Batch Release

## CERTIFICATE OF ANALYSIS

Product: DANILON EQUI. DCP 60SOBX10G-H+RO+SLO  
Date of manufacture: 04-02-2016

Date of analysis: 13-09-2016

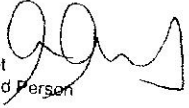
Batch: 161918

TEST	SPECIFICATIONS	RESULTS
Description	Yellow granules	
Uniformity of mass		Complies
Units		
Mean	10 or 30	10
Standard Deviation		100,5 %
Acceptance Value		1,7
CCF Identification	Complies Ph. Eur.	4,1
Suxibuzone	Positive	Positive
Saccharin sodium	Positive	Positive
Quinoline yellow (E-104) (UV)		
Identification	Positive	Periodic Test
Particulate size		
> 0,21 mm	Less than 10 %	6 %
Dissolution rate Suxibuzone (UV)		
Units	6, 12 or 24	6
Mean	Q = 75 % in 45 min	99 %
Minimum		96 %
Complies acceptance criteria	Complies Ph. Eur./USP	Complies
Suxibuzone HPLC		
Identification	Positive	Positive
Assay	95,0 - 105,0 % (0,143 - 0,158 g/g)	96,9 %
Assay/sachet	95,0 - 105,0 % (1,425 - 1,575 g/sachet)	97,4 %
Degradation product		
Phenylbutane	Not more than 1,0 %	0,05 %
Microbiological purity	Complies Eur. Ph. 6.3 chapter 5.1.4	Complies

This is to certify that the product has been manufactured and tested according to current EC Good Manufacturing Practices (GMP) and other rules governing Medicinal Products in the EU at the above mentioned in accordance with the specifications in the Marketing Authorisation.

Signed and dated:

13 SEP 2016



C. Colet  
Qualified Person

AN: 040000067904,890000093829