

ESTEVE

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CERTIFICATE OF ANALYSIS	Date of analysis: 19-08-2015
Product: DANILON EQUI DCP 60S08X10G-H+RO+SLO	Batch: 151265
Date of manufacture: 18-12-2014	

TEST	SPECIFICATIONS	RESULTS
Description	Yellow granules	Complies
Uniformity of mass		
Units	10 or 30	10
Mean		102,8 %
Standard Deviation		1,3
Acceptance Value	Complies Ph. Eur.	4,4
CCF Identification		
Suxibuzone	Positive	Positive
Saccharin sodium	Positive	Positive
Quinoline yellow (E-104) (UV)		
Identification	Positive	Periodic Test
Particulate size		
> 0,21 mm	Less than 10 %	10 %
Dissolution rate Suxibuzone (UV)		
Units	6, 12 or 24	6
Mean	Q = 75 % in 45 min	100 %
Minimum		99 %
Complies acceptance criteria	Complies Ph. Eur./JSP	Complies
Suxibuzone HPLC		
Identification	Positive	Positive
Assay	95.0 - 105.0 % (0.143 - 0.158 g/g)	99,6 %
Assay/sachet	95.0 - 105.0 % (1.425 - 1.575 g/sachet)	98,4 %
Degradation product		
Phenylbutane	Not more than 1.0 %	0,08 %
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:

E. Rodón
 Qualified Person



AN: 040000056580,890000077110

