

vetoquinol
 ACHIEVE MORE TOGETHER
CERTIFICATE OF ANALYSIS

FINISHED PRODUCT	
Product name :	CLAVASEP 500MG CPR100 GR HU RO
Analytical code :	456279
Pharmaceutical form :	Not Coated Tablets
Manufacturing date :	27/05/2019
Batch number :	9C1620E
Expiration date :	26/05/2022

TESTS	SPECIFICATIONS	RESULTS
Characteristics of tablets	Oblong, off-white to brownish sprinkled, scored	Pass
Length	>= 20.8 <= 21.2 mm	21.1 mm
Average weight and uniformity of weight	>= 904.4 <= 999.6 mg	951.6 mg
Average mass	Complies with Eur. Ph. 2.9.5	Pass
Uniformity of mass		
Resistance to crushing	>= 80 N	130 N
Disintegration test	<= 15 min	<4 min
Equilibrium relative humidity	<= 15 %	9 %
Amoxicillin dissolution rate	Q=85% within 30 min	102 %
Amoxicillin dissolution compliance	Complies with Eur. Ph. 2.9.3	Pass
Clavulanic ac dissolution rate	Q=80% within 30 min	102 %
Clavulanic ac dissolution compliance	Complies with Eur. Ph. 2.9.3	Pass
Amoxicillin and clavulanic acid identity (HPLC)		
Amoxicillin UV spectrum	Positive	Pass
Amoxicillin retention time	Positive	Pass
Clavulanic acid UV spectrum	Positive	Pass
Clavulanic acid retention time	Positive	Pass
Amoxicillin and clavulanic acid assay		
Amoxicillin content	>= 380 <= 420 mg/tab	406 mg/tab.
Clavulanic acid content	>= 97.8 <= 108.2 mg/tab	101.1 mg/tab.

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LABORATOIRE PHARMACEUTIQUE VÉTÉRIINAIRE

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 VETOQUINOL S.A. au Capital de 29.704.750 € - SIRET 676 250 111 (0117) RCS VESOUL GRAY B 676 250 111

Meoszám:191100009MBT

6.pld. Kiadva: 19/11/27

MEDIMPEX GYÓGYSZER ZRT.
 1158 Bp., Rákospalotai határút 2.
 Nagyné Kári Éva
 Minőségbiztosítási szisztem
 Csak a piros szín hiteles

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TESTS	SPECIFICATIONS	RESULTS
Amoxicillin purity (HPLC)		
Imp.A	<= 1.0 %w/w	
Imp.B	<= 1.0 %w/w	<0.3 % w/w
Imp.C	<= 1.0 %w/w	<0.3 % w/w
Imp.D	<= 1.0 %w/w	<0.3 % w/w
Imp.E	<= 1.0 %w/w	0.3 % w/w
Imp.F	<= 1.0 %w/w	<0.3 % w/w
Imp.G	<= 1.0 %w/w	0.3 % w/w
Imp.K	<= 1.0 %w/w	0.6 % w/w
Any unidentified degradation product	<= 1.0 %w/w	<0.3 % w/w
Total unidentified degradation products	<= 2.0 %w/w	<0.3 % w/w
Total degradation products	<= 3.0 %w/w	<0.3 % w/w
Clavulanic acid impurities (HPLC)		1.3 % w/w
Any unspecified degradation product	<= 1.0 %area	<0.3 % area
Total degradation products	<= 2.0 %area	<0.3 % area

I hereby certify that this lot is manufactured, packed and labelled under conditions as stated in the current European Union rules for Good Manufacturing Practice for medicinal products for veterinary use, and that the finished product in all aspects answers the requirements laid down in the marketing authorisation.

Released by Qualified Person : **Paul-Adrien MATHON**
 Date : 10/10/2019 14:27:13
 Decision : FULL RELEASE

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