

Certificate of Analysis  
 Bulletin d'Analyse

BA 5 Edition 15 du 09/05/2014

**CEPRAVIN DRY COW**

**3 G**

**BATCH N° 5511**

**CSSA**

Specification Reference: Dossier HRA/88/035

<u>TEST</u>	<u>METHOD</u>	<u>RESULT</u>	<u>ACCEPTANCE LIMITS AT RELEASE</u>
Appearance	Q-PF- CEPRAVIN	Conform	White to cream coloured suspension
Identity	Colorimetric Q-PF- CEPRAVIN	Conform	Complies with test
Assay	UV/ HPLC Q-PF- CEPRAVIN	0.248	0.243 to 0.283 g/syringe (MFW)
Sterility	BP Vet Q-PF- CEPRAVIN	Conform	Complies with the BP (Vet)
Uniformity of fill weight /extractable weight	In a random sample of 10 applicators Q-PF- CEPRAVIN	Conform	NMT 1 in 10 shall deviate from the mean fill weight by more than +/- 10% and none shall deviate by more than +/- 20%
Related substances	HPLC Q-PF- CEPRAVIN	0.8 0.2 1.1 1.9	Isonicotinamide ≤ 2% Any other individual impurity ≤ 1% Total unknown impurities ≤ 3% Total impurities ≤ 5%

MFW : Mean Fill Weight

Manufacturing Date : February 18<sup>th</sup> 2015

Expiry Date : January 2018

This batch has been manufactured and released in accordance with GMP requirements and according to the Marketing Authorisation.

Release date : March 18<sup>th</sup> 2015

  
**Mikael CAUCHE**  
 Qualified Person

R0407