

EMA and VICH Guidelines

Here you can find a list of [quality guidelines for veterinary medicinal products](#) as published by European Medicines Agency (EMA) and Veterinary International Co-operation on Harmonization (VICH). These guidelines together with requirements of the current version of [European](#) and Czech Pharmacopoeia should be taken into consideration for the preparation of the quality part of a registration dossier.

Apart from the following guidelines, EMA also publishes and continuously updates a list of questions and answers ([Part 1](#) and [Part 2](#)) concerning interpretation and implementation of the guidelines.

1. Development pharmaceuticals

Name	Number	Effective from	Source	Note
Development pharmaceuticals	EMA/CVMP/315/98	March 2000	EMA	
Sterilisation of the medicinal products	EMA/CVMP/CVMP/OWB/BWP/850/2015	September 2015	EMA	primary
Decision trees for selection of analytical methods	EMA/CVMP/1065/99	September 2000	EMA	for guidance on development pharmaceuticals

2. Manufacture of medicinal product

Name	Number	Effective from	Source	Note
Manufacture of the finished product	EMA/CVMP/1126/95	June 1996	EMA	G
Start of shelf-life of the finished product	EMA/CVMP/453/01	December 2001	EMA	Annex 1 for guidance on the manufacture of the finished product
Process validation for finished products	EMA/CVMP/CVMP/OWB/BWP/850/2015	September 2015	EMA	Annex 1 for guidance on the manufacture of the finished product

Limitations to the use of ionizing radiation in the manufacture of medicinal products

EMA

Guideline

Use of ionizing radiation in the manufacture of medicinal products

Eudralex

Gui

3. Active substance

Name	Number	Effective from	Source	Note
Active substance master file procedure	EMA/CVMP/134/02	October 2012	EMA	Gu
Chemistry of active substances	EMA/CVMP/70736/02	2018	EMA	Gui
Summary of requirements for active substances in the part of the dossier	EMA/CVMP/1039/02	February 2005	EMA	Gui
Template for the qualified person declaration on GMP manufacturing practice compliance	EMA/19629/2014	June 2014	EMA	Chai
Chemical structure and impurity data for the evaluation of new active substances	EMA/CVMP/1629/2016	October 2017	EMA	Ref
Use of cocrystals of active substances	EMA/CVMP/26400/2015		EMA	Ref

4. Impurities

Name	Number	Effective from	Source	Note
Implementation of risk assessment for impurities in veterinary medicinal products	EMA/CVMP/QWP/60101/2017	2018	EMA	Ch
Assessment and control of impurities in veterinary medicinal products	EMA/CVMP/QWP/377245/2016		EMA	Dr
Control of impurities of pharmaceuticals in accordance with the European Pharmacopoeia	EMA/CVMP/059/04	March 2014	EMA	Page
Setting specifications for related compounds	EMA/CVMP/QWP/15925/06	2013	EMA	Gui
VICH GL10 Impurities in New Veterinary Substances	EMA/VICH/397/00	September 2008	VICH	Gui
VICH GL11 Impurities in New Veterinary Products	EMA/VICH/398/00	September 2008	VICH	Gui
VICH GL18 Residual solvents in primary packaging of medicinal products, active substances and excipients	EMA/VICH/502/00	July 2012	VICH	Gui
Annexes to VICH GL 18	EMA/CVMP/511/03	March 2013	EMA	Ann
Application of VICH GL 18 to veterinary medicinal products containing existing	EMA/CVMP/420/01	May 2001	EMA	Ca

5. Excipients

Name	Number	Effective from	Source	Note
Excipients in the dossier for marketing authorisation for veterinary medicinal products	EMA/CVMP/004/98	June 1999	EMA	6

PART II Quality

Written by Administrator - Last Updated Wednesday, 06 March 2019 06:24

Inclusion of antioxidants and antioxidants in preservation in medicinal products	EMA/CVMP/QWP/015/95	January 1998	EMA	Guideline
Quality of water for pharmaceuticals	EMA/CVMP/QWP/158/01	June 2002	EMA	Guideline
Water for injection prepared by conventional methods	EMA/CVMP/QWP/287/08	March 2008	EMA	Reference

6. Packaging

Name	Number	Effective from	Source	Note
Plastic primary packaging for medicinal products	EMA/CVMP/205/04	December 2005	EMA	Guideline

7. Specifications, analytical procedures and analytical validation

Name	Number	Effective from	Source	Note
Parametric release	EMA/CVMP/QWP/339588/2005	May 2007	EMA	Guideline
Specifications and control tests on the finished product	3AQS1A	1992	Eudralex	Guideline
VICH GL1 Validation of analytical procedures: definitions and terminology	EMA/VICH/QWP/590/98	October 1999	VICH	Guideline
VICH GL2 Validation of analytical procedures: methodology	EMA/VICH/QWP/591/98	October 1999	VICH	Guideline
VICH GL39 Test procedures and acceptance criteria for veterinary drug substances and new formulations	EMA/CVMP/VICH/810/06	November 2006	VICH	Guideline
VICH GL40 Test procedures and acceptance criteria for biological veterinary products	EMA/CVMP/VICH/811/06	November 2006	VICH	Guideline
Dissolution specification for medicinal products	EMA/CVMP/QWP/339588/2007	May 2007	VICH	Reference

8. TSE

Name	Number	Effective from	Source	Note
Minimising the risk of transmitting a prion agent from encephalopathy agents via human and animal products	EMA/142011	2011	EMA	Guideline

9. Stability

PART II Quality

Written by Administrator - Last Updated Wednesday, 06 March 2019 06:24

Name	Number	Effective from	Source	Note
Declaration of storage conditions for medicinal products of pharmaceutical veterinary medicinal products	EMA/CVMP/112/00	October 2003	EMA	Annex
In-use stability testing of medicinal products	EMA/CVMP/624/01	September 2002	EMA	Guidance
Maximum shelf-life for sterile ophthalmics after opening or following reconstitution	EMA/CVMP/198/99	February 2001	EMA	Guidance
Stability testing for applications for variation of product	EMA/CVMP/101/00	February 2001	EMA	Guidance
Stability testing of existing medicinal products and new products	EMA/CVMP/846/99	September 2001	EMA	Guidance
VICH GL3 Stability testing of drug substances and medicinal products	CVMP/VICH/399/99	January 2000	VICH	Guidance
VICH GL4 Stability testing of dosage forms	CVMP/VICH/600/99	May 2000	VICH	Guidance
VICH GL5 Stability testing of veterinary drug substances and medicinal products	CVMP/VICH/110/00	May 2000	VICH	Guidance
VICH GL8 Stability testing of mixtures	CVMP/VICH/186/99	December 2000	VICH	Guidance
VICH GL45 Bracketing and matrixing	EMA/CVMP/158/01	April 2001	VICH	Guidance
VICH GL51 Quality: stability	EMA/CVMP/158/01	April 2001	VICH	Guidance

10. Herbal medicinal products

Name	Number	Effective from	Source	Note
Declaration of herbal substances	EMA/CVMP/101/00	February 2001	EMA	Guidance
Quality of combination herbal medicinal products	EMA/CVMP/101/00	February 2001	EMA	Guidance
Quality of herbal medicinal products	EMA/CVMP/101/00	February 2001	EMA	Guidance
Specifications: test procedures	EMA/CVMP/101/00	February 2001	EMA	Guidance

11. Specific veterinary dosage forms

Name	Number	Effective from	Source	Note
Additional quality requirements for medicinal products intended for incorporation into animal feeding-stuffs (medicinal products)	EMA/CVMP/101/00	February 2001	EMA	Guidance
Quality aspects of pharmaceutical products for administration in drinking water	EMA/CVMP/101/00	February 2001	EMA	Guidance
Quality aspects of single-dose veterinary products	EMA/CVMP/101/00	February 2001	EMA	Guidance
Quality of modified release veterinary products	EMA/CVMP/101/00	February 2001	EMA	Guidance
Maximum in-use shelf-life of veterinary products	EMA/CVMP/101/00	February 2001	EMA	Guidance
Premixes for medicated feed	EMA/CVMP/101/00	February 2001	EMA	Guidance

12. Minor uses / minor species (MUMS)

Name	Number	Effective from	Source	Note
Quality data requirements for CVMP medicinal products intended for minor uses or minor species	EMA/CVMP/QWP/1287/2004	1 January 2007	EMA	CVMP
CVMP guidelines on data requirements for medicinal products intended for minor uses or minor species	EMA/CVMP/1367/2005	1 January 2007	EMA	CVMP