28 Novembre 2024 EMA/899164/2022 Rev.2 Human Medicines Division Veterinary Medicines Division

# Abbreviations used in EMA scientific committees & CMD documents and in relation to EMA's regulatory activities

Abbreviation <sup>1</sup>	
1S1A	One substance, one assessment (see EU chemicals assessment reform)
3Rs	3Rs principles -Replace, Reduce and Refine- for the ethical use of animals in
	medicine testing across the European Union (see also Joint 3Rs WP)
AA	Accelerated Assessment
ACT EU initiative	Accelerate Clinical Trials in the EU (see <u>ACT EU</u> )
ADI	Acceptable Daily Intake
ADR(s)	Adverse Drug Reaction(s) (see <u>GVP</u> annex I)
AE(s)	Adverse Event(s) (see <u>GVP</u> annex I)
AEFI(s)	Adverse Event(s) Following Immunisation (see <u>GVP</u> annex I)
AER	Adverse Event Report
AESI	Adverse Event of Special Interest
AHEG	(EMA) Ad Hoc Expert Group
AI	Artificial Intelligence
AM	Additional Monitoring
AMEG	(EMA CHMP/CVMP) Antimicrobial Advice Ad Hoc Expert Group (see <u>AMEG</u> )
AMR	Antimicrobial resistance (see Antimicrobial resistance)
ANVISA	Brazilian health regulatory agency (see International agreements)
API	Active Pharmaceutical Ingredient (see International collaboration on GMP
	inspections)
API	Application Programming Interface (see substance and product data
	management under SPOR)
AR(s)	Assessment Report(s)
ARSP	Assessment Report Summary for the Public (see <u>EU herbal monographs</u> )
ASMF WG	(Joint EMA/HMA) Active Substance Master File Working Group (see ASMF WG)
ASU	Antimicrobial sales and use

 $^1$  Acronyms are abbreviations that can be pronounced as a word (e.g. 'CAT') whereas initialisms are abbreviations for which each letter is pronounced separately (as in 'SME')

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ATC(/DDD)	Anatomical Therapeutic Chemical classification system, maintained by WHO
	(with Defined Daily Doses)
ATD	(EMA) Access to Documents (see <u>Access to documents</u> )
ATD	Anti-Tampering Device (see Falsified medicines: overview)
ATMP(s)	Advanced Therapy Medicinal Product(s) (i.e. gene, cell and tissue engineering
AWP	products) (EMA CVMP) Antimicrobials Working Party (see <u>AWP</u> )
BA	Bioavailability
BE	Bioequivalence (see also Country codes: BE = Belgium)
BEMA	Benchmarking of European Medicines Agencies (see <u>Integrated quality</u>
	management system)
ВСР	(EMA) Business Continuity Planning
BDSG	(HMA-EMA) Big Data Steering Group (see <u>HMA-EMA joint BDSG</u> )
BMWP	(EMA CHMP) Biosimilar Medicinal Products Working Party
B/R	Benefit/Risk (in B/R assessment, B/R balance, B/R profile)
BWP	(EMA CHMP) Biologics Working Party
CAMD	Competent Authority for Medical Devices (see <u>CAMD</u> )
CAP(s)	Centrally Authorised Product(s)
CAPA plan	Corrective and preventive action plan
CAR-T cell	Chimeric antigen receptor T cell
САТ	(EMA) Committee for Advanced Therapies
СВМР	Cell-based Medicinal Product
CBRN	Chemical, Biological, Radiological and Nuclear (see EU CBRN risk mitigation)
CCDS	Company Core Data Sheet (see <u>GVP</u> annex I)
CCI	Commercially Confidential Information
CCRVDF	Codex Committee on Residues of Veterinary Drugs in Foods (see <u>Codex</u>
	<u>Alimentarius</u> )
CCSI	Company Core Safety Information (see <u>GVP</u> annex I)
CdT	Centre de Traduction (see <u>Translation Centre for the bodies of the EU</u> )
CDx	Companion Diagnostics
CDP	(EMA) Clinical Data Publication (see <u>Clinical data publication</u> )
CE mark	Conformité Européenne = European conformity mark (see <u>Medical Devices</u> )
CECP	Clinical Evaluation Consultation Procedure (see <u>Medical Devices</u> )
CEP(s)	Certificate(s) of Suitability to the monographs of the European Pharmacopoeia (see <u>EDQM- Certification of suitability</u> )
СНМР	(EMA) Committee for Medicinal Products for Human Use (previously: CPMP)
CI	Confidence interval
CI	Contraindication
CIA	Critically Important Antimicrobials
CIOMS	Council for International Organizations of Medical Sciences
СМА	Conditional Marketing Authorisation
CMDh	Coordination Group for Mutual Recognition and Decentralised Procedures
	(human)
CMDv	Coordination Group for Mutual Recognition and Decentralised Procedures for
	Veterinary Medicinal Products
CMDS	Critical Medical Devices Shortage
СМО	Contract Manufacturing Organisation

CMS(s)	Concerned Member State(s)
CNSWP	(EMA CHMP) Central Nervous System Working Party (see <u>CNSWP</u> )
СОМР	(EMA) Committee for Orphan Medicinal Products
Corr.	Corrigendum
CP	Centralised Procedure (see <u>Applying for EU marketing authorisation</u> )
CP	Concept Paper (see <u>Scientific quidelines</u> )
CPAR	Consultation Procedure public Assessment Report (see <u>CHMP opinions on</u> <u>consultation procedures</u> )
СРМР	Committee for Proprietary Medicinal Products, former name of CHMP
CQA	Critical Quality Attribute
CRM	Customer Relationship Management
CRO	Contract Research Organisation
CSP(s)	Core Safety Profile(s)
CSR	Clinical Study Report
СТ	Clinical Trial (see <u>Clinical trials</u> )
СТА	Clinical Trial Application
CTCG	Clinical Trial Coordination Group (see <u>HMA CTCG</u> )
CTD	Common Technical Document – see eCTD
CTIS	Clinical Trials Information System (see <u>CTIS</u> )
CTR	Clinical Trial Regulation (see <u>Clinical trials human medicines</u> )
CTS	Communication and Tracking System (see HMA – CMDh CTS Working Group)
CV	Curriculum vitae
СУМР	(EMA) Committee for Veterinary Medicinal Products
CVSWP	(EMA CHMP) Cardiovascular Working Party (see <u>CVSWP</u> )
DARWIN EU®	Data Analysis and Real World Interrogation Network (see DARWIN EU)
DCP	Decentralised Procedure (see <u>Applying for EU marketing authorisation</u> )
DDCs	Drug-Device Combination(s)
DDD	Defined Daily Dose (see ATC)
DDI	Drug-Drug Interaction
DER	Drug Extract Ratio (see <u>HMPC scientific guidelines</u> )
DG	Directorate-General (at the European Commission)
DG(s)	(EMA) Drafting Group(s) (see <u>Working parties and domains</u> )
DHPC	Direct Healthcare Professional Communication (see GVP annex I)
DIA	Drug Information Association
DIBD	Development International Birth Date (see <u>GVP</u> annex I)
DILI	Drug Induced Liver Injury
DFS	Disease-free survival
DLP	Data Lock Point
DMP	Development Medicinal Product (see <u>EudraVigilance</u> medicinal product dictionary)
DoI	Declaration of Interests (see <u>Handling competing interests</u> )
DoC	Declaration of conformity
DPO	Data Protection Officer
DPC	Data Protection Coordinator
DSMB	Data Safety Monitoring Board

DUS	Drug Utilisation Study
eAF	electronic Application Form
EC	European Commission ( <u>http://ec.europa.eu/index_en.htm</u> )
ECDC	European Centre for Disease Prevention and Control
	(https://www.ecdc.europa.eu/en)
ECHA	European Chemicals Agency ( <u>https://echa.europa.eu/</u> )
eCTD	electronic Common Technical Document (see <u>eSubmission website's section on</u>
	eCTD)
EDPB	European Data Protection Board (see EDPB)
EDPS	European Data Protection Supervisor (see Data protection and privacy)
EDQM	European Directorate for the Quality of Medicines (see EDQM of the Council of
	Europe)
EEA	European Environment Agency ( <u>https://www.eea.europa.eu/</u> )
EEA-EFTA states	European Economic Area – European Free Trade Association states
EFS	Event-free survival
EFSA	European Food Safety Authority ( <u>http://www.efsa.europa.eu</u> )
EHDS	European Health Data Space ( <u>https://health.ec.europa.eu/ehealth-digital-</u>
	health-and-care/european-health-data-space_en)
EMA/CAT-NB	Ad hoc European Medicines Agency/Committee for Advanced Therapies and
	Medical Devices Notified Body Collaboration Group – see EMA/CAT-NB
EMANS	European Medicines Agencies Network Strategy (see <u>EMANS</u> )
EMCDDA	was European Monitoring Centre for Drugs and Drug Addiction - see EUDA
EMEA	Old acronym for: European Medicines Agency; use: EMA
EMR	Electronic Medical Records
EMRN	European Medicines Regulatory Network (see EMRN)
EMT	(EMA) Experts Management Tool (see <u>European experts</u> )
ENCePP	European Network of Centres for Pharmacoepidemiology and Pharmacovigilance
	(https://www.encepp.eu/)
Enpr-EMA	European network of paediatric research at EMA (see Enpr-EMA)
ENPRS	European Network for Partnership in Regulatory Science
EoI	Extension of Indication
EP	European Parliament ( <u>http://www.europarl.europa.eu/</u> )
EPAR	European Public Assessment Report
e-PI	electronic Product Information
EPITT	European Pharmacovigilance Issues Tracking Tool
EPMAR	European Public MRL Assessment Report (see <u>Maximum residue limit</u>
	assessment reports)
ERA	Environmental Risk Assessment
ERAWP	(EMA CVMP) Environmental Risk Assessment Working Party (see <u>ERAWP</u> )
ERMS	European Risk Management Strategy (see <u>ERMS</u> )
eRMR	electronic Reaction Monitoring Report
ESEC(s)	(EMA) European Specialised Expert Community(ies) (see <u>Working parties and</u>
	domains)
ESMP	European Shortages Monitoring Platform (see <u>Availability of medicines</u> )
ESUAvet	European Sales and Use of Antimicrobials for Veterinary Medicine (see <u>ESUAvet</u>
50/40	Working Group)
ESVAC	European Surveillance of Veterinary Antimicrobial Consumption (see ESVAC)

ETF	(EMA) Emergency Task Force (see <u>ETF</u> )
EU	European Union
EUAN	European Union Agencies Network (see <u>EUAN</u> )
EU-ADR Project	Exploring and Understanding Adverse Drug Reactions by Integrative Mining of
-	Clinical Records and Biomedical Knowledge (formerly known as ALERT)
EUDA	European Union Drugs Agency (formerly known as European Monitoring Centre
	for Drugs and Drug Addiction – see <u>EUDA</u> )
EUDAMED	European database on medical devices (see <u>EUDAMED</u> )
Eudra-	European Union Drug Regulating Authorities
EudraCT	European Union Drug Regulating Authorities Clinical Trials database:
	see EudraCT and EU Clinical Trials Register
EU-IN	(Joint HMA/EMA) EU Innovation Network (see EU-IN)
EU-M4all	EU Medicines for all: see Medicines for use outside the European Union
	(formerly known as 'Article 58 procedure')
EUnetHTA	European Network for Health Technology Assessment
EU-NTC	EU Network Training Centre (see <u>EU-NTC</u> )
EU PAS Register	EU Post-Authorisation Study register
EURD list	List of EU Reference Dates and frequency of PSUR submission (see EURD list)
EURL ECVAM	European Union Reference Laboratory for alternatives to animal testing (see
	Ethical use of animals in medicine testing)
EURORDIS	European Organisation for Rare Diseases ( <u>http://www.eurordis.org/</u> )
EUTCT	European Union Telematics Controlled Terms – has been replaced by RMS
EU IVMAB	EU Immunisation and Vaccine Monitoring Board
EV	EudraVigilance (see EudraVigilance: electronic reporting)
EVDAS	EudraVigilance Data Analysis System
EVVet	EudraVigilance Veterinary
EV-EWG	EudraVigilance Expert Working Group (see <u>EV-EWG</u> )
EVMPD	EudraVigilance Medicinal Products Dictionary
EWP-V	(EMA CVMP) Efficacy Working Party (see <u>EWP-V</u> )
fAR	final Assessment Report
FDA	Food and Drug Administration (US) (see International agreements)
FDC	Fixed Dose Combination
FDHA	Federal Department of Home Affairs (Switzerland) (see International
	agreements)
FIM	First-In-Man
FMD	Falsified Medicines Directive (see Falsified medicines: overview)
FUQ	Follow-up questionnaire
fvAR	final variation Assessment Report
FWG	(EMA CHMP) Formulation Working Group (see <u>FWG</u> )
GACP	Good Agricultural and Collection Practice (see <u>HMPC GACP guideline</u> )
GCG	(EMA CHMP) Guideline Consistency Group (see <u>GCG</u> )
GCP	Good Clinical Practice (see GCP)
GCP IWG	Good Clinical Practice Inspectors Working Group (see <u>Compliance: overview</u> )
GDP	Good Distribution Practice (see GDP)
GDPR	General Data Protection Regulation (see Workshop on GDPR and secondary use
	of data for medicines and public health purposes)
GEG	(EMA CHMP) Geriatric Expert Group (see <u>GEG</u> )

GLP	Good Laboratory Practice (see <u>GLP</u> )
GMA	Global Marketing Authorisation
GMO	Genetically Modified Organism
GMP	Good Manufacturing Practice (see <u>GMP</u> )
GMDP IWG	Good Manufacturing Practice/Good Distribution Practice Inspectors Working
	Group (see <u>Compliance: overview</u> )
GPAG	(EMA PRAC) Granularity and Periodicity Advisory Group
GSPR	General Safety and Performance Requirements (see <u>Medical Devices</u> )
GTMP	Gene Therapy Medicinal Product
GVP	Good Pharmacovigilance Practices (see <u>GVP</u> )
HaDEA	European Health and Digital Executive Agency (see <u>HaDEA</u> )
HAEMWP	(EMA CHMP) Haematology Working Party (see <u>HAEMWP</u> )
HBD	Harmonised Birth Date
НС	Health Canada (see International agreements)
HCP(s)	Healthcare Professional(s)
HCPWP	(EMA) Healthcare Professionals' Working Party (see <u>HCPWP</u> )
HERA	Health Emergency Preparedness and Response Authority (see <u>HERA</u> )
НМА	Heads of Medicines Agencies (formerly: HoA) – see HMA
- HMA-Joint	with three groups: HMA-Joint, HMA-Human and HMA-Veterinary
- HMA(h)	
- HMA(v)	
НМР	Herbal Medicinal Product (see <u>EU herbal monographs</u> )
НМРС	(EMA) Committee on Herbal Medicinal Products
НоА	was: Heads of Agencies, use: HMA
HP	Herbal preparation (see EU herbal monographs)
	equivalent to 'Herbal drug preparation' in Ph. Eur. monographs
HR	Hazard Ratio
HRQoL	Health-related quality of life
HS	Herbal substance (see EU herbal monographs)
	equivalent to 'Herbal drug' in Ph. Eur. monographs
HTA	Health Technology Assessment
HTAb	Health Technology Assessment body (see <u>HTA Bodies</u> )
HTACG	Member State Coordination Group on HTA (see <u>HTACG</u> )
HTAR	Health Technology Assessment Regulation (EU) 2021/2282
HTD	Health Technology Developer
IBD	International Birth Date (see <u>GVP</u> annex I)
ICH	International Conference on Harmonisation of Technical Requirements for
	Registration of Pharmaceuticals for Human Use
ICMRA	International Coalition of Medicines Regulatory Authorities (see <u>ICMRA</u> )
ICSR(s)	Individual Case Safety Report(s) (see <u>GVP</u> annex I)
ICTPR	(WHO) International Clinical Trials Registry Platform
IDWP	(EMA CHMP) Infectious Diseases Working Party (see <u>IDWP</u> )
IDMC	Independent Data Monitoring Committee
IEC	Independent Ethics Committee
IFU	Instructions For Use (see <u>Medical Devices</u> )
IGDRP	International Generic Drug Regulators Programme
IHSI	International Horizon Scanning Initiative (see IHSI)

im	intramuscular
IMP	Investigational Medicinal Product
IMP	(EU Regulatory Network) Incident Management Plan (see <u>IMP</u> )
INN	International Nonproprietary Name (see <u>WHO/INN</u> )
IPs	Interested Parties
IPRP	International Pharmaceutical Regulators Programme (see <u>IPRP</u> )
IR	Inspection Report
IRB	Institutional Review Board
IRCH	International Regulatory Cooperation for Herbal Medicines (under WHO)
IIR	Integrated Inspection Report
IRIS	Not an abbreviation. Refers to the regulatory & scientific information
1110	management platform between EMA and stakeholders (NCAs, industry)
IRN	(EU Regulatory Network) Incident Review Network (see IMP)
ISO IDMP	Internal Organization for Standardization for the Identification of Medicinal
100 10111	Products (see <u>ISO IDMP standards</u> ) – implementation through the following EMA
	services:
	- OMS = Organisation Management Service
	- PMS = Product Management Service
	- RMS = Referentials Management Service
	- SMS = Substance Management Service
ISRR	Immunisation Stress-Related Response
ITF	(EMA) Innovation Task Force (see <u>Innovation in medicines</u> )
ITT	Intention-To-Treat (analysis)
iv	intravenous
IVD	In vitro Diagnostics
IVDR	(EU) In vitro Diagnostic medical devices Regulation (see <u>Medical Devices</u> )
IVMAB	(ECDC/EMA) Immunisation and Vaccine Monitoring Advisory Board
IVMP	Immunological Veterinary Medicinal Product
IWP	(EMA CVMP) Immunologicals Working Party (see IWP)
JAMS	Joint Action on Market Surveillance of Medical Devices (see <u>JAMS</u> )
JAP	(HMA/EMA) Joint Audit Plan
JECFA	Joint FAO/WHO Expert Committee on Food Additives
JIACRA	Joint Inter-agency Antimicrobial Consumption and Resistance Analysis (see
	Analysis of antimicrobial consumption and resistance)
Joint 3Rs WP	(EMA CHMP/CVMP) Joint 3Rs Replacement, Reduction and Refinement Working
	Party (see <u>3Rs principles</u> )
JSA	Joint Scientific Assessment
JSC	Joint Scientific Consultation (see Parallel joint scientific consultation with
	regulators and HTA bodies)
KPI	Key Performance Indicator
LE	List entry (see EU herbal monographs and list entries)
LM	Limited Markets
LoI	Letter of Intent
LoOI	List of Outstanding Issues
LoQ	List of Questions
LTT	Lines to take [internal EMA document usually not for publication]
MA	Marketing Authorisation

MAA	Marketing Authorisation Application
MAH	Marketing Authorisation Holder
MAWP	Multi-annual Work Plan
MB	(EMA) Management Board
MCMN (trial)	Multicenter/multinational (trial)
MD	Medical Device
MDCG	(EU) Medical Device Coordination Group
MDIG	(EMA) Medical Devices Implementation Group
MDR	(EU) Medical Devices Regulation (see <u>Medical Devices</u> )
MDSSG	(EMA) Medical Devices Shortages Steering Group
MedDRA	Medical Dictionary for Regulatory Activities – organised in a hierarchical
Ticabiot	structure characterised by different levels:
	- SOC = System Organ Class
	- HLGT = High Level Group Term
	- HLT = High Level Term
	- PT = Preferred Term
	- LLT = Lowest Level Term
MIC	Minimum Inhibitory Concentration
MIDD	Model-Informed Drug Development
MLM	Medical literature monitoring
MLWP	Monographs and List entries Working Party (former HMPC working party)
MNAT	Multinational Assessment Team (see <u>Multinational assessment team concept</u> )
МО	Major Objection
MoU	Memorandum of Understanding
MR	Mutual Recognition
MRA	Mutual Recognition Agreement (see <u>MRA</u> )
MRL	Maximum Residue Limit (see <u>Maximum residue limits</u> )
MRP	Mutual Recognition Procedure (see <u>Applying for EU marketing authorisation</u> )
MS(s)	Member State(s) of the European Union
MSP	multi-stakeholder platform (see <u>ACT-EU</u> )
MSSG	(EMA) Medicines Shortages Steering Group (see MSSG)
MUMS	Minor Use, Minor Species
MWP	(EMA CHMP) Methodology Working Party (see <u>MWP</u> )
NAMs	New Approach Methodologies
NAP(s)	Nationally Authorised Product(s)
NAS	New Active Substance
NB	Notified Body (see High-risk medical devices: consultation procedures and
	advice   European Medicines Agency (EMA))
NcWP	(EMA) Non-clinical Working Party (see <u>NcWP</u> )
NCA(s)	National Competent Authority(ies)
NCD(s)	Non-communicable disease(s) (see <u>EU Public Health NCDs</u> )
NfG	Note for Guidance
NRG	(EMA) [Invented] Name Review Group (see <u>NRG</u> )
NtA	Notice to Applicants (see <u>Eudralex – Volume 2</u> )
NTWP	(EMA CVMP) Novel Therapies and Technologies Working Party (see <u>NTWP</u> )
NUI	Non-Urgent Information (see also RA/NUI System)
OD	Orphan Designation (see <u>Orphan designation: Overview</u> )
00	orphan Designation (see <u>orphan designation. Overview</u> )

OE	Oral Explanation
OECD	Organisation for Economic Co-operation and Development
OEG(s)	(EMA) Operational Expert Group(s) (see <u>Working parties and domains</u> )
	- BOEG = Biostatistics Operational Expert Group
	- MSOEG = Modelling and Simulation Operational Expert Group
	- RWDOEG = Real World Data Operational Expert Group
OIE	World Organisation for Animal Health, based on its original name Office
	International des Epizooties – see also WOAH
OLAF	European Anti-Fraud Office
OMCL(s)	Official Medicines Control Laboratory(ies) ( <u>https://www.edqm.eu/en/omcl-</u>
	background-and-mission)
OMS	see ISO IDMP
ONCWP	(EMA CHMP) Oncology Working Party (see <u>ONCWP</u> )
OPEN initiative	Opening our Procedures at EMA to Non-EU authorities - see OPEN Pilot: one-
	year review and recommendations
ORGAM	Organisational Matters (see PROM; see also <u>HMPC</u> )
OS	Overall survival
ОТС	Over-the-counter
PA	Protocol Assistance (see <u>Scientific advice and protocol assistance</u> )
PaedPAR	Paediatric Public Assessment Report
PAES	Post-Authorisation Efficacy Study (see <u>PAES Q&amp;A</u> )
PAM(s)	Post Authorisation Measure(s) categorised as follows in EMA's product and
	procedure tracking database – see PAMs Q&A
	ANX = Annex II condition
	LEG = Legally Binding Measure
	MEA = Additional PhV activity in the RMP
	SOB = Specific Obligation
	REC = Recommendation
pAR	preliminary Assessment Report
PAS	Post-Authorisation Safety
PASS	Post-Authorisation Safety Study (see GVP annex I)
PBRER	Periodic Benefit-Risk Evaluation Report
PBT	Persistent Bioaccumulative Toxic (chemical)
PCO	Patients' and Consumers' Organisations
PCWP	(EMA) Patients' and Consumers' Working Party (see <u>PCWP</u> )
PCU	Population Correction Unit
PD	Personal Data
PD	Pharmacodynamic(s)
PD	(EMA) Parallel Distribution (see <u>Parallel distribution</u> )
PdAR	Paediatric Assessment Report
PDCO	(EMA) Paediatric Committee
PECP	Performance Evaluation Consultation Procedure (see <u>Medical Devices</u> )
PED	Patient Experience Data
PEM (study)	Prescription-Event Monitoring (study)
PFS	Progression-free survival
PHE	Public Health Emergency
Ph.Eur.	European Pharmacopoeia ( <u>https://www.edqm.eu/en/european-pharmacopoeia</u> )
rii.Lui.	European Filannacopoela ( <u>https://www.euqin.eu/en/european-pharmacopoela</u> )

PhV	Pharmacovigilance
PhV IWG	Pharmacovigilance Inspectors Working Group (see <u>Compliance: overview</u> )
PhVWP	Pharmacovigilance Working Party (working party that preceded the PRAC)
PhVWP-V	(EMA CVMP) Pharmacovigilance Working Party (see <u>PhVWP-V</u> )
PI	Product Information (see <u>Product Information requirements for human</u>
	medicines and Product Information requirements for veterinary medicines)
PICO	Population, Intervention, Comparator, Outcome
PIC/S	Pharmaceutical Inspection Co-operation Scheme (see PIC/S)
PIL	Patient Information Leaflet
PIP(s)	Paediatric Investigation Plan(s) (see <u>PIPs</u> )
PK	Pharmacokinetic(s)
PL	Package Leaflet
PL	(EMA) Product Lead
PLD	Patient Level Data
PLM	Product Lifecycle Management
PMDA	Pharmaceuticals and Medical Devices Agency (Japan) (see <u>International</u>
	agreements)
PMF	Plasma Master File (see <u>PMF certification</u> )
PMS	Post-Marketing Surveillance (see also under ISO IDMP)
POM	Prescription-only Medicine
PP	Per Protocol (analysis)
PPD	Protected Personal Data
PPP	Pregnancy Prevention Programme
PPP	Public-Private Partnership
PRA	Preliminary Risk Analysis (see IMP)
PRAC	(EMA) Pharmacovigilance Risk Assessment Committee
PRIME	(EMA) Priority Medicines scheme (see <u>PRIME</u> )
PRISMA	(EMA) PRAC Risk Minimisation Alliance
PRO	Patient-Reported Outcome (see HTA)
PROM	Patient-Reported Outcome Measure (see HTA)
PROM	(EMA CHMP) Preparatory and Organisational Matters (see <u>CHMP</u> – <i>formerly</i>
TROM	known as ORGAM)
PRP	Preliminary Risk Profiling (see <u>Use of antimicrobials in animals</u> )
PRR	Proportional Reporting Ratio
PSA	Parallel Scientific Advice
PSMF	Pharmacovigilance System Master File (for human medicines: see <u>GVP</u> annex I;
	for veterinary medicines: see <u>VGVP</u> )
PSUFU	PSUSA Follow-Up
PSUR	Periodic Safety Update Report (see GVP annex I)
PSUSA	PSUR Single Assessment
PUMA	Paediatric Use Marketing Authorisation (see <u>PUMA</u> )
QIG	(EMA CHMP/CVMP) Quality Innovation Group (see <u>QIG</u> )
QMS	Quality Management System
QoL	Quality of Life
Qol QoNM	Qualification of Novel Methodologies
QP	Qualified Person
QP QPPV	
QPPV	Qualified Person responsible for Pharmacovigilance

QRD-WG	(EMA) Working Group on Quality Review of Documents (see <u>QRD</u> )
QWP	(EMA CHMP/CVMP) Quality Working Party (see <u>QWP</u> )
RA	Rapid Alert – see also RA/NUI System
RA	Reference Authority
RA	Regulatory Affairs
rAAV	recombinant adeno-associated viral vector
RA/NUI System	Rapid Alert/Non-Urgent Information System
RCT(s)	Randomised Controlled Trial(s)
R&D	Research and Development
REA	Relative Effectiveness Assessment
REMS	Risk Evaluation & Mitigation Strategies
RFI	(EMA) Request for Information
RfR	Report for Release
RIWP	(EMA CHMP) Rheumatology/Immunology Working Party (see <u>RIWP</u> )
RMAT	Regenerative Medicine Advanced Therapy
RMM(s)	Risk Minimisation Measure(s) / Risk Mitigation Measure(s)
RMP or RefMP	Reference Medicinal Product
RMP	Risk Management Plan (see <u>GVP</u> annex I)
RMR	Reaction Monitoring Report
RMS or RefMS	Reference Member State (see also 'RMS' under ISO IDMP)
RMS	Risk Management System
ROG	Regulatory Optimisation Group (see <u>HMA ROG</u> )
RPCs	Regional Pharmacovigilance Centres
RPI	Research Product Identifier (see <u>Requesting SA or PA from EMA</u> )
RRR	Relative Risk Reduction
RSI	Request for Supplementary Information
RSS	(EMA) Regulatory Science Strategy (see <u>RSS</u> )
RCT	Randomised Clinical Trial
RUP	Repeat Use Procedure (see <u>CMDh MRP/RUP</u> )
RWD	Real World Data
RWE	Real World Evidence
SA	Scientific Advice
SAE	Serious Adverse Event
SAG(s)	(EMA) Scientific Advisory Group(s)
SAP	Statistical Analysis Plan
SAWP	(EMA CHMP) Scientific Advice Working Party (see <u>SAWP</u> )
SAWP-V	(EMA CVMP) Scientific Advice Working Party (see <u>SAWP-V</u> )
SB	Significant Benefit
SBP(s)	Similar Biotherapeutic Product(s) (WHO term for biosimilars)
SC	subcutaneous
SCAR	Serious Cutaneous Adverse Reaction
SEND	Standard for Exchange of Nonclinical Data
SFDA	State Food and Drug Authority (China) (see International agreements)
SmAR	Summary Assessment Report
	Builling / lobebolliene report
SMEs	Small and Medium-sized Enterprises (see <u>Support to SMEs</u> )
SMEs SME	

SmPC SMQs SMS SNSA SoC SOC SOC SOH SOH SOP SPC SPOC SPOR SRLM
SMS SNSA SoC SOC SOH SoHo SOP SPC SPOC
SNSA SoC SOC SOH SoHo SOP SPC SPOC
50C 50C 50H 50H0 50P 5PC 5POC
SOC SOH SoHo SOP SPC SPOC
SOH SoHo SOP SPC SPOC
SoHo SOP SPC SPOC
SOP SPC SPOC
SPC SPOC SPOR
SPOC
SPOR
אררה
SSR
STAMP
SUSAR
Swissmedic
SWP-V
ГСМ
DGs
TDD
ГGA
ГНМР
TMF
ГоС
ГоС
ГоD
ГТР
ГU
VICH
VICH
TMF ToC ToC ToD TTP TU TUR JDI JI JMN JPD JPhV JSR VarWP

VMP	(ECDC/EMA) Vaccine Monitoring Platform (see Vaccine Monitoring Platform)	
VMP	Veterinary Medicinal Product	
VNeeS	Veterinary Non-eCTD Electronic Submission	
VNRA	Variation Not Requiring Assessment	
VRA	Variation Requiring Assessment	
VWP	(EMA CHMP) Vaccines Working Party (see <u>VWP</u> )	
WEU	Well-established use	
WG	Working Group	
WHO	World Health Organization (see <u>WHO</u> )	
WHO-UMC	WHO-Uppsala Monitoring Centre	
WOAH	World Organisation for Animal Health	
WP	Working party (see Working parties and domains)	
WS	Work Sharing	

#### **Country codes of EU/EEA Countries<sup>2</sup>**

Country (short name in English)	Country Code	Agency	Acronym
Austria	AT	Austrian Agency for Health and Food Safety	AGES
Belgium	BE	Federal Agency for Medicines and Health Products	FAMHP
Bulgaria	BG	Bulgarian Drug Agency	BDA
Bulgaria (V)	BG	Bulgarian Food Safety Authority	BFSA
Croatia	HR	Agency for medicinal products and medical devices of Croatia	HALMED
Croatia (V)	HR	Ministry of Agriculture - Veterinary and food safety directorate	MPS
Cyprus	CY	Ministry of Health -Pharmaceutical Services	мон
Cyprus (V)	CY	Veterinary Services, Ministry of Agriculture, Natural Resources and Environment	MOA
Czechia	CZ	State Institute for Drug Control	SUKL
Czechia (V)	CZ	Institute for State Control of Veterinary Biologicals and Medicines	USKVBL
Denmark	DK	Danish Medicines Agency	DKMA
Estonia	EE	State Agency of Medicines	SAM
Finland	FI	Finnish Medicines Agency	FIMEA
France	FR	National Agency for the Safety of Medicines and Health Products	ANSM

<sup>&</sup>lt;sup>2</sup> Sources: <u>https://ec.europa.eu/eurostat/statistics-explained/index.php?title=Glossary:Country\_codes</u>; <u>https://commission.europa.eu/strategy-and-policy/policies/eu-enlargement\_en</u>; <u>Tutorial:Country\_codes and protocol order</u> <u>- Statistics Explained</u>; <u>Online Browsing Platform (OBP)</u>

France (V)	FR	French Agency for Food, Environmental and Occupational Health & Safety	ANSES
Germany (H+V)	DE	Federal Institute for Drugs and Medical Devices	BfArM
Germany (H+V)	DE	Paul Ehrlich Institute	PEI
Greece	GR (ISO) EL <sup>2</sup>	National Organization for Medicines	EOF
Hungary	HU	National Centre for Public Health and Pharmacy	NNK
Hungary (V)	HU	Directorate of Veterinary Medicinal Products	NEBIH
Iceland	IS	Icelandic Medicines Agency	IMA
Ireland	IE	Health Products Regulatory Authority	HPRA
Italy	IT	Italian Medicines Agency	AIFA
Italy (V)	IT	Ministry of Health	
Latvia	LV	State Agency of Medicines	ZVA
Latvia (V)	LV	Food and Veterinary Service	PVD
Liechtenstein	LI	Office of Health/ Department of Pharmaceuticals	LLV
Lithuania	LT	State Medicines Control Agency	VVKT
Lithuania (V)	LT	State Food and Veterinary Service	VMVT
Lithuania (V)	LT	National Food and Veterinary Risk Assessment Institute	NMVRVI
Luxembourg	LU	Ministry of Health	MS
Malta	МТ	Malta Medicines Authority	ММА
Malta	MT	Veterinary and Phytosanitary Regulation Department	
Netherlands	NL	Medicines Evaluation Board	CBG-MEB
Norway	NO	Norwegian Medicinal Products Agency	DMP
Poland	PL	Office for Registration of Medicinal Products, Medical Devices and Biocidal Products	URPL
Portugal	PT	National Authority of Medicines and Health Products	INFARMED
Portugal (V)	PT	National Authority for Animal Health	DGAV
Romania	RO	National Agency for Medicines and Medical Devices of Romania	ANM
Romania (V)	RO	Institute for Control of Biological Products and Veterinary Medicines	ICBMV
Slovakia	SK	State Institute for Drug Control	SUKL
Slovakia (V)	SK	Institute for State Control of Veterinary Biologicals and Medicaments	USKVBL

Slovenia	SI	Agency for Medicinal Products and	JAZMP
		Medical Devices of the Republic of	
		Slovenia	
Spain	ES	Spanish Agency of Medicines and	AEMPS
		Medical Devices	
Sweden	SE	Swedish Medical Products Agency	MPA

### Country Codes of EU candidate countries<sup>2</sup>

Country	Country Code
Bosnia and Herzegovina	BA
Montenegro	ME
Moldova	MD
North Macedonia	МК
Georgia	GE
Albania	AL
Serbia	RS
Turkey	TR
Ukraine	UA

## **Country Codes of Other European Countries**<sup>2</sup>

Country	ISO Country Code
Andorra	AD
Armenia	AM
Azerbaijan	AZ
Belarus	BY
Holy See (Vatican City State)	VA
Козоvо	ХК
Monaco	MC
Russia	RU
San Marino	SM
Switzerland	СН
Vatican City State	See Holy See

## Other Country Codes<sup>2</sup>

Country	ISO Country Code
Australia	AU
Canada	CA
China	CN
Japan	JP
New Zealand	NZ
United States (of America)	US(A)