

# **GLOSSARY OF REGULATORY HEALTHCARE ACRONYMS & ABBREVIATIONS**

www.topra.org/glossary

Note: Medical prescription abbreviations can be found at www.abbreviations.com/acronyms/PRESCRIPTION

1-1-1 - One dossier, one European scientific assessment, one decision for marketing authorisation **3Rs** – Replacement, refinement and reduction (in research using animals) 510(k) – Medical device premarket notification (US FDA) **AA** – Accelerated assessment/approval AAC - Accelerated Access Collaborative (UK) AADA – Abbreviated antibiotic drug application AAP - Accelerated approval pathway (US) - and also: **AAP** – Accelerated assessment procedure (EU) AAPS - American Association of Pharmaceutical Scientists AAR – Accelerated access review **AAS** – Atomic absorption spectroscopy AAV - Adeno-associated virus **ABHI** – Association of British Healthcare Industries (medical devices sector) **ABPI** – Association of the British Pharmaceutical Industry A-CASI – Audio computer-assisted self-interviewing ACO – Addendum to clinical overview ACRP - Association of Clinical Research Professionals ACSS - Australia, Canada, Singapore, Switzerland Consortium **ACT** – Artemisinin-based combination therapy ACTD - ASEAN common technical dossier (see ASEAN) ACVM - Agricultural Compounds and Veterinary Medicines (New Zealand) ADA - Anti-drug antibodies ADaM – Analysis data model ADC – Additional data collection – and also: **ADC** – Antibody–drug conjugate ADCC – Antibody-dependent cellular cytotoxicity **ADE** – Adverse device event (AE judged to be related to the medical device) ADEC – Australian Drug Evaluation Committee ADI - Acceptable daily intake ADME - Absorption, distribution, metabolism and excretion/elimination (also AME - absorption, metabolism, excretion/elimination) **ADR** – Adverse drug reaction ADROIT - Adverse Drug Reactions On-Line Tracking System **ADVAC** – Ad hoc group on veterinary vaccine availability (CVMP) **ADVENT** – Ad Hoc Expert Group on Veterinary Novel Therapies AE - Adverse event AEFI - Adverse event following immunisation AEGIS - Adverse Experience Gathering Information System **AEM** – Agencia Espanola Medicamento (Spain) AEMPS – Agencia Española de Medicamentos y Productos Sanitarios (Spain) AEPAR - Associación Española de Profesionales de Actividades de Registro (Spanish Regulatory Affairs Association) AERS - Adverse event reporting system (US FDA) AESGP – Association Européenne des Spécialitiés Pharmaceutiques Grand Public (Association of the European Self-Medication Industry) AF - Application Form AFAR – Association Française des Affaires Reglémentaires (French Regulatory Affairs Association) AFDO - Association of Food and Drug Officials (US) AFMPS – Agence Fédérale des Médicaments et des Produits de Santé (Belgium) Afssaps – former French regulatory agency (Agence Française de Sécurité Sanitaire des Produits de Santé) – replaced by **ANSM** in 2012 (see below) AGES PharmMED – Osterreichische Agentur fur Gesundheit und Ernahrungssicherheit GmbH (Austria's medicines & devices agency) AHSC - Academic Health Science Centre (UK) **AHWP** – Asian Harmonisation Working Party AI – Adverse incident (medical devices sector) – and also:



**AI** – Artificial intelligence **AIFA** – Agenzia Italiana del Farmaco (Italy's health authority) **AIM** – Active ingredient manufacturer **AIMD** – Active implantable medical device AITS – Adverse Incident Tracking System (medical devices sector) **AKP** – Alkaline phosphatase ALARP – As low as reasonably practical ALATF - As low as technically feasible (terminology superseded by "ALARP" - see above) ALIMS - Medicines and Medical Devices Agency (Serbia) ALL – Acute lymphocytic leukaemia ALT – Alanine aminotransferase (ALT = SGPT) AM – Agence du Medicament (France) AMA – American Medical Association AMEG – AntiMicrobial advice ad hoc Expert Group AMI – Acute myocardial infarct AML - Acute myeloid leukemia **AMM** – Autorisation de mise sur le marché (France) = Product licence **AMP** – Authorised medicinal product – **and also:** AMP - Auxiliary medicinal product (formerly non-investigational medicinal product, NIMP, **AMR** – Antimicrobial resistance **AMRH** – African Medicines Regulatory Harmonisation **ANADA** – Abbreviated New Animal Drug Application (US) **ANDA** – Abbreviated new drug application **ANDS** – Abbreviated new drug submission (Canada) **ANMV** – Agence nationale du médicament vétérinaire (French vet medicines agency) ANOVA - Analysis of Variance **ANPR** – Advanced notice of proposed rulemaking (US) ANSES – Agence Francaise de Securite Sanitaire des Aliments Agence nationale due medicament veterinaire **ANSM** – French regulatory agency (Agence nationale de sécurité du médicament et des produits de santé) [formerly Afssaps] **ANZTPA** – Australia New Zealand Therapeutic Products Agency (scheduled to come into force in 2016 – replacing Australia's TGA and New Zealand's Medsafe) AO – Auditing organisation AOAC – Association of Official Analytical Chemists (US) **AOB** – Any other business AP - Accredited person - and also: **AP** – Adaptive pathway **APEC** – Asia-Pacific Economic Cooperation **APHIS** – Animal and Plant Health Inspection Service (US) **API** – Active pharmaceutical ingredient **APIC** – Active Pharmaceutical Ingredients Committee APLB - Advertising and Promotional Labeling Branch (FDA's CBER) **APMA** – Australian Pharmaceutical Manufacturers Association APVA – Additional pharmacovigilance activities **APVMA** – Australian Pesticides and Veterinary Medicines Authority (Australia) AQL - Acceptable quality level AR – Adverse reaction – and also: AR - Assessment Report (EU) - and also: AR – Authorised representative ARfD – Acute reference dose (veterinary) **ARMAs –** Additional risk minimisation activities **ARMMs** – Additional risk minimisation measures AS – Active Substance **ASAP** – Accelerated Stability Assessment Program ASCII – American Standard Code for Information Interchange Quality Assurance ASDI – Acceptable single-dose intake **ASEAN** – Association of Southeast Asian Nations **ASMF** – Active Substance Master File **ASMF WG** – Working Group on Active Substance Master File procedures **ASPR** – Anonymised single patient report (formerly ASPP – anonymised single patient printout) **ASR** – Annual safety report **AST** – Aspartate aminotransaminase (AST = SGOT)



ATA – Alternatives to antibiotics ATC - Anatomical - therapeutic - chemical (WHO) - and also: ATC – Animal Test Certificate (UK) – and also: ATC Code - Anatomical Therapeutic Chemical Code ATC Vet Code - Anatomical Therapeutic Chemical Veterinary Code ATC(/DDD) - Anatomical Therapeutic Chemical classification system (with Defined Daily Doses) **ATD** – Access to documents (EMA policy) – and also: ATD - Anticipated therapeutic dose - and also: ATD - Anti-tampering device **ATECT** – Advanced T-cell Engineering for Cancer Therapy ATF – Alcohol – Tobacco and Firearms (Bureau of) (US) ATMPs - Advanced therapy medicinal products (aka "advanced therapies") **ATU** – Authorisation for temporary use  $AUC_{\infty}$  \_ Area under the concentration time curve between zero and infinity AUCx – Area under the curve during a given time AVEG – AIDS Vaccine Evaluation Group AWP – Antimicrobials Working Party **AXREM –** Association of X-ray Equipment Manufacturers AYA – Adolescents and young adults BBB **BA** – Bioavailability **BA/BE** – Bioavailability/bioequivalence **BACPAC** – Bulk active chemical post approval changes (US) BAI - Breath actuated inhaler **BAID** – Batch identifier **BAN** – British Approved Name BAP - Biotechnology Action Programme/Biosimilars Action Plan BARQA - British Association of Research Quality Assurance BCS – Biopharmaceutics Classification System **bd/bid** – twice a day (Latin: bis in die) **BDA** – Bulgarian drug agency **BE** – Bioequivalence **BEMA** – Benchmarking of European Medicines Agencies BfArM – Federal Institute for Drugs and Medical Devices (Bundesinstituts für Arzneimittel und Medizinprodukte) (Germany's regulatory authority) BGMA - British Generic Manufacturers Association **BIND** – Biological investigational new drug **BIO** – Biotechnology Industry Organization (US) **BLA** – Biologics license application (US) BM – Bone marrow **BMA** – British Medical Association **BMD** – Bone mineral density **BMG** – Bundesministerium für Gesundheit = Federal Ministry of Health (Germany) **BMGF** – Bundesministerium fuer Gesundheit und Frauen (Austrian agency) **BMWP** – Biosimilar Medicinal Products Working Party **BNF** – British National Formulary BoH – Board of Health BOS - Break-out session **BP** – Blood pressure – **and also:** BP – British Pharmacopoeia BPC - British Pharmacopoeia Commission - and also: **BPC** – Bulk pharmaceutical chemicals **BPCA** – Best Pharmaceuticals in Children Act (US) BPG – Best Practice Guide BPI - Bundesverband der Pharmazeutischen Industrie (German pharmaceutical industry trade association) **BPR** – Biocidal Products Regulation **BPWP** – Blood Products Working Party (EMA) Br – Barrier reared (in older reports – 'Brown') **BRAS** – Belgian Regulatory Affairs Society BRAT – Benefit–Risk Action Team



BRIC - Brazil, Russia, India & China BRICK- Brazil, Russia, India, China & (South) Korea BRICS - Brazil, Russia, India, China & South Africa BROMI - Better Regulation of Over the Counter Medicines Initiative **BSE** – Bovine Spongiform Encephalopathy **BTD** – Breakthrough therapy designation (US) **BTDR** – Breakthrough therapy designation request BTF – Brexit Task Force **BWP** – Biotech Working Party (EMA) CCC **C&P** – Chemistry and Pharmacy CA - Commercial appraisal - and also: CA - Competent authority **CAC** – Codex Alimentarius Commission (veterinary sector) CAD - Coronary artery disease **CADREAC** – Collaboration agreement between drug regulatory authorities of European Union associated countries (also **nCADREAC** – new Collaboration Agreement) **CADTH** – Canadian Agency for Drugs and Technologies in Health (formerly CCOHTA) **CAMD** – Competent Authorities for Medical Devices **CAMS** – Chinese Academy of Medical Sciences **CANDA** – Computer assisted new drug application **CAO** – Central Agricultural Office (Hungary) **CAP** – Centrally authorised product **CAPA** – Corrective action and preventive action CAPA plan - Corrective and preventive action plan **CAPLA –** Computer Assisted Product Licence Application **CAPRA –** Canadian Association of Pharmaceutical Regulatory Affairs **CAR** – Chimeric antigen receptor **CARPHA** – The Caribbean Public Health Agency **CAS** – Central alerting system (UK) – and also: **CAS** – Chemical abstract systems **CAT** – Committee for Advanced Therapies (EMA) CATMP - Combined Advanced Therapy Medicinal Product **CAVDRI** - Collaboration agreement between veterinary drug registration institutions **CAVOMP** – Clinical added value orphan medicinal product **CBER** – Center for Biologics Evaluation and Research (US FDA) **CBG/MEB** – Medicines Evaluation Board (the Netherlands) **CBP** – Corticoid binding protein CC – Candidate country (EU) **CCDP** – Complete clinical data package CCDS – Company core data sheet CCG - Clinical Commissioning Group (UK NHS) CCG IAC - Clinical Commissioning Group Indicator Advisory Committee CGTPs - Cell and gene therapy products **CCI** – Commercially confidential information CCRB - Change control review board **CCSI** – Company core safety information CD - Caesarean derived - and also: CD - Controlled drug CDA – China Drug Administration **CDC** – Centers for Disease Control and Prevention (US) **CDDD** – Clinical dossier of drug development (Brazil) CDE - Center for Drug Evaluation (China) **CDEC** – Canadian Drug Expert Committee (Canada) **CDER** – Center for Drug Evaluation and Research (US FDA) **CDISC** – Clinical Data Interchange Standards Consortium **CDMA** – Canadian Drug Manufacturers Association **CDR** – Common Drug Review (Canada) **CDRH** – Center for Devices and Radiological Health (US FDA) CDS – Clinical decision support



**CDSCO** – Central Drug Standard Organization (India's clinical trials licensing authority) **CDSM** – Committee on Dental and Surgical Materials (UK) **CDx** – Companion Diagnostics **CE Mark** – Conformité European (approval for EU medical devices) **CEA** – Cost-effectiveness analysis **CEC** – Central ethics committee – and also: **CEC** – Commission of the European Communities **CED** – Coverage with evidence development **CEE** – Central and Eastern Europe **CEEC** – Central and Eastern European Countries **CEFTA –** Central Europe Free Trade Area CEN – Comité Européan des Normes – European Committee for Standardization CEP - Central enquiry point (MHRA) - and also: **CEP** – Certificate of European Pharmacopoeia (aka Certificate of Suitability) CER – Clinical evaluation report – and also: **CER** – Comparative effectiveness research **CESP** – Common European submission portal **CF** – Cystic fibrosis CFC - Chlorofluorocarbons CFDA - China Food and Drug Administration (formerly State FDA - SFDA) **CFR** – Code of Federal Regulations (US) CFS - Certificate of Free Sale **CFSAN** – Center for Food Safety and Applied Nutrition (US) **cGLP** – Current good laboratory practice **cGMP** – Current good manufacturing practice CGP – Clinical Guidance Panel (Canada) **CH** – Clinical hold CHAI - Commission for Healthcare Audit and Inspection (UK) CHC - Consumer healthcare **CHMB** - Creatine kinase Muscle Brain CHMP - Committee for Medicinal Products for Human Use (EMA) CHMP - Committee for Medicinal Products for Human Use (previously: CPMP) CHO – Chinese hamster ovary cells CHPA – Consumer Healthcare Products Association CI - Confidence Interval, and also: **CI** – Contraindication **CIA** – Corporate Integrity Agreement (US) **CIOMS** - Council for International Organizations of Medical Sciences (WHO) **CIRS** – Centre for Innovation in Regulatory Science CIS (countries) - Commonwealth of Independent States (members are former Soviet Republic countries, currently including Armenia, Azerbaijan, Belarus, Kazakhstan, Kyrgyzstan, Moldova, Russia, Tajikistan, Usbekistan, Turkmenistan, Ukraine CK – Creatine kinase **CI** – Total body clearance Class Im – Class I with measuring function (medical devices) **CLIA** – Clinical Laboratory Improvement Amendments (US) CLL – Chronic lymphocytic leukaemia **CLO** – Clinical overview **CLP** – Classification, labelling and packaging (medical devices) **CLS** – Clinical summary Cm or Cmax – Maximum plasma concentration at steady state **CMA** – Conditional marketing authorisation (US) **CMC** – Chemistry, manufacturing, and controls CMDCAS - Canadian Medical Devices Conformity Assessment System CMDh - Co-ordination Group for Mutual Recognition and Decentralised Procedures - Human (EMA) **CMDR** – Canadian Medical Device Regulation CMDv - Co-ordination Group for Mutual Recognition and Decentralised Procedures - Veterinary (EMA) CMN – Comité de Moléculas Nuevas" (New Molecules Committee) (Mexico) CMP - Certificate of Medicinal Product - and also: CMP – Common product model CMR - Carcinogenic, mutagenic or reprotoxic [toxic to reproduction] - and also:



**CMR** – Centre for Medicines Research **CMS** – Concerned member state (EU) CMT – Convergent medical technologies COA/CofA - Certificate of analysis CoAg - Cooperative Agreement COE - Council of Europe **COMET –** Core Outcome Measures in Effectiveness Trials **COMP** – Committee for Orphan Medicinal Products (EMA) **COREPER** – Committee of Permanent Representatives to the Community **COSHH –** Control of Substances Hazardous to Health **COSTART** – Coding Symbols for a Thesaurus of Adverse Reaction Terms CoU - Context of Use CP - Centralised procedure (EU) - and also: CP - Comparability protocol (US) **CPAC** – Central Pharmaceutical Affairs Council (Japan) **CPC** – Combination Products Coalition **CPD** – Continuing professional development **CPI** – Critical Path Initiative (US) **CPMP** – Committee for Proprietary Medicinal Products (EMA) **CPP** – Certificate of pharmaceutical product – and also: **CPP** – Critical process parameter **CPQ** – Costs per quality-adjusted life year **CPR** – Cosmetic Products Regulation **CPRD** – Clinical Practice Research Datalink (MHRA) **CPS** – Chemistry – Pharmacy and Standards Subcommittee of the CSM (UK) – and also: **CPS** – Clinical performance study **CPSP** – Clinical performance study plan **CPU** – Clinical pharmacology unit **CPWP** – Cell-based Products Working Party (EMA) CQA – Clinical quality assurance – and also: CQA – Critical quality attribute CR – Computed radiology – and also: **CR** – Controlled releasse CRF - Case report form CRG – Clinical reference group (UK) CRO - Clinical Research Organisation CRP - Canadian reference product (WHO) - and also: **CRP** – Collaborative registration procedure CRS – The Caribbean Regulatory System – and also: **CRS** – Cytokine release syndrome **CS** – Clinically significant – **and also: CS** – Common specifications CSA - Controlled Substances Act CSI - Core safety information CSM - Centralised statistical monitoring - and also: **CSM** – Committee on Safety of Medicines (UK) CSO - Consumer Safety Officer (US) CSP - Core safety profile CSR – Clinical study report (EU) **CSV** – Comma-separated values CT – Clinical trial – and also: **CT** – Computed tomography **CTA** – Clinical trial application – **and also:** CTA – Clinical trial assay – and also: CTA – Clinical trial authorisation CTAG – Clinical Trials Action Group (Australia) – and also: **CTAG** – Clinical Trials Coordination and Advisory Group **CTC** – Clinical trial certificate (Hong Kong, Singapore) CTD – Clinical Trials Directive – and also: CTD – Common technical document\* [\*Although 'dossier' has become commonplace – the correct term is

`document']

**CTEG** – Clinical Trials Expert Group



- **CTFG** Clinical Trials Facilitation Group
- CTIS Clinical Trials Information System (formerly the EU clinical trial portal and database, EudraCT)
- **CTMP** Cell therapy medicinal product
- CTMS Clinical trial management system
- **CTN** Clinical trial notification (Australia)
- CTOC Comprehensive Table of Contents Headings and Hierarchy
- **CTR** Clinical Trial Regulation
- CTS Common technical specification and also:
- **CTS** Communication Tracking System (formerly Eudratrack)
- CTTI Clinical Trials Transformation Initiative
- CTU Clinical trials unit
- CTX Clinical trial exemption (UK)
- **CUA** Cost utility analysis
- **CUP** Compassionate use programme
- **CV** Controlled vocabulary
- **CVM** Center for Veterinary Medicine (US)
- **CVMP** Committee for Medicinal Products for Veterinary Use (EMA)
- CVO Chief Veterinary Officer
- **CVS** Cardiovascular system
- CVZ Dutch Health Care Insurance Board
- **CWoW** Combined Ways of Working
- CZ Climatic zone

## DDD

- **DAB** German Pharmacopoeia (Deutsches Arznei Buch)
- **DAC** Data analysis centre
- DACS Detailed and critical summary
- **DAE –** Discontinuation due to an adverse event
- DAL Defect action level (US)
- DAMOS Drug application methodology with optical storage
- **DB** Device Bulletin (MHRA)
- DCGI Drugs Controller General of India
- **DCGI** India's regulatory authority (Directorate General of Health Services in the Ministry of Health and Family Welfare)
- DCP Decentralised procedure (EU)
- **DCTs** Decentralised clinical trials
- **DD** District Director (US)
- **DDC(P)** Drug-device combination (product)
- **DDD** Defined daily dose
- DDMAC Division of Drug Marketing, Advertising and Communications (CDER)
- DDPS Detailed description of pharmacovigilance system
- **DDX** Doctors' and dentists' exemption (UK)
- **DE** Designated examination
- **DEA** Drug Enforcement Agency (US)
- **DEREK** Deductive estimate of risk from existing knowledge
- DES Data exchange standard (EU) and also:
- **DES** Drug eluting stent
- **DESI** Drug efficacy study implementation (US)
- DG Directorate-General (at the European Commission)
- **DGEM** Disease-gene expression matching
- **DGV** Direccao Geral de Veterinaria (Veterinary Medicines Agency) (Portugal)
- **DH** Department of Health (UK) **and also:**
- DH Digital healthcare
- **DHHS** Department of Health and Human Services (US)
- DHPC Direct healthcare professional communication (formerly 'Dear Doctor Letter')
- **DIA** Drug Information Association (US)
- **DIBD** Development international birth date
- **DID** Design inputs document
- **DIMDI** Deutsches Institut für Medizinische Dokumentation und Information (Germany)
- DKMA Lægemiddelstyrelsen/Danish Medicines Agency (Denmark)
- **DLP** Data lock point



**DMF** – Drug master file **DMPK** – Drug metabolism and pharmacokinetics DMRC – Defective Medicines Report Centre (MHRA) **DMS** – Document management system **DMT** – Disease modifying therapy **DOE** – Design of experiments **DoR** – Duration of Response **DP** – Drug product **DPI** – Dry powder inhaler **DPIA –** Data protection impact assessment DPO – Data Protection Officer **DPR** – Data Protection Representative – and also: **DPR** – Dual Pack import Registration DR - Deliberate release - and also: **DR** – Digital radiology **DRA** – Drug Regulatory Authority **DRF(S)** – Dose range finding (study) **DRMP** – Developmental risk management plan DRR – Drug Registration Regulation (China) – and also: **DRR** – Durable response rate **DS** – Drug substance **DSC** – Differential scanning calorimetry **DSMC** – Data safety monitoring committee **DSRU** – Drug Safety Research Unit (EMA) **DSUR –** Development safety update report DTaP - Diphtheria, tetanus and pertussis DTC - Direct-to-consumer **DTD** – Document type definition **DUNS** – Data universal numbering system **DUS** – Drug utilisation study **DVPHNFS** – Department for Veterinary Public Health, Nutrition and Food Safety (Italy) **DWH** – Data warehouse **Dx** – Diagnostic EEE **EA** – Environmental assessment **EAC** – East African Community **eAF** – electronic Application Form EAI – Estimated acute intake EAMS - Early Access to Medicines Scheme (UK) **EBE** – European Biopharmaceutical Enterprises EbM – Evidence-based medicine EC - Established conditions (ICH Q12 Guideline) - and also: EC – Ethics committee – and also: EC – European Commission – and also: **EC** – Exceptional circumstances **ECDC** – European Centre for Disease Prevention and Control ECG – Electrocardiogram ECHAMP – European Coalition on Homoeopathic and Anthroposophic Medicinal Products ECHR - European Court of Human Rights ECJ – European Court of Justice **ECPHIN** – European Community Pharmaceutical Information Network **ECRAB** – European Committee on Regulatory Aspects of Biotechnology (EBCG) eCRF – electronic case report form eCTD – electronic common technical document [not dossier\*] \*Although 'dossier' has become commonplace – the correct term is 'document' ED – Early dialogue EDA – Egyptian Drug Authority **EDC** – electronic data capture EDMF - European drug master file eDMS - electronic document management system



**EPDB** – European Data Protection Board EDQM – European Directorate for the Quality of Medicines EDQM – European Directorate for the Quality of Medicines | EDT – Electronic data transfer ED<sub>X</sub> – Effective dose at X% EEA - European Economic Area (comprising the EU countries, plus Iceland, Liechtenstein and Norway) **EEC** – European Economic Community **EEG** – Electroencephalogram eERA – extended Environmental Risk Assessment **EEU** – Eurasian Economic Union **EFA** – European Federation of Allergy and Airways Diseases Patients' Associations **EFPIA** – European Federation of Pharmaceutical Industries and Associations (http://www.efpia.eu) **EFPIA** – European Federation of Pharmaceutical Industries and Associations **EFQM** – European Foundation for Quality Management EFSA – European Food Safety Authority **EFTA** – European Free Trade Association EGA – European Generic medicines Association – Name changed 10 March 2016 to "Medicines for Europe" **EGGVP** – European Group for Generic Veterinary Products EGP – Economic Guidance Panel (Canada) EHR – Electronic health record EIA – Environmental Impact Assessment **EINECS** – European Inventory of Existing Chemical Substances ELA – Establishment license application (US) EMA – European Medicines Agency (formerly European Medicines Evaluation Agency – EMEA) **EMACOLEX** – European Medicines Agencies Co-operation of Legal and Legislative Issues **EMCDDA** – European Monitoring Centre for Drugs and Drug Addiction EMEA – Europe, Middle East & Africa EMEA - see above - and also: EMEAA – Europe, Middle East, Africa & Asia **EMR** – Electronic medical records **EMRC** – European Medical Research Councils (a unit of the **ESF** – see below) **EMVO** – European Medicines Verification Organisation **EMVS** – European Medicines Verification System ENCePP – European Network of Centres for Pharmacoepidemiology and Pharmacovigilance eNDA – Electronic New Drug Application **ENDS** – Electronic nicotine delivery system **ENP** – European Neighborhood Policy Enpr-EMA - European Network of Paediatric Research at the European Medicines Agency **ENS** – Early notification system EOF – Ethnikos Organismos Farmakon – aka National Organization for Medicines (Greece's regulatory agency) EoP – End of Procedure EOP1 – End of Phase 1 (US) EOP2 – End of Phase 2 (US) EOQ – European Organization for Quality EP – European Parliament – and also: EP/Ph Eur – European Pharmacopoeia (aka Pharm Eur) **EPA** – Environmental Protection Agency (US) and (Ireland) EPAA - European Partnership for Alternative approaches to Animal testing **EPAD** – European Prevention of Alzheimer's Dementia **EPADES** – European Parliament Document Exchange Server **EPAR** – European public assessment report **EPC** – European Pharmacopoeia Commission **EPHA** – European Public Health Alliance ePI – Electronic product information **EPI** – Essential Program for Immunisation EPID – Extended (also Expanded) Public Information Document **EPITT** – European Pharmacovigilance Issues Tracking Tool **EPL** – Effective patent life **EPO** – European Patent Office EPPOSI - European Platform for Patients' Organisation - Science & Industry **EPPV** – Early post-marketing phase vigilance (eg, in Japan)



EPRG – European Pharmacovigilance Research Group **EPRUMA** – European Platform for the Responsible Use of Medicines in Agriculture EPS - Eco-Pharmaco-Stewardship **ePSUR** – electronic periodic safety update report **EQM** – Equivalence margin ERs - Essential requirements (devices) ERA – Environmental risk assessment – and also: ERA - European regulatory affairs ERB - Ethical review board eRMR – electronic Reaction Monitoring Report ERMS - European risk management strategy **ERMS-FG** – European Risk Management Strategy Facilitation Group (HMA) **ERP** – European Reference Medicinal Product ESF - European Science Foundation ESG - Electronic submissions gateway (FDA) ESM – European stakeholder model ESPAR - Executive Summary Pharmacovigilance Assessment Report (EU) ESR - Erythrocyte Sedimentation Rate ESRA – European Society of Regulatory Affairs **ESTRI** – Electronic Standards for the Transfer of Regulatory Information **ESVAC** – European Surveillance of Veterinary Antimicrobial Consumption **ETASU** – Elements to ensure safe use (US) eTMF - electronic Trial Master File ETOMEP - European Technical Office for Medical Products (within EMA) EU - European Union EU5 - Group of countries comprising Germany, France, Italy, Spain and the UK **EUA** – Emergency use authorisation EU-ADR – Exploring and Understanding Adverse Drug Reactions by Integrative Mining of Clinical Records and Biomedical Knowledge (formerly known as ALERT) (EU) **EUBAN –** European Borderline Assessment Network **EU-IN** – EU Innovation Network **EUCERD –** EU Committee of Experts on Rare Diseases **EUCOMED** – European Confederation of Medical Device Associations **EUDAMED** – European Databank on Medical Devices **EUDRA** – European Union Drug Regulatory Authorities EudraCT - European Union Drug Regulatory Authorities Clinical Trials database EudraNet - European Union Drug Regulatory Authorities Network EudraSmPC – Summary of Product Characteristics **EUnetHTA** – European Network for Health Technology Assessment EU-NTC – EU Network Training Centre **EUPATI** – European Patients' Academy on Therapeutic Innovation **EUPD –** EU Portal and Database **EuPFI** – European Paediatric Formulation Initiative EURD - European Union reference date **EUREC** – European Network of Research Ethics Committees **EURL** – EU reference laboratory **EUR-OP** – EU Office for Publications **EUTCT** – European Union Telematics Controlled Terms **EUTMB** – EU Telematics Management Board EV – EudraVigilance – European Union Drug Regulating Authorities Pharmacovigilance **EVALI** – e-cigarette or vaping product use-associated lung injury **EVCTM** – EudraVigilance Clinical Trial Module EV-EWG – EudraVigilance Expert Working Group **EVIDENT** - Evidence Database on New Technologies **EVM** – European Vaccine Manufacturers **EVMPD** – EudraVigilance medicinal products dictionary **EVPM** – EudraVigilance post-authorisation module **EVPRM** – EudraVigilance product report message **EWG** – Expert Working Group **EWP** – Efficacy Working Party (EMA)



FACC – Food Additives and Contaminants Committee (UK) FAGG – Federaal Agentschap voor Geneesmiddelen en Gezondheidsproducten (Belgium) FAIR (data) - Findable, accessible, interoperable and reusable (data) FAMHP – Federal Agency for Medicines and Healthcare Products (Belgium) FAR – Final assessment report Farmindustria – Association of Italian Pharmaceutical Manufacturers (Italy) FCC – Food and Chemical Codex FDA – Food and Drug Administration (the US regulatory authority) FDAAA - FDA Amendments Act FDAMA – FDA Modernization Act FDASIA – Food and Drug Administration Safety and Innovation Act FDC – Fixed dose combination FDC Act - Food - Drug and Cosmetic Act (US) FDF – Finished dosage form FIH - First-in-human (aka FIM - first-in-man; and FTIM - first-time-in-human) FIM – First-in-man FIM-A - Federal Institute for Medicines (Austria) **FIMEA** – Finnish Medicines Agency (Finland) FIP - International Pharmaceutical Federation **FMD** – Falsified Medicines Directive (EU) FMEA – Failure mode and effect analysis FMECA – Failure Modes Effects and Criticality Assessment FNOM-CeO – Federazione Nazionale degli Ordini dei Medici-Chirurghi e degli Odontoiatri (IT) = Italian organisation of doctors and dentists FOB – Follow-on biologic FOFI – Federazione Ordini Farmacisti Italiani (IT) = Italian Organisation of Pharmacists FOI Act – Freedom of Information Act (US) FOM - Francophone Overseas Markets FONSI – Finding of no significant impact FOP – Follow-on protein FPA – Food producing animal FPFV – First patient first visit **FPIF** – Finnish Pharmaceutical Industry Association FPP – Finished pharmaceutical product FPRC – Final product release control FPRR – Final product release responsibility FQA – Full quality assurance FR – Federal Register (US) **FRPs** – Facilitated regulatory pathways FrP – French Pharmacopoeia (Pharmacopée Française, aka PF) FSCA – Field safety corrective action (medical devices sector) **FSIS** – Food Safety and Inspection Service (US) FSN – Field safety notice (medical devices) FTA – Fault tree analysis FTC – Federal Trade Commission (US) FTD – Fast track designation (US) FTE – Full Time Equivalent (employee) FTIM – First-time-in-human FTIR - Fourier Transform infra-red FU – Farmacopea Ufficiale – the Italian Pharmacopoeia FUM – Follow-up measures FVAR – Final Variation Assessment Report FY - Fiscal year GGG **GAIN Act** – Generating Antibiotic Incentives Now Act (US) GATT - General Agreement on Tariffs and Trade GCC (region) – Gulf Cooperation Council (region)

GCC-DR – Gulf Central Committee for Drug Registration

- **GCD** Global clinical development
- GCG Global Cooperation Group (ICH)
- GCP Good clinical practice



**GCPv** – Good Clinical Practice (Veterinary) **GDP** – Good distribution practice **GDPR** – General Data Protection Regulation **GDUFA** – Generic Drug User Fee Amendments (FDA) **GEG** – Geriatrics Expert Group GEP - Good epidemiological practice **GGP** – Good guidance practice **GHTF** – Global Harmonisation Task Force **GIVIMP** – Good in vitro method practices GLC – Gas liquid chromatography **GLP** – Good laboratory practice **GLPMA** – Good Laboratory Practice Monitoring Authority (UK) **GMA** – Global marketing authorisation GMC - General Medical Council (UK) **GMDN** – Global medical device nomenclature (medical devices sector) **GMiA** – Generic Medicines industry Association (Australia) **GMO** – Genetically modified organism **GMP** – Good management practice GMP - Good manufacturing practice - and also: **GNA** – Grounds for non-acceptance **GPAG** – Granularity and Periodicity Advisory Group **GPhP** – Good Pharmacopoeial Practices **GPIA** – Generic Pharmaceutical Industry Association (US) **GPMSP** – Good postmarketing surveillance practice (Japan) GPP - Good paediatric practice - and also: GPP - Good pharmacoepidemiology practice **GPP2** – Good publication practice **GPSP** – Good Post-marketing Study Practice **GpvP** – Good pharmacovigilance practice GQCLP – Good Quality Control Laboratory Practice **GQP** – Good quality practice GRAS - Generally Recognised as Safe (US) **GRB** – Global Regulatory Board GRP - Good regulatory practice - and also: **GRP** – Good review practice (US) **GSL** – General sales list **GSP** – Good statistics practice – and also: **GSP** – Good storage practice **GSPRs** – General Safety and Performance Requirements **GTI** – Genotoxic impurity **GTMP** – Gene therapy medicinal product **GTP** – Gene therapy product **GTWP** – Gene Therapy Working Party GVD - Global value dossier GvHD – Graft versus Host Disease **GVP** – Good pharmacovigilance practice **GxP** – general term for "good practice" quality guidelines and regulations, where "x'' is the symbol for the variable descriptor HHH

**HA** – Health authority **HACCP** – Hazard analysis critical control point (inspection technique) (US) **HAI** – Health Action International HAS - Haute Autorité de santé (French health authority) Hb - Haemoglobin HBD – Harmonised Birth Date HCD - Historical control data HCP - Healthcare professional **HCPWP** – Healthcare Professionals Working Party (EMA) **HCR** – Holder of certificate of registration (South Africa) **HCRW** – Health and Care Research (Wales)

HCT – Haematocrit



HCT/P – Human cells, tissues, and cellular and tissue-based products HDE – Humanitarian device exemption HDI – Human development index **HE** – Hospital exemption HEOR - Health economics and outcomes research HEW - Health, Education and Welfare (US) **HFE** – Human factors engineering HGAC - Human Genetics Advisory Committee **HGPRT** – Hypoxanthine-guanine-phosphoribasyltransferase activity HHMG – Human Harmonisation Maintenance Group HHS – US Department of Health and Human Services HIC - High income countries С **HIMA** – Health Industry Manufacturers Association (US) HL7 - Health Level Seven HLGT – High level group term (in MedDRA) **HLT** – High level term (in MedDRA) **HMA** – Heads of Medicines Agencies (Human and Veterinary) (EU) **HMO** – Health Maintenance Organisation (US) HMPC - Committee on Herbal Medicinal Products (EMA) **HMR** – Human Medicines Regulations **HNSTD** – Highest Non Severely Toxic Dose HoA – Heads of Agencies HPB - Health Protection Board (Canada) **HPLC** – High performance liquid chromatography HPRA – Health Products Regulatory Authority (formerly Irish Medicines Board) HR - Heart rate **HRA** – Health Research Authority (UK) HRB - Health Research Board HREC – Human Research Ethics Committee HRQoL – Health-related quality of life **HRT** – Hormone replacement therapy HSA – Human serum albumin **HSC** – Haematopoietic stem cells HSE - Health and Safety Executive (UK) HST - Highly specialised technologies HTA – Health technology assessment HTS - High-throughput screening HV - Healthy volunteer III I&AC - Imaging and acute care (medical devices sector) **IAM** – Identity and Access Management IAPO - International Alliance of Patients' Organisations IB – Investigator's brochure

- **IBD** International Birth Date
- **IBMS** Institute of Basic Medical Sciences (China)
- IC Informed consent
- ICD Informed consent document and also:
- **ICD** International Classification of Diseases
- ICDRA International Conference of Drug Regulatory Authorities
- ICER Incremental cost-effectiveness ratio
- **ICF** Informed consent form
- **ICH** International Council for Harmonisation (formerly International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use
- ICI Immune checkpoint inhibitor
- **ICMJE** International Committee of Medical Journal Editors
- **ICMRA** International Coalition of Medical Regulatory Authorities
- ICP-MS Inductively coupled plasma mass spectrometry
- ICSR Individual case safety report
- **ICT** Information and communications technology
- **ICTRP** International Clinical Trials Registry Platform (WHO)



IC<sub>X</sub> – Inhibition concentration at X% **IDE** – Investigational Device Exemption IDMP – Identification of medicinal products – and also: **IDMP** – Infectious diseases management program (US) IDR – Idiosyncratic drug reaction IDRAC – International Drug Registration Assisted by Computer IEC - Independent ethics committee **IFAH** – International Federation for Animal Health **IFPMA** – International Federation of Pharmaceutical Manufacturers and Associations IFU – Instructions for use **IGDG** – Informal Generic drug Discussion Group **IGDRP** – International Generic Drug Regulators Pilot **IGPA** – International Generic Pharmaceutical Alliance **IGZ** – the Netherlands Healthcare Inspectorate IIG – Inactive ingredient guide (US FDA) IIS – Investigator initiated study IM – Intramuscular – and also: IM – Issue management IM(ER)R – Ionising radiation (medical exposure) regulations IMA – Lyfjastofnun/Icelandic Medicines Agency (Iceland) **IMB** – Irish Medicines Board [name changed in July 2014 to **HPRA** – Health Products Regulatory Authority] **IMCA** – Lyfjastofnun/Icelandic Medicines Control Agency (Iceland) **IMD** – Implantable medical device **IMDA** – Irish Medical Device Association **IMDRF** – International Medical Device Regulators Forum IME – Important medical event **IMI** – Innovative Medicines Initiative **IMM** – Irreversible morbidity or mortality **IMP** – Investigational medicinal product **ImPACT** – Imaging performance assessment of CT scanner **IMPD** – Investigational medicinal product dossier **IMRDF** – International Medical Device Regulatory Forum **IMS** – Information management strategy **INADA** – Investigational new animal drug application **IND** – Investigational new drug (US) **INDA** – Investigational new drug application (US) **INDC** – Investigational New Drug Committee **INFARMED** – Instituto Nacional da Farmacia e do Medicamento (Portugal's regulatory agency) **INN** – International nonproprietary name **IO** – Immune-oncology IP – Intellectual property – and also: IP – Interested Parties – and also: **IP** – Intraperitoneal **IPAC** – International Pharmaceutical Aerosol Consortium IPC – International Pharmaceuticals Council **IPCs** – In-process controls IPD – Individual Patient Data **IPEC** – International Pharmaceutical Excipients Council **IPI** – International Pricing Index **iPiE** – Intelligence-led assessment of Pharmaceuticals in the Environment **IPO** – Intellectual Property Office IPR – Intellectual property rights **IPRF** – International Pharmaceutical Regulators Forum iPSP – initial Paediatric Study Plan **IPU** – Irish Pharmaceutical Union **IQM** – Integrated quality management IR – Infra-red – and also: IR (tablets) – Immediate release **IRAS** – Integrated Research Application System **IRB** – Institutional review board (aka Independent Ethics Committee (**IEC**) or Ethical Review Board (**ERB**)) **IRC** – Institutes Review Committee



**IRN** – Incident Review Network IRP - Independent review panel **IRR** – Ionising radiation regulation **IRT** – Interactive response technology – and also: IRT – Interdisciplinary Review Team (US) IS – Information science/systems – and also: IS - Internal standard **ISA** – Integrated scientific advice **ISCT** – In silico clinical trial **ISE** – Integrated summary of efficacy **ISI** – Integrated summary of immunogenicity ISS – Integrated summary of safety **ISO** – International Standards Organisation **ISRB** – Integrated summary of risk benefit **ISS** – Integrated summary of safety **IT** – Information technology ITF - Innovation Task Force (EMA) ITT - Intent-to-treat IU – International unit **IUPAC** – International Union of Pure and Applied Chemistry **IV** – Intravenous IVD - in vitro (medical) device; and also:

**IRDiRC** – International Rare Diseases Research Consortium

IVD - III VILIO (Illeuical) device

**IVD** – *in vitro* diagnostics

**IVDR** – In Vitro Diagnostic Regulation

**IRD** – International registration document

**IVIVC** – *in vitro in vivo* correlation **IVMP** – Immunological veterinary medicinal product

**IVRS** – Interactive voice response system

**IWG** – Implementation working group

**IWP** – Immunologicals Working Party (EMA)

JJJ

JAN – Japanese Approved Name

JAZMP – Javna agencija Republike Slovenije za zdravila in medicinske pripomočke (Slovenia's regulatory agency) JFDA – Jordan Food & Drug Administration

JIACRA – Joint Interagency Antimicrobial Consumption and Resistance Analysis

JNDA – Japanese New Drug Application

JP – Japanese Pharmacopoeia

JPMA – Japan Pharmaceutical Manufacturers Association

**J-RMP** – Japanese risk management plan (template)

## KKK

KAS – Known active substance

**KFDA** – Korean Food and Drug Administration

**KIT** – Key intelligence topic

KM – Knowledge management

KOL – Key opinion leader

**KOM –** Kick-off meeting

### LLL

LABST – Laboratory animal batch safety testing

LAT – Light authoring tool (EU)

**LCM** – Lifecycle management

**LD<sub>50</sub>** – Lethal dose required to kill 50% of the study population

LDH – Lactate dehydrogenase

LEC – Local ethics committee

LED - Least Effect Dose

**LEEM –** Les Entreprises du Médicament (French Pharmaceutical Industry Association)

LFT - Liver function test

LICT – Low-intervention clinical trial



LM – Limited markets (veterinary) LMA - Limited marketing authorisation LMICs - Low and middle income countries LoA - Letter of access (China) LOD – Loss on drying **LOI** – Letter of intent (US) LoNR - Letter of non-repudiation agreement (FDA) LoOI – List of Outstanding Issues LoQ – List of Questions LPLV – Last patient last visit LSIF – Life Sciences Innovation Forum LT (stability) – Long term LTT - Lines to take [document usually not for publication] (EMA) LVP - Large volume parenterals МММ M&S - Modelling and simulation MA – Marketing authorisation MAA – Marketing authorisation application (EU) MABEL – Minimal anticipated biological effect level MAD – Multiple ascending dose (study), and also: MAD - Mutual acceptance of data (OECD Council Decision) **MAFF** – Ministry of Agriculture, Forestry and Fisheries (Japan) MAH - Marketing authorisation holder **MAID** – Manufacturers, authorised representatives, importers and distributors MALAM - Medical Lobby for Appropriate Marketing Mane – Morning MANSEV - Marketing Authorisation by Network Submission and Evaluation **MAPPs** – Medicines adaptive pathways to patients MAUDE – Manufacturer and User Facility Device Experience (US) MAWP - Multi-Annual Work Plan (HMA) MaxSPRT – Maximised sequential probability ratio test MB – Management Board MCC – Medicines Control Council (South Africa) MCDA – Multi-criteria decision analysis MCH - Mean cell haemoglobin concentration MCPC – Major contribution to patient care MCV – Mean cell volume **MD** – Medical device MDA – Medical device alert MDCG – Medical Device Coordination Group MDD - Medical Device Directive - and also: MDD – Medical Devices Directorate MDDS – Medical device data systems **MDEG** – Medical Devices Expert Group MDEG-BC - Medical Devices Expert Group on Borderline and Classification MDI – Metered dose inhaler MDLO – Medical Device Liaison Officer MDR – Medical Device Regulation – and also: **MDR** – Medical device reporting – and also: **MDR** – Multi-drug resistant MDSAP - Medical Device Single Audit Program (US, Canada) **MDV** – Medical device vigilance MEB - Medicines Evaluation Board (the Netherlands) - also known as Dutch College MedDevs - Guidances outlining the requirements of the Medical Device Directive MedDRA - Medical Dictionary for Regulatory Activities **MEDEV** – Medicine Evaluation Committee (EU) **MEDSAFE** – New Zealand Medicines and Medical Devices Safety Authority

LIF – Läkemedelsindustriföreningen (Swedish Pharmaceutical Industry Association)

**LLL** – Lifelong learning

**MENA** – Middle East and North Africa

**MERS** – Multi-agency electronic regulatory system



MFDS – Ministry of Food and Drug Safety (Korea) **MgSzH** – Mezogazdasagi Szakigazgatasi Hivatal Dictorate of Veterinary Medicinal Products (Hungary) MHRA – Medicines and Healthcare products Regulatory Agency **MHW** – Ministry of Health and Welfare (Japan) MIA - Manufacturing and Importation Authorisation MIA(IMP) - Manufacturer's Authorisations for IMPs MIDD - Model-informed drug development (US) MIMS - Monthly Index of Medical Specialities (UK) **MINE –** Medicines Information Network for Europe MIR – Manufacturer incident report MISG – Ministerial industry strategy group ML – Machine learning – and also: ML – Manufacturer's licence MLD - Minimal lethal dose MLM – Medical literature monitoring MMA - Malta Medicines Authority - and also: **MMA** – Mobile medical app **MNAT –** Multinational Assessment Team MO - Major Objection MoA – Mechanism of action – and also: **MOA** – Ministry of Agriculture MoCA – Mechanism of Coordinated Access MOD 1 – Module One (laboratory facility) (US) **MOD 2** – Module Two (laboratory facility) (US) MORE - Manufacture's Online Reporting Environment (MHRA) (medical devices sector) mOS - median Overall Survival **MOU** – Memorandum of Understanding MPA - Medical Products Agency - Sweden MPD – Medicinal Products Directive MPID – Medicinal product identifier MQAS – Model Quality Assurance System MQSA – Mammography Quality Standards Act of 1992 (US) **MR** – Mutual Recognition MRA – Mutual recognition agreement MRAs - Medicines regulatory authorities - and also: MRC – Medical Research Council MRD – Multiple rising dose **MRFG** – Mutual Recognition Facilitation Group (EMA) **MRH** – Medicines regulatory harmonisation MRI (scan) – Magnetic resonance imaging (scan) – and also: **MRI** – Mutual recognition information MRL – Maximum residue limit **MRP** – Mutual recognition procedure (EU) MRSD - Maximum recommended safe dose MRTP – Modified risk tobacco product MRU – Medicines Regulatory Unit (Health Division Malta) MS – Mass spectrometry – and also: MS - Member state/s (EU) **MSWG** – Modelling and Simulation Working Group MTD – Maximum tolerated dose **MTS** – Medicines testing scheme (MHRA) **MUMS** – Minor use and minor species (veterinary) NNN **N&ET** – New and emerging technologies (see also: NET WG) N-11 – Next 11 (group of countries comprising Bangladesh, Egypt, Indonesia, Iran, Korea, Mexico, Nigeria, Pakistan, Philippines, Turkey and Vietnam) NAD - No abnormality detected **NADA** – New animal drug application (US) **NAFDAC** – National Agency for Food and Drug Administration and Control (Nigeria) **NAFTA** – North American Free Trade Association (US) NAI - No action indicated



NAO – National Audit Office (UK) **NAP** – Nationally authorised product NAS – New active substance **NB** – Notified body (EU) **NBE** – New biological entity **NBIC** – Nanotechnology, biotechnology, information science and cognitive science **NBO** – Notified body opinion **NBOG** – Notified Body Operations Group (EU) NC3Rs - National Centre for the Replacement, Refinement and Reduction of Animals in Research (UK). NCA – National competent authority NCAS – New chemical active substance NCD – Non-communicable diseases NCE – New chemical entity NCI - National Cancer Institute (US) - and also: NCI - National Coordinating Investigator NCO - Non clinical overview NCS – Non clinical summarv **NCTR** – National Center for Toxicological Research (US) **NDA** – New drug application (US) NDAC – New Drug Advisory Committee (India) NDMA – Non-Prescription Drug Manufacturers Association (US) **NDS** – New drug submission (Canada) NED - Non effect dose **NeeS –** Non eCTD electronic submission **NEFARMA** – Netherlands Pharmaceutical Industries Association **NET WG –** New & Emerging Technologies Working Group **NF** – National Formulary NfG - Note for Guidance (EU) NGS – Next generation sequencing NHL - non-Hodgkin's lymphoma **NHP** – Non-human primate NHS – National Health Service **NHV** – Normal healthy volunteer NIAID – National Institute of Allergy and Infectious Diseases NIBSC – National Institute for Biological Standards Control (UK) NICE – National Institute for Health and Care Excellence (formerly 'Clinical' Excellence) NICHD – National Institute of Child Health and Human Development (US) NIH – National Institutes of Health (US) NIHR - National Institute for Health Research (UK) **NIMP** – Non-investigational medicinal product (but see AMP – Auxiliary medicinal product) NIR – near infrared (spectroscopy) – and also: NIR – Non-interventional research NIS - Non-interventional study NK cells – Natural killer cells NLEA – Nutrition Labelling and Education Act of 1990 (US) NLN – Nordic Council on Medicines NMA – National Medicines Agency (Romania) NMCA – Norwegian Medicines Control Agency (aka SLK) **NME** – New molecular entity **NMFS** – National Marine Fisheries Service (US) NMPA - National Medical Products Administration (China) (国家药品监督管理局) (formerly CFDA) **NMR** – Nuclear magnetic resonance **NMRAs –** National Medicines Regulatory Authorities **NMVO** – National Medicines Verification Organisation NMVRVI – Nacionalinis Maistro Ir Veterinarijos Rizikos Vertinimo Institutas (National Food and Veterinary Risk Assessment Institute) (Lithuania) NOAEL – No observable adverse effect level **NOAH** – National Office of Animal Health (UK) **NOAL** – No observed adverse effect level **NOC** – Notice of Compliance (Canada) **NOC/c** – Notice of Compliance with Conditions (Canada)



Nocte – Night NOEL - No observable effect level **NoMA** – Norwegian Medicines Agency NPCB - National Pharmaceutical Control Bureau (Malaysia) **NPP** – Named patient product **NPRM** – Notice of Proposed Rulemaking NPT - Near-patient test **NRA** – National regulatory authority NRG - (invented) Name Review Group **NSA** – National Security Agency (US) **NSAID** – Nonsteroidal anti-inflammatory drug NSB - National Standards Body - and also: NSB - Non-similar biologic NSCLC – Non-small cell lung cancer **NSF** – No biologically significant finding (may be used in older reports) **NSN** – New substances notification (Canada) **NSR** – Non-significant risk **NSVA** – National Sanitary Veterinary Agency (Romania) NtA – Notice to applicants (EC) **NTD** – Neglected tropical disease NTE – No toxic effect level **NTI** – Narrow therapeutic index NUI - Non-urgent information (aka "Infofax") (EU) **NWIP** – New work item proposal (EU) 000 O/E – Observed versus expected [analysis] oab - On anhydrous basis oasfb - On anhydrous solvent free basis **OBL** – Own brand labelling **OBP** – On-boarding partner **OC** – Office of the Commissioner (US) **OCA** – Office of Consumer Affairs (US) **OCABR –** Official control authority batch release **OCI** – Office of Criminal Investigation (US) **OCP** – Office of Combination Products (US FDA) od – once a day [Latin: omne in die] – and also: **OD** – Orphan drug **ODA** – Orphan Drugs Act (US) **ODC** – Optimal diagnostic concentration (used on allergy products) **ODD** – Orphan drug designation **OE** – Oral explanation **OECD** – Organisation for Economic Co-operation and Development **OEI** – Official establishment inventory (US) **OEM** – Original equipment manufacturer **OES** – Original equipment supplier **OGTR** – Office of the Gene Technology Regulator (Australia) **OGYI/NIP** – National Institute of Pharmacy (Hungary) OH - Oral Hearing **OHDSI –** Observational Health Data Science and Informatics OIA – Official action indicated **OIE** – World Organisation for Animal Health **OJ/OJEC** – Official Journal of the European Communities OLE (study) - Open label extension (study) **OMAR –** Orphan Maintenance Assessment Report **OMCL** – Official Medicines Control Laboratories (part of **EDQM**) **OMP** – Orphan medicinal product **OMS** – Organisations data management service **OOPD** – Office of Orphan Products Development (US FDA) **OOS** – Out of specification **OPA** – Office of Public Affairs (US) **OPD** – Original pack dispensing

OPDP – Office of Prescription Drug Promotion (FDA's CDER)
OPE – Office of Planning and Evaluation (US)
ORA – Office of Regulatory Affairs (US FDA)
ORGAM – Organisational Matters
ORR – Overall response rate
OS – Overall survival
OTAT – Office of Tissues and Advanced Therapies (US CBER)
OTC – Over-the-counter

# PPP

P - Pharmacy only (ie, medicinal product dispensed by a pharmacist) P to GSL – Pharmacy to General Sales List P&L – Packaging and labelling P&R – Pricing and reimbursement **PA** – Product authorisation – and also: **PA** – Protocol assistance **PAB** – Pharmaceutical Affairs Bureau (Japan) **PAC-ATLS** – Post Approval Change – Analytical Testing Laboratory Site (US) **PACMP** – Post-approval change management protocol **PAD** – Pharmacologically active dose PaedPAR – Paediatric Public Assessment Report **PAES** – Post authorisation efficacy study PAGB – Proprietary Association of Great Britain **PAI** – Pre-approval inspection **PAL** – Pharmaceutical Affairs Law (Japan) **PAM** – Patient activation measure (UK) **PAM(s)** – Post Authorisation Measure(s) **PAO** – Period after opening (cosmetic products) **PAR –** Preliminary assessment report PAR – Public Assessment report **PARENT** – Patient Registries Initiative (EU) PAS – Patient Affairs Staff, and also: **PAS** – Public Affairs Specialist (US) **PASS** – Post authorisation safety study **PAT –** Priority Action Team (EFPIA) **PAT –** Process analytical technology – and also: PBAC - Pharmaceutical Benefits Advisory Committee (Australia) PBI - Protein-bound iodine **PBPK** – Physiologically based pharmacokinetic modelling PBRER – Periodic benefit-risk evaluation report PBS – Pharmaceutical Benefit Scheme (Australia) **PBT** – Persistent, bioaccumulative and toxic (biocidal products) PC – Packaged commodities (India) PCA – Perception, cognition, action PCG - Product Coordination Group (EU) PCID - Package indentifier pCODR - pan-Canadian Oncology Drug Review PCORI - Patient-Centered Outcomes Research Institute PCPA – Pan-Canadian Pricing Alliance PCT – Primary care trust (UK) PCWP – Patients' and Consumers' Working Party PD - Parallel distribution, and also: **PD** – Pharmacodynamics PdAR – Paediatric Assessment Report **PDCO** – Paediatric Committee (EMA) **PDE** – Permitted daily exposure **PDG** – Pharmacopoeial discussion group **PDMA** – Prescription Drug Marketing Act (US) **PDP** – Product development protocols (for medical devices) (US) **PDPs** – Product development partnerships PDR – Physician's desk reference PDS – Public disclosure synopsis/system



**PDUFA** – Prescription Drug User Fee Act (US) **PDX** – Patient-derived xenograft PE – Pharmacoeconomics **PEAG** – Pharmacovigilance Expert Advisory Group (MHRA) PEC - Patient Engagement Collaborative, and also: PEC - Predicted environmental concentration **PECA** – Protocol to the Europe Agreement on Conformity Assessment and Acceptance of industrial products **PED** – Patient experience data **PEFR** – Peak expiratory flow rate **PEFRAS** – Pan European Federation of Regulatory Affairs PEI – Paul-Ehrlich-Institut (Federal Institute for Vaccines and Biomedicines (one of the two German regulatory agencies) PEM (study) - Prescription-event monitoring (study) PER – Pharmaceutical evaluation report **PeRC –** Paediatric Review Committee (US) **PERF** – Pan European Regulatory Forum **PET/CT** – Positron emission tomography and computerised tomography pfa (or b) - pure free acid (or base) **PFDD** – Patient-focused drug development **PFI** – Pediatric Formulation Initiative (US) **PFMD** – Patient Focused Medicine Development **PFS** – Progression-free survival PGD - Patient group directions (written instructions) **PGENI** – Pharmacogenetics for Every Nation Initiative **PGI** – Potentially genotoxic impurity **PgWP** – Pharmacogenomics Working Party **PGx** – Pharmacogenomics Ph Eur – European Pharmacopoeia **PHA** – Preliminary hazard analysis **PHARE** – Poland and Hungary; aid of the Restructure of the Economy; Now the Phare programme is one of the three pre-accession instruments financed by the European Communities to assist the applicant countries of central Europe in their preparations for joining the EU **PHARMO** – Institute for Drug Outcomes Research (the Netherlands) PHC – Personalised healthcare PhI – Pharmacological intelligence **PhPID** – Pharmaceutical product identifiers (EU) PhRMA – Pharmaceutical Research and Manufacturers of America PHS - Public Health Service (US) PhV – pharmacovigilance (aka PV) PhV WSP WP - Pharmacovigilance Procedures Work Sharing Working Party **PhVIWG** – Pharmacovigilance Inspectors Working Group PhVWP - Pharmacovigilance Working Party (EMA) **PhVWP-V** - Pharmacovigilance Working Party - Veterinary PI - Package insert - and also: PI - Parallel import - and also: PI – Prescribing information – and also: PI – Principal investigator – and also: PI - Production information - and also: PI - Protease inhibitor **PIA –** Pharmaceutical Industries Association **PIC** – Pharmaceutical Inspection Convention (EU) PIC/S - Pharmaceutical Inspection Convention and Pharmaceutical Inspection Co-operation Scheme **PICO** – Population, intervention, comparator, outcome(s) **PICS** – Pharmaceutical inspection cooperation scheme (EU) PIE - Pharmaceuticals in the environment PIIGS - Portugal, Ireland, Italy, Greece and Spain **PIL** – Patient information leaflet PIL-ICF - Patient information leaflet-informed consent form **PIM** – Product information management (EMA) – and also: **PIM** – Promising innovative medicine **PIN** – Patient identification number **PIP** – Paediatric investigation plan – and also:



**PIP** – Poly Implant Prothèse (breast implant) PIQ – Product Information Quality Review Group **PK** – Pharmacokinetics **pKa** – acid dissociation constant **PKWP** – Pharmacokinetic Working Party PL - Package leaflet - and also: PL - Product license (US) **PLA** – Product license application (for biologics) (US) **PLLR** – Pregnancy and Lactation Labeling Rule (US) **PLPI** – Parallel import licence [product licence parallel import] PLR – Physician Labeling Rule (US) PLR - Product license renewal (US) - and also: **PLT** – Platelet count PMA - Pre-market approval (application for medical devices) (US) PMC - Postmarketing commitments (US) **PMCF** – Post-market clinical follow-up (studies) PMDA – Japan's regulatory agency – the Pharmaceutical and Medical Devices Agency (within the Ministry of Health, Labor and Welfare - MHLW) **PMDI** – Pressurised metered dose inhaler **PMDL** – Pharmaceutical and Medical Device Law (Japan) **PMF** – Plant master file (US and Canada) PMI – Pharmacological, metabolic and immunological **PMN** – Pre-market notification **PMOA** – Primary mode of action PMPF - Post Market Performance Follow-up PMPRB - Patented Medicines Prices Review Board (Canada) **PMR** – Postmarketing requirements (US) PMS – Postmarket(ing) surveillance – and also: **PMS** – Product data management service/product management services **PMS study** – Post-marketing safety study **PMTA** – premarket tobacco application **PNC** – Pre-notification consultation (Canada) **PNEC –** Predicted no-effect concentration **po** – by mouth/orally [Latin: per os] **POC** – Proof of concept **POCA** – Phonetic and Orthographic Computer Analysis **POM –** Prescription-only medicine **POM to P** – Prescription-only medicine to pharmacy **PONV** – Post-operative nausea and vomiting **POP db** – Planned and Ongoing Projects database (an EUnetHTA database) **popPK** – Population PK PPA – Parallel production authorisation **PPD** – Protected personal data PPI - Patient and Public Involvement (UK) - and also: **PPI** – Patient package insert (US) **PPP** – Pregnancy Prevention Programme **PPP** – Public-private partnership **PPRS** – Pharmaceutical Price Regulation Scheme **PPSR** – Proposed Paediatric Study Request (US) **PQP** – Prequalification of Medicines Programme (WHO) PQR - Product quality review PQS – Pharmaceutical quality system PR – Pulse rate PRAC – Pharmacovigilance Risk Assessment Committee (EMA) PRAG – PSUR Repository Advisory Group PrAR – Preliminary Assessment Report **PRD-PRV** – Pediatric rare disease priority review voucher (US) **PREA** – Paediatric Research Equity Act (US) **PREG** – Pandemic Response Expert Group **PRIME** – Priority medicines scheme P-RMS – PSUR reference member state (also see PSUR)



**prn** – as needed (Latin: pro re nata) **PRO** – Patient reported outcome **PRO-AE** – Patient-reported outcomes in adverse event reporting **PROM –** Patient-relevant outcome measure **PROSPER** – Patient-reported outcomes safety event reporting **PROTECT –** Pharmacoepidemiological Research on Outcomes of Therapeutics **PRR** – Proportional reporting ratio **PRRC** – Person responsible for regulatory compliance PRS - PIM review system (EU) - also see PIM **PRSPH** – Potential serious risk to public health **PSA –** Parallel scientific advice **PSBGL(s)** – Product-specific bioequivalence guideline(s) **PSD** – Particle size distribution **PSM –** Pre-submission meeting **PSMF** – Pharmacovigilance system master file PSP - Paediatric study plan - and also: **PSP** – Patient Support Programme **PSR** – Periodic summary report – and also: **PSR –** Product safety reference **PSRPH** – Potential Serious Risk to Public Health **PSS** – Personal social services **PSUR** – Periodic safety update report PSUSA – PSUR single assessment PT – Preferred term – and also: PT time - Prothrombin time PtC - Points to consider. PTD – Protection of technical documentation **PTE –** Patent term extension **PuAR –** Public assessment report PUL module – Performance of the Upper Limb module **PUMA** – Paediatric-use marketing authorisation **PV** – Pharmacovigilance **PVAR** – Preliminary Variation Assessment Report **PXRD** – Powder xray diffraction 000 (Q)SAR – Quantitative structure activity relationships **QA** – Quality assurance QALY – Quality-adjusted life year **QbD** – Quality by design QC – Quality control **qd** – once a day [Latin: quaque die] **qds/qid** – four times a day [Latin: quater die sumendum/quater in die] **QIDP** – Qualified infectious disease product (US) QMS – Quality management system QOF – Quality and Outcomes Framework (NICE, UK) QOL – Quality of life QoS – Quality overall summary QP – Qualified person **QPPV** – Qualified person for pharmacovigilance **QR(C)** – Quick response (code) (EU) **QRD** – Quality review of documents [template] **QRM** – Quality risk management QS – Quality system **QSE** – Quality, safety and efficacy **QSIT -** Quality System Inspection Technique (US FDA) QTPP - Quality target product profile **OUAMED –** Quality Medicines for All **QWP** – Quality Working Party (EMA) RRR

R&D - Research & development



R4BP – Register for Biocidal Products RA – Rapid alert – and also: **RA** – Regulatory affairs RA/NUI System - Rapid Alert/Non-Urgent Information System RADAR – Risk assessment of drugs analysis and response **RAMA** – Remote access for marketing authorisations (MHRA) RAPS – Regulatory Affairs Professionals Society (US) RAS – Rapid alert system **RAT** – Regenerative advanced therapy RBC - Red blood cell count **RBI** – Risk-based inspection **RBM** – Risk-based monitoring **RCB** – Registered certification body (Japan) **RCFID** – Registration Certificate for Import of Drug RCH - Remove clinical hold RCP – Royal College of Physicians (UK) RCT – Randomised controlled trial RCTP - Regenerative and cellular therapy product **RDE** – Remote data entry RDI – Research, development and innovation **RDP** – Regulatory data protection **RDS** – Repeat dose study **RDT** – Rising-dose tolerance **REA –** Relative effectiveness assessment **REACH** – Registration, evaluation, authorisation and restriction of chemicals REC – Research Ethics Committee RefMP(s) - Reference Medicinal Product(s), see also RMP(s) **REMS** – Risk evaluation and mitigation strategy (US) RfD – Reference dose (veterinary) **RFDD** – Regional Food and Drug Director (US) **RFI** – Request for information RfMs – Requests for modifications RH – Relative humidity RHSC – Regulatory Harmonisation Steering Committee **RI** – Regulatory intelligence **RIM** – Regulatory information management **RING** – Regulatory Intelligence Network Group (EU) rINN – Recommended international non-proprietary name **RiskMAP** – Risk minimisation action plan **RLD** – Reference listed drug (US) RMM – Risk minimisation materials – and also: **RMM** – Risk minimisation measure **RMP** – Reference medicinal product – and also: **RMP** – Risk management plan RMR – Reaction monitoring report – and also: RMR – Risk management report RMS - Reference member state (Europe) - and also: **RMS** – Referentials data management service rMS - Reporting member state (Europe) **ROG** – Regulatory Optimisation Group **RoHS** – Restriction of hazardous substances (Directive) **ROI** – Residues on ignition – and also: ROI – Return on investment RONAFA - Reduction of need for antimicrobials in food-producing animals RoW - Rest of (the) World **RP** – Responsible person **RPA** – Robotic process automation **RPI** – Research Product Identifier (formerly called 'Unique Product Identifier, UPI) **RPS** – Regulated product submission **RPSGB** – Royal Pharmaceutical Society of Great Britain **RQA** – Research quality assurance RR - Relative risk - and also:



RR – Respiratory rate – and also:
RR – Risk ratio
RRI – Regional regulatory initiatives
RRR – Relative risk reduction
RSA – Risk share agreement
RSI – Reference safety information – and also:
RSI – Request for supplementary information (EU)
RTF – Refusal-to-file (US)
RTI – Respiratory tract infection
RTQ – Response to questions
RTT – Real time release testing
RTT – Right to Try
RU–MRP – Repeat use mutual recognition procedure
RWD – Real world data

- **RWE** Real word evidence
- **Rx** Prescription

## SSS

S+T – Sampling and testing **SA** – Scientific advice **SAARC** – South Asia Association for Regional Cooperation SaaS – Software as a service SABS – Safety alert broadcast system **SAD** – Single ascending dose (study) SADR – Serious adverse drug reaction SAE – Serious adverse event SAG – Scientific Advisory Group SAL - Sterility assurance level SaMD – Software as a Medical Device **SAMM** – Safety assessment of marketed medicines (US) SANDS – Supplemental abbreviated new drug submission (Canada) SAP – Scientific advice procedure – and also: SAP – Statistical analysis plan SAR – Safety assessment report – and also: SAR – Serious adverse reaction **SAT** – Special Action Team (EFPIA) SAWP – Scientific Advice Working Party SBA/SBOA – Summary basis of approval (US) **SBP** – Similar biotherapeutic product (WHO) sc – subcutaneou (aka sq) SCB – Scientific Coordination Board SCCS – Self-controlled case series design **SCF** – Scientific Committee for Food (UK) **SCOTT** – Ethics and Standing Committee on Therapeutic Trials (Australia) SCT - Stem cell transplant **sCTMP** – somatic Cell Therapy Medicinal Product SD - Standard deviation **SLDC** – Software development lifecycle SDR – Statistic of disproportionate reporting SDRG – Study data reviewer's guide **SDTM** – Study Data Tabulation Model (US) SE – Standard error – and also: SE – Substantially equivalent/substantial equivalence SEAR - Safety, Efficacy and Adverse Reactions (sub-committee of CSM) **SEB** – Subsequent entry biologic SEED Consortium - Shaping European Early Dialogues Consortium SEND - Standard for exchange of nonclinical data SFDA - Formerly China's State Food and Drug Administration (now CFDA) and also: SFDA – Safety Features Delegated Act – and also: **SFDA** – Saudi Food & Drug Authority SFFC medicines – Spurious/falsely-labelled/falsified/counterfeit medicines (US)



SGML – Standard general mark-up language **SGOT** – Serum glutamic oxalo-acetic acid transaminase (SGOT = AST) **SGPT** – Serum glutamic pyruvic transaminase (SGPT = ALT) SHBG - Sex-hormone-binding globulin SI - Statutory instrument SKU - stock-keeping unit **SLA –** Service level agreement SLK/NMCA - Statens legemiddelverk/Norwegian Medicines Control Agency SmAR – Summary Assessment Report SMC – Scottish Medicines Consortium **SMDA** – Safe Medical Devices Act (US) SME - Significant medical event - and also: SMEs - Small and medium-sized enterprises SMF - Site master file **SMO** – Site management organisation SmPAR – Summary Pharmacovigilance Assessment Report (EU) SmPC - Summary of product characteristics (aka SPC in veterinary sector) SMQ - Standardised MedDRA query SMS - Substance data management service **SNDA** – supplemental new drug application (US) **SNDS** – Supplemental new drug submission (Canada) **SNIF** – Summary Notification Information Format **SO** – Scientific opinion SOC - Standard of care - and also: **SOC** – System organ class SOCMA – Society of Chemical Manufacturers and Affiliates SOCRA – Society of Clinical Research Associates (US-based) **SOP** – standard operating procedure **SOUP** – Software of unknown pedigree SPA – Special protocol assessment **SPC** – Summary of product characteristics (typically for veterinary sector) – and also: **SPC** – Supplementary protection certificate (EU) **SPECT** – Single photon emission computed tomography **SPIN –** Special interest network SPL – Structured product labelling (US) SPOR data - Substance, product, organisation and referential data **SPS** – Summary of Pharmacovigilance Systems **sq** – subcutaneous (aka **sc**) **SQP** – Suitably gualified person **SR** – Significant risk SRAs – Stringent regulatory authorities SRM – Specified risk materials c - and also: SRN - Stroke Research Network (part of NIHR, UK) SSC – Scientific Steering Committee SSCP – Summary of safety and clinical performance SSFFC – Substandard, spurious, falsely labelled, falsified and counterfeit (medical products) SSRI – Selective serotonin reuptake inhibitor SSU - Study start up STAMP – Safe and timely access to medicines for patients **stat** – immediately [Latin: statim] STD - Severely toxic dose STED – Summary technical documentation [for Demonstrating Conformity to the Essential Principles of Safety and Performance of Medical Devices Safety and Performance of Medical Devices] **StEM** – Stakeholder engagement meeting (MHRA) STF - Study tagging files STR – Stirred tank bioreactors **STRPC** – Scientific, Technical and Regulatory Policy Committee (EFPIA) SUD – Single use device – and also: SUD – Sudden unexpected death **SUE –** Serious undesirable effect SUKL – State Institute for Drug Control (Czech Republic and Slovakia)



**SUPAC** – Scale-up and post-approval changes **SUPAC-IR** – Scale up and post approval changes – immediate release **SUPAC-MR** – Scale up and post approval changes – modified release SUSAR – Suspected unexpected serious adverse reaction SWOT (analysis) - Strengths, weaknesses, opportunities, threats **SWP** – Safety Working Party (CHMP) Sx - Symptoms TTT  $t_{1/2}$  -Terminal half-life of elimination TA - Targeted assessment - and also: TA – Therapeutic area **TABST** – Target animal batch safety testing TAG – Technical Advisory Group (UK's NICE) – and also: TAG – Therapeutic Advisory Group TAS (studies) - Target animal safety (studies) TATFAR - TransAtlantic Task Force on Antimicrobial Resistance TBC - The Biomarker Consortium **TBG** – Thyroid binding globulin TCA - Tricyclic antidepressant TCM – Traditional Chinese medicine TCP – Target candidate profile TCT – Toxicity, Clinical Trials and Therapeutic Efficacy Subcommittee of the CSM (UK) **TDD** – Transdermal drug delivery **TD-PRV** – Tropical disease priority review voucher (US) **TDR** – Totally drug-resistant tds/tid - three times a day [Latin: ter die sumendum/ter in die] TE – Therapeutic equivalence **TEP** – Tissue engineered product **TESS** – Tamper evident security seal TFEU - Treaty on the Functioning of the European Union **TFM** – Tentative final monograph (US) TGA – Therapeutic Goods Administration (Australia's regulatory agency) – and also: **TGA** – Thermogravimetric analysis THMP – Traditional herbal medicinal product THMPD - Traditional Herbal Medicinal Products Directive THMRS - Traditional Herbal Medicines Registration Scheme **THR** – Traditional herbal registration TIGes - Telematic Implementation Group-electronic submissions **TIND** – Treatment IND (see **IND**) TK – Thymidine kinase – and also: **TK** – Toxicokinetics **TLC** – Thin layer chromatography TLV – Threshold limit value TMF - Trial Master File TOC – Table of contents TOD - Table of decisions **TOM** – Target operating model **TOPRA** – The Organisation for Professionals in Regulatory Affairs **TOPS** – The Over-volunteering Prevention System (database) **TPP** – Target product profile **TRF** – Tamper-resistant formulation TRIPS - Trade Related Aspects of Intellectual Property Rights **TRK** – Tropomyosin receptor kinase TRL – Total residue level (veterinary) **TSA –** Therapeutic Substances Act **TSE** – Transmittable spongiform encephalopathy **TTC** – Threshold of toxicological concern **TUBITAK** – Scientific and Technological Research Council of Turkey



**UAT** – User acceptance testing UCN - Unique carton number **UDI** – Unique device identification **UI** – Unique Identifier (according to the FMD) ULTRA - Unlocking Lifesaving Treatments for Rare-Diseases Act (US) UMBRA - Unified Methodologies for Benefit-Risk Assessment **UMP** – Beijing Union Medical and Pharmaceutical General Corp (the innovative arm of the Chinese Academy of Medical Sciences) **UOUP** – User Interface of Unknown Provenance **UPS-NF** – United States Pharmacopeia and National Formulary **USAN –** United States Approved Name **USC** – United States Code **USDA** – United States Department of Agriculture USKVBL – Ustav pro Statni Kontrolu Veterinarnich Biopreparatu a Leciv (Institute for State Control of Veterinary Biologicals and Medicines) (Czech Republic) - and also: USKVBL – Ústav štátnej kontroly veterinárnych biopreparátov a liečiv (Department of State Control of Veterinary Biologicals and Medicaments) (Slovenia) **USP** – United States Pharmacopoeia **USP-DI** – United States Pharmacopeia-Drug Information **USPI –** United States Product Insert **USP-NF** – United States Pharmacopeia-National Formulary **USR** – Urgent safety restriction **UTI** – Urinary tract infection **UUP** – Urgent union procedure (European Commission) VVV **VAERS** – Vaccine adverse event reporting system (US) VAESCO – Vaccine adverse event surveillance & communication VAF - Virus antibody free VAI – Voluntary action indicated VAMF – Vaccine antigen master file **VAR** – Variation assessment report VarWP – Working Party on Variation Regulation, also: Variation Working Party VBA – Value-based assessment VBP - Value-based pricing VCS – Viral challenge study **VDD** – Veterinary Drugs Directorate (Canada) VeDDRA – Veterinary Dictionary for Drug Related Affairs VF – Ventricular failure VHP – Voluntary harmonisation procedure VICH – International Cooperation on Harmonization of Technical Requirements for Registration of Veterinary Products **VIPP** – Verified internet pharmaceutical practice site (US) VMD – Veterinary Medicines Directorate VMP – Veterinary medicinal product VMRFG – Veterinary Mutual Recognition Facilitation Group **VNeeS** – Veterinary non-eCTD electronic submission **VPC** – Veterinary Products Committee (UK) **VPN** – Virtual private network vPvB – Very persistent and very bioaccumulative (biocidal products) VSI – Validation Supplementary Information VTE – Venous thromboembolism VWP - Vaccines Working Party WWW WBC - White blood cell **WC** – Written confirmation (issued by competent authority) WCPB - Women of childbearing potential WDA – Wholesale dealer's licence

WEBAE - Web adverse event(s)

WEB-RADR (project) – Recognising Adverse Drug Reactions

WEU – Well-established use



WG - Working Group
WGEO - Working Group of Enforcement Officers (HMA)
WHO - World Health Organization
WL - Warning letter - and also:
WL - Wholesale dealer's licence
WOCBP - Women of child-bearing potential
WoE - Weight of evidence
WP - Working Party
WRAC - Worldwide Regulatory Affairs Committee
WS - Work sharing
WTO - World Trade Organisation

## ххх

XEVIMPD – Extended EudraVigilance Investigational Medicinal Product Dictionary
 XEVMPD – Extended EudraVigilance medicinal products dictionary
 XEVPRM – Extended EudraVigilance product report message
 XML – Extensible Markup Language
 XRF – X-ray fluorescence

## ZZZ

ZAPI – Zoonosis Anticipation and Preparedness InitiativeZVA – Zalu Valsts Agentura (State Agency for Medicines) (Latvia)

[Last updated July 2020]