

Evaluation of certain veterinary drug residues in food

Seventy-eighth report of the Joint
FAO/WHO Expert Committee on
Food Additives



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Evaluation Of Certain Veterinary Drug Residues In Food

Padhraic Smyth



Evaluation Of Certain Veterinary Drug Residues In Food:

Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives. Meeting, World Health Organization, Food and Agriculture Organization of the United Nations, 2009 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of veterinary drugs within the terms of reference of the Joint FAO WHO Expert Committee on Food Additives JECFA including a hypothesis driven decision tree approach for the safety evaluation of residues of veterinary drugs comments on the Committee for Veterinary Products for Medicinal Use reflection paper on the new approach developed by JECFA for exposure and maximum residue limit MRL assessment of residues of veterinary drugs in honey and possible approaches to derive MRLs for this commodity comments on a paper entitled Risk assessment policies Differences among jurisdictions and the use of no observed effect level NOEL and no observed adverse effect level NOAEL in JECFA assessments Summaries follow of the Committee's evaluations of toxicological and residue data on a variety of veterinary drugs three antimicrobial agents avilamycin tilmicosin tylosin one authentic triclabendazole one production aid melengestrol acetate two antimicrobial agents and production aids monesin and narasin a glucocorticosteroid dexamethasone and an antimicrobial agent and contaminant malachite green Annexed to the report is a summary of the Committee's recommendations on these drugs including acceptable daily intakes ADIs and proposed MRLs

Toxicological evaluation of certain veterinary drug residues in food, 2021-04-12 The monographs contained in this volume were prepared at the eighty eighth meeting of the Joint Food and Agriculture Organization of the United Nations FAO World Health Organization WHO Expert Committee on Veterinary drugs JECFA which met at WHO headquarters in Rome Italy 22-31 October 2019 These monographs summarize the data on selected veterinary drugs reviewed by the Committee

Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives. Meeting, World Health Organization, 2006 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of veterinary drugs within the terms of reference of JECFA including compounds without an ADI or MRL recommendations on principles and methods in derivation of MRLs including a new procedure for estimating chronic dietary intakes the use of a spreadsheet based procedure for the statistical evaluation of residue depletion data a revised approach for the derivation of microbiological ADIs and the Committee's review of and comments on documents provided by the Codex Committee on Residues of Veterinary Drugs Summaries follow of the Committee's evaluations of toxicological and residue data on a variety of veterinary drugs three antimicrobial agents colistin erythromycin flumequine two production aids melengestrol acetate

ractopamine hydrochloride an insecticide trichlorfon metrifonate and an anthelmintic triclabendazole In addition the attempt by the Committee to use tylosin as an example to investigate if evaluations are possible based on published data in the absence of data submissions from sponsors is described Annexed to the report is a summary of the Committee's recommendations on these drugs including acceptable daily intakes and proposed maximum residue limits *Evaluation of Certain Veterinary Drug Residues in Food* Joint FAO/WHO Expert Committee on Food Additives, World Health Organization, 2001-05 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of veterinary drugs within the terms of reference of JECFA including compounds without an ADI or MRL recommendations on principles and methods in derivation of MRLs including a new procedure for estimating chronic dietary intakes the use of a spreadsheet based procedure for the statistical evaluation **Evaluation of Certain Veterinary Drug Residues in Food** World Health Organization, 1989

Toxicological evaluation of certain veterinary drug residues in food World Health Organization, Food and Agriculture Organization of the United Nations, 2023-04-28 *Toxicological evaluation of certain veterinary drug residues in food: prepared by the eighty-eighth meeting of the Joint FAO/WHO Expert Committee on Food Additives (JECFA)* World Health Organization, Food and Agriculture Organization of the United Nations, 2024-05-29 The report contains the monographs that were prepared at the eighty eighth meeting of the Joint FAO WHO Expert Committee on Food Additives JECFA which took place at the headquarters of the Food and Agriculture Organization of the United Nations FAO in Rome on 22-31 October 2019 Author These monographs summarize the data on the safety of residues in food of selected veterinary drugs reviewed by the JECFA Author *Evaluation of certain veterinary drug residues in food: Ninety-eighth report of the Joint FAO/WHO Expert Committee on Food Additives* World Health Organization, Food and Agriculture Organization of the United Nations, United Nations Children's Fund, 2024-06-17 This report represents the conclusions of the Joint FAO WHO Expert Committee on Food Additives JECFA convened to evaluate the safety of veterinary drug residues in food clopidol imidacloprid and fumagillin Annexed to this report is a summary of the Committee's recommendations on these drugs discussed at the ninety eighth meeting including acceptable daily intakes ADIs acute reference doses ARfDs estimated dietary exposure and proposed maximum residue limits MRLs Summaries are also provided for general considerations around the matters of interest arising from previous sessions of the Codex Committee on Residues of Veterinary Drugs in Foods CCRVDF update of Guidance for the Safety Evaluation of Residues of Veterinary Drugs with incomplete data packages and release of JECFA Toolbox for Veterinary Drug Residues Risk Assessment This volume and others in the WHO Technical Report Series contain information that is useful to those who produce and use veterinary drugs and those involved with controlling contaminants in food government and food regulatory officers industrial testing laboratories toxicological

laboratories and universities

Toxicological evaluation of certain veterinary drug residues in food World Health Organization, Food and Agriculture Organization of the United Nations, 2025-02-26 The monographs contained in this volume were prepared at the ninety eighth meeting of the Joint FAO WHO Expert Committee on Food Additives JECFA which met at the headquarters of the Food and Agriculture Organization of the United Nations FAO in Rome Italy on 20 29 February 2024 These monographs summarize the data on the safety of residues in food of selected veterinary drugs reviewed by the Committee namely Clopidol Fumagillin dicyclohexylamine and Imidacloprid

Toxicological Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives, Joint FAO/WHO Expert Committee on Food Additives. Meeting, 2005 The monographs in this volume summarize data on the veterinary drug residues that were evaluated toxicologically by the Committee which included three antimicrobial agents cefuroxime flumequine and pirlimycin two insecticides cyhalothrin and cypermethrin and alpha cypermethrin and one production aid ractopamine The Committee also evaluated the safety of low levels of the antimicrobial agent chloramphenicol in animal products This volume and others in the WHO Food Additives Series contain information that is useful to those who produce and use food additives and veterinary drugs and those involved with controlling contaminants in food government and food regulatory officers industrial testing laboratories toxicological laboratories and universities

Evaluation of certain veterinary drug residues in food World Health Organization, Food and Agriculture Organization of the United Nations, 2022-09-05

Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives. Meeting, 2000 This report presents the conclusions of a Joint FAO WHO Expert Committee on Food Additives which was convened to evaluate the safety of residues of certain veterinary drugs The first part of the report considers an approach to assessing the safety of antimicrobial drug residues and their effects on the human intestinal microflora and other general issues relation to the recommendation of Maximum residue Levels MRLs for veterinary drug residues in food This is followed by a summary of the Committee s evaluations of toxicological and residue data on a number of veterinary drugs a beta adrenoceptor blocking agent four antimicrobial agents two insecticides four production aids and one tranquillising agent The recommendations on these drugs including MRLs and Acceptable Daily Intakes are given as appendices

Toxicological Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives. Meeting, World Health Organization, 2003-01-01

Evaluation of Certain Veterinary Drug Residues in Food , 2003

Evaluation of Certain Veterinary Drug Residues in Food World Health Organization, 2016-02-22 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of residues of veterinary drugs within the terms of reference of the Joint FAO WHO Expert Committee on Food Additives JECFA including MRLs for generic fish species acute reference doses ARfDs for veterinary drugs an approach for

dietary exposure assessment of compounds used for multiple purposes i.e. veterinary drugs and pesticides dietary exposure assessment for less than lifetime exposure and the assessment of short term 90 day and 12 month studies in dogs Summaries follow of the Committee's evaluations of toxicological and residue data on a variety of veterinary drugs two insecticides diflubenzuron and teflubenzuron an antiparasitic agent ivermectin an ectoparasiticide sisapronil and a B2 adrenoceptor agonist zilpaterol hydrochloride In addition the Committee considered issues raised in concern forms from the Codex Committee on Residues of Veterinary Drugs in Foods on lasalocid sodium an antiparasitic agent Annexed to the report is a summary of the Committee's recommendations on these drugs including acceptable daily intakes ADIs ARfDs and proposed MRLs

Evaluation of Certain Veterinary Drug Residues in Food Joint FAO/WHO Expert Committee on Food Additives. Meeting, World Health Organization, 2014 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of residues of veterinary drugs within the terms of reference of the Joint FAO WHO Expert Committee on Food Additives JECFA including extrapolation of maximum residue limits MRLs to minor species MRLs for veterinary drug residues in honey MRLs relating to fish and fish species dietary exposure assessment methodologies the decision tree approach to the evaluation of residues of veterinary drugs and guidance for JECFA experts Summaries follow of the Committee's evaluations of toxicological and residue data on a variety of veterinary drugs two anthelmintic agents derquantel monepantel three antiparasitic agents emamectin benzoate ivermectin lasalocid sodium one antibacterial antifungal and anthelmintic agent gentian violet and two production aids recombinant bovine somatotropins zilpaterol hydrochloride Annexed to the report is a summary of the Committee's recommendations on these drugs including acceptable daily intakes ADIs and proposed MRLs

Technical Report Series, 1950 Toxicological evaluation of certain veterinary drug residues in food: prepared by the eighty-fifth meeting of the Joint FAO/WHO Expert Committee on Food Additives (JECFA) World Health Organization, Food and Agriculture Organization of the United Nations, 2024-05-29 This volume contains monographs prepared at the eighty fifth meeting of the Joint FAO WHO Expert Committee on Food Additives JECFA which met in Geneva Switzerland from 17 to 26 October 2017 Author The toxicological monographs in this volume summarize data on the veterinary drug residues that were evaluated toxicologically by the Committee amoxicillin ampicillin ethion flumethrin halquinol and lufenuron Author Annexed to the report is a summary of the Committee's recommendations on these and other drugs discussed at the eighty fifth meeting including acceptable daily intakes ADIs acute reference doses ARfDs and proposed maximum residue limits MRLs Author This volume and others in the WHO Food Additives Series contain information that is useful to those who produce and use food additives and veterinary drugs and those involved with controlling contaminants in food government and food regulatory officers industrial testing laboratories toxicological laboratories and universities Author Toxicological

evaluation of certain veterinary drug residues in food World Health Organization, Food and Agriculture Organization of the United Nations, 2019-07-17 **Evaluation of certain veterinary drug residues in food: ninety-fourth report of the Joint FAO/WHO Expert Committee on Food Additives** World Health Organization, Food and Agriculture Organization of the United Nations, 2022-10-05 This report represents the conclusions of a Joint FAO WHO Expert Committee convened to evaluate the safety of residues of certain veterinary drugs in food and to recommend maximum levels for such residues in food The first part of the report considers general principles regarding the evaluation of residues of veterinary drugs within the terms of reference of the Joint FAO WHO Expert Committee on Food Additives JECFA It covers topics such as the parallel review process estimation of dietary exposure to veterinary drug residues a risk based decision tree approach for safety evaluation assessment of the potential effects of residues on the human intestinal microbiome Summaries follow the Committee s evaluations of toxicological and residue data on a variety of veterinary drugs two antiparasitic agents imidacloprid ivermectin and one coccidiostat nicarbazin Additionally further evaluation of the parasiticide selamectin is included as part of a pilot in support of the proposed parallel review process Annexed to the report is a summary of the Committee s recommendations on these drugs including acceptable daily intakes and proposed maximum residue limits

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