
Council Regulation EEC No 2309/93 of 22 July 1993 laying down Community procedures for the authorization and supervision of medicinal products for human and veterinary use and establishing a European Agency for the Evaluation of Medicinal Products.


Guide to Good Distribution Practice (94/C 63/03).

Current version of the Guide to Good Manufacturing Practice, Rules Governing Medicinal Products in the European Community, Volume IV.

2. For the United States:

[Copies of FDA documents may be obtained from the Government Printing Office, 1510 H St. NW., Washington, DC 20005. FDA documents, except the FDA Compliance Program Guidance Manual, may be viewed on FDA’s Internet web site at http://www.fda.gov.]


APPENDIX B TO SUBPART A OF PART 26—LIST OF AUTHORITIES

1. For the United States: In the United States, the regulatory authority is the Food and Drug Administration.

2. For the European Community: In the European Community, the regulatory authorities are the following:

   Belgium: Inspection générale de la Pharmacie, Algemene Farmaceutische Inspectie.
   Denmark: Laegemiddelstyrelsen.
   Spain: For medicinal products for human use: Ministerio de Sanidad y Consumo, Subdirección General de Control Farmaceutico. For medicinal products for veterinary use: Ministerio de Agricultura, Pesca y Alimentación (MAPA), Dirección General de la Producción Agraria.
   Ireland: Irish Medicines Board.
   Italy: For medicinal products for human use: Ministero della Sanità, Dipartimento Farmaci e Farmacovigilanza. For veterinary products for veterinary use: Ministero della Sanità, Dipartimento alimenti e nutrizione e sanità pubblica veterinaria-Div. IX.
   Luxembourg: Division de la Pharmacie et des Médicaments.
   Netherlands: Staat der Nederlanden.
   Austria: Bundesministerium für Arbeit, Gesundheit und Soziales.
   Portugal: Instituto da Farmácia e do Medicamento (INFARMED).
   Finland: Lääkölaitos/Läkemedelsverket (National Agency for Medicines).
   Sweden: Läkemedelsverket-Medical Products Agency.
   European Community: Commission of the European Communities. European Agency for the Evaluation of Medicinal Products (EMEA).

APPENDIX C TO SUBPART A OF PART 26—INDICATIVE LIST OF PRODUCTS COVERED BY SUBPART A

Recognizing that precise definition of medicinal products and drugs are to be found in the legislation referred to above, an indicative list of products covered by this arrangement is given below:

—human medicinal products including prescription and nonprescription drugs;
—human biologicals including vaccines, and immunologicals;
—veterinary pharmaceuticals, including prescription and nonprescription drugs, with the exclusion of veterinary immunologicals (Under 9 CFR 101.2 “veterinary immunologicals” are referred to as “veterinary biologicals”);
—premixes for the preparation of veterinary medicated feeds (EC), Type A medicated articles for the preparation of veterinary medicated feeds (EC).