# Directive 2001/83/EC of the European Parliament and of the Council of 6 November 2001 on the Community code relating to medicinal products for human use

# TITLE I

### DEFINITIONS

## Article 1

For the purposes of this Directive, the following terms shall bear the following meanings:

[ <sup>F2</sup> 2.Medicinal product		(a)	Any substance or combination of substances presented as having properties for treating or preventing disease in human beings; or				
		(b)	Any substance or combination of substances which may be used in or administered to human beings either with a view to restoring, correcting or modifying physiological functions by exerting a pharmacological, immunological or metabolic action, or to making a medical diagnosis.]				
3. Substance		Any matter irrespective of origin which may be: — human, e.g.					
		—					
		_	human blood and human blood products; animal, e.g.				
		_	micro-organisms, whole animals, parts of organs, animal secretions, toxins, extracts, blood products; vegetable, e.g.				
		_	micro-organisms, plants, parts of plants, vegetable secretions, extracts; chemical, e.g.				
			elements, naturally occurring chemical materials and chemical				
[ <sup>F3</sup> 3a.Active substance	:	products obtained by chemical change or synthesis. Any substance or mixture of substances intended to be used in the manufacture of a medicinal product and that, when used in its production, becomes an active ingredient of that product intended to exert a pharmacological, immunological or metabolic action with a view to restoring, correcting or modifying physiological functions or to make a medical diagnosis.					
3b.Excipient	:	Any constituent of a medicinal product other than the active substance and the packaging material.] Any medicinal product consisting of vaccines, toxins, serums or allergen products:					
4. Immunological medicinal product	:						
		(a)	vaccines, toxins and serums shall cover in particular:				

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			(i)	agents used to produce active immunity, such as cholera vaccine, BCG, polio vaccines, smallpox vaccine;			
			(ii)	agents used to diagnose the state of immunity, including in particular tuberculin and tuberculin PPD, toxins for the Schick and Dick Tests, brucellin;			
			(iii)	agents used to produce passive immunity, such as diphtheria antitoxin, anti-smallpox globulin, antilymphocytic globulin;			
		(b)	intended	product' shall mean any medicinal product which is to identify or induce a specific acquired alteration in inological response to an allergizing agent.			
[ <sup>F4</sup> 4a. <i>Advanced</i> <i>therapy medicinal</i> <i>product</i> [ <sup>F2</sup> 5.Homeopathic medicinal product	:	A product as defined in Article 2 of Regulation (EC) No 1394/2007 of the European Parliament and of the Council of 13 November 2007 on advanced therapy medicinal products <sup>(1)</sup> .] Any medicinal product prepared from substances called homeopathic stocks in accordance with a homeopathic manufacturing procedure described by the European Pharmacopoeia or, in the absence thereof, by the pharmacopoeias currently used officially in the Member States. A homeopathic medicinal product may contain a number of principles.]					
(							
6.	:			duct which, when ready for use, contains one or more			
Radiopharmaceutica	al			oactive isotopes) included for a medicinal purpose.			
7. Radionuclide	:			porating a fixed parent radionuclide from which is			
generator		<b>.</b>	•	er radionuclide which is to be obtained by elution or of and used in a radiopharmaceutical.			
8. [ <sup>F2</sup> Kit]	:	* * *		be reconsitituted or combined with radionuclides in maceutical, usually prior to its administration.			
9. Radionuclide	:	Any othe	er radion	uclide produced for the radio-labelling of another			
precursor		substance	e prior to	administration.			
10. Medicinal	:	Medicina	al product	s based on blood constitutents which are prepared			
products derived from human blood				ublic or private establishments, such medicinal g, in particular, albumin, coagulating factors and			
or human plasma				of human origin.			
[ <sup>F5</sup> 11.Adverse reaction	:			edicinal product which is noxious and unintended.]			
12. Serious	:	An adver	se reactio	n which results in death, is life-threatening, requires			
adverse reaction	•	inpatient results in	hospitali n persiste	sation or prolongation of existing hospitalisation, ent or significant disability or incapacity, or is a y/birth defect.			
13. Unexpected adverse reaction	:			on, the nature, severity or outcome of which is not summary of product characteristics.			
E.E	:						
[ <sup>F5</sup> 15.Post- authorisation safety study	:	the aim of confirmin	of identifing the saf	to an authorised medicinal product conducted with ying, characterising or quantifying a safety hazard, ety profile of the medicinal product, or of measuring f risk management measures.]			
16. Abuse				dic, intentional excessive use of medicinal products			
of medicinal products	•			nied by harmful physical or psychological effets.			

17. Wholesale : distribution of medicinal products	All activities consisting of procuring, holding, supplying or exporting medicinal products, apart from supplying medicinal products to the public. Such activities are carried out with manufacturers or their depositories, importers, other wholesale distributors or with pharmacists and persons authorized or entitled to supply medicinal products to the public in the Member State concerned.			
[ <sup>F3</sup> 17a.Brokering : of medicinal products	All activities in relation to the sale or purchase of medicinal products, except for wholesale distribution, that do not include physical handling and that consist of negotiating independently and on behalf of another legal or natural person.]			
18. Public service : obligation	The obligation placed on wholesalers to guarantee permanently an adequate range of medicinal products to meet the requirements of a specific geographical area and to deliver the supplies requested within a very short time over the whole of the area in question.			
[ <sup>F7</sup> 18aRepresentative: of the marketing authorisation holder	The person, commonly known as local representative, designated by the marketing authorisation holder to represent him in the Member State concerned.]			
19. Medicinal :	Any medicinal prescription issued by a professional person qualified to			
Prescription	do so.			
[ <sup>F2</sup> 20.Name of the : medicinal product	The name, which may be either an invented name not liable to confusion with the common name, or a common or scientific name accompanied by a trade mark or the name of the marketing authorisation holder.]			
21. Common :	The international non-proprietary name recommended by the World			
name 22. Strength of the :	Health Organization, or, if one does not exist, the usual common name. The content of the active substances expressed quantitatively per dosage			
medicinal product	unit, per unit of volume or weight according to the dosage form.			
23. Immediate :	The container or other form of packaging immediately in contact with			
packaging	the medicinal product.			
24. Outer :	The packaging into which is placed the immediate packaging.			
packaging				
25. Labelling:26. Package:	Information on the immediate or outer packaging.			
leaflet	A leaflet containing information for the user which accompanies the medicinal product.			
[ <sup>F8</sup> 26a.Variation :	An amendment to the contents of the particulars and documents referre			
or variation to	to in:			
the terms of	(a) Article 8(3) and Articles 9 to 11 of this Directive and Annex			
a marketing authorisation	I thereto, Article 6(2) of Regulation (EC) No 726/2004 and in Article 7 of Regulation (EC) No 1394/2007; and			
	(b) the terms of the decision granting the marketing authorisation for a medicinal product for human use, including the summary of the product characteristics and any conditions, obligations, or restrictions affecting the marketing authorisation, or changes to the labelling or the package leaflet related to			
[ <sup>F2</sup> 27.Agency :	changes to the summary of the product characteristics.] The European Medicines Agency established by Regulation (EC) No 726/2004 <sup>(2)</sup> .]			
[ <sup>F2</sup> 28.Risks related :	— any risk relating to the quality, safety or efficacy of the			
to use of the	medicinal product as regards patients' health or public health;			
medicinal product	— any risk of undesirable effects on the environment.			

28a.Risk-benefit	:	An evalu	uation of the positive therapeutic effects of the medicinal					
balance		product i	n relation to the risks as defined in point 28, first indent.]					
[ <sup>F9</sup> 28b.Risk	:		pharmacovigilance activities and interventions designed to					
management		identify,	characterise, prevent or minimise risks relating to a medicinal					
system			including the assessment of the effectiveness of those activities ventions.					
28c.Risk	:		d description of the risk management system.					
management plan								
28d.Pharmacovigila	nnce		used by the marketing authorisation holder and by Member					
system		States to	fulfil the tasks and responsibilities listed in Title IX and					
		designed	to monitor the safety of authorised medicinal products and					
		detect an	y change to their risk-benefit balance.					
28e.Pharmacovigila	nce	ed description of the pharmacovigilance system used by the						
system master file			g authorisation holder with respect to one or more authorised il products.]					
[ <sup>F10</sup> 29.Traditional			l medicinal product that fulfils the conditions laid down in					
herbal medicinal	•	Article 1						
product			(1).					
30.Herbal		Any med	licinal product, exclusively containing as active ingredients one					
medicinal product	•		herbal substances or one or more herbal preparations, or one					
medicinal product		or more such herbal substances in combination with one or more such						
			eparations.					
31.Herbal			ly whole, fragmented or cut plants, plant parts, algae, fungi,					
substances	•		an unprocessed, usually dried, form, but sometimes fresh.					
substances			xudates that have not been subjected to a specific treatment are					
			sidered to be herbal substances. Herbal substances are precisely					
			by the plant part used and the botanical name according to the					
			system (genus, species, variety and author).					
32.Herbal			ons obtained by subjecting herbal substances to treatments					
preparations	•		extraction, distillation, expression, fractionation, purification,					
proparations			ation or fermentation. These include comminuted or powdered					
			bstances, tinctures, extracts, essential oils, expressed juices and					
			d exudates.]					
[ <sup>F3</sup> 33.Falsified		<b>1</b>	licinal product with a false representation of:					
medicinal product	•	Any mee	nemai product with a faise representation of.					
medicinal product		(a)	its identity, including its packaging and labelling, its name or					
			its composition as regards any of the ingredients including					
			excipients and the strength of those ingredients;					
		(1-)	ite second including ite menufactures ite secondary of					
		(b)	its source, including its manufacturer, its country of					
			manufacturing, its country of origin or its marketing					
			authorisation holder; or					
		(c)	its history, including the records and documents relating to the					
			distribution channels used.					
	This definition does not include unintentional quality defects a							
		without p	prejudice to infringements of intellectual property rights.]					

#### **Textual Amendments**

**F1** Deleted by Directive 2004/27/EC of the European Parliament and of the Council of 31 March 2004 amending Directive 2001/83/EC on the Community code relating to medicinal products for human use.

- **F2** Substituted by Directive 2004/27/EC of the European Parliament and of the Council of 31 March 2004 amending Directive 2001/83/EC on the Community code relating to medicinal products for human use.
- **F3** Inserted by Directive 2011/62/EU of the European Parliament and of the Council of 8 June 2011 amending Directive 2001/83/EC on the Community code relating to medicinal products for human use, as regards the prevention of the entry into the legal supply chain of falsified medicinal products (Text with EEA relevance).
- F4 Inserted by Regulation (EC) No 1394/2007 of the European Parliament and of the Council of 13 November 2007 on advanced therapy medicinal products and amending Directive 2001/83/EC and Regulation (EC) No 726/2004 (Text with EEA relevance).
- **F5** Substituted by Directive 2010/84/EU of the European Parliament and of the Council of 15 December 2010 amending, as regards pharmacovigilance, Directive 2001/83/EC on the Community code relating to medicinal products for human use (Text with EEA relevance).
- F6 Deleted by Directive 2010/84/EU of the European Parliament and of the Council of 15 December 2010 amending, as regards pharmacovigilance, Directive 2001/83/EC on the Community code relating to medicinal products for human use (Text with EEA relevance).
- **F7** Inserted by Directive 2004/27/EC of the European Parliament and of the Council of 31 March 2004 amending Directive 2001/83/EC on the Community code relating to medicinal products for human use.
- F8 Inserted by Regulation (EU) 2019/5 of the European Parliament and of the Council of 11 December 2018 amending Regulation (EC) No 726/2004 laying down Community procedures for the authorisation and supervision of medicinal products for human and veterinary use and establishing a European Medicines Agency, Regulation (EC) No 1901/2006 on medicinal products for paediatric use and Directive 2001/83/EC on the Community code relating to medicinal products for human use (Text with EEA relevance).
- F9 Inserted by Directive 2010/84/EU of the European Parliament and of the Council of 15 December 2010 amending, as regards pharmacovigilance, Directive 2001/83/EC on the Community code relating to medicinal products for human use (Text with EEA relevance).
- **F10** Inserted by Directive 2004/24/EC of the European Parliament and of the Council of 31 March 2004 amending, as regards traditional herbal medicinal products, Directive 2001/83/EC on the Community code relating to medicinal products for human use.

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- (1) [<sup>F4</sup>OJ L 324, 10.12.2007, p. 121.]
- (2) [<sup>F2</sup>OJ L 136, 30.4.2004, p. 1.]

#### **Textual Amendments**

- **F2** Substituted by Directive 2004/27/EC of the European Parliament and of the Council of 31 March 2004 amending Directive 2001/83/EC on the Community code relating to medicinal products for human use.
- F4 Inserted by Regulation (EC) No 1394/2007 of the European Parliament and of the Council of 13 November 2007 on advanced therapy medicinal products and amending Directive 2001/83/EC and Regulation (EC) No 726/2004 (Text with EEA relevance).