CHAPTER 20

Model declaration

Declaration for the import from third countries and for the transit through the European Union of intermediate products to be used for the manufacture of medicinal products, veterinary medicinal products, medical devices, in vitro diagnostics and laboratory reagents

COUNTRY: UNITED STATES

Veterinary certificate to EU

	I.1. Consignor Name	I.2. Certificate reference	No	l.2.a.	
	Address				
		I.3. Central competent au	thority		
	Tel.	I.4. Local competent auth	ority		
	I.5. Consignee Name Address	I.6. Person responsible fo Name Address	or the load in E	U	
ient	Postal code Tel.	Postal code Tel.			
Part I : Details of dispatched consignment	I.7. Country of ISO code I.8. Region of Code origin		ISO I. code	10. Region of Code destination	
Itche	I.11. Place of origin	I.12. Place of destination			
ls of dispa	Name Approval number Address	Name Address	Custom warehouse Approval number		
Detail		Postal code			
Par	Name Approval number Address Name Approval number Address				
	I.13. Place of loading	I.14. Date of departure			
	I.15. Means of transport	I.16. Entry BIP in EU			
	Aeroplane Ship Railway wagon				
	Road vehicle Other Other	I.17.			
	Identification				
	Documentation references				
	I.18. Description of commodity	I.19. Commodity code	(HS code)		
			I.20. Quan	tity	
	I.21. Temperature of product Ambient Chilled	Frozen 🗆	I.22. Numl	per of packages	
	I.23. Seal/Container No		I.24. Type	of packaging	
	Page of				

	1.2.	Certificate reference	No	l.2.a.		
I.25. Commodities certified for:			~~~~			
Technical use 🗆						
I.26. For transit through EU to third country		I.27. For import or admission into EU				
Third country ISO code						
I.28. Identification of the commodities		1				
SpeciesApproval number of establishments(Scientific name)Manufacturing plant		Net weight	Batch number			

	н.	Health ii	nformation	II.a.	Certificate reference No	II.b.			
1	DECLARATION								
		into tl			product referred to above is intended to for in point 35 of Annex I of Commission I				
_	(1) ⁽²⁾ either	it is intended for the manufacture of:							
ior		[-	medicinal products,]						
cat	⁽²⁾ and/or	[-	veterinary medicinal products,]						
Tifi	⁽²⁾ and/or	[-	medical devices,]						
Cel	⁽²⁾ and/or	[-	active implantable medical devices,]						
≓	⁽²⁾ and/or	[-	in vitro diagnostic medical devices,]						
Part II: Certification	⁽²⁾ and/or [- laboratory reagents;]								
-	(2) its design, transformation and manufacturing stages have been sufficiently completed in order to qualify the material directly or as a component of a product intended for that purpose, except for the fact that it requires further handling or transformation such as mixing, coating, assembling, packaging or labelling to make it suitable for placing on the market or putting into service as medicinal products, veterinary medicinal products, active implantable medical device, medical devices or in vitro diagnostic medical device in accordance with the Union legislation ^(1b) applicable to those products or as laboratory reagents;								
	(3)	it has been derived from the following material which may have originated from animals which have been submitted to illegal treatment as defined in Article 1(2)(d) of Council Directive 96/22/EC or Article 2(b) of Council Directive 96/23/EC ⁽²⁾ :							
⁽²⁾ either [- carcases and parts of animals slaughtered or, in the case of game, bodies or parts of an and which are fit for human consumption in accordance with Union legislation, but are not human consumption for commercial reasons;]									
	⁽²⁾ and/or [- carcases and the following parts originating either from animals that have been slaughtered slaughterhouse and were considered fit for slaughter for human consumption following an ante-me inspection or bodies and the following parts of animals from game killed for human consumption accordance with Union legislation:				ing an ante-mortem				
					imals which are rejected as unfit for hun but which did not show any signs of dise				
			(ii) heads of poultry;						
	 (iii) hides and skins, including trimmings and splitting thereof, horns and fe phalanges and the carpus and metacarpus bones, tarsus and metatarsus b other than ruminants; 								
			(iv) pig bristles;						
	⁽²⁾ and/or	[-	animals obtained from animals other	er than for sla	signs of disease communicable through the ruminants that have been slaughtered in ughter for human consumption followin tion;]	n a slaughterhouse			
	⁽²⁾ and/or	[-			tion of products intended for human con separator sludge from milk processing;]	sumption, including			
	⁽²⁾ and/or	[-	products of animal origin, or foodstuffs containing products of animal origin, which are no longer intended for human consumption for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arise;]						
	⁽²⁾ and/or	[-	products, which are no longer inte	s of animal origin, or feedingstuffs containing animal by-products or derived longer intended for feeding for commercial reasons or due to problems of ging defects or other defects from which no risk to public or animal health					
⁽²⁾ and/or [- blood, placenta, wool, feathers				ir, horns, hoof cuts and raw milk originating from live animals that did nmunicable through that product to humans or animals;]					

Page ____ of ____

Н.	Health ii	nformation II.a. Certificate reference No II.b.						
(2) and/or	[-	aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;]						
⁽²⁾ and/or	[-	animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;]						
⁽²⁾ and/or	[-	 the following material originating from animals which did not show any signs of disease communicable through that material to humans or animals: (i) shells from shellfish with soft tissue or flesh; (ii) the following originating from terrestrial animals: hatchery by-products, eggs, egg by-products, including egg shells; (iii) day-old chicks killed for commercial reasons;] 						
⁽²⁾ and/or	[-	animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;]						
⁽²⁾ and/or	[-							
⁽²⁾ and/or	[-	products derived from or generated by:						
		 aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of disease communicable to humans or animals, 						
		- aquatic or terrestrial invertebrates other than species pathogenic to humans or animals,						
		 animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) and Category 2 material as referred to in Article 9(a) to (g) of Regulation (EC) No 1069/2009;] 						
⁽²⁾ and/or	[-	animals and parts of animals, other than those referred to in Article 8 or Article 10 of Regulation (EC) No 1069/2009,						
		 that die other than by being slaughtered or killed for human consumption, including animals killed for disease control purposes; 						
		 (ii) foetuses; (iii) oocytes, embryos and semen which are not destined for breeding purposes; and (iv) dead-in-shell poultry;] 						
(2)and/or	[-	animal by-products other than Category 1 material or Category 3 material;]						
(4)	MEDI DEVI	uter packaging is labelled 'FOR MEDICINAL PRODUCTS / VETERINARY MEDICINAL PRODUCTS / ICAL DEVICES/ ACTIVE IMPLANTABLE MEDICAL DEVICES / IN VITRO DIAGNOSTIC MEDICAL CES/ LABORATORY REAGENTS ONLY' and it is not intended to be diverted at any stage within the in for any other use;						
(5)		onsignment will be transported directly to the place of destination as indicated under point I.12 of this ration, that is:						
	-	an establishment or plant for the production of medicinal products, veterinary medicinal products, medical devices, in vitro diagnostics or laboratory reagents, which has been registered in accordance with Article 23 of Regulation (EC) No 1069/2009,						
	-	an establishment or plant which has been approved in accordance with Article 24(1)(i) of Regulation (EC) No 1069/2009, from where they shall only be dispatched to an establishment or plant referred to in the preceding subpoint of (5).						
Notes	_							
(10)								
(11)		2.2011, p. 1. 01/02/EC of the European Darliament and of the Council of 6 Nevember 2001 on the Community and						
Directive 2001/82/EC of the European Parliament and of the Council of 6 November 2001 on the Community code relating to veterinary medicinal products (OJ L 311, 28.11.2001), 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 (OJ L 311, 28.11.2001, p. 67), Council Directive 93/42/EEC of 14 June 1993 concerning medical devices (OJ L 169, 12.7.1993, p. 1.) and Directive 98/79/EC of the European Parliament and the Council of 27 October 1998 on in vitro								

Page ____ of ____

II.	Health information	II.a.	Certificate reference No	II.b.				
(2)	diagnostic medical devices (OJ L 331, 7.12.1998, p. 1.), as appropriate.							
Nar	importer ne (in capital letters): Iress:							
Dat	e: Signature:							