Export Health Certificate

	I.1. Consignor				I.2. IMSOC Ref	erence			
	Name				I.2.a. Local Reference				
	Address								
	Country ISO Code								
	I.5. Consignee				L3. Central co	mpetent authority			
Ļ	Name					petent authority			
en	Address								
띩	Country ISO Code								
igi	I.7. Country of origin ISO Code				I.9. Country of destination ISO Co			ISO Code	
Suc	1.7. Country of orig	3111			1.3. Country of destination				130 Code
Part I : Details of consignment	IO Danian of animi			Cada	I 10 Danian at				
S O	L8. Region of origin Code L11. Place of Dispatch				I.10. Region of destination I.12. Place of destination				
ail	Name				Name				
ĕ	Name Address				Address				
:	Approval Number	r			Approval Number				
ᇳ	Country		ISO Code		Country ISO Code				
Pa	I.13. Place of Loadi	ing			I 14 Date and	time of departure			
	Name	шв			1.14. Date and	unie of departure			
	Address								
	Approval Number	ſ							
	Country		ISO Code						
	I.15. Means of Trai	acnort			I.16 Entry Poi	<u></u>			
	Mode	International	Identification		1.10 EIIII y Poli	ц			
	Wode	transport	identification						
		document							
	I.18. Transport con		<u>_</u>	_	I.17. Accompanying documents				
	Chilled ☐ Controlled Frozen ☐ Ambient ☐ temperature ☐			nbient \square	Accompanyi				
		temperature =	•		ng Date of issu		issue		
					reference Place of			.f	
					Country		issue	1	
	I.19. Container No	/ Seal No							
	I.20. Certified as								
	Technical use \square		Slaughter \square		Production of petfood \square		Production \square		
	Human consumption \square		Other 🗆		Relaying 🗆		Animal Feedingstuff \square		
	Breeding \square Fattening \square			Pharmaceutical use Breeding and production		ion 🗆			
	Artificial reproduction								
	I.21. For transit through a third country				I.22. For transit through Member State(s)				
	Country	-	ISO Code						
	EU Entry Authority BC		BCP code				ISO Code		
			BCP code						
							105 7		
			I.24. Total quantity		I.25. Total net weight		1.25. 10	tal gross weigh	ıı
	I.28. Description of consignment								
	1. 04 DAIRY PRODUCE; BIRDS' EGGS; NATURAL HONEY; EDIBLE PRODUCTS OF ANIMAL ORIGIN, NOT ELSEWHERE SPECIFIED OR INCLUDED								LUDED
	0407 Birds' eggs	0407 Birds' eggs, in shell, fresh, preserved or cooked							
	Commodity	Spe	cies	Quantity		Net weight		Package count	
	Identification nun	nber			Identification system				

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	II. Health information							
	DECLARATION							
	i, the undersigned, declare that the intermediate product referred to above is intended to be imported by me into or							
	to be transited through Great Britain, Channel Islands or Isle of Man and satisfies the definition of an intermediate product provided for in point 35 of Annex I to Commission Regulation (EU) No 142/2011, and in particular that:							
_	(1)	it is intended for the manufacture	of:					
ion		(2) □ either [- medicinal products	,]					
Part II: Certification		(2) □ and/or [- veterinary medicin	nal products,]					
		(2) \square and/or [- medical devices fo	r medical and veterinary pur	poses,]				
		(2) □ and/or [- active implantable	medical devices,]					
		(2) \square and/or [- in vitro diagnostic	medical devices for medical a	and veterinary purposes,]				
		(2) □ and/or [- laboratory reagent	rs,]					
		(2) □ and/or [- cosmetic products;]					
	(2)	its design, transformation and ma order to qualify the material direct purpose, except for the fact that it as mixing, coating, assembling or p putting into service as a medicinal for medical and veterinary purpos diagnostic medical device for medi- accordance with the retained EU la reagent;	tly or as a component of a pro requires further manufacturi packaging to make it suitable of product, veterinary medicinal es, an active implantable medical and veterinary purposes	oduct intended for that ng or transformation such for placing on the market or al product, medical device dical devices, an in vitro or a cosmetic product in				
	(3)	it has been derived from:						
		(2) ☐ either [- material which material treatment as defined in Article 1(2) Council Directive 96/23/EC;]	•	_				
		(2) □ and/or [- carcases and parts parts of animals killed, and which retained EU law, but are not intended.	are fit for human consumptio	on in accordance with				
		(2) □ and/or [- carcases and the form been slaughtered in a slaughterhor consumption following an ante-more animals from game killed for hum	use and were considered fit fo ortem inspection or bodies an	or slaughter for human d the following parts of				
		human consumption in	and parts of animals which ar a accordance with retained EV ase communicable to humans	J law, but which did not				
		(ii) heads of poultry;						
		including the phalange	cluding trimmings and splitti is and the carpus and metaca nimals other than ruminants;	rpus bones, tarsus and				
		(iv) pig bristles;						
	(v) feathers;]						
		(2) □ and/or [-blood of animals w through blood to humans or animals been slaughtered in a slaughterhold human consumption following an law;]	als obtained from animals oth use after having been conside	er than ruminants that have red fit for slaughter for				
		(2) □ and/or [-animal by-products human consumption, including de from milk processing;]						

EUROPEAN UNION (GB) Intermediate products GBHC144E II. Health information (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 [- petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises;] (2) and/or [- blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals;] (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) \square and/or [- animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;] (2) \square 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) \square and/or [- animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;] (2) \square and/or [- 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) \square 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 1069/2009:1 (2) \square and/or [- animals and parts of animals, other than those referred to in Article 8 or Article 10 of Regulation (EC) No 1069/2009, (i) that died 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; (iv) dead-in-shell poultry;] (2) ☐ and/or [- animal by-products other than Category 1 material or Category 3 material;] (4) its outer packaging is labelled 'FOR MEDICINAL PRODUCTS / VETERINARY MEDICINAL PRODUCTS /

MEDICAL DEVICES FOR MEDICAL AND VETERINARY PURPOSES / ACTIVE IMPLANTABLE MEDICAL DEVICES / IN VITRO DIAGNOSTIC MEDICAL DEVICES FOR MEDICAL AND VETERINARY PURPOSES / LABORATORY REAGENTS / COSMETIC PRODUCTS ONLY' and it is not intended to be diverted at any stage within Great Britain, Channel Islands or Isle of Man for any other use;

EUROPEAN UNION

	N UNION		(GD) Internit	emate products GBHC144E			
II. Health i	information						
(5)	the consignment will Islands or Isle of Man	Great Britain, Channel is:					
ation	(2) o either	[an establishment or plant for the production of medicinal products, veterinary medicinal products, medical devices for medical and veterinary purposes, active implantable medical devices, in vitro diagnostic medical devices for medical and veterinary purposes, laboratory reagents or cosmetic products, which has been registered in accordance with Article 23 of Regulation (EC) No 1069/2009],					
Part II: Certification	(2) ∘ or	24(1)(i) of Regulation (I	olant which has been approve EC) No 1069/2009, from where Lishment or plant referred to i	e they may only be			
Notes							
	e countries subject to the ; Iceland and Switzerland	_	ngements include: an EU men	nber State; Liechtenstein;			
been ret	ained in Great Britain (re	tained EU law as defined	ficate are references to direct l in the European Union (With				
Reference			anel Islands and Isle of Man.				
			e end of the transitional stagi				
	 Box reference I.19: use appropriate Harmonised System (HS) code in accordance with Commission Decision 2007/275/EC of 17 April 2007 concerning lists of animals and products to be subject to controls at border control posts in accordance with Council Directives 91/496/EEC and 97/78/EC 						
_	Box reference I.25: te	chnical use: any use othe	er than for animal consumption	on.			
(1b) 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, 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, Council Directive 93/42/EEC of 14 June 1993 concerning medical devices Directive 98/79/EC of the European Parliament and the Council of 27 October 1998 on in vitro dia medical devices, Regulation (EC) No 1223/2009 of the European Parliament and of the Council of November 2009 on cosmetic products, as appropriate.							
(2)	Delete as appropriate						
Certifying	Officer		Qualification and title				
Name (in Date of signal Stamp	capital letters) gnature						

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