EUROPEAN UNION

	I.1. Consignor			I.2. IMSOC Reference		
	Name			I.2.a. Local Reference		
	Address					
	Country		ISO Code			
	I.5. Consignee			I.3. Central competent authority	I.3. Central competent authority	
nt	Name			I.4. Local competent authority	I.4. Local competent authority	
ne	Address					
Eng	Country		ISO Code			
consi	I.7. Country of ori	gin	ISO Code	I.9. Country of destination	ISO Code	
Part I : Details of consignment	I.8. Region of origi	in	Code	I.10. Region of destination		
	I.11. Place of Dispa	atch		I.12. Place of destination		
	Name			Name		
Ď	Address			Address		
t I :	Approval Numbe Country	r	ISO Code	Approval Number Country	ISO Code	
Par	country		150 Coue	country	130 Code	
H	I.13. Place of Load	ling		I.14. Date and time of departure		
	Name					
	Address	7				
	Approval Numbe Country	r	ISO Code			
	-		100 0000			
	I.15. Means of Tra	T	T.J (*C*	I.16 Entry Point		
	Mode	International transport	Identification			
		document		_		
				_		
	I.18. Transport co			I.17. Accompanying documents		
	Frozen Controlled Ambient Chilled Lemperature			Accompanying document reference		
		-		Date of issue		
				Country Place of issue		
	I.19. Container No	/ Seal No		Thee of issue		
	I.20. Certified as					
	Pharmaceutical us	se 🗆	Other 🗆	Slaughter 🗆	Relaying 🗆	
	Pharmaceutical use □ Artificial reproduction □ Production of petfood □		Fattening 🗆	Production	Breeding	
			Breeding and production \Box	Animal Feedingstuff	Human consumption	
	Technical use					
	I.21. For transit th	rough a third cour	itry	I.22. For transit through Member State(s)		
	Country		ISO Code	Country	ISO Code	
	EU Exit		BCP code			
	Authority EU Entry		BCP code			
	Authority			—		
	I.23. Total number	r of packages	I.24. Total quantity	I.25. Total net weight	I.25. Total gross weight	
	I.28. Description of consignment					
	1. 35 ALBUMINOI	DAL SUBSTANCES;	MODIFIED STARCHES; GLUES; ENZ	ZYMES		
	3501 Casein, ca	seinates and other	casein derivatives; casein glues			
	#1. Commodity		Quantity	Net weight	Package count	
	Species		Identification number	Identification system		
				1		

EUROPEAN UNION

	II. Health info	rmation							
	II.1.	Animal Health Attestation							
	11.1.				o that I am awaro of the roles	ant provisions of Directive			
		I, the undersigned official veterinarian, declare that I am aware of the relevant provisions of Directive 2002/99/EC and of Regulation (EC) No 853/2004 and hereby certify that the \Box colostrum/ \Box colostrum-based products (1) described in Part I:							
	have been	obtained or	manufactu	red from colostrum obta	ained from animals:				
апо		(i)	under the o	control of the official ve	terinary service;				
Part II: Certification		(ii)	which were in a third country or part thereof that has been free of foot-and-mouth disease and of rinderpest for a period of at least 12 months prior to the date of this certificate, and where vaccination against foot-and-mouth disease has not been carried out during that period;						
		(iii)	belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest, and						
		(iv)	subject to regular veterinary inspections to ensure that they satisfy the animal health conditions laid down in Chapter I of Section IX of Annex 3 to Regulation (EC) No 853/2004 and in Directive 2002/99/EC.						
	II.2.	Public Hea	Public Health Attestation						
		I, the undersigned official inspector, declare that I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EU) 2019/627 and hereby certify that the \Box colostrum/ \Box colostrum-based products made with colostrum(1) described in Part I were produced in accordance with those provisions, and in particular that:							
		(a)							
			(i)		lings registered in accordanc in accordance with Articles 49				
			(ii)	-	ollected, cooled, stored and tr tions laid down in Chapter I o /2004;	-			
			(iii)	provided by the monitor	ne guarantees on the residues oring plans for the detection o ce with Directive 96/23/EC, an	of residues or substances			
			(iv)	food business operator of Chapter I of Section with the maximum res	ing for residues of antibacter in accordance with the requi- IX of Annex III to Regulation idue limits for residues of ant d down in the Annex to Regul	rements of point 4 in Part III (EC) No 853/2004, complies ibacterial veterinary			
			(v)	maximum residue leve	ced under conditions guarant els for pesticides laid down in or contaminants laid down in	Regulation (EC) No 396/2005,			
		(b) they come from an establishment principles in accordance with Reg			based on the HACCP				
		(c)	-	-	wrapped, packaged and label Annex III to Regulation (EC)				
		(d)	Regulation	(EC) NO 853/2004 and the	nts laid down in Chapter II, Se he relevant microbiological cr nicrobiological criteria for foo	riteria laid down in			
		(e)			als and products thereof prov active 96/23/EC, and in particu				

EUROPEAN UNION

	II. Health info	rmation							
	been retair	References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018) and can be viewed on the UK legislation website (legislation.gov.uk).							
	References	eferences to Great Britain in this certificate include Channel Islands and Isle of Man.							
ification	This certificate is intended for colostrum or colostrum-based products from third countries or parts thereof authorised in column A as set out in a document relating to 'milk and milk products' published on gov.uk in accordance with Regulation (EU) No 605/2010. (2) Part I:								
Part II: Certification		Box reference I.7:		country or part thereof as appearing in a document s' published on gov.uk in accordance with Commission					
ה	Box reference I.11: Name, address and app			oval number of the establishment of dispatch.					
		(aircraft) o	nce I.15: Registration number (railw r name (ship). In case of unloading s st of introduction into Great Britain.	and reloading, the consignor	-				
		Box reference I.19:	For containers or boxes, the contai included.	iner number and the seal number (if applicable) should be					
		Box reference I.25:	Indicate total gross weight and tota	al net weight.					
	Box reference I.28:		Use the appropriate Harmonised System (HS) code under the following headings: 04.01; 04.02; 04.03; 04.04; 04.05; 04.06; 04.10; 15.17; 17.02; 19.01; 21.05; 21.06; 22.02; 28.35; 30.01; 35.01; 35.02 or 35.04.						
		Box reference I.28:	Manufacturing plant: introduce the collection centre or standardizatio						
	Part II:								
	(2) A d		Keep as appropriate.						
			A document relating to 'milk and milk products' for EU and EFTA states published by the Secretary of State, with the consent of the Scottish and Welsh Ministers, may be found here:						
	EU and EFT	[A states ap]	states approved to export animals and animal products to Great Britain - data.gov.uk						
		The colour of the signature shall be different to that of the printing. The same rule applies to stamps other than hose embossed or watermark.							
	Certifying Officer Name (in capital letters) Date of signature Stamp		Qualification and title Signature						