EUROPEAN UNION

	I.1. Consignor				I.2. IMSOC Reference						
	Name				I.2.a. Local Reference						
	Address										
	Country		ISO Coo	de							
	I.5. Consignee						I.3. Central co	mpetent au	ithority		
	Name					I.4. Local com					
En.	Address							,			
nm	Country ISO Code										
Part I : Details of consignment	I.7. Country of orig				ISO Code	I.9. Country of	Country of destination ISO Code				
of c	I.8. Region of origi				Code	I.10. Region of destination					
ls (I.11. Place of Dispa					I.12. Place of destination					
tai	Name					Name					
a	Address					Address					
	Approval Number					Approval Number					
art	Country							Country ISO Code			
Pŝ	I.13. Place of Load	ing					I.14. Date and	time of de	parture		
	Name										
	Address										
	Approval Number	Approval Number									
	Country ISO Code										
	I.15. Means of Trai	nsport					I.16 Entry Point				
	Mode	Internation	nal	Identificati	on						
		transport document									
							-				
							-				
	I.18. Transport cor	nditions					I.17. Accompanying documents Commercial document Date of issue reference Place of				
	Frozen	Controlled	_	Ambient 🗆] Ch	illed 🗆					
		temperatur	re 🗆								
							Country Place of sissue				
	I.19. Container No	/ Seal No									
	I.20. Certified as										
	Human consumpti	ion 🗆									
	.21. For transit through a third country						I.22. For transit through Member State(s)				
	Country			ISO Code							
	EU Exit Authority	EU Exit BCD code					Country ISO Code				
	EU Entry			BCP code							
	Authority BCP code I.23. Total number of packages I.25. Total net weight								I.25. Total gross we	ight	
	1.23. 10tal liuliber	of package.	5		1.23. 10ta	i net weight			1.23. 10tal gross we	igitt	
	I.28. Description of	-									
	 35 ALBUMINOIDAL SUBSTANCES; MODIFIED STARCHES; GLUES; ENZYMES 3502 Albumins (including concentrates of two or more whey proteins, containing by weight more than 80 % whey proteins, calculated on the dry matter), albuminates and other albumin derivatives 										
	matter), albumins	oncent	rates of two bumin deriv	or more v vatives	vhey proteins, o	containing by w	eight more	e than 80 % whey p	roteins, calcula	ted on the dry	
	Commodity Package count Species							Net weigł	ıt	Manufacturing	g plant
	Batch number										

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	II. Health info	rmation								
Part II: Certification										
	I, the under	I, the undersigned Official veterinarian hereby certify that:								
	1.									
		a)	under the control of the official ver	terinary service,						
		b)	which were in a 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,							
		c)	belonging to holdings which were rinderpest	e not under restrictions due to foot-and-mouth disease or						
		d)	subject to regular veterinary inspections to ensure that they satisfy the animal health requirements of the EU, and							
		e)	which were fed with feed produced in compliance with Regulation (EC) No 1829/2003 on genetically modified food and feed.							
	2.	It has undergone pasteurisation or been produced from raw milk which has been submitted to a pasteurisation treatment fulfilling the provisions of Regulation (EC) No 853/2004 laying down specific rules or requirements for heat treatment.								
	3.	It was man	ufactured from raw milk:							
		a)	which comes from holdings registered in accordance with Regulation (EC) No 852/2004 and checked in accordance with Article 49 and Article 50 of Regulation (EU) 2019/627,							
		b)	which was produced, collected, cooled, stored and transported in accordance with the hygiene conditions laid down in Annex III to Regulation (EC) No 853/2004,							
		c)	which meets the plate and somatic cell count criteria laid down in Annex III to Regulation (EC) No 853/2004,							
		d)	which, pursuant to testing for residues of antibacterial drugs carried out by the food business operator in accordance with the requirements of the Regulation (EC) No 853/2004, it complies with the maximum residue limits for residues of antibacterial veterinary medicinal products laid down in Commission Regulation (EU) No 37/2010,							
		e)	which has been produced under conditions guaranteeing compliance with the maxim residue levels for pesticides in accordance with the requirements of the EU, and							
		f)	which has been produced under co Directive 96/22/EC concerning the j substances having a hormonal or t	prohibition on the use in stoc	k farming of certain					
	4.	It comes from an establishment implementing a programme based on the HACCP princip accordance with Regulation (EC) No 852/2004.								
	5.		been processed, stored, wrapped, packaged and transported in accordance with the relevant e requirements of the EU.							
	6.	It meets the	he relevant microbiological criteria of Commission Regulation (EC) No 2073/2005.							
	7.		antees covering live animals and products thereof provided by the residue plans submitted in nce with the requirements of Regulation (EU) 2017/625 are fulfilled.							
	Notes:									
	Part I:									
	Box I.11: The approval number of the entity should be indicated, if applicable.									
	Box I.12: The approval number of the entity should be indicated, if applicable.									
	Box I.19: Either seal- or container number or both are to be indicated in this box.									
	Box I.28: "CN code": use the appropriate Harmonized System (HS) code of the World Customs Organisation:04.01; 04.02; 04.03; 04.04; 04.05; 04.06; 15.17; 17.02; 19.01; 21.05; 21.06; 22.02; 28.35; 35.01; 35.02 or 35.04.									
	Signature and stamp must be different color that in the printed certificate.									
	The certificate must be provided in at least the English language									
	Certifying Officer									

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II. Health information		
Name (in capital letters) Date of signature Stamp	Qualification and title Signature	