Export Health Certificate

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	I.1. Consignor					I.2. IMSOC Reference				
						I.2.a. Local Reference				
	Name					1.2.a. Local Reference				
	Address									
	Country ISO Code									
	100 code				<u> </u>					
	I.5. Consignee					I 3 Central competent authority				
	_					I.3. Central competent authority				
坦	Name	Name				I.4. Local competent authority				
Part I: Details of consignment	Address									
اغ	Country		ISO Cod	io.						
딤	Country		150 000	ie						
. <u>p</u> o										
S	I.7. Country of origin			I	ISO Code	I.9. Country of destination ISO Code			ISO Code	
o										
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j	I.8. Region of origin Code				I.10. Region of	destinatio	n			
S	I.11. Place of Dispatch					I.12. Place of destination				
ij	1.11. Place of Dispatch				1.12. Place of destination					
ta	Name	Jame				Name				
<u>e</u>	Address					Address				
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Н	Approval Number					Approval Nur	nber			
ピ	Country		ISO	Code		Country			ISO Code	
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щ	I.13. Place of Loading					I.14. Date and	time of de	parture		
	_							F		
	Name									
	Address									
	Approval Number									
	Country		ISO	Code						
	I.15. Means of Transp	ort				I.16 Entry Poir	nt			
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		ternational	Identification	on						
		ansport								
	αο	cument				-				
						-				
	I.18. Transport condit	ione				I.17. Accompanying documents				
				,		1.17. Accompa	nymg doct	intents		
	Chilled Cor	ntrolled	Ambient \Box	Froz	zen 🗆	Commercial				
	ten	nperature \square				document		Date o	f issue	
	reference									
							Country Place of			
						Country		Place o	of	
						Country		Place o issue	of	
	I 10 Container No / Se	nal No				Country			of	
	I.19. Container No / Se	eal No				Country			of	
		eal No				Country			of	
	I.19. Container No / Se	eal No				Country			of	
	I.20. Certified as					Country			of	
						Country			of	
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	I.20. Certified as Human consumption		try ISO Code				it through	issue		
	I.20. Certified as Human consumption I.21. For transit through Country EU Exit		ISO Code				it through	issue		
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EUROPEAN UNION

EU	ROPEAN U	JNION				consumpti	on from the EU to Georgia		
	II. Health information								
	I, the undersigned Official veterinarian hereby certify that:								
	1.	The dairy product described above, which is exported to Georgia, has been obtained from animals:							
on	a)	under the control of the official veterinary service,							
	b)	belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpoand							
	c)	subject to regular veterinary inspections to ensure that they satisfy the animal health requirements of the EU.							
	d)	which were fed with feed produced in compliance with Regulation (EC) No 1829/2003 on genetically modified food and feed.							
		o (1)either	2.	It was made from raw milk sourced from confiderable of the species Camelus dromedarius, and he			-		
			o (1)either	(i)	value equal to or greater				
			o (1)or	(ii)		gh temperature (UHT) treatment on with a suitable holding tim			
			○ (1)or	(iii)	°C for 15 se than 7.0 ac	perature-short time pasteuris conds applied twice to milk w hieving, where applicable, a r tosphatase test, applied imme	vith a pH equal to or greater negative reaction to an		
			o (1)or	(iv)	achieving,	t with an equivalent pasteuris where applicable, a negative i se test, applied immediately a	reaction to an alkaline		
			o (1)or	(v)	a HTST trea	atment of milk with a pH belo	w 7.0;		
				(vi)		atment combined with anothe			
				o (1)either	(1)	a sterilisation process, to ach greater than three;	ieve an F0 value equal to or		
				o (1)or	(2)	additional heating equal to o combined with desiccation.	r greater than 72 °C,		
		o (1)or	2.			milk sourced from animals ot the species Camelus dromeda			
			o (1)either	(i)	a sterilisati than three	on process, to achieve an F0 v	alue equal to or greater		
			o (1)or	(ii)		gh temperature (UHT) treatmon with a suitable holding tim			
	3.	It was manufactured from raw milk:							
 a) which comes from holdings registered in accordance with Regulation (EC) No 852/2004 and check accordance with Article 49 and Article 50 to 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; 						No 852/2004 and checked in			
						ce with the hygiene			
						I to Regulation (EC) No			
	d) which, pursuant to testing for residues of antibacterial drugs carried out by the food business operaring accordance with the requirements of the Regulation (EC) No 853/2004, it complies with the maximal residue limits for residues of antibacterial veterinary medicinal products laid down in Commission Regulation (EU) No 37/2010;						complies with the maximum		
	e)					taranteeing compliance with the nents of the EU;	the maximum residue levels		
f) which has been produced under conditions guaranteeing compliance with the Cour 96/22/EC concerning the prohibition on the use in stock farming of certain substance hormonal or thyrostatic action and of beta-agonists.									

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EUROPEAN UNION

	II Health infor	emation										
	II. Health infor	manon										
	4.	It comes from an establishment implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004.										
	5.	It has been processed, stored, wrapped, packaged and transported in accordance with the relevant hygiene requirements of the EU.										
_	6.	It meets the relevant microbiological criteria o	f the EU Regulation.									
Part II: Certification	7.	The guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with the requirements of the Regulation (EU) 2017/625 are fulfilled.										
Ιij	Notes:											
ပ္ပ	Part I:	Part I:										
tΪ	Box I.11: Th	Box I.11: The approval number of the entity should be indicated, if applicable.										
Par	Box I.12: Th	ne approval number of the entity should be ind	icated, if applicable.									
	Box I.19: Ei	ther seal- or container number or both are to b	e indicated in this box.									
	04.02; 04.03	N code": use the appropriate Harmonized Systos; 04.04; 04.05; 04.06; 15.17; 17.02; 19.01; 21.05; 2										
	Part II:											
	(1)Delete as	s appropriate										
	Signature a	and stamp must be different color that in the pr	inted certificate.									
		ate must be provided in at least the English lan	guage.									
	Certifying Offi		0. 115 11									
	Name (in cap Date of signat Stamp		Qualification and title Signature									

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