Export Health Certificate

	I.1. Consignor					I.2. IMSOC Reference				
	Name					I.2.a. Local Reference				
	Name Address				1.2.a. Local Reference					
	Address Country ISO Code									
	Country		130 CC	Jue						
	I.5. Consignee				I.3. Central competent authority					
ţ	Name				I.4. Local competent authority					
uə	Address					•				
ŭ	Country ISO Code									
g	- Country - Coun									
ß	I.7. Country of origin ISO Code				I.9. Country of	destination	n		ISO Code	
Part I : Details of consignment										
J(I.8. Region of origin Code				I.10. Region of	destinatio	n			
S	I.11. Place of Dispatch				I.12. Place of destination					
ail										
et	Name					Name				
Q:	Address					Address	,			
Ι	Approval Number		100	20.1.		Approval Nu	mper		100 0 1	
art	Country		150	O Code		Country			ISO Code	
P	I.13. Place of Loading	g				I.14. Date and	time of de	parture		
	Name	0				In In Batto and	arno or ac	purture		
	Address									
	Approval Number		***							
	Country		ISC	O Code						
	I.15. Means of Transp	nort				I.16 Entry Poi	nt			
	1	nternation	al Identificat	tion		1				
	ll tr	ransport	lai lueitillicat	11011						
	de	locument				-				
						-				
						-				
						-				
	I.18. Transport condi		_	_	_	I.17. Accompanying documents				
	Chilled Controlled Ambient Frozen				Commercial					
	temperature				1 .					
	te	emperatur	е Ш			document reference		Date o	of issue	
	te	emperatur	е Ш			reference		Date o		
	te	emperatur	e 🗀							
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	I.19. Container No / S		е Ц			reference		Place		
	I.19. Container No / S I.20. Certified as	Seal No	e 🗀			reference		Place		
	I.19. Container No / S	Seal No	e 🗀			reference		Place		
	I.19. Container No / S I.20. Certified as Human consumption	Seal No				reference Country	it through	Place issue	of	
	I.19. Container No / S I.20. Certified as Human consumption I.21. For transit throu	Seal No	d country			reference Country	it through	Place		
	I.19. Container No / S I.20. Certified as Human consumption I.21. For transit throu	Seal No	d country			reference Country	it through	Place issue	of	
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	I.19. Container No / S I.20. Certified as Human consumption I.21. For transit throu Country EU Exit Authority EU Entry	Seal No	d country ISO Code BCP code			reference Country I.22. For trans	it through	Place issue	of	
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	I.19. Container No / S I.20. Certified as Human consumption I.21. For transit throu Country EU Exit Authority EU Entry Authority I.23. Total number of	Seal No n ugh a third	d country ISO Code BCP code BCP code		l net weight	reference Country I.22. For trans	it through	Place issue Member State(s)	of	
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EUROPEAN UNION

EUROPEAN UNION consumption from the EU to Ge							on from the EU to Georgia		
	II. Health information								
	I, the unde	rsigned Off	icial veterin	arian hereb	y certify tha	t:			
	1.	The dairy product described above, which is exported to Georgia, has been obtained from animals:							
	a)	under the control of the official veterinary service,							
ification	b)	belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest, and							
	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.							
		(1)either of th			nade from raw milk sourced from cows, ewes, goats, buffaloes or, camels pecies Camelus dromedarius, and has undergone:				
			o (1)either	(i)	a sterilisation process, to achieve an F0 value equal to or greater than three;				
			o (1)or	(ii)		gh temperature (UHT) treatmon with a suitable holding time			
			○ (1)or	(iii)	°C for 15 se than 7.0 ac	perature-short time pasteurism conds applied twice to milk w hieving, where applicable, a r cosphatase 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		
				atment of milk with a pH belo	w 7.0;				
				(vi)	a HTST trea	atment combined with anothe	er physical treatment by:		
				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	value equal to or greater		
			o (1)or	(ii)		gh temperature (UHT) treatmon on with a suitable holding time			
	3.	It was manufactured from raw milk:							
	a)	accordance with Article 49 and Article 50 to Regulation (EU) 2019/627;							
	b)								
	c)						I to Regulation (EC) No		
d) which, pursuant to testing for residues of antibacterial drugs carried out by in accordance with the requirements of the Regulation (EC) No 853/2004, it residue limits for residues of antibacterial veterinary medicinal products la Regulation (EU) No 37/2010;					complies with the maximum				
	e)	which has been produced under conditions guaranteeing compliance with the maximum residue le for pesticides in accordance with the requirements of the EU;					the maximum residue levels		
f) which has been produced under conditions guaranteeing compliance with the Council Directi 96/22/EC concerning the prohibition on the use in stock farming of certain substances having hormonal or thyrostatic action and of beta-agonists.									

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EUROPEAN UNION

			<u> </u>	U						
	II. Health infor	rmation								
	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.								
rtif	Notes:									
: Ce	Part I:	Part I:								
tΠ	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	CN code": use the appropriate Harmonized Systems; 04.04; 04.05; 04.06; 15.17; 17.02; 19.01; 21.05; 2								
	Part II:									
		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 Offic		Ovalification and title							
	Name (in cap Date of signat Stamp		Qualification and title Signature							

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