**Export Health Certificate** 

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	I.1. Consignor					I.2. IMSOC Reference			
	Name					I.2.a. Local Reference			
	Address								
	Country ISO Code								
	LE Consigned					I 2 Control competent outh			
	I.5. Consignee				I.3. Central competent authority				
ıt	Name					I.4. Local competent authority			
er	Address								
Part I : Details of consignment	Country ISO Code								
g		<u> </u>							
si	I.7. Country of origin ISO Code					I.9. Country of destination ISO Code		ISO Code	
on									
C									
of	I.8. Region of origin Code					I.10. Region of destination			
ls	I.11. Place of Dispatch					I.12. Place of destination			
ai	Name						Name		
et	Address								
D	Address					Address			
Ι:	Approval Number	r				Approval Number			
rt	Country		ISO	Code		Country		ISO Code	
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I	I.13. Place of Load	ing				I.14. Date and time of de	parture		
	Name								
	Address								
	Approval Number	r							
	Country		ISO	Code					
	7.47.76					7.40 T			
	I.15. Means of Trai	nsport				I.16 Entry Point			
	Mode	Internationa	al Identification	on					
		transport							
		document				-			
	I.18. Transport cor	nditions	·			I.17. Accompanying doc	iments		
			Ambient 🗆	Chilled			uments		
	Frozen 🗀	Controlled	Ambient L	1 Chilled	ш	Commercial	Data	ficeno	
		temperature $\square$				document Date of issue			
						l reference			
						reference	Place	nf	
						reference Country	Place o	of	
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	I.19. Container No	/ Seal No						of	
		/ Seal No						of	
	I.20. Certified as							of	
								of	
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## **EUROPEAN UNION**

EUROPEAN UNION consumption from the EU to								
	II. Health info	ormation						
	I, the unde	rsigned Off	icial veterinarian hereby certify that	:				
	1.	The dairy	product described above, which is ex	xported to Georgia, has been	obtained from animals:			
Part II: Certification		a)	under the control of the official vet	erinary service,				
		b)	which were in a country or part thereof that has been free of foot-and-mouth disease and 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 not under restrictions due to foot-and-mouth disease crinderpest					
		d)	subject to regular veterinary inspect requirements of the EU, and	ctions to ensure that they sati	isfy the animal health			
		e)	which were fed with feed produced genetically modified food and feed.		ion (EC) No 1829/2003 on			
	2.	pasteurisa	t 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 mar	nufactured from raw milk:					
		a)	which comes from holdings registe checked in accordance with Article	_				
		b)	which was produced, collected, coo hygiene conditions laid down in Ar					
		c)	which meets the plate and somatic (EC) No 853/2004,	cell count criteria laid down	in Annex III to Regulation			
		d)	which, pursuant to testing for reside business operator in accordance with the maximum resimedicinal products laid down in Co	ith the requirements of the R due limits for residues of ant	egulation (EC) No 853/2004, ibacterial veterinary			
		e)	which has been produced under coresidue levels for pesticides in accoresidue.					
		f)	which has been produced under co Directive 96/22/EC concerning the p substances having a hormonal or the	prohibition on the use in stoc	k farming of certain			
	4.	It comes from an establishment implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004.						
	5.		as been processed, stored, wrapped, packaged and transported in accordance with the relevant giene requirements of the EU.					
	6.	It meets th	ne relevant microbiological criteria of Commission Regulation (EC) No 2073/2005.					
	7.		ntees covering live animals and proc e with the requirements of Regulatio					
	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 r	nust be different color that in the pri	nted certificate.				
	The certifi							
Certifying Officer								

## **EUROPEAN UNION**

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