# CHAPTER 2(A)

## Health certificate

# For milk, milk-based products and milk-derived products not intended for human consumption for dispatch to or transit through(<sup>2</sup>) the European Union

#### COUNTRY: UNITED STATES

Veterinary certificate to EU

		2			
	I.1. Consignor Name Address	I.2. Certificate reference No I.2.a.			
		I.3. Central competent authority APHIS-VS			
	Tel.	I.4. Local competent authority			
	I.5. Consignee Name Address	I.6. Person responsible for the load in EU Name Address			
	Post code Tel.	Post code Tel.			
gnment					
ed consi	I.7. Country of ISO code I.8. Region of Code origin origin	I.9. Country of ISO I.10. Region of Code destination code destination			
ch	I.11. Place of origin	I.12. Place of destination			
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse       Name       Address			
I : Detai		Postal code			
Part					
	Name Approval number Address				
	Name Approval numb <del>er</del> Address				
	I.13. Place of loading	I.14. Date of departure			
	I.15. Means of transport	I.16. Entry BIP in EU			
	Aeroplane Ship Railway wagon				
	Road vehicle  Other  Identification	1.17.			
	Documentation references				
	I.18. Description of commodity	I.19. Commodity code (HS code)			
		I.20. Quantity			
	I.21. Temperature of product	I.22. Number of packages			
	Ambient Chilled Frozen				
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#### COUNTRY: UNITED STATES

### Milk, milk-based products and milk-derived products not for human consumption

		1.2.	Certificate reference No		l.2.a.
I.23. Seal/Container No		1.24	Type of packaging		
			. The or hearing		
I.25. Commodities certified for: Animal feedingstuff  ☐ Further proces Technical use  ☐					
I.26. For transit through EU to third co	untry	1.27.	For import or admission into E	U	
Third country	ISO code				
I.28. Identification of the commodities					
Species (Scientific name)	Approval number of establish Manufacturing plant	iments	Net weight	Batch number	

COUNTRY: UNITED STATES Milk, milk-based products and milk-derived products not for human consumption

	П.	Health information		II.a. Certificate reference No	II.b.	
		No 1069/2009 of the and Commission R Annex X, and Chapt	European Parliament ar egulation (EU) No 142/2 er I of Annex XIV thereto	are that I have read and understood Regund of the Council( <sup>1a</sup> ), and in particular Article 2011( <sup>1b</sup> ), and in particular Section 4 of Ch, and certify that the milk( <sup>2</sup> ), the milk-based pox I.28 comply with the following conditions:	10 thereof, apter II of products( <sup>2</sup> )	
Part II: Certification	II.1.	they were produced and derived in <i>(insert name of exporting country)</i> ( <sup>3</sup> ), <i>(insert name of region)</i> ( <sup>3</sup> ), which is listed in Part I of Annex II to Commission Regulation (EU) No 605/2010( <sup>4</sup> ), and which has been free from foot-and-mouth disease (FMD) and rinderpest for a period of 12 months immediately prior to export and has not practised vaccination against rinderpest during that period;				
Part II:	II.2.	they were produced clinical signs of any kept for a period of a	hey were produced from raw milk derived from animals which at the time of milking did not show linical signs of any disease transmissible through milk to humans or animals, and which had been ept for a period of at least 30 days prior to production on holdings that were not subject to official estrictions due to foot-and-mouth disease or rinderpest;			
	II.3.		rgone one of the treatme	ents or combinations thereof described in po		
			hey was collected from	als of species susceptible to foot-and-mout milk subjected to one of the treatments de		
		•		at least 16 hours after clotting and has a pH	below 6;]	
( <sup>2</sup> )( <sup>5</sup> )or [the whey has been produced at least 21 days before the shippi that period no cases of FMD have been detected in the exporting			luced at least 21 days before the shipping MD have been detected in the exporting co	and_during untry;]		
( <sup>2</sup> )( <sup>5</sup> )or [the whey has been produced on this date, in consideration of the foreseen voyage duration, being at days before the consignment is presented to a border inspection po European Union;]]			at least 21			
	II.4.	• •	ect to one of the followin	-		
		equivalent		teurisation at 72°C for at least 15 secor - a negative reaction to a phosphatase tes		
( <sup>2</sup> )either [a subsequent second high temperature short time pasteurisation at least 15 seconds or an equivalent pasteurisation which itself a negative reaction to a phosphatase test in bovine milk;]		equivalent pasteurisation which itself a				
	( <sup>2</sup> )or [a subsequent drying process that in the case of milk intended for fe combined with additional heating to 72°C or higher;]		-			
		ł	nour at a level below 6;]	ey which the pH is reduced and kept for a		
	(2)(5) or [the condition that the milk/milk product has been produced at least prior to the date of shipping and during that period no cases of FMD ha detected in the exporting country;]				st 21 days have been	
		<del>(²)(⁵)or [</del> - € ŧ	the milk/milk consideration of the fore the date that the consigr European Union;]	product has been produce (insert the date), this seen voyage duration, being at least 21 da nment is presented to a border inspection	<del>s date, in</del> <del>iys prior to</del>	
			sterilisation at a level of		on with	
( <sup>2</sup> )either [a subsequent dryin		a subsequent drying pro	: 132°C for at least one second in combinati pcess that in the case of milk intended for beating to 72°C or higher:1			
	combined with additional heating to 72°C or higher;] ( <sup>2</sup> )or [a subsequent process by which the pH is reduced and kept for at lea hour at a level below 6;]		t least one			

Ithe condition that the milk/milk product has been produced at least 21 days (<sup>2</sup>)(<sup>5</sup>)or prior to the date of shipping and during that period no cases of FMD has been detected in the exporting country:] (2)(5)or milk/milk product [the has been produced on ..... (insert the date), this date, in consideration of the foreseen voyage duration, being at least 21 days prior to the date that the consignment is presented to a border inspection post of the European Union;]] 11.5. every precaution was taken to avoid contamination of the milk/milk-based product/milk-derived product after processing; II.6. the milk/milk-based product/milk-derived product was packed: <sup>(2</sup>)either [in new containers;] (2)or fin vehicles or bulk containers disinfected prior to loading using a product approved by the competent authority;] the containers are marked so as to indicate the nature of the milk/milk-based and product/milk-derived product and bear labels indicating that the product is Category 3 material and not intended for human consumption; 11.7. the milk, milk-based products and milk-derived products described above: (2)either [does not contain milk or milk products of ovine or caprine animal origin or is not intended for feed for farmed animals, other than fur animals.] fcontains milk or milk products of ovine or caprine animal origin and is intended for <del>(²)or</del> feed for farmed animals, other than fur animals, and the milk or milk products: <del>(a)</del> are derived from ovine and caprine animals which have been kept continuously since birth in a country where the following conditions are fulfilled: classical scrapie is compulsorily notifiable; (i) an awareness, surveillance and monitoring system is in place for <del>(ii)</del> classical scrapie: official restrictions apply to holdings of ovine or caprine animals in (;;;) the case of a suspicion of TSE or the confirmation of classical scrapie; ovine and caprine animals affected with classical scrapie are killed and destroyed: the feeding to ovine and caprine animals of meat-and-bone meal or greaves, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health (OIE), of ruminant origin has been banned and effectively enforced in the whole country for a period of at least the preceding seven years; originate from holdings where no official restrictions are imposed due to a suspicion of TSE; originate from holdings where no case of classical scrapie has been (c)diagnosed during a period of at least the preceding seven years or, following the confirmation of a case of classical scrapie: (2)either [all ovine and caprine animals on the holding have been killed and destroyed or slaughtered, except for breeding rams of the ARR/ARR genotype, breeding ewes carrying at least one ARR allele and no VRQ allele and other ovine animals carrying at least one ARR allele;] [all animals in which classical scrapie was confirmed have been killed (<sup>2</sup>)or and destroyed, and the holding has been subjected for a period of at least two years since the date of confirmation of the last classical scrapie case to intensified TSE monitoring, including testing with

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II.a. Certificate reference No

negative results for the presence of TSE in accordance with the laboratory methods set out in point 3.2 of Chapter C of Annex X to

Πb

П.

Health information

11.	Health information	II.a. Certificate reference No II.b.			
		gulation (EC) No 999/2001( <sup>6</sup> ), of all of the following animals whi			
		• over the age of 18 months, except ovine animals of the ARR/AF notype:			
	9 <del>0</del>	animals which have been slaughtered for hum			
consumption; and					
	_	animals which have died or been killed on the holding to which were not killed in the framework of a disea eradication campaign.]]			
Notes					
Part I					
	be filled in only if it is a certificat	nsible for the load in the European Union: this box is required e for a commodity to be transited through the European union s for a commodity to be imported into the European Union.			
_	<ul> <li>Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transi commodity.</li> </ul>				
_	<ul> <li>Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In the case of unloading and reloading, the consignor must inform the border inspection post of the European Union.</li> </ul>				
_		propriate Harmonised System (HS) code of the World Custor 3; 04.04; 23.09.10, 23.09.90, 35.01, 35.02 or 35.04.			
_	must be included.	ainers, the container number and the seal number (if applicab			
_	animals, and the production or n				
_		according to whether it is a transit or an import certificate. uring plant': provide the registration number of treatment			
Part II	:				
• •	OJ L 300, 14.11.2009, p. 1.				
• •	OJ L 54, 26.2.2011, p. 1.				
• •	Delete as appropriate. For completion if the authorisation to import into or transit through the European Union is restricted to				
(	certain regions of the third country concerned.				
• •	OJ L 175, 10.7.2010, p. 1.				
	this condition applies only to third countries listed in column 'A' of Annex I to Regulation (EU) No 605/2010.				
	OJ L 147, 31.5.2001, p. 1.				
_		st be in a different colour to that of the printing.			
		or the consignment in the European Union: This certificate is or accompany the consignment until it reaches the border inspecti			

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II. Health information	II.a. Certificate reference No	II.b.
Official veterinarian/Official inspector		
Name (in capital letters):	Qualification and title:	
Date:	Signature:	
	Stamp:	
		/

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