#### **CHAPTER 2**

#### Health certificate

For milk, mill	k products,	colostrum	and o	colostrum	products	not i	ntended	for human
consun	ption for a	dispatch to	or tre	ansit throu	$\frac{1}{1} (2)$ the	Euro	pean Ur	<i>iion</i>

CO	UNTRY:	Veterinary certificate to EU				
	I.1. Consignor Name	I.2. Certificate reference number I.2.a				
ent	Address	I.3. Central Competent Authority APHIS-VS				
mm	Tel.N°	I.4. Local Competent Authority				
nsig	I.5. Consignee	I.6. Person responsible for the load in EU				
d coi	Name	Name				
cheo	Address Postal code	Address				
pate	Tel.N°	Postal code Fet.N°				
dis	1.7.Country of origin US US US-0	I.9. Country of destination ISO code I.10. Region of destination Code				
s of	I.11. Place of origin	I.12. Place of destination				
Part I : Details of dispatched consignment	Name Approval number	Custom warehouse				
<b>D</b>	Address	Name Approval number				
rt I		Address				
Pa		Postal code				
	I.13. Place of loading	I.14. Date of departure				
	1.15. Means of transport	I.16. Entry BIP in EU				
	Aeroplane Ship Railway wagon Road vehicle Other					
	Identification:	I.17. No.(s) of CITES				
	Documentary references: 1.18. Description of commodity	I.19. Commodity code (HS code)				
		L20.Quantity				
	I.21 Temperature of product	1.00 Number of a strange				
	Ambient Chilled	I.22. Number of packages				
	I.23. Identification of container/Seal number	I.24. Type of packaging				
	I.25. Commodities certified for:	•				
	Animal feedingstuff	Technical use D Other				
	I.26. For transit through EU to 3rd Country	1.27. For import or admission into EU				
	3rd country ISO code					
	1.28. Identification of the commodities	I				
	1 11	er of establishments Net weight Batch number uring plant				

COUNTRY

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# Milk, milk products, colostrum and colostrum products not for human consumption

II.	Health information	II.a. Certificate reference number II.b.				
	I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) N $1774/2002^{(1)}$ , and in particular Article 6 and Chapter V of Annex VII thereto, and certify that the milk <sup>(2)</sup> , the milk products <sup>(2)</sup> the colostrum <sup>(2)</sup> or the colostrum products <sup>(2)</sup> referred to in box I.28 comply with the following conditions:					
II.1.	they were produced and derived in the <u>United States</u> ( <i>insert name of exporting country</i> ) <sup>(3)</sup> , ( <i>insert name of region</i> ) <sup>(3)</sup> , which is listed in the Annex to Decision 2004/438/EC, and which has from foot-and-mouth disease (FMD) and rinderpest for 12 months immediately prior to export an practiced vaccination against rinderpest during that period;					
II.2.	they were produced from raw milk or colostrum derived from animals which at the time of milking did show clinical signs of any disease transmissible through milk or colostrum to humans or animals, and wh had been kept for at least 30 days prior to production on holdings that were not subject to official restrict due to foot-and-mouth disease or rinderpest;					
II.3.	they are milk or milk produc					
		eatments or combinations thereof described in point II. 4]				
<sup>(2)</sup> or	collected from milk subjecte	animals of species susceptible to foot-and-mouth disease, and that whey we do one of the treatments described in point II. 4 and				
		ollected at least 16 hours after clotting and has a pH below 6;]				
	of FMD have b	een produced at least 21 days before the shipping and during that period no cas een detected in the exporting country;]				
		been produced on//, this date, in consideration of the foreseen voya at least 21 days before the consignment is presented to a Border Inspection Po- Union;]]				
II.4.	they have been subject to on	of the following treatments:				
<sup>(2)</sup> either		ne pasteurisation at 72°C for at least 15 seconds, or an equivalent pasteurisati to a phosphatase test in bovine milk, in combination with:				
	15 seconds or	second High Temperature Short Time pasteurisation at 72°C for at lea an equivalent pasteurisation which itself achieves a negative reaction to t in bovine milk;]				
	<sup>(2)</sup> or [a subsequent-	lrying process that in the case of milk intended for feeding is combined with a second to the second s				
		rocess by which the pH is reduced and kept for at least one hour at a level bek				
	<sup>(2)(4)</sup> or [the condition t	hat the milk/milk product has been produced at least 21 days before the shippi period no cases of FMD have been detected in the exporting country;]				
	<sup>(2)(4)</sup> or [the milk/milk- voyage-duration Inspection Post	product has been produced on//, this date, in consideration of the forese n, being at least 21 days before the consignment is presented to a Borc of the European Union;]				
	<sup>(2)</sup> or [sterilisation at	a level of at least F <sub>0</sub> 3;]]				
<sup>(2)</sup> or		tment at 132°C for at least one second in combination with:				
	additional heat	rying process that in the case of milk intended for feeding is combined with a second to 72°C or higher,]				
	<del>6;]</del>	rocess by which the pH is reduced and kept for at least one hour at a level belo				
		nat the milk/milk product has been produced at least 21 days before the shippi period no cases of FMD has been detected in the exporting country;]				
		period no cases of 1 MiD has been detected in the exporting country,				
	voyage duration	n, being at least 21 days before the consignment is presented to a Bord of the European Union;]]				
II.5.	they are colostrum or colostr	im products of bovine animals that have been subject to High Temperature Sho for at least 15 seconds, or an equivalent pasteurisation achieving a negati				

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# Milk, milk products, colostrum and colostrum products not for human consumption

II.	Health in	formation	II.a.	Certificate reference nu	imber II.b.		
	reaction to	a phosphatase test in	bovine milk. in o	combination with:			
				or colostrum products have	been produced at least 21 d	dave	
	enner	before the shipping	g and in this pe	riod no cases of FMD have	been detected in the export	<del>ting</del> :	
	<sup>(2)(4)</sup> 0r	of the foreseen voy	rage duration, be	ts have been produced on/ wing at least 21 days before the bean Union:1			
and	Border Inspection Post of the European Union;] have been obtained from animals subject to regular veterinary inspections to ensure that animals come fror holdings on which all bovine herds are:						
	<sup>(2)(4)</sup> either		cially tuberculos	is and brucellosis free <sup>(5)</sup> ,]			
			er the national le	egislation of the third country	r of origin regarding eradica	tior	
and	(2)(4) <u>either</u>			vine-leukosis free <sup>(5)</sup> ;]			
	(2)(4)			the control of enzootic boving	e leukosis and there has beer	n nc	
				boratory testing of this diseas			
II.6.	every prec after proce		avoid contamina	ation of the milk/milk produ	ict/colostrum/colostrum proc	duct	
II.7.	-	ilk product/colostrum	/colostrum prod	uct was packed:			
	<sup>(2)</sup> either	[in new containers,	·	1			
	<sup>(2)</sup> <i>or</i>	L ,	lk containers dis	sinfected prior to loading us	ing a product approved by	-the	
and	the containers are marked so as to indicate the nature of the milk/milk product/colostrum and bear labels indicating that the product is Category 3 material and not intended for human consumption.						
Notes Part I:	•						
•			ible for the load	in EU: this box is to be fille	d in only if it is a certificate	e fo	
•	Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity.						
•	Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the Border Inspection Post of the European Union.						
•	Box reference I.19: Use the appropriate Harmonised System (HS) code of the World Customs Organisation 23.09.10, 23.09.90, 35.01, 35.02 or 35.04.						
•	Box reference I.23: For bulk containers, the container number and the seal number (if applicable) must b included.					t b	
•	Box reference I.26 and I.27: Fill in according to whether it is a transit or an import certificate.						
•	Box reference 1.28: "Manufacturing plant": provide the registration number of treatment or processing establishment.						
Part I							
(1)		).10.2002. n. 1					
(2)	OJ L 273, 10.10.2002, p. 1. Delete as appropriate.						
(3)	For completion if the authorisation to import into the European Union is restricted to certain regions of the third country concerned.						
(4)	-		d countries lister	l in column "A" of Annex I to	Decision 2004/438/FC		
(5)				as laid down in Annex A to		EC	

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# Milk, milk products, colostrum and colostrum products not for human consumption

II.	Health information	II.a.	Certificate reference number	II.b.				
	– The signature and the seal must be	in a differen	nt colour from that of the printing.					
	<ul> <li>Note for the importer: this certificate is only for veterinary purposes and must accompany the consignment until it reaches the Border Inspection Post of the European Union.</li> </ul>							
Offic	Official veterinarian							
Name (in capital letters):		Qualification and title:						
	Date:		Signature:					
	Stamp:							