					_		
	I.1. Consignor			I.2. IMSOC Reference			
	Name Address			I.2.a. Local Reference			
	Country		ISO Code				
	country		100 couc				
	I.5. Consignee				I.3. Central competent authority		
님	Name				I.4. Local competent authority		
ne	Address						
빏	Country		ISO Code				
S	I.7. Country of ori	gin	ISO Co	I.9. Country of destination		ISO Code	
읽	, ,	5					
ы	I.8. Region of origi	in	Code		I.10. Region of destination		
Part I : Details of consignment	I.11. Place of Dispa	atch			I.12. Place of destination		
<u>[a</u> ]	Name				Name		
al	Address				Address		
ï	Approval Numbe	r			Approval Number		
턻	Country		ISO Code		Country	ISO Code	
പ്	I.13. Place of Load	ling			I.14. Date and time of departure		
	Name						
	Address						
	Approval Numbe	r					
	Country		ISO Code				
	I.15. Means of Tra	nsport			I.16 Entry Point		
	Mode	International	Identification				
		transport document					
					-		
					-		
					-		
		,					
	I.18. Transport co	nditions Controlled	_	_	I.17. Accompanying documents Accompanying document		
	Frozen 🗆	temperature	Ambient Chille	dЦ	reference		
		I I I I I I I I I I I I I I I I I I I			Date of issue		
					Country		
	I.19. Container No	/ Soal No			Place of issue		
		7 Sear No					
	I.20. Certified as	_			_	_	
	Pharmaceutical u		Other 🗆		Slaughter 🗆	Relaying 🛛	
	Artificial reprodu	ction $\Box$	Fattening 🗆		Production $\Box$	Breeding $\Box$	
	Production of petf	food 🗆	Breeding and productior	ι 🗆	Animal Feedingstuff 🗖	Human consumpt	tion 🗆
	Technical use 🛛						
	104						
	I.21. For transit through a third country				I.22. For transit through Member S		
	Country		ISO Code		Country	ISO Code	
	EU Exit Authority		BCP code				
	EU Entry BC Authority		BCP code				
			-			105 m - 1	• • •
			I.24. Total quantity		I.25. Total net weight	I.25. Total gross w	reight
	I.28. Description o	f consignment					
			MODIFIED STARCHES; GI				
	3502 Albumins	(including concent	rates of two or more whe	y proteins, c	ontaining by weight more than 80 9	% whey proteins, calo	culated on the dry
	#1. Commodity	mates and other al	Quantity		Net weight	Package count	
					Identification system		
	Species Identification number				accontinuation system		

# (GB) Milk, milk-based products and milk-derived products not for human consumption GBHC091E

	II. Health infor	rmation						
	I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 1069/2009 of the European Parliament and of the Council , and in particular Article 10 thereof, and Commission Regulation (EU) No 142/2011 , and in particular Section 4 of Chapter II of Annex X, and Chapter I of Annex XIV thereto, and certify that the milk(2) , the milk-based products (2) and milk-derived products(2) referred to in box I.28 comply with the following conditions:							
Part II: Certification		they were produced and derived in (insert name of exporting country)(3), (insert name of region)(3),which is listed in Part I of Annex II to Commission Regulation (EU) No 605/2010,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:		clinical sign for a period	y were produced from raw milk derived from animals which at the time of milking did not show ical signs of any disease transmissible through milk to humans or animals, and which had been kept a period of at least 30 days prior to production on holdings that were not subject to official rictions due to foot-and-mouth disease or rinderpest;					
	II.3.	they are mi	ilk or milk p	products that:				
	(2)	$\circ$ either	[have unde	rgone one of the treatm	ents or combinations thereof	described in point II.4;]		
	(2)			whey to be fed to animals of species susceptible to foot-and-mouth disease, and was collected from milk subjected to one of the treatments described in point II.4				
		(2)	$\circ$ either	[the whey was collected	l at least 16 hours after clottin	g and has a pH below 6;]		
		(2)(5)			duced at least 21 days before have been detected in the exp			
		(2)(5)		before the consignment	duced on / f the foreseen voyage duratio t is presented to a border cont mel Islands or Isle of Man;]]	<b>u</b>		
	II.4.	they have <b>k</b>	oeen subject	t to one of the following	treatments:			
	(2)		[high temperature short time pasteurisation at 72oC for at least 15 seconds, or an equivale pasteurisation achieving a negative reaction to a phosphatase test in bovine milk, in combination with:					
		(2)			igh temperature short time pa quivalent pasteurisation whic ise test in bovine milk;]			
		(2)			rocess that in the case of milk nal heating to 72oC or higher;]			
		(2)		[a subsequent process b at a level below 6;]	py which the pH is reduced an	d kept for at least one hour		
		(2)(5)			milk/milk product has been pr ping and during that period n ng country;]			
		(2)(5)		in consideration of the the date that the consig	has been produced on//(i foreseen voyage duration, bei nment is presented to a borde , Channel Islands or Isle of Ma	ng at least 21 days prior to er control post of the point of		
		(2)	$\circ$ or	[sterilisation at a level of	of at least F03;]]			
	(2)	$\circ$ or	[ultra high	temperature treatment	at 132oC for at least one secor	nd in combination with:		
		(2)		[a subsequent drying process that in the case of milk intended for feeding is combined with additional heating to 72'C or higher;]				
		(2)		[a subsequent process b at a level below 6;]	by which the pH is reduced an	d kept for at least one hour		
		(2)(5)			milk/milk product has been pr ping and during that period n			

						numan consumption objective		
	II. Health ii	nformation						
				detected ir	ا h the exportin	g country:]		
		(2)(5)	∘ or	[the milk/milk product has been produced on//(insert the date), this date, in consideration of the foreseen voyage the duration, being at least 21 days prior to the date that consignment is presented to a border control post of the point of entry into Great Britain, Channel Islands or Isle of Man;]]				
Certification	II.5.		caution wa		oid contamin	ation of the milk/milk-based product/milk- derived		
tific	II.6.	-	-	•	-derived prod	luct was packed:		
Cer	(2)	$\circ$ either		- containers;]	-	-		
Part II:	(2)	$\circ$ or		les or bulk co nt authority;]		fected prior to loading using a product approved by the		
Р			nd bear la	hiners are marked so as to indicate the nature of the milk/milk-based product/milk-derived bear labels indicating that the product is Category 3 material and not intended for human				
	II.7.	the milk,	milk-based	d products and milk-derived products described above:				
	(2)	∘ either			contain milk or milk products of ovine or caprine animal origin or is not intended or farmed animals, other than fur animals.]			
	-				-	ovine or caprine animal origin and is intended for feed animals, and the milk or milk products:		
			(a)			and caprine animals which have been kept continuously where the following conditions are fulfilled:		
				(i)	classical s	crapie is compulsorily notifiable;		
				(ii)	an awarer classical s	ness, surveillance and monitoring system is in place for crapie;		
				(iii)		strictions apply to holdings of ovine or caprine animals in f a suspicion of TSE or the confirmation of classical		
				(iv)	ovine and and destro	caprine animals affected with classical scrapie are killed oyed;		
				(v)	greaves, a World Org has been l	g to ovine and caprine animals of meat-and-bone meal or s defined in the Terrestrial Animal Health Code of the ganisation for Animal Health (OIE), of ruminant origin banned and effectively enforced in the whole country for f at least the preceding seven years;		
			(b)	originate f suspicion (	-	where no official restrictions are imposed due to a		
						where no case of classical scrapie has been diagnosed st the preceding seven years or, following the f classical scrapie:		
			(2)	○ either	destroyed genotype,	and caprine animals on the holding have been killed and or slaughtered, except for breeding rams of the ARR/ARR breeding ewes carrying at least one ARR allele and no and other ovine animals carrying at least one ARR		
			(2)	° or	killed and period of a last classic testing wit with the la Annex X to	ls in which classical scrapie was confirmed have been destroyed, and the holding has been subjected for a at least two years since the date of confirmation of the cal scrapie case to intensified TSE monitoring, including th negative results for the presence of TSE in accordance aboratory methods set out in point 3.2 of Chapter C of the Regulation (EC) No 999/2001, of all of the following hich are over the age of 18 months, except ovine animals		

	II. Health info	rmation							
	of the ARR/ARR genotype:								
	- animals which have been slaughtered for human consumption; and								
cation	- animals which have died or been killed on the holding but which were not killed in the framework of a disease eradication campaign.]]								
irtifi	Notes	Notes							
Part II: Certification	(*) Those countries subject to the transitional import arrangements include: an EU member State; Liechtenstein; Norway; Iceland and Switzerland.								
Paı	References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018).								
	References	to Great Br	itain in this certificate include Char	nnel Islands and Isle of Man.					
	Part I:								
	-	BoxPerson responsible for the load in Great Britain, Channel Islands or Isle of Man: this box isreferencerequired to be filled in only if it is a certificate for a commodity to be transited throughI.6:Great Britain, Channel Islands or Isle of Man; it may be filled in if the certificate is for a commodity to be imported into the Great Britain, Channel Islands or Isle of Man.							
	-	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 the case of unloading and reloading, the consignor must inform the border control post of the point of entry into Great Britain, Channel Islands or Isle of Man .						
	-	Box reference I.16:	Do not use this box until the end of the transitional staging period.						
	-	Box reference I.19:	use the appropriate Harmonised System (HS) code of the World Customs Organisation: 04.01; 04.02; 04.03; 04.04; 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 be included.						
	-	Box reference I.25:	echnical use: any use other than feeding of farmed animals, other than fur animals, and the oroduction or manufacturing of pet food.						
	-	Box reference I.26 and I.27:	fill in according to whether it is a transit or an import certificate.						
	-	Box reference I.28:	'Manufacturing plant': provide the establishment.	registration number of treat	nent or processing				
	Part II:								
(2) Delete as appropriate.									
	(3) For completion if the authorisation to import into or transit through Great Britain, Channel								
		slands or Isle of Man is restricted to certain regions of the third							
	country concerned.								
	(5) this condition applies only to third countries listed in column 'A' of Annex I to Regulation (EU) No								

	II. Health information		
	605/2010.		
	- The signature and the stamp must be in a diffe	erent colour to that of the prin	ting
	<ul> <li>Note for the person responsible for the consignation</li> </ul>		
	This certificate is only for veterinary purposes	s and must accompany the cor	signment until it reaches
ų	the border control post.		
atio	Certifying Officer		
ific	Name (in capital letters) Date of signature	Qualification and title Signature	
Cert	Stamp		
Part II: Certification			
art			
Ч			