	COUNTRY: Country	Health certificate to Great Britain,							
		port arrangements	(*)			nnel Isl	ands and	Isle of Man	
П	I.1. Consignor			I.2. Cert			I.2.a.		
consignment	Name			reference number					
шe	Address								
臣		I.3. Central Competent Authority							
i g									
13	Plana			I.4. Local Competent Authority					
or	Phone								
ט	I.5. Consignee			I.6.					
Þ	Name								
ъе Те	Address					_			
ପ୍ର									
at	Postal Code								
dispatched	Phone I.7. Country ISO code I.8. Region Code								
1.	I.7. Country of origin	code I.8. Region of origin	Code	I.9. Coun of destin	try ation	ISO code	I.10.		
ם	1	=	ı	01 0000111			_		
of									
O	I.11. Place of origin			1.12.					
Ω	Name	Approval number							
ij	Address	Address							
d d									
Detail				_					
А									
: H	I.13. Place of load:	ing		I.14. Dat	e of dep	arture			
	I.15. Means of trans	sport		I.16. Ent	rv BCP i	n Great Br	itain, Chan	nel Islands	
rt				I.16. Entry BCP in Great Britain, Channel Islands or Isle of Man					
Part	Aeroplane								
щ	Road vehicle	□ Other □		I.17.					
	Identification:								
	Documentary references:								
	I.18. Description of commodity			I.19. Commodity code (HS code)					
	I.21. Temperature if products			I.20. Quantity		I.22. Number of packages			
	Ambient ☐ Chilled ☐ Frozen ☐								
	I.23. Seal/Container No.					I.24. Type of packaging			
	1.23. Seal/Container No.			1.21. Type of packaging			,1119		
	I.25. Commodity certified for:								
	Human Consumption								
		•							
	I.26.	I.27. For import or admission into Great Britain,							
				Channel Islands or Isle of Man			Man 📙		
	I.28. Identification	•							
	Manufacturing	Number of packages	Species		Net we	ight	Batch	Number	
	plant		(scienti	fic name)					
					 				

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COUNTRY: Countries subject to transitional import arrangements (*)

Milk-HTB - Dairy products derived from milk of cows, ewes, goats and buffaloes for human consumption from third countries authorised in column B

II. Health information	II.a. Certificate reference	II.b.
	number	

II.1. Animal Health Attestation

I, the undersigned official veterinarian, declare that I am aware of the relevant provisions of Directive 2002/99/EC and of Regulation (EC) No 853/2004 and hereby certify that the dairy product described above:

- (a) has been obtained from animals:
 - (i) under the control of the official veterinary service,
 - (ii) which were in a country or part thereof that has been free of foot-and-mouth disease and of 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,
 - (iii) belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest, and,
 - (iv) subject to regular veterinary inspections to ensure that they satisfy the animal health conditions laid down in Chapter I of Section IX of Annex 3 to Regulation (EC) No 853/2004 and in Directive 2002/99/EC,
- (b) has undergone or been produced from raw milk which has been submitted to a pasteurisation treatment involving a single heat treatment with a heating effect at least equivalent to that achieved by a pasteurisation process of at least 72°C for 15 seconds and, where applicable, sufficient to ensure a negative reaction to an alkaline phosphatase test applied immediately after the heat treatment.

II.2. Public Health attestation

- I, the undersigned official inspector, declare that I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EU) 2019/627 and hereby certify that the dairy product made with raw milk described above was produced in accordance with those provisions, in particular that:
- (a) it was manufactured from raw milk:
 - (i) which comes from holdings registered in accordance with Regulation (EC) No 852/2004 and checked in accordance with Article 49-50 of Regulation (EU) 2019/,
 - (ii) which was produced, collected, cooled, stored and transported in accordance with the hygiene conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004,
- (iii) which meets the plate and somatic cell count criteria laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004,
- (iv) which complies with the guarantees on the residues status raw milk provided by the monitoring plans for the detection of residues substances submitted in accordance with Council Directive 96/23/EC, and in particular, Article 29 thereof,
- (v) which, pursuant to testing for residues of antibacterial drugs carried out by the food business operator in accordance with the requirements of Annex III, Section IX, Chapter I, Part III, point 4 of Regulation (EC) No 8532004, it complies with the maximum residue limits for residues of antibacterial veterinary medicinal products laid down in the Annex to Regulation (EU) No 37/2010,
- (vi) which has been produced under conditions guaranteeing compliance with the maximum residue levels for pesticides laid down in Regulation (EC) No 396/2005, and maximum levels for contaminants laid down in Regulation (EC) No 1881/2006.
- (b) it comes from an establishment implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004,

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COUNTRY: Countries subject to

Milk-HTB - Dairy products derived from COUNTRY: Countries subject to Milk-HTB - Dairy products derived from transitional import arrangements (*) milk of cows, ewes, goats and buffaloes for human consumption from third countries authorised in column B

TT Wes	1+h	information	II.a. Certificate	II.b.					
II. IICa	II CII	IIIOImacion	reference number	11.0.					
	, ,								
	(c) it has been processed, stored, wrapped, packaged and transported in accordance								
	with the relevant hygiene conditions laid down in Annex II to Regulation (EC) No 852/2004 and Chapter II of Section IX of Annex III 3 to Regulation (EC) No								
	853/2004;								
	(4)	(d) mosts the relevant switchis laid days in Chapter Il of Costion IV of Apper III to							
	(u)	(d) meets the relevant criteria laid down in Chapter Il of Section IX of Annex III to Regulation (EC) No 853/2004 and the relevant microbiological criteria laid down in							
	Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs,								
	(a) the greenteed governor live enimals and products thereof provided by the regidue plans								
	(e) the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are								
	fulfilled.								
Notes									
(*)Thos	se co	ountries subject to the transitional imp	ort arrangements include:	an EU member State;					
		ein; Norway; Iceland and Switzerland.							
D - £		to Tourney Wiley Lord Land of the Control of the Control		name to discount TITE					
		to European Union legislation within th which has been retained in Great Brita							
) Act 2018).	(
D - 6		to Court Duitain in this mortificate in	alada Obassal Talasda and	Talla of Man					
Reieren	ıces	to Great Britain in this certificate in	clude Channel Islands and	isle of Man.					
Part I:									
_		reference I.7: Provide name and ISO coo	de of the country or part t	hereof as appearing in					
	Ann	ex 1 to Regulation (EU) No 605/2010.							
_	Box reference I.11: Name, address and approval number of the establishment of dispatch.								
_	Box	reference I.15: Registration number (ra	ailway wagons or container	and road vehicles),					
		ght number (aircraft) or name (ship). In	_						
		ber of containers and their registration							
		l it must be indicated in box I.23. In t t inform the border control post of into							
-	Box reference I.16: Do not use this box until the end of the transitional staging period.								
-		reference I.19: Use the appropriate Har							
	headings: 04.01; 04.02; 04.03; 04.04; 04.05; 04.06; 15.17; 17.02; 21.05; 22.02; 28.35; 35.03 or 35.04.								
	33.	02 01 33.01.							
_	Box reference I.20: Indicate total gross weight and total net weight.								
_	Box	reference T.23: For containers or boxes	s, the container number and	the seal number (if					
	Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.								
	_								
_		reference I.28: Manufacturing plant: in		er of the treatment and/or					
processing establishment(s) approved for export to Great Britain.									
Part II:									
The colour of the signature shall be different to that of the printing. The same rule applies									
to stamps other than those embossed or watermark.									
Official Veterinarian									
Name (in ca		pital letters):	Qualification and t	itle:					
Date:			Signature:						
Ob /									
Stamp:									

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