Official certificate to the EU

	I.1. Consignor					I.2. Certifica	te reference		I.2.a. Il	MSOC refere	nce		
	Name						I.3. Central competent authority Specimen not to be used for imports into the				to the		
	Address					I.4. Local con	mpetent autho	ority	EU				
	Country			ISO Code									
	I.5. Consignee					I 6 Responsi	ible for the co	neignm	ont in F	et i			
	Name					Name	ible for the cor	nsignin	CIII III I	.0			
int	Address					Address							
וונ	Country			ISO Code		Country				ISO Code			
gn	Country	150 code								100 0000			
: Details of consignment	I.7. Country of original	in ISO Co	de	I.8. Region of origin	Code	I.9. Country destination	of	ISO Co	de I.1	0. Region of	destinat	ion	Code
כ	I.11. Place of dispatch					I.12. Place of	f destination						
Ö	Name					Name							
Ë	Address					Address							
בונ	Approval Number					Approval N	umber						
1	Country			ISO Code		Country				ISO Cod	.e		
raiti.	-												
מדו	I.13. Place of loading	g				I.14. Date an	d time of depa	arture					
1	Name												
	Address												
	Approval Number												
	Country			ISO Code									
	145 M					IAC File D	CD.						
	I.15. Means of Trans	-		*1		I.16. Entry B	CP						
		International transport		Identification		Authority Country							
	(document				Country							
	I.18. Transport cond Chilled \square	Frozen		Ambien	t 🗆	Type	oanying docun	nents					
	I.19. Container No /	Seal No				Number							
L20 Contified as													
	I.20. Certified as												
	Conn Cloud Anny		Pet	ts Tech Quar Fat	te Anim Artifi g al cial	i Phar Regis mace tered	Furth Gam	Hum an	Prod uctio	Relay Man	Trad	Petfo	Bree ding/
		Circu Othe s/exh r l ibitio n l	Pet	ts Tech Quar Fat nical antin nin Use e	te Anim Artifi g al cial Feedi repro ngstu ducti ff μη	i Phar Regis mace tered o utical equi	Furth Gam er e proc resto ess cking	Hum an Cons umpt ion	Prod uctio n of petfo od	Relay Man ing ufac ure of petfo	les -	Petfo	Bree ding/ prod uctio n 🗀
	Cann Slaug Appring hter oved indus Bodi	ibįtiρ	Pet	ts Tech Quar Fat nical antin pin Use e	ngstu ducti	i Phar Regis mace tered o utical equi use dae		umpt	petfo	of	les -	Petfo	Bree ding/ prod uctio n 🏻
	Cann Slaug Appring hter oved indus Bodi	ibįtiρ	Pet	ts Tech Quar Fat nical antin pin Use e	ngstu ducti	i use dae		umpt	petfo	of petfo	les -	Petfo	Bree ding/ prod uctio n 📙
	Cann Slaug Appring hter oved indus Bodi	ibįtiρ	Pet	ts Tech Quar Fat nical antin nin Use e	ngstu ducti	i use dae	ernal market	umpt	petfo	of petfo	les -	Petfo	Bree ding/ prod uctio n
	Cann Islaug Appring oved indus Bodi es I.21. For transit	ibįtiρ	Pet		ngstu ducti	I.22. For inte	ernal market entry	umpt	petfo	of petfo	les -	Petfo	Bree ding/ prod uctio n
	Cann ing indus Herrican vived indus Herrican Bodi es I.21. For transit Non-EU	ibitio n 🔲	Pet		ngstu ducti	I.22. For inte	ernal market entry	umpt	petfo	of petfo	les -	Petfo	Bree ding/ prod uctio n
	Cann ing indus Herrican vived Bodi es I.21. For transit Non-EU I.25. Quantity	consignment	Pet		ngstu ducti	I.22. For inte	ernal market entry	umpt	petfo	of petfo	les -	Petfo	Bree ding/prod uction
	Cann indus Literal Provided in State Control	consignment	Pet		ngstu ducti	I.22. For inte	ernal market entry	umpt	petfo	of petfo	les -	Petfo	Bree ding/ prod uctio n
	Cann indus Herrican Slaug Appring Step 1.21. For transit Non-EU I.25. Quantity I.27. Description of 0.1.01 LIVE ANIMAL 0102 Live bovine	consignment .S		ISO Code	ngstu ducti	I.22. For inte I.23. For re-e I.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/ prod uctio n
	I.21. For transit Non-EU I.27. Description of o	consignment .S	ecies	ISO Code	ngstu ducti	I.22. For inte I.23. For re-e I.26. Total gr	ernal market entry	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved Bodi es I.21. For transit Non-EU I.25. Quantity I.27. Description of or 1. 01 LIVE ANIMAL 0102 Live bovine	consignment .S		ISO Code	ngstu ducti	I.22. For inte I.23. For re-e I.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uctio
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction
	Cann Slaug Appring ved indus Slaug Bodi es S	consignment .S		ISO Code	ngstu ducti	1.22. For inte 1.23. For re-6 1.26. Total gr	ernal market entry coss weight	umpt ion	petfo	of petfo		Petfo	Bree ding/prod uction

	II. Health information		
١.			
<u>.</u>			
t			
問			
[등			
۱ü			
Part II: Certification			
Pa			
		/ / / /	
		>	
	5		

en

COUNTRY 206/2010 (2021/403) Model BO						
	II. Health information					
	II.1. Public hea	alth attestati	on [*to delet	te when the Union is not the final destination of the animals]		
				by certify, that the animals described in this certificate:		
	II.1.1.	have not r				
		-	any stilber	ne or thyrostatic substances,		
fication		-		c, androgenic, gestagenic or beta-agonist substances for purposes therapeutic or zootechnical treatment (as defined in Council Directive		
Part II : Certification	II.1.2.	plans subr	mitted in acc l animals are	overing live animals and products thereof provided by the residue cordance with Article 29 of Council Directive 96/23/EC and the e listed in Commission Decision 2011/163/EU for the concerned		
Pa	II.1.3.	with regar	d to bovine	spongiform encephalopathy (BSE):		
		(a)		ls are identified by a permanent identification system enabling them d back to the dam and herd of origin, and they are not:		
			(i)	BSE cases;		
			(ii)	bovine animals which, during their first year of life, were reared with BSE cases during their first year of life, and which an investigation has shown that they have consumed the same potentially contaminated feed during that period, or		
	and		(iii)	if the results of the investigation referred to in indent (ii) are inconclusive, bovine animals which, during their first year of life, were reared with BSE cases during their first year of life, or were born in the same herd as, and within 12 months preceding or following the date of the birth of, the BSE cases;		
	(1)	either ○ [(b)	(i)	the animals were born and continuously reared in a country or region or countries or regions classified in accordance with Commission Decision 2007/453/EC as countries or regions posing a negligible BSE risk;		
		Ç	(ii)	if there have been BSE indigenous cases in the country concerned, the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health, was effectively enforced, or they were born after the date of birth of the last BSE indigenous case if born after the date of the feed ban.]		
	(1)	or ○ [(b)	(i)	the country or region of origin of the animals is classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk;		
			(ii)	the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health, was effectively enforced, or they were born after the date of birth of the last BSE indigenous case if born after the date of the feed ban.]		
	(1)	or ○ [(b)	(i)	the country or region of origin of the animals is classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk;		
			(ii)	the feeding of ruminants with meat-and-bone meal and greaves from ruminants, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health, has been banned and the ban has been effectively enforced in the country or region of origin;		

_	UNIKI						
	II. Health info	ormation					
11				(iii)	the ban on greaves der Animal Hea was effectiv	s were born at least two years after the date from which the feeding of ruminants with meat-and-bone meal and rived from ruminants, as defined in the Terrestrial alth Code of the World Organisation for Animal Health, wely enforced, or they were born after the date of birth of indigenous case if born after the date of the feed ban.]	
מחנ	II.2.	Animal he	ealth attest	ation			
	I, the unde	ersigned off	icial veteri	narian, her	eby certify that	t the animals described in Part I:	
raitii. eeimicadon		II.2.1.	this certi	ficate is aut	thorised for en		
TOT		II.2.2.	have ren	nained cont	inuously:		
			(i)			in point II.2.1. since birth or for a period of time of at the date of their dispatch to the Union, and	
			(ii)	days pri period n	or to the date o to bovine anim	Forigin since birth or for a period of time of at least 40 of their dispatch to the Union, into which during this als and no animals of other species listed for the same nals have been introduced.	
		II.2.3.			animals of a lo	ower health status since birth or at least for 30 days prior Union.	
		II.2.4.	the relev	ant listed d		al programme for the eradication of diseases, including d to in Annex I to Commission Delegated Regulation (EU	
	(1)	either o [II.2.5.			ed directly fron other establis	n their establishment of origin to the Union without hment].	
	(1)	or ○ [II.2.5.	have undergone one single assembly operation in the zone of origin fulfilling the following requirements:				
			(a)	the asse	mbly operation	took place in an establishment:	
				(i)	competent	or conducting assembly operations of ungulates by the authority in the third country or territory in accordance of Delegated Regulation (EU) 2019/2035;	
				(ii)		an unique approval number assigned by the competent f the third country or territory;	
				(iii)	country or	nat purpose by the competent authority of the third territory of dispatch with the information set out in f Delegated Regulation (EU) 2019/2035;	
				(iv)	•	e requirements provided for in Article 8 of Delegated (EU) 2020/692.	
			(b)	the asse	mbly operation	n in the assembly centre took no longer than 6 days.]	
		II.2.6.	in point l loaded fo	I.2.11. since or dispatch	e they were dis	ce that does not comply with the requirements laid down patched from their establishment of origin until they are nd during that period they have not been in contact with	
		II.2.7.	transpor	t which wa	s cleaned and d	n on (dd/mm/yyyy)(3) in a means of disinfected prior to loading with a disinfectant authorise third country or territory and constructed in such a way	
			(i)	animals	cannot escape	or fall out;	
			(ii)	visual in	spection of the	e space where animals are kept is possible;	
1			(iii)	the esca	no of animal or	screments, litter or feed is prevented or minimized.	

en 4/ 10

II. Health information II.2.8. have been subjected to a clinical inspection within the 24 hour period prior to loading for dispatch to the Union, carried out by an official veterinarian in the third country or territory of origin, who did not detect signs indicative of the occurrence of diseases, including the relevant listed diseases referred to in Annex I to Delegated Regulation (EU) 2020/692 and emerging diseases. II.2.9. have not been vaccinated against: (i) foot and mouth disease, infection with Rift Valley fever virus, infection with Mycoplasma mycoides subsp. mycoides SC (Contagious bovine pleuropneumonia), Mycobacterium tuberculosis complex (M.bovis, M.caprae and M.tuberculosis) and infection with Brucella abortus, B. melitensis and B. suis, and (ii) infection with bluetongue virus (serotypes 1-24) with a live vaccine during 60 days prior to their dispatch to the Union. II.2.10. come from a zone: II.2.10.1. in which: (i) foot and mouth disease has not been reported for: either o [at least 24 months prior to the date of dispatch of the animals to the Union] (1) (dd/mm/yyyy)](1)(4) or o [since (ii) vaccination against foot and mouth disease has not been carried out for at least 12 months prior to the date of dispatch of the animals to the Union, and no animals vaccinated against foot and mouth disease have been introduced during that period. in which infection with lumpy skin disease virus has not been reported for at II.2.10.2. least 12 months prior to the date of dispatch of the animals to the Union. in which infection with rinderpest virus, infection with Rift Valley fever virus II.2.10.3. and infection with Mycoplasma mycoides subsp. mycoides SC (Contagious bovine pleuropneumonia) has not been reported for at least 12 months prior to the date of dispatch of the animals to the Union and during that period: (i) vaccination against these diseases has not been carried out, and (ii) animals vaccinated against these diseases have not been introduced. either o which is free from infection with bluetongue virus (serotypes 1-24)] (1)(5) [II.2.10.4. $or \circ$ which is seasonally free from infection with bluetongue virus (serotypes 1-24): [II.2.10.4. for at least 60 days prior to the date of dispatch of the animals to the [II.2.10.4.1 Union.](1)(6) for at least 28 days prior to the date of dispatch of the animals to the $or \circ$ [II.2.10.4.1 Union and the animals have been subjected to a serological test in accordance with Article 9(b) of Delegated Regulation (EU) 2020/692, with negative results, carried out on samples collected at least 28 days following the date of entry of the animal into the seasonally free zone.](1)(6) $or \circ$ for at least 14 days prior to the date of dispatch of the animals to the [II.2.10.4.1 Union and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of entry of the animal in the seasonally free zone.](1)(6) is not free from infection with bluetongue virus (serotypes 1-24) and the animals $or \circ$ have been vaccinated against all the serotypes (1 to 24) of bluetongue virus [II.2.10.4. reported during the past 2 years in that zone and are still within the immunity period of time guaranteed in the specifications of the vaccine and

COUNTRY 206/2010 (2021/403) Model BOV-X II. Health information have been vaccinated more than 60 days prior to the date of dispatch [II.2.10.4.1 of the animals to the Union.]](1) or \circ have been vaccinated with an inactivated vaccine and were [II.2.10.4.1 subjected to a PCR test, with negative results on samples collected at least 14 days after the onset of the immunity protection set in the Part II : Certification specifications of the vaccine.]](1) $or \circ$ is not free from infection with bluetongue virus (serotypes 1-24) and the animals [II.2.10.4. have been subjected with positive results to a serological test able to detect specific antibodies against all serotypes (1 to 24) of bluetongue virus reported during the past 2 years in that zone and: the serological test has been carried out on samples collected at least [II.2.10.4.1 60 days prior to the date of dispatch of the animals to the Union.]](1) or \circ the serological test has been carried out on samples collected at least [II.2.10.4.1 30 days prior to the date of dispatch of the animals to the Union and the animals were subjected to a PCR test, with negative results, carried out on samples collected not earlier than 14 days prior to the date of dispatch of the animals to the Union.]](1) is free from enzootic bovine leukosis](1)(7) either o [II.2.10.5. is not free from enzootic bovine leukosis and the disease has not been reported $or \circ$ [II.2.10.5. in the establishment of origin of the animals during at least the 24 months prior to the date of dispatch of the animals to the Union, and the animals of the consignment over 24 months of age: [II.2.10.5.1 have been kept in isolation from the other bovine either 🌢 [II.2.10.5.1 animals kept in the same establishment prior to dispatch to the Union and during the period of isolation have been subjected to a laboratory examination for enzootic bovine leukosis using one of the diagnostic methods referred to in Article 9(b)(i) of Delegated Regulation (EU) 2020/692, with negative results, carried out on samples taken on two occasions at an interval of at least 4 months.]](1) have been subjected to a laboratory examination for $or \circ$ [II.2.10.5.1 enzootic bovine leukosis using one of the diagnostic methods referred to in Article 9(b)(i) of Delegated .1. Regulation (EU) 2020/692, with negative results, carried out on a sample taken during the 30 day period prior to the date of their dispatch to the Union and all bovine animals over 24 months of age kept in the establishment of origin have been subjected to a laboratory examination for enzootic bovine leukosis with one of the diagnostic methods referred to in Article 9(b)(i) of Delegated Regulation (EU) 2020/692, carried out, with negative results, on samples taken on two occasions at an interval of not less than 4 months during the 12 month period prior to the date of dispatch of the animals to the

en 6/10

Union.]](1)

CO	UNTRY				206/201	l0 (2021/403) Model BOV-X	
	II. Health information						
cation			□ [II.2.10.5.2 ·	born to dar examinatio methods re 2020/692, w occasions a	s of the consignment younger ns which have been subjected n for enzootic bovine leukosi ferred to in Article 9(b)(i) of I with negative results, carried of t an interval of not less than od prior to the date of dispato	d to a laboratory s with one of the diagnostic Delegated Regulation (EU) out on samples taken on two 4 months during the 12	
ij	II.2.11.	come from	an establisl	nment:			
Part II: Certification		II.2.11.1.	third coun	competent authority of the in place to maintain for at ion regarding:			
			(1)	establishme		ithication of animals on the	
			(ii)	movements	s of animals into and out of th	ne establishment;	
			(iii)	_	the establishment.		
		II.2.11.2.	of the deter diseases, in Delegated	ction of, and ncluding the Regulation (I	animal health visits from a v information on, signs indicate relevant listed diseases refer EU) 2020/692 and emerging distance by the establishment.	tive of the occurrence of	
		II.2.11.3.	which was not subject to national restriction measures for animal health reasons, including the relevant listed diseases referred to in Annex I to Delegated Regulation (EU) 2020/692 and emerging diseases, at the time of dispatch of the animals to the Union.				
		in and around which, in an area of 10 km radius, including where appropriate the territory of a neighbouring country, none of the following listed diseases has been reported for at least 30 days prior to the date of dispatch of the animals to the Union: foot and mouth disease, infection with rinderpest virus, infection with Rift valley fever virus, infection with Mycoplasma mycoides subsp. mycoides SC (Contagious bovine pleuropneumonia) and infection with lumpy skin disease virus.					
	either o [II.2.11.5.		appropriat disease has	e the territo	n an area with a 150 km radio ry of a neighbouring country, ported for at least 2 years be n.](1)	, epizootic haemorrhagic	
		or o [II.2.11.5.	which is lo disease.](1)		ne seasonally free of epizooti	ic haemorrhagic	
		II.2.11.6.			h Mycobacterium tuberculosi losis) as regards bovine anim		
			either o [II.2.11.6.1		zone free from the disease we is not practiced.](1)(10)	vhere vaccination against	
			or o [II.2.11.6.1	provided for infection caprae and	s have been tested with one o or in Article 9(b)(i) of Delegate n with Mycobacterium tubero M. tuberculosis), with negati r to the date of dispatch of th	ed Regulation (EU) 2020/692 culosis complex (M. bovis, M. ve results, during the 30 day	
			or ○ [II.2.11.6.1	the animals	s are less than six weeks old.]	(1)	
		II.2.11.7.		nfection wit	h Brucella abortus, B. meliter	nsis and B. suis as regards	

bovine animals(9), and

7/10

or ○ [II.2.1: (1)(12) □ the an pustul either [II.2.1:		
(1)(12)		
(1)(12)		a zone free from the disease where vaccination against e is not practiced.](1)(11)
(1)(12)	[II.2.11.7.1 provided for for infection negative rethe date of post-parture	s have been tested with one of the diagnostic methods or in Article 9(b)(i) of Delegated Regulation (EU) 2020/692 on with Brucella abortus, B. melitensis and B. suis, with esults, on a sample taken during the 30 day period prior to dispatch of the animals to the Union, and in the case of crient females, the test is carried out on a sample taken at ys after parturition.](1)
(1)(12)	or \circ the animals [II.2.11.7.1 .	s are less than 12 months old.](1)
(1)(12)	or \circ the animals [II.2.11.7.1	s are castrated.](1)
(1)(12)	1.8. in which infection with prior to dispatch of the	n rabies virus has not been reported for at least 30 days
(1)(12)	 in which anthrax has n dispatch of the animals 	not been reported for at least 15 days prior to the date of s to the Union.
(1)(12)	7.	nosoma evansi) has not been reported for at least 2 years patch of the animals to the Union.](1)
[III.2.12. pustul either [II.2.12 or o [II.2.13] (1)(12)	1.10. prior to the date of disp was reported in the est of dispatch of the anim restriction until the inf the remaining animals to a test for surra (Tryp Delegated Regulation (1	nosoma evansi) has not been reported for at least 30 days patch of the animals to the Union and when the disease tablishment of origin during the 2 years prior to the date hals to the Union, the establishment remained under fected animals were removed from the establishment and son the establishment were subjected with negative result panosoma evansi) as described in Article 9(b)(i) of EU) 2020/692 carried out on samples taken at least 6 ted animals were removed from the establishment.](1)
(1)(12) ☐ [II.2.13. the an	nimals have not been vaccina lar vulvovaginitis, and	ited against infectious bovine rhinotracheitis/infectious
(1)(12) □ [II.2.13. the an		country or territory or zone thereof free from Infectious / infectious pustular vulvovaginitis.]](1)(13)
	2.1. the animals to the Unic of antibodies against w diagnostic methods ref 2020/692, with negative	rantine for at least 30 days prior to the date of dispatch of on and have undergone a serological test for the detection whole bovine herpes virus-1 (BoHV-1) with one of the ferred to in Article 9(b)(i) of Delegated Regulation (EU) e results, on a sample taken within 15 days prior to the animals to the Union.]](1)
aithar	nimals have not been vaccina	ited against bovine viral diarrhoea, and:
[II.2.13	9	country or territory or zone thereof free from bovine vira
or ○ [II.2.13	3.1. of the diagnostic method	ovine viral diarrhoea virus antigen or genome using one ods provided for in Part 6 of Annex I to Commission EU) 2020/688 with negative results, and
		kept in a quarantine establishment for a period of at least or to their dispatch to the Union.]]](1)

en 8/ 10

II. Health information or o the animals are pregnant dams and have been kept in a quarantine [II.2.13.1.1 establishment for a period of at least 21 days prior to their dispatch to the Union and have been subjected to a serological test for the detection of antibodies against bovine viral diarrhoea virus using one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688 with negative results carried out Part II : Certification on samples taken not less than 21 days after the commencement of the quarantine.]]](1) have been subjected to serological test for the detection of antibodies or o [II.2.13.1.1 against bovine viral diarrhoea virus using one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, with positive results, carried out on samples taken prior to their dispatch to the Union.]]](1) the animals are pregnant dams that have been subjected to or \circ [II.2.13.1.1 serological test for the detection of antibodies against bovine viral diarrhoea virus using one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688 with positive results, carried out on samples taken before insemination preceding the current gestation.]]] (1)

9/10 en

COUNTRY

				.0 (2022, 100) 1110 401 20 1 11					
	II. Health info	rmation							
	Notes:								
		cate is intended for entry into the Union of boy of the animals.	ine animals, including when t	he Union is not the final					
icatio	In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland / Northern Ireland in conjunction with Annex 2 to that Protocol, references to European Union in this certificate include the United Kingdom in respect of Northern Ireland.								
II : Cer	This animal health/official certificate shall be completed according to the notes for the completion of certificates provided for in Chapter 4 of Annex I to Commission Implementing Regulation (EU) 2020/2235.								
art]	Part I:								
	Box reference I.27:	"Identification system and identification number tattoo, transponder etc., from the list in Annex individual identification codes of the animals in (EU) 2020/692.	III to Delegated Regulation (F	EU) 2019/2035) and the					
	Part II:								
	(1)	Keep as appropriate.							
	(2)	Code of the zone as it appears in Column 2 of F [C(2021)1800].	Part 1 of Annex II to Implemen	nting Regulation (EU)					
	(3)	Date of loading: it cannot be a date prior to the Union, or a date in a period when restriction n of these animals from this zone.							
	(4)	Only for zones with opening date in accordance Regulation (EU) 2021/404.	e with column 8 in part 1 of A	Annex II to Implementing					
	(5)	For zones with entry BTV in column 7 of Part 1	nes with entry BTV in column 7 of Part 1 of Annex II to Implementing Regulation (EU) 2021/404.						
	(6) For zones with entry SF-BTV in column 7 of Part 1 of Annex II to Implementing Regulation (EU) 2021/404.								
	(7)	For zones with entry EBL in column 7 of Part 1	of Annex II to Implementing	Regulation (EU) 2021/404.					
	(8)								
	(9)	In accordance with Article 10 of Delegated Reg	gulation (EU) 2020/692.						
	(10)	For zones with entry TB for bovine animals in column 7 of Part 1 of Annex II to Implementing Regulation (EU) 2021/404.							
	(11)	For zones with entry BRU for bovine animals in column 7 of Part 1 of Annex II to Implementing Regulation (EU) 2021/404.							
	Only applicable when the Member State of destination or Switzerland, in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002), either has disease-free status or an approved eradication programme for the diseases mentioned in point II.2.12 and II.2.13 (infectious bovine rhinotracheitis/infectious pustular vulvovaginitis and bovine viral diarrhoea).								
	(13)	For zones with entry IBR in column 7 of Part 1	of Annex II to Implementing	Regulation (EU) 2021/404.					
	(14)	For zones with entry BVD in column 7 of Part 2	l of Annex II to Implementing	Regulation (EU) 2021/404.					
	Official veterin	narian or Official inspector							
	Name (in cap Date of signar Stamp		Qualification and title Signature						