	UNITED KINGDOM					ealth certificate to the EU	
	I.1 Consignor/Exporter			I.2 Certificate ref	ference	I.2a	
	Name				······		
	Address		ŀ	I.3 Central Comp	petent Authority	7 /	
				FOOD & RURAL			
´			ļ	I.4 Local Compet	tent Authority] /	
	Country	ISO co	untry code	ANIMAL AND P	LANT HEALTH AGENCY		
	1.5 Consignee/Importer			I.6 Operator resp	oonsible for the consignmen	t	
	Name	!	Name				
별●	Address		!	Address			
Part I: Description of consignment	. 6						
n	Country	ISO cor	untry code	Country	I	SO country code	
riptio	I.7 Country of origin	ISO cor	untry code	I.9 Country of de	estination	ISO country code	
: Desc	I.8 Region of origin	Code		I.10 Region of de		Code	
Part I	I.11 Place of dispatch	Regis	stration/Approval No	I.12 Place of dest	ination	Registration/Approval No	
	Name	1		Name			
	Address			Address			
		ISO 22			,	20	
ļ	Country	130 00	untry code	Country		SO country code	
	I.13 Place of loading			I.14 Date and tim			
	I.15 Means of transport		ļ	I.16 Entry Borde	er Control Post		
	☐ Aircraft ☐	Vessel		I.17 Accompanyi	ng documents		
			!	Type	Co	ode	
	□ Railway □	☐ Road veh	icle	Country		O country code	
	Identification		!	Commercial documents	ment reference		
-	I.18 Transport conditions	□ An	nbient	☐ Chilled	☐ Froz	en	
	I.19 Container number/Seal number				•		
	Container No			Seal No	*		
	I.20 Certified as or for						
	☐ Germinal products						
	I.21		1	I.22 🗆 Fo	or internal market		
	Third country	ISO c	country code	I.23			
	I.24 Total number of packages		I.25 Total quantity		I.26		

HAITT	ED KINCDOM			II.a Certific	ate reference
I.27	ED KINGDOM Description of cons	signment			
1	CN code	Species	Subspecies/Category	Identification number	Quantity
•	2. Coac	Species	Sucception carried by		
	Туре	Approval or registration number of plant/establishment/centre	Identification mark	Date of collection/production	
2	CN code	Species	Subspecies/Category	Identification number	Quantity
3	Туре	Approval or registration number of plant/establishment/centre	Identification mark	Date of collection/production	
3	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration number of plant/establishment/centre	Identification mark	Date of collection/production	
4	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration number of plant/establishment/centre	Identification mark	Date of collection production	
5					
	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration number of plant/establishment/centre	Identification mark	Date of collection/production	

II.a	1 (Certifi	icate r	eferenc	ce		

II. Health information

I, the undersigned official veterinarian, hereby certify that:

- II.1. The germinal product storage centre ⁽¹⁾ described in box I.11 at which the [semen] ⁽²⁾ [oocytes] ⁽²⁾ [in vivo derived embryos] ⁽²⁾ [in vitro produced embryos] ⁽²⁾ [micromanipulated embryos] ⁽²⁾ to be dispatched to the Union was/were stored:
 - II.1.1. is located in a third country or territory, or zone thereof:
 - II.1.1.1. authorised for the entry into the Union of [semen] (2) [cocytes] (2) [embryos] (2) of [ovine] (2) [caprine] (2) animals and listed in Annex X to Commission Implementing Regulation (EU) 2021/404;
 - **either [II.1.1.2. where foot and mouth disease was not reported for at least 24 months immediately prior to the date of [collection] (2) [production] (2) of the [semen] (2) [oocytes] (2) [embryos] (2) and until the date of its/their dispatch;]
 - where foot and mouth disease was not reported for a period starting on the date / / (ad/mm/yyyy) immediately prior to the date of [collection] (2) [production] (2) of the [semen] (2) [oocytes] (2) [embryos] (2) and until the date of its/their dispatch;]
 - II.1.3. where infection with rinderpest virus, infection with Rift Valley fever virus, infection with peste des petits ruminants virus, sheep pox and goat pox and contagious caprine pleuropneumonia were not reported for at least 12 months immediately prior to the date of [collection] (2) [production] (2) of the [semen] (2) [oocytes] (2) [embryos] (2) and until the date of its/their dispatch;
 - II.1.1.4. where no vaccination against infection with rinderpest virus, infection with Rift Valley fever virus, infection with peste despetits ruminants virus, sheep pox and goat pox and contagious caprine pleuropneumonia has been carried out for at least 12 months immediately prior to the date of [collection] (2) [production] (2) of the [semen] (2) [cocytes] (2) [embryos] (2) and until the date of its/their dispatch, and no vaccinated animals entered into the third country or territory, or zone thereof during that period, and
 - (2) either [no vaccination against foot and mouth disease has been carried out for the same period, and no vaccinated animals entered into the third country or territory, or zone thereof during that period;]
 - (2) or [vaccination against foot and mouth disease has been carried out for the same period, or vaccinated animals entered into the third country or territory, or zone thereof during that period;]
 - II.1.2. is approved and listed by the competent authority of the third country or territory;
 - II.1.3. complies with requirements as regards responsibilities, operational procedures, facilities and equipment set out in Part 5 of Annex I to Commission Delegated Regulation (EU) 2020/686.]
- II.2. The [semen] (2) [oocytes] (2) [embryos] (2) described in Part I is/are intended for artificial reproduction and
 - II.2.1. has/have been [collected] (2) [produced] (2), [processed] (2) [stored] (2) [in a semen collection centre] (2) (4) [by an embryo collection team] (2) (4) [by an embryo production team] (2) (4) [and] (2) [processed] (2) [stored] (2) [in a germinal product processing establishment] (2) (4) and stored in a germinal product storage centre (4) complying with requirements set out in [Part 1] (2) [Part 2] (2) [Part 3] (2) [Part 4] (2) [Part 5] (2) Annex I to Delegated Regulation (EU) 2020/686, and:
 - (2) either [located in the third country or territory of dispatch to the Union;]
 - [located in ______, and has/have been introduced into the third country or territory of dispatch to the Union under conditions at least as strict as for the entry into the Union of [semen] (2) [cocytes] (2) [embryos] (2) of [ovine] (2) [caprine] (2) animals in accordance with Regulation (EU) 2016/429 and Commission Delegated Regulation (EU) 2020/692;]
 - II.2.2. was/were moved to the germinal product storage centre described in box I.11 under conditions at least as strict as described in:
 - (2) either [Model OV/CAP-SEM-A-ENTRY (6);]
 - (2) and/or [Model OV/CAP-SEM-B-ENTRY (6);]
 - (2) and/or [Model 1 in Section A of Part 2 of Annex II to Decision 2010/472/EU (6);]
 - (2) and/or [Model 2 in Section B of Part 2 of Annex II to Decision 2010/472/EU (6);]
 - (2) and/or [Model OV/CAP-OOCYTES-EMB-A-ENTRY (6);]
 - (2) and/or [Model OV/CAP-OOCYTES-EMB-B-ENTRY (6);]
 - (2) and/or [Model OV/CAP-GP-PROCESSING-ENTRY (6);]
 - (2) and/or [Model OV/CAP-GP-STORAGE-ENTRY (6);]
 - II.2.3. has/have been collected, processed and stored in accordance with animal health requirements set out in Annex III to Delegated Regulation (EU) 2020/686;
 - II.2.4. is/are placed in straws or other packages on which the mark is applied in accordance with requirements provided for in Article 83 point (a), of Delegated Regulation (EU) 2020/692 and that mark is indicated in box I.27;
 - II.2.5. is/are transported in a container which
 - II.2.5.1. was sealed and numbered prior to the date of dispatch from the germinal product storage centre under responsibility of the centre veterinarian, or by an official veterinarian, and the seal bears the number as indicated in box I.19;
 - II.2.5.2. has been cleaned and either disinfected or sterilised before use, or is single-use container;
 - (2) (7) [II.2.5.3. has been filled in with a cryogenic agent which has not been previously used for other products;]
 - (2) (8) [II.2.6. is/are placed in straws or other packages which are securely and hermetically sealed;

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II.2.7. is/are transported in a container where the different types are separated from each other by physical compartments or by being placed in secondary protective bags.]

Notes

This animal health certificate is intended for the entry into the Union of semen, oocytes and embryos of ovine and caprine animals, including when the Union is not the final destination of the semen, oocytes and embryos.

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 Windsor Framework (see Joint Declaration No 1/2023 of the Union and the United Kingdom in the Joint Committee established by 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 of 24 March 2023, OJ L 102, 17.4.2023, p. 87) in conjunction with Annex 2 to that Framework, references to the Union in this animal health certificate include the United Kingdom in respect of Northern Ireland.

This animal health certificate shall be completed in accordance with the notes for the completion of certificates provided for in Chapter 4 of Annex I to Commission Implementing Regulation (EU) 2020/2235.

Part I.

Box reference I.11:

"Place of dispatch": Indicate the unique approval number and the name and address of the germinal product storage entre of dispatch of the consignment of semen, oocytes and/or embryos. Only germinal product storage centres listed in accordance with Article 233(3) of Regulation (EU) 2016/429 on the Commission website:

http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm.

"Place of destination": Indicate the address and unique registration or approval number of the establishment of Box reference I.12:

destination of the consignment of semen, oocytes and/or embryos.

Box reference I.17:

"Accompanying documents": Number(s) of related original animal health certificate(s) shall correspond to the serial number of the individual official document(s) or animal health certificate(s) that accompanied the semen, oocytes and/or embryos described in Part I from the semen collection centre where the semen was collected, and/or from the embryo collection team and/or the embryo production team by which the oocytes and/or embryos were collected or produced, and/or from the germinal product processing establishment where the semen, oocytes and/or embryos were processed and stored, and/or from the germinal product storage centre where the semen, oocytes and/or embryos were stored, to the germinal product storage centre described in box I.11. The original(s) of those document(s) or those animal health

certificate(s) or the officially endorsed copies thereof shall be attached to this animal health certificate.

Box reference I.19: Seal number shall be indicated.

Box reference I.24: Total number of packages shall correspond to the number of containers.

"Species": indicate "Ovis aries" and/or "Capra hircus" as appropriate. Box reference I.27:

"Type": Specify if semen, in vivo derived embryos, in vivo derived occytes, in vitro produced embryos or

micromanipulated embryos.

"Identification number": Indicate identification number of each donor animal

"Identification mark": Indicate mark on the straw or other packages where semen, oocytes and/or embryos of the consignment are placed.

"Date of collection/production": Indicate the date on which semen, oocytes and/or embryos of the consignment was/were collected or produced.

"Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the semen collection centre, where the semen of the consignment was collected, and/or of the embryo collection team or the embryo production team by which oocytes, in vivo derived embryos or in vitro produced embryos of the consignment were collected or produced.

"Quantity": Indicate number of straws or other packages with the same mark.

Part II:

- Only germinal product storage centres listed in accordance with Article 233(3) of Regulation (EU) 2016/429 on the Commission websites http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm.
- (2) Delete if not applicable.
- (3) Only for a third country or territory, or zone thereof with an opening date in accordance with column 9 of the table in Part 1 of Annex II to Implementing Regulation (EU) 2021/404.
- Only approved germinal product establishments listed in accordance with Article 233(3) of Regulation (EU) 2016/429 on the Commission website: http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm.
- (5) Only a third country or territory, or zone thereof listed in Annex X to Implementing Regulation (EU) 2021/404 and Member States.
- The original(s) of the document(s) or the animal health certificate(s) or the officially endorsed copies of thereof that accompanied the semen, oocytes and/or embryos described in Part I from the semen collection centre where the semen was collected, and/or from the embryo collection team or the embryo production team by which the oocytes and/or embryos were collected or produced, and/or from the germinal product processing establishment where the semen, oocytes and/or embryos were processed and stored, and/or from the germinal product storage centre

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11.a	Certificate reference

where the semen, oocytes and/or embryos were stored, to the germinal product storage centre of dispatch of the semen, oocytes and/or embryos described in box I.11 shall be attached to this animal health certificate.

- Applicable for frozen semen, oocytes or embryos.
- Applicable for consignments where semen, oocytes, in vivo derived embryos, in vitro produced embryos and micromanipulated embryos of

ovine and/or caprine animals are placed and transported in one container.	
Official veterinarian	
Name (in capital letters) Oate Qualification and title	
Stamp	