EUROPEAN UNION INTRA

	I.1. Consignor				I.2. IMSOC ref	erence	I.2.a. Local reference			
	Name						I.3. Central Competent Authority			
	Address						I.4. Local Competent Authority			
	Country		ISO Code							
_	I.5. Consignee				I.6. Operator	I.6. Operator conducting assembly operations independently of an				
Part I: Description of consignment	Name				establishmen	Ċ				
Ĕ	Address				Name					
롡	Country		ISO Code		Address					
딿	Country		130 code		Approval Nu	mber				
Ë					Country		ISO Code			
ဗ					country		100 couc			
H	I.7. Country of orig	zin		ISO Code	I.9. Country of	f destination	ISO Code			
=	1177 00 411(1) 01 0116	,			1.01 00 01101) 0.	i doctifiation				
2										
ದ	I.8. Region of origi	n		Code	I.10. Region of	f destination	Code			
Ę	I.11. Place of dispa	tch			I.12. Place of o	destination				
န္တ	Nama									
ڇ	Name				Name					
∺	Address				Address					
E	Approval Number	ſ				Approval Number				
ā	Country		ISO Cod	ie	Country		ISO Code			
_										
	I.13. Place of loadi	ng			I.14. Date and	I.14. Date and time of departure				
	Name									
	Address									
	Approval Number	r								
		L	ICO C	J_						
	Country		ISO Coo	ie						
	I.15. Means of Trai	nenort			I.16. Transpor	rtor				
			- 1			ter				
	Mode	International transport	Identification		Name					
		document			Address					
					Activity ID					
					Country		ISO Code			
					I.17. Accompa	nying documents	;			
						Commercial				
					document reference		Date of issue			
					reference		D1 C			
					Country		Place of issue			
	I.18. Transport conditions									
	-									
	I.19. Container No	/ Seal No								
	I.20. Certified as									
	פר המודוזים) ()/ []									
			l Travelling circ	rus/animal act \square	Quarantine or establishmen	r similar	Exhibition \square			
	Event or activity n	ear borders \square	Travelling enre			+				
	Event or activity n		· ·							
	Event or activity n Release into the w		Other 🗆		Further keepi		Confined establishment \square			
	Event or activity n		· ·							
	Event or activity n Release into the w Slaughter	ild □	Other 🗆			ing 🗆				
	Event or activity n Release into the w Slaughter I.21. For transit th	ild □	Other 🗆		Further keepi					
	Event or activity n Release into the w Slaughter I.21. For transit the Third country	ild □	Other 🗆		Further keepi	ing 🗆				
	Event or activity n Release into the w Slaughter I.21. For transit th	ild □	Other 🗆		Further keepi	ing 🗆				
	Event or activity n Release into the w Slaughter I.21. For transit the Third country	ild □	Other 🗆		Further keepi	ing 🗆				
	Event or activity n Release into the w Slaughter I.21. For transit the Third country Exit point	ild □ rough a third c	Other ountry		Further keepi ISO Code BCP code	ing 🗆				
	Event or activity not	ild □ rough a third c	Other ountry	_	ISO Code BCP code BCP code I.23. For expo	ing 🗌 💮	Confined establishment			
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	Event or activity n Release into the w Slaughter I.21. For transit the Third country Exit point Entry point I.22. For transit the Member State	ild □ rough a third c rough Member	Other ountry State(s)	_	ISO Code BCP code BCP code I.23. For expo Third country Exit point I.25. Journey I	rt /	Confined establishment ISO Code			
	Event or activity not	ild □ rough a third c rough Member	Other ountry State(s)	_	ISO Code BCP code BCP code I.23. For expo Third country Exit point	rt /	Confined establishment ISO Code			
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	II. Health info	rmation								
	I, the undersigned official veterinarian, hereby certify that:									
	II.1. The porcine animals(1) of the consignment described in Part I meet the following requirements:									
		II.1.1.	They are id	hey are identified as provided for in Article 52 or 54(2) of Commission Delegated egulation (EU) 2019/2035.						
Part II: Certification		II.1.2.	They, for a	t least the 3		l prior to the departure of the consignment, or since birth,				
			II.1.2.1.		•	y resident in the esta	ablishmeı	nt of origin;		
			II.1.2.2.	have not b	een in conta	eact with kept porcine animals of a lower health status or restrictions for animal health reasons;				
			II.1.2.3.	the Union		country or territory	_	animals that have entered he 30 day period prior to the		
		II.1.3.	They have not shown clinical signs or symptoms of diseases listed for porcine animals during the clinical examination which was carried out, within the 24 hour period prior to departure of consignment, on (insert date dd/mm/yyyy).							
(2) [II.1.4. They come from one or more holdings officially recognised as applying of conditions in accordance with Article 8 of Commission Implementing Re 2015/1375 and have not passed through an establishment approved for a in accordance with Article 99(3) of Regulation (EU) 2016/429 that does not requirements set out in Chapter I(A)(j) of Annex IV of Implementing Reg 2015/1375.]							enting Regulation (EU) ved for assembly operations t does not meet the			
	II.2.	According requirement	g to official information, the animals described in Part I meet the following health							
		II.2.1.	They do not come from establishments subject to movement restrictions affecting the species or situated in a restricted zone established for reasons of diseases listed for porcanimals.							
		II.2.2.	They come from establishments in which infection with rabies virus in kept terrestrial animals has not been reported during the 30 day period prior to departure.							
		II.2.3.	They come from establishments in which anthrax in ungulates has not been reported during the 15 day period prior to departure.							
		II.2.4.	They come from establishments in which infection with Brucella abortus, B. melitensis and B. suis in porcine animals has not been reported during the last 42 days prior to departure, and in which during at least the 12 month period prior to departure							
	(2)		either \square [II.2.4.1.			tigating measures se Regulation (EU) 202		Article 19(1)(f)(i) of we been introduced;]		
	(2)		and/or surveillance for infection with Brucella abortus, B. melitensis and B. suis has been carried out on the porcine animals kept on the establishments in accordance with Article 19(1)(f)(ii) of Delegated Regulation (EU) 2020/688.]							
		II.2.5.	They come from establishments in which infection with Aujeszky's disease virus has not been reported during the 30 day period prior to departure of the consignment.							
	(2)	○ [II.2.6.	2.6. They are moved to a Member State or zone thereof with the status free from infection with Aujeszky's disease virus and have not been vaccinated against infection with Aujeszky's disease virus, and either come from establishments free from infection with Aujeszky's disease virus, an [II.2.6.1.							
	(2)									
	(2)			either □ [II.2.6.1.1.				in a Member State or zone Aujeszky's disease virus;]]		
[II.2.6.1.2. test for virus w						detection of antibod one of the diagnostic Delegated Regulation sample taken durin	lies again c methods n (EU) 202	en subjected to a serological st whole Aujeszky's disease s provided for in Part 7 of 20/688(3)(4), with a negative day period prior to		

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II. Health info	rmation						
(2)		and/or □ [II.2.6.2.	come from	ı establishme	ents not free from infection w	rith Aujeszky's disease virus,	
		-	-		kept in an approved quaranti t least 30 days; and	ne establishment for a	
(2)			-	antibodies method pro (EU) 2020/6 on two occa	subjected to a serological test against whole Aujeszky's dise ovided for in Part 7 of Annex 88, with a negative result, car asions at an interval of not le en during the 15 day period p	ease virus with the diagnostic I to Delegated Regulation cried out on samples taken ss than 30 days, the last	
(2)	○ [II.2.6.	-	y are moved to a Member State or zone thereof with an approved eradication gramme for infection with Aujeszky's disease virus, and				
(2)		either \square [II.2.6.1.	come from	ı establishme	ents free from infection with	Aujeszky's disease virus, and	
(2)			either □ [II.2.6.1.1.		hments of origin are situated h the status free from infectio		
(2)			and/or □ [II.2.6.1.2.	thereof wit	hments of origin are situated h an approved eradication pr disease virus;]]		
(2)			and/or □ [II.2.6.1.3.	test for the virus or an where appl Part 7 of Ar	s in the consignment have be detection of antibodies again tibodies against Aujeszky's di icable, with one of the diagnomex I to Delegated Regulation sult, on a sample taken durin	st whole Aujeszky's disease sease virus-gE protein, ostic methods provided for in n (EU) 2020/688(4), with a	
(2)		and/or \square [II.2.6.2.	come from virus, and	ı an establish	nment not free from infection	with Aujeszky's disease	
			-		kept in an approved quaranti t least 30 days; and	ne establishment for a	
			-	antibodies method pro (EU) 2020/6 on two occa	subjected to a serological test against whole Aujeszky's dise ovided for in Part 7 of Annex 88, with a negative result, can asions at an interval of not les en during the last 15 days pri	ease virus with the diagnostic I to Delegated Regulation cried out on samples taken ss than 30 days, the last	
II.3.		t of my knowledge and as declared by the operator, the animals come from establishments re were no abnormal mortalities with an undetermined cause.					
(2) \square [II.4.	According	to official information and as declared by the operator, they are semen donor animals, and					
	II.4.1.	they come from a semen collection centre and will be transported directly to another semen collection centre in accordance with Article 19 of Commission Delegated Regulation (EU) 2020/686; and					
(2)	either o	they were continuously resident since the date of their admission at the semen collection centre and were subjected, with negative results, to all compulsory routine tests referred to in point 2 of Chapter I of Part 2 of Annex II to Delegated Regulation (EU) 2020/686 in the period of the preceding 12 months prior to date of that movement; and]					
(2)	or 0 [II.4.2.	they were subjected, with negative results, to all tests referred to in point 1(b) and (c) of Chapter I of Part 2 of Annex II to Delegated Regulation (EU) 2020/686, required before admission to a semen collection centre carried out during the period immediately preceding quarantine and during the quarantine period; and]					
	II.4.3.			ie centre vete operator; and	erinarian of the semen collect	tion centre of destination has	
		DCCII ODIUI	incu by the t	sperator, and	4		

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ъ	JROPEAN (INION	2023/594 (2021/403) MODEL POR-INTRA-X						
	II. Health info	rmation							
	II.5.	Arrangements are made to transport the consignment in accordance with Article 4 of Delegated Regulation (EU) 2020/688.							
	II.6.		rtificate is valid for 10 days from the date of issuing. In the case of transport by waterway/sea of s, the period of validity of the certificate may be extended by the duration of the journey by ay/sea.						
Part II: Certification	(2)(5) □ [II.7.		ing their establishments of origin and before arriving to this establishment approved for operations, none of the animals of the consignment has undergone more than two assembly is, and						
ert	(2)	either \circ [they come from their esta	ablishments of origin.]]						
rt II: C	(2)	or \circ [at least one of the animals of on an approved establishment.]]	the consignment has undergone one assembly operation						
Pa	(2)	or \circ [at least one of the animals of on approved establishments.]]	the consignment has undergone two assembly operations						
	Animal we	lfare attestation							
	At the time of inspection, the animals covered by this health certificate were fit to be transported in accordance with the provisions of Council Regulation (EC) No 1/2005 on the intended journey due to start on (insert date) (6)(7).								
	(2) \square [II.8.	The animals meet the additional guarantees for	r:						
	(2)	either \circ [II.8.1. Porcine animals ke	pt in a restricted zone I in compliance with the special to African swine fever laid down in Commission						
	(2)		n a restricted zone II in compliance with the special to African swine fever laid down in Commission 3/594.]						
	(2)		n a restricted zone III in compliance with the special to African swine fever laid down in Commission 23/594.]]						

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EUROPEAN UNION 2023/594 (2021/403) MODEL POR-INTRA-X II. Health information Notes: 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. Certification This animal health certificate shall be completed according to the notes for the completion of certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235. Part I: Ħ Box "Place of dispatch": Indicate an establishment of the origin of the animals in the consignment or an refer I.11: reference establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. Box "Place of destination": Indicate an establishment of the final destination of the consignment or an reference establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. I.12: Box "Accompanying documents": In case the animals are dispatched from an establishment approved for assembly operations in the Member State of origin, the reference number(s) of the official document(s), reference based on which the animal health certificate for this consignment is issued in this establishment I.17: approved for assembly operations, may be indicated. In case the animals are dispatched from an establishment approved for assembly operations in the Member State of passage, the reference number(s) of the certificate(s), based on which the animal health certificate for this consignment is issued in this establishment approved for assembly operations, must be indicated. "Identification number": Indicate identification codes of the animals in the consignment identified in Box reference accordance with Article 52 or 54(2) of Delegated Regulation (EU) 2019/2035. 1.30: Part II: (1) There can be one or more animals in the consignment. (2) Delete if not applicable. (3) For porcine animals less than four months old born to dams vaccinated with a gE-deleted vaccine, the diagnostic method for the detection of antibodies against Aujeszky's disease virus gE protein provided for in Part 7 of Annex I to Delegated Regulation (EU) 2020/688 may be used. (4)The number of porcine animals tested must allow at least for the detection of 10% seroprevalence of the consignment with 95% confidence. (5) Applicable in case the consignment is dispatched from the establishment approved for assembly operations. (6) In the case where a consignment is grouped in an establishment approved for assembly operations and comprises animals that were loaded on different dates, the date which the journey commenced for the whole consignment is considered to be the earliest date when any part of the consignment left the establishment of origin. (7)This statement does not exempt transporters from their obligation in accordance with Union rules in force in particular regarding the fitness to be transported. Certifying Officer/Official veterinarian

Name (in capital letters) Authority name Signature Date of signature Stamp

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