

I. rész: Bemutatott szállítmány adatai	I.1. Feladó		I.2. Bizonyítvány hivatkozási száma		I.2.a Helyi hivatkozási szám:										
	Név														
	Cím		I.3. Központi illetékes hatóság												
			I.4. Helyi illetékes hatóság												
	Ország														
	I.5. Címzett		I.6. Kapcsolódó eredeti bizonyítványok száma(i) Kisérő okmányok száma(i)												
	Név														
	Cím														
			I.7. Kereskedő												
	Ország		Név		Jóváhagyási szám										
	I.8. Származási ország		ISO kód	I.9. Származási régió	Kód	I.10. Rendeltetési ország	ISO kód	I.11. Rendeltetési régió	Kód						
	I.12. Származási hely/Betakarítás helye								I.13. Rendeltetési hely						
Telep <input type="checkbox"/> Gyűjtő központ <input type="checkbox"/> Kereskedő telephelye <input type="checkbox"/>								Telep <input type="checkbox"/> Gyűjtő központ <input type="checkbox"/> Kereskedő telephelye <input type="checkbox"/>							
Engedélyezett intézmény <input type="checkbox"/> Termékenyítő központ <input type="checkbox"/> Engedélyezett aquakultúra telep <input type="checkbox"/>								Engedélyezett intézmény <input type="checkbox"/> Termékenyítő központ <input type="checkbox"/> Engedélyezett aquakultúra telep <input type="checkbox"/>							
Embrióátültető csoport <input type="checkbox"/> Létesítmény <input type="checkbox"/> Egyéb <input type="checkbox"/>								Embrióátültető csoport <input type="checkbox"/> Létesítmény <input type="checkbox"/> Egyéb <input type="checkbox"/>							
Név								Név							
Jóváhagyási szám								Jóváhagyási szám							
Cím								Cím							
Írányítószám								Írányítószám							
I.14. Berakodás helye								I.15. Indulás dátuma és időpontja							
Írányítószám															
I.16. Szállítóeszköz								I.17. Szállító							
Repülőgép <input type="checkbox"/> Hajó <input type="checkbox"/> vagon <input type="checkbox"/>								Név							
Tehergépkocsi <input type="checkbox"/> Egyéb <input type="checkbox"/>								Jóváhagyási szám							
Azonosítás::								Cím							
Szám(ok):								Írányítószám							
I.21. Termékek hőmérséklete								I.20. Szám/Mennyiség							
Környezeti hőmérséklet <input type="checkbox"/> Hűtött <input type="checkbox"/> Fagyasztott <input type="checkbox"/>								I.22. Csomagok száma							
I.23. Konténer azonosítása/Plomba száma															
I.25. Az állatok/termékek felhasználási célja::															
I.26. Árutovábbítás harmadik országon keresztül <input type="checkbox"/>								I.27. Árutovábbítás tagállamon keresztül <input type="checkbox"/>							
Kilépési pont								Kód							
Belépési pont								BIP egység száma:							
I.28. Kivitel								I.29. Szállítás becsült időtartama							
Harmadik ország <input type="checkbox"/> ISO kód															
Kilépési pont								Kód							
I.30. Útvonalterv															
Igen <input type="checkbox"/> Nem <input type="checkbox"/>															
I.31. Az állatok azonosítása															

II. Egészségügyi információk		II.a. Bizonyítvány hivatkozási száma	II.b. Helyi hivatkozási szám
I, the undersigned official veterinarian, hereby certify that:			
II.1. The ovine/caprine animals(1) of the consignment described in Part I meet the following requirements:			
II.1.1. They are identified as provided for in Article 45(2) or (4) or Article 46(1) of Commission Delegated Regulation (EU) 2019/2035.			
II.1.2. They, for at least the 30 day period prior to the departure of the consignment, or since birth, if they are younger than 30 days of age,			
II.1.2.1. have been continuously resident in the establishment of origin;			
II.1.2.2. have not been in contact with kept ovine or caprine animals of a lower health status or subject to movement restrictions for animal health reasons;			
II.1.2.3. have not been in direct or indirect contact with kept animals that have entered the Union from a third country or territory during the 30 day period prior to the departure of the animals.			
II.1.3. They have not shown clinical signs or symptoms of diseases listed for ovine/caprine animals during the clinical examination which was carried out, within the 24 hour period prior to departure of the consignment, on (insert date dd/mm/yyyy).			
II.2. According to official information, the animals described in Part I meet the following health requirements:			
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 ovine/caprine animals.			
(2)	either [II.2.2.	They come from establishments free from infection with Brucella abortus, B. melitensis and B. suis without vaccination regarding ovine and caprine animals, and	
(2)		either [the establishments of origin are situated in a Member State or zone thereof with the status free from infection with Brucella abortus, B. melitensis and B. suis regarding the ovine and caprine population;]	
(2)		and/or [they have been subjected to a test for infection with Brucella abortus, B. melitensis and B. suis with one of the diagnostic methods provided for in Part 1 of Annex I to Commission Delegated Regulation (EU) 2020/688, carried out, with negative results, on a sample taken during the 30 day period prior to departure, and in the case of post-parturient females taken at least 30 days after parturition;]	
(2)		and/or [they are less than 6 months old;]	
(2)		and/or [they are castrated.]	
(2)	or [II.2.2.	They come from establishments free from infection with Brucella abortus, B. melitensis and B. suis with vaccination regarding ovine and caprine animals and they are moved to a Member State or zone thereof without the status free from infection with Brucella abortus, B. melitensis and B. suis regarding ovine and caprine animals.]	
(2)	either [II.2.3.	They are kept ovine animals and come from establishments in which infection with Mycobacterium tuberculosis complex (M. bovis, M. caprae and M. tuberculosis) has not been reported during the last 42 days prior to departure.]	
(2)	and/or [II.2.3.	They are kept caprine animals and come from establishments in which surveillance for infection with Mycobacterium tuberculosis complex (M. bovis, M. caprae and M. tuberculosis) has been carried out on the caprine animals kept on the establishments during at least the 12 month period prior to departure, as referred to in Article 15(3) of Delegated Regulation (EU) 2020/688.]	
II.2.4. 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.2.5. They come from establishments situated in an area of at least 150 km radius around those establishments in which infection with epizootic haemorrhagic disease virus has not been reported in kept animals of listed species for that disease during the last 2 years prior to departure.			
II.2.6. They come from establishments in which anthrax in ungulates has not been reported during the 15 day period prior to departure.			
II.2.7. They come from establishments in which surra (Trypanosoma evansi) has not been reported during the 30 day period prior to departure, and			
(2)		either [surra has not been reported in the establishments during the last 2 years prior to their departure.]	
(2)		or [surra has been reported during the last 2 years prior to departure, following the last outbreak the affected establishments have remained under movement restrictions until:	
		– the infected animals have been removed from the establishments, and	
		– the remaining animals on the establishments have been subjected to a test for surra (Trypanosoma evansi) with one of the diagnostic methods provided for in part 3 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken at least 6 months after the infected animals have been removed from the establishments.]	
(2)	[II.2.8.	They are kept uncastrated male ovine animals, and	
	–	come from establishments in which ovine epididymitis (Brucella ovis) has not been reported during the 12 month period prior to departure, and	
	–	have been subjected to a serological test for ovine epididymitis (Brucella ovis), carried out, with negative results, on a sample taken during the 30 day period prior to departure.]	
(2)	either [II.2.9.	They originate from a Member State or a zone free from infection with bluetongue virus (serotypes 1-24), where no case of infection with bluetongue virus (serotypes 1-24) has been confirmed during the last 24 months in the targeted animal population and have not been vaccinated with a live vaccine against infection with bluetongue virus (serotypes 1-24) in the 60 day period before the date of movement and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled.]	
(2)	and/or [II.2.9.	They originate from a Member State or a zone covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled, and they	
(2)	either [II.2.9.1.	have been kept in a Member State or zone seasonally free from infection with bluetongue virus (serotypes 1-24) in accordance with Article 40(3) of Commission Delegated Regulation (EU) 2020/689	
(2)		either [II.2.9.1.1. for at least 60 days prior to the date of movement]]	
(2)		and/or [II.2.9.1.2. for at least 28 days prior to the date of movement and have been subjected to a serological test, with negative results, carried out on samples collected at least 28 days following the entry date of the animal into the Member State or zone seasonally free from infection with bluetongue virus (serotypes 1-24)]]	
(2)		and/or [II.2.9.1.3. for at least 14 days prior to the date of movement and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the entry date of the animal into the Member State or zone seasonally free from infection with bluetongue virus (serotypes 1-24);]]	
(2)	and/or [II.2.9.2.	have been protected against attacks by the vectors during transportation to the place of destination and have been kept protected against attacks by vectors in a vector protected establishment	
(2)		either [II.2.9.2.1. for at least 60 days prior to the date of movement]]	
(2)		and/or [II.2.9.2.2. for at least 28 days prior to the date of movement and have been subjected to a serological test, with negative results, carried out on samples collected at least 28 days following the date of the commencement of the period of protection against attacks by vectors]]	

II: Egészségügyi információk		II.a. Bizonyítvány hivatkozási száma	II.b. Helyi hivatkozási szám
II. rész: Bizonyítvány	(2)	and/or [II.2.9.2.3.	for at least 14 days prior to the date of movement and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of the commencement of the period of protection against attacks by vectors;]]]
	(2)	and/or [II.2.9.3.	have been vaccinated against those serotypes from 1 to 24 of infection with bluetongue virus which were reported during the past 2 years in that Member State or zone and are within the immunity period guaranteed in the specifications of the vaccine and
	(2)	either [II.2.9.3.1.	have been vaccinated more than 60 days before the date of movement]]
	(2)	and/or [II.2.9.3.2.	have been vaccinated with an inactivated vaccine and subjected to a PCR test, with negative results on samples collected at least 14 days after the onset of the immunity set in the specifications of the vaccine;]]]
	(2)	and/or [II.2.9.4.	have been subjected with positive results to a serological test able to detect specific antibodies against all serotypes 1-24 of infection with bluetongue virus reported during the past 2 years in that Member State or zone and
	(2)	either [II.2.9.4.1.	the serological test has been carried out on samples collected at least 60 days before the date of movement]]
	(2)	and/or [II.2.9.4.2.	the serological test has been carried out on samples collected at least 30 days before the date of the movement and the animal has been subjected to a PCR test, with negative results, carried out on samples collected not earlier than 14 days before the date of movement;]]]
	(2)	and/or [II.2.9.	They originate from a Member State or a zone neither free from infection with bluetongue virus (serotypes 1-24) nor covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled, and they
	(2)	either [II.2.9.1.	have been protected against attacks by the vectors during transportation to the place of destination and have been kept protected against attacks by vectors in a vector protected establishment
	(2)	either [II.2.9.1.1.	for at least 60 days prior to the date of movement]]
	(2)	and/or [II.2.9.1.2.	for at least 28 days prior to the date of movement and have been subjected to a serological test, with negative results, carried out on samples collected at least 28 days following the date of the commencement of the period of protection against attacks by vectors]]
	(2)	and/or [II.2.9.1.3.	for at least 14 days prior to the date of movement and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of the commencement of the period of protection against attacks by vectors;]]]
	(2)	and/or [II.2.9.2.	have been kept for the 60 day period prior to departure in an establishment situated in a Member State or in an area of at least 150 km radius centred on the establishment, where surveillance in compliance with the requirements set out in Sections 1 and 2 of Chapter 1 of Part II of Annex V to Delegated Regulation (EU) 2020/689 has been carried out during that period, and
	(2)	either [II.2.9.2.1.	the animals have been vaccinated against those serotypes from 1 to 24 of infection with bluetongue virus which were reported during the past 2 years in an area of at least 150 km radius centred on the place where the animals were kept and are within the immunity period guaranteed in the specifications of the vaccine and
	(2)	either [II.2.9.2.1.1.	have been vaccinated more than 60 days before the date of movement]]]
	(2)	and/or [II.2.9.2.1.2.	have been vaccinated with an inactivated vaccine and subjected to a PCR test, with negative results on samples collected at least 14 days after the onset of the immunity set in the specifications of the vaccine;]]]]]
	(2)	and/or [II.2.9.2.2.	the animals have been immunised against those serotypes from 1 to 24 of infection with bluetongue virus which were reported during the past 2 years in an area of at least 150 km radius centred on the place where the animals were kept, and
	(2)	either [II.2.9.2.2.1.	the animals have been subjected with positive results to a serological test carried out on samples collected at least 60 days before the date of movement]]]
	(2)	and/or [II.2.9.2.2.2.	the animals have been subjected with positive results to a serological test carried out on samples collected at least 30 days before the date of the movement and to a PCR test, with negative results, carried out on samples collected not earlier than 14 days before the date of movement;]]]
	(2)	and/or [II.2.9.	They do not fulfil the requirements laid down in points 1 to 3 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689 and the competent authority of the Member State of origin authorised movement of those animals to another Member State or zone thereof
	(2)	either [II.2.9.1.	with the status free from infection with bluetongue virus (serotypes 1-24) and the Member State of destination has informed the Commission and the other Member States that such movement is authorised subject to the conditions referred to in Article 43(2)(a), (b) and (c) of Delegated Regulation (EU) 2020/689, and
	(2)	either [II.2.9.1.1.	point 5 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.1.2.	point 6 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.1.3.	point 7 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.1.4.	point 8 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
			the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]

II. Egészségügyi információk		II.a. Bizonyítvány hivatkozási száma	II.b. Helyi hivatkozási szám
II. rész: Bizonyítvány	(2)	and/or [II.2.9.2.	with an approved eradication program for infection with bluetongue virus (serotypes 1-24) and the Member State of destination has informed the Commission and the other Member States that such movement is authorised under the conditions referred to in Article 43(2)(a), (b) and (c) of Delegated Regulation (EU) 2020/689, and
	(2)		
	(2)	either [II.2.9.2.1.	point 5 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.2.2.	point 6 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.2.3.	point 7 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
	(2)	and/or [II.2.9.2.4.	point 8 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]
	(2)	and/or [II.2.9.3.	neither free from infection with bluetongue virus (serotypes 1-24) nor covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the Member State of destination has informed the Commission and the other Member States that such movement is authorised
	(2)	either [II.2.9.3.1.	without any conditions, and
	(2)	and/or [II.2.9.3.2.	subject to the conditions referred to in point 5 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
	(2)	and/or [II.2.9.3.3.	under the conditions referred to in point 6 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
	(2)	and/or [II.2.9.3.4.	under the conditions referred to in point 7 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
	(2)	and/or [II.2.9.3.5.	under the conditions referred to in point 8 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]
	(2)	either [II.2.10.	The animals are intended for a Member State or zone of a Member State listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 of the European Parliament and of the Council as having a negligible risk status for classical scrapie or for a Member State listed in point 3.2. of that Section as having an approved national scrapie control programme, and
	(2)		
	(2)	either [come from a holding situated in a Member State or zone of a Member State listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie.]	
	(2)	and/or [come from a holding recognised as having a negligible risk of classical scrapie in accordance with point 1.2. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with point 1.1. of that Section.]	
	(2)	and/or [come from a holding not subject to the measures laid down in points 3 and 4 of Chapter B of Annex VII to Regulation (EC) No 999/2001 and the animals are of the ovine species and are of the ARR/ARR prion protein genotype, or the animals are of the caprine species and carry at least one of the K222, D146 or S146 alleles.]	
	(2)	and/or [come from and are destined for an approved body, institute or centre as defined in Article 2(1)(c) of Council Directive 92/65/EEC.]	
	(2)	or [comply with the conditions set out in point 4.1.(d) of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001.]]	
	(2)	or [II.2.10.	The animals are for breeding and are intended for a Member State or zone of a Member State other than those listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie or other than those listed in point 3.2. of that Section as having an approved national scrapie control programme, and
(2)			
(2)	either [come from a holding situated in a Member State or zone of a Member State listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie.]		
(2)	and/or [come from a holding recognised as having a negligible risk of classical scrapie in accordance with point 1.2. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with point 1.1. of that Section.]		
(2)	and/or [come from a holding recognised as having a controlled risk of classical scrapie in accordance with point 1.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with point 1.1. of that Section.]		
(2)	and/or [come from a holding not subject to the measures laid down in points 3 and 4 of Chapter B of Annex VII to Regulation (EC) No 999/2001 and the animals are of the ovine species and are of the ARR/ARR prion protein genotype, or the animals are of the caprine species and carry at least one of the K222, D146 or S146 alleles.]		
(2)	and/or [come from and are destined for an approved body, institute or centre as defined in Article 2(1)(c) of Directive 92/65/EEC.]		
(2)	or [comply with the conditions set out in point 4.1.(d) of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001.]]		
(2)	or [II.2.10.	The animals are not for breeding and are intended for a Member State or zone of a Member State other than those listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie or other than those listed in point 3.2. of that Section as having an approved national scrapie control programme.]	
II.3.	To the best of my knowledge and as declared by the operator, the animals come from establishments where there were no abnormal mortalities with an undetermined cause.		
(2) [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 [II.4.2.	they were continuously present 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 1 of Part 3 of Annex II to Delegated Regulation (EU) 2020/686 in the period of the preceding 12 months prior to date of that movement; and]	

II. rész: Bizonyítvány

II. Egészségügyi információk		II.a. Bizonyítvány hivatkozási száma	II.b. Helyi hivatkozási szám
(2)	or [II.4.2. they were subjected, with negative results, to all tests referred to in point 1(c) and (d) of Chapter I of Part 3 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. the prior consent of the centre veterinarian of the semen collection centre of destination has been obtained by the operator; and		
	II.4.4. the means of transport used have been cleansed and disinfected before use.]		
II.5.	Arrangements are made to transport the consignment in accordance with Article 4 of Delegated Regulation (EU) 2020/688.		
II.6.	This certificate is valid for 10 days from the date of issuing. In the case of transport by waterway/sea of animals, the period of validity of the certificate may be extended by the duration of the journey by waterway/sea.		
(2)(3) [II.7.	Since leaving their establishments of origin and before arriving at this establishment approved for assembly operations, none of the animals of the consignment has undergone more than two assembly operations, and		
(2)	either [they come from their establishments of origin.]]		
(2)	or [at least one of the animals of the consignment has undergone one assembly operation on an approved establishment.]]		
(2)	or [at least one of the animals of the consignment has undergone two assembly operations on approved establishments.]]		
Animal welfare 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) (4)(5).			
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.			
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:			
Box reference I.11:	“Place of dispatch”: Indicate an establishment of the origin of the animals in the consignment or an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429 of the European Parliament and of the Council.		
Box reference I.12:	“Place of destination”: Indicate an establishment of the final destination of the consignment or an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429.		
Box reference I.17:	“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), based on which the animal health certificate for this consignment is issued in this establishment 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.		
Box reference I.30:	“Identification number”: Indicate identification codes of the animals in the consignment identified in accordance with Article 45(2) or (4) or Article 46(1) of Delegated Regulation (EU) 2019/2035.		
Part II:			
(1)	There can be one or more animals in the consignment.		
(2)	Delete if not applicable.		
(3)	Applicable in case the consignment is dispatched from the establishment approved for assembly operations.		
(4)	This statement does not exempt transporters from their obligation in accordance with Union rules in force in particular regarding the fitness to be transported.		
(5)	To be completed in case of consignment grouped in an establishment approved for assembly operations located in the Member State of transit.		
Hatósági állatorvos vagy hatósági ellenőr			
Név (nagybetűkkel):		Képesítés és beosztás:	
Helyi állat-egészségügyi egység:		A kapcsolódó helyi állat-egészségügyi egység száma::	
Dátum:		Aláírás:	
Pecset			

III. rész: Ellenőrzés

III.1. Az ellenőrzés dátuma

III.3. Iratellenőrzés:

Nem

Igen

EU sztandard

Megfelelő

Nem megfelelő

Kiegészítő garanciák

Megfelelő

Nem megfelelő

Nemzeti szabályozás

Megfelelő

Nem megfelelő

III.5. Fizikai ellenőrzés:

Nem

Az összes állapot ellenőrzve

Megfelelő

Nem megfelelő

III.7. Állatvédelmi ellenőrzés

Nem

Igen

Megfelelő

Nem megfelelő

III.8. Az állatvédelmi jogszabályok megsértése:

III.8.1. Szállító engedélye érvénytelen

III.8.2. Nem megfelelő szállítóeszköz

III.8.3. Rakodási sűrűség túllépése

III.8.4. Szállítási idő túllépése

III.8.5. Nem megfelelő itatás és takarmányozás

III.8.6. Rossz vagy hanyag bánásmód az állatokkal

III.8.7. Kiegészítő intézkedések nagy távolságra történő szállítás esetén

III.8.8. A járművezetők képzési bizonyítványa

III.8.9. A menetlevélben rögzített adatok

III.8.10. Egyéb

III.8.10.a Az utazás nem EU-s részének megfelelő megtervezése

III.8.10.b Szélsőséges hőmérsékletekkel

Átlagos terület

III.9. Az egészségügyi jogszabályok megsértése

III.9.1. Hiányzó/Érvénytelen bizonyítvány

III.9.2. Eltérés az okmányoktól

III.9.3. Nem engedélyezett ország

III.9.4. Nem engedélyezett régió/övezet

III.9.5. Tiltott állatfaj

III.9.6. Kiegészítő biztosítékok hiánya

III.9.7. Nem engedélyezett gazdaság

III.9.8. Beteg vagy betegsége gyanús állatok

III.9.9. Nem megfelelő vizsgálati eredmények

III.9.10. Hiányzó vagy nem jogszerű azonosítás

III.9.11. Nemzeti követelmények be nem tartása

III.9.12. rendeltetési hely címe érvénytelen

III.9.13. Egyéb

III.11. Korrekciós intézkedések

III.11.1. Késleltetett indulás

III.11.2. Átrakodási eljárás

III.11.3. Karantén

III.11.4. Levágás/kíméletes leölés

III.11.5. Tetemek/Termékek megsemmisítése

III.11.6. Szállítmány visszafordítása

III.11.7. Termékek kezelése

III.11.7.7. Termék egyéb célra történő felhasználása

Azonosítás:

III.12. Karanténból történő elbocsátás

III.12.1. Levágás/kíméletes leölés

III.12.2. Karanténból történő elbocsátás

III.13. Az ellenőrzés helye

Létesítmény

Telep

Gyűjtő központ

Kereskedő telephelye

Engedélyezett intézmény

Termékenyítő központ

Kikötő

Repülőtér

Kilépési pont

Útvonal

Egyéb

[hu] Control post

III.10. Szállítás hatása az állatokra

Elhullott állatok száma:

Értékelés:

Szállításra alkalmatlan állatok száma:

Értékelés:

Ellések vagy vetélések száma:

III.14. hatósági állatorvos vagy hatósági ellenőr

Helyi állat-egészségügyi egység

A kapcsolódó helyi állat-egészségügyi egység száma:

Név (nagybetűkkel):

Dátum:

Képesítés és beosztás

Aláírás:

hu

6/ 7

TERVEZÉS

1.1. SZÁLLÍTÁSSZERVEZŐ neve és címe (a) (b)		1.2. A szállításért az út során felelős személy neve			
		1.3. Telefon / Fax			
2. TELJES VÁRHATÓ IDŐTARTAM (órák / napok) / 2. VÁRHATÓ ÖSSZIDŐTARTAM (órák / napok)					
3.1. INDÍTÁSI hely és ország		4.1. RENDELTETÉSI hely és ország			
3.2. Dátum	3.3. Idő	4.2. Dátum	4.3. Idő		
5.1. Állatfaj	5.2. Állatok létszáma	5.3. Állat-egészségügyi bizonyítvány(ok) száma(i)			
5.4. A szállítmány becsült összsúlya (kg-ban megadva)		5.5. A szállítmány tervezett összterülete (m²-ben megadva)			
6. TERVEZETT PIHENTETÉSI, SZÁLLÍTÁSI VAGY KILÉPTETÉSI PONTOK LISTÁJA					
6.1. Azon helyek, ahol az állatok pihentetésre, pihenésre vagy átrakodásra kerülnek (beleértve a kiléptetési pontokat)	6.2. Érkezés		6.3. Időhossz (órákban mérve)	6.4. Szállítmányozó neve és engedélyezési száma (amennyiben az különbözik a szállításszervezőétől)	6.5. Azonosítás
	dátum	Idő			
Alulírott, szállításszervező kijelentem, hogy a fent említett szállítás lebonyolításáért vállalom a felelősséget, és az 1/2005/EK tanácsi rendelet rendelkezéseinek megfelelően megtettem a szükséges előkészületeket az állatok jólétének a szállítás teljes időtartama alatt történő biztosítására.					
8. Szállításszervező aláírása					

(a) "Szállításszervező": a fogalom-meghatározást lásd a 1/2005/EK tanácsi rendelet 2. cikkének (q) pontjában

(b) Amennyiben a szállításszervező egyben a szállítmányozó, akkor az engedélyének számát/ a jóváhagyási számot meg kell adni