EUROPEAN UNION INTRA

_							
	I.1. Consignor			I.2. IMSOC reference	I.2.a. Local reference		
	Name				I.3. Central Competent Authority		
	Address				nor contrar competent rationally		
	Country		ISO Code		I.4. Local Competent Authority		
of consignment	I.5. Consignee Name Address			establishment Name			
igis	Country		ISO Code	Address Approval			
Suc				Number			
JC				Country	ISO Code		
	I.7. Country of orig	gin	ISO Code	I.9. Country of destination	ISO Code		
I: Description	I.8. Region of origi	in	Code	I.10. Region of destination	Code		
Se(	I.11. Place of dispa	atch		I.12. Place of destination			
Ι:Ι	Name			Name			
Part	Address			Address			
P	Approval Number			Approval Number			
	Country		ISO Code	Country	ISO Code		
	-						
	I.13. Place of loadi	ing		I.14. Date and time of departure			
	Name Address						
	Approval						
	Number						
	Country		ISO Code				
	I.15. Means of Tra	nsport		I.16. Transporter			
	Mode	International transport document	Identification	Name Address			
				Approval Number			
				Country	ISO Code		
				I.17. Accompanying documents	150 couc		
				1 . , ,	Document Type Accompanying document reference Date of issue Country Place of issue		
				,			
	I.18. Transport co	nditions					
	Ambient 🗆		Chilled □	Frozen	Frozen 🗆		
	I.19. Container No	o / Seal No					
	I.20. Certified as	7	~ · □	m			
	Transformation L		Processing	Treatment $\square$			
	I.21. For transit th Third country	rough a third cour	try				
	Exit point			ISO Code BCP code			
	Entry point			BCP code			
	I.22. For transit th	rough Member Sta	te(s)	I.23. For export			
	Member State		ISO Code	Third country	ISO Code		
				Exit point	BCP code		
	I.24. Estimated jou	urney time		I.25. Journey Log			
	I.26. Total number	r of packages	I.27. Total quantity	I.28. Total net weight	I.28. Total gross weight		
	I.30. Description o	of consignment		l .	•		
	1. 30 PHARMACEUTICAL PRODUCTS						
	<b>3002</b> Human blood; animal blood prepared for therapeutic, prophylactic or diagnostic uses; antisera and other blood fractions and modified immunological products, whether or not obtained by means of biotechnological processes; vaccines, toxins, cultures of micro-organisms (excluding yeasts) and similar products						
	<b>300290</b> Other	L					

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	30029030 Animal blood prepared for therapeutic, prophylactic or diagnostic uses								
	#1. Commodity	Nature of commodity	Category	Manufacturing plant	Quantity				
	Species	Batch number	Package count	Net weight					
Ħ									
mei									
g									
nsi									
3									
סכ									
tio									
rip									
esc									
<u>:</u>									
Part I: Description of consignment									
Pē									

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## **EUROPEAN UNION**

EU —	ROPEAN UNION		esta	EDIIShed for	r the prevention and contr	of of certain fisted diseases
	II. Health information					
	I, the undersigned offi	icial veterin	arian. herek	ov certify the	at:	1
	• (1)either [II.1					kept animals:
	<ul> <li>(1)either [II.1 The animal by-products described in Part I were obtained from kept animals:</li> <li>(1)either [killed for the purpose of the prevention and control of:</li> </ul>					
		- (1)010101	<ul><li>(1)</li></ul>	l Larbone	_	
Part II: Certification	. *11			[ (introduce the name of the relevant category A disease) following the instructions of the competent authority in accordance with Regulation (EU) 2020/687 and are intended for processing by		
Sertif				o (1) either	[Method 1 to 5.]]]	
ij				o (1) or	[incineration.]]]	
art				<ul><li>(1) or</li><li>(1) or</li></ul>	[co-incineration.]]]	
Д			o (1)or	[ following	(introduce the name of the the instructions of the compe	,
				with emergency measures adopted by the Commission pursuant to Article 259 of Regulation (EU) 2016/429 and are intended for processing by		
				o (1) either	[Method 1 to 5.]]]	
				o (1) or	[incineration.]]]	
				o (1) or	[co-incineration.]]]	
o (1)or [not subject to killing by the competent authority for the and control of category A diseases or emerging diseases located in restricted zones established for the prevent diseases in accordance with			eases, kept in establishments ention and control of animal			
				o (1) either	[Delegated Regulation (EU) 2	2020/687,]
				o (1) or	[temporary special disease of the control of the co	control measures referred to EU) 2016/429,]
				o (1) or	[emergency measures adopt accordance with Article 259	ted by the Commission in of Regulation (EU) 2016/429,]
	and the animal by-products are moved from that restricted zones in compliance with the conditions set out in (2), for the					
				o (1) either	from aquatic animals as set	as set out in Chapter II of ilage of by-products obtained out in point K of Section 2 of egulation (EU) No 142/2011.]]]
				o (1) or	[processing or treatment by Annex X, Annex XI or Annex 142/2011.]]]	
				o (1) or	[production of processed pe referred to in Annex XIII to 142/2011.]]]	tfood, other than raw petfood Regulation (EU) No
				o (1) or	[transformation into compo Section 1 of Chapter III of An 142/2011.]]]	st or biogas referred to in nnex V to Regulation (EU) No
	o (1) or [II.1	The animal by-products described in Part I were obtained from wild animals of listed species found dead or killed for the purpose of the prevention and control of (introduce the name of the relevant category A disease) following the instructions of the competent authority (3) in accordance with Article 64(2)(c) of Delegated Regulation (EU) 2020/687 and are intended for processing by				
		o (1) either	[Method 1	to 5.]]		

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## 2023/1521 (2021/1699) Animal by-products from restricted zones established for the prevention and control of certain listed diseases $\,$

	II. Health information								
			o (1) or	[incineration.]]					
Part II: Certification			o (1) or	[co-incineration.]]					
	(1)	either	□ [II.2 Unprocessed animal by-products other than hides and skins, hides and skins, colostrum, milk and dairy products, indicate as appropriate) obtained from bovine animals kept in vaccination zone I in relation to emergency protective vaccination against lumpy skin disease, in compliance with Article 13(3) of, and Annex IX, Part 3, points (3.5) and (3.7), to Commission Delegated Regulation (EU) 2023/361.]						
	(1)	or	□ [II.2 Unprocessed animal by-products other than hides and skins, hides and skins, colostrum, milk and dairy products, indicate as appropriate) obtained from bovine animals kept in vaccination zone II in relation to emergency protective vaccination against lumpy skin disease, in compliance with Article 13(3) of, and Annex IX, Part 3, points (3.6) and (3.7), to Commission Delegated Regulation (EU) 2023/361.]						
	Notes								
	Part I:								
	•	Box refere	nce I.9 and	I.11: delete as appropria	ate.				
	•	Box refere	nce I.12, I.1	3 and I.17: approval nui	mber or registration number.				
	•	Box refere	nce I.14: co	mplete if different from	'I.1. Consignor'.				
	•	Box refere	nce I.25: for	r 'processing', 'treatmen	t', 'transformation'				
	•			iture of commodity: 'AB Category 1','Category 2',	Ps referred to in Article 22(5) o 'Category 3'	of Delegated Regulation (EU)			
	Part II:								
	(1)	Delete as a	ppropriate.						
	(2)	Insert the number of the relevant article(s), the title and date of publication in the Official Journal of the European Union of the relevant legal act adopted by the Commission providing those conditions or the reference to the legal act or instruction approved and made public by the competent authority providing for those conditions.							
	(3)	See special legislation on the prevention of transmissible diseases.							
		icer/Official ve	terinarian						
	Name (in capi Date of signate Stamp				Qualification and title Signature				
						1			

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