	I.1. Consignor						I.2. IMSOC Ref	erence			
	Name						I.2.a. Local Reference				
	Address										
	Country ISO Code										
ľ	I.5. Consignee					I.3. Central co	mpetent a	uthority			
Ľ	Name						I.4. Local com	petent aut	hority		
<u>e</u>	Address										
E	Country			ISO Coc	le						
Part I : Details of consignment	I.7. Country of orig	gin				ISO Code	I.9. Country of	f destinatio	on		ISO Code
ы Ц	I.8. Region of origi	n				Code	I.10. Region of	destinatio	on		
s							I.12. Place of destination				
tai	Name						Name				
Pe	Address						Address				
	Approval Number	r					Approval Nu	mber			
Ľ	Country			ISO	Code		Country			ISO Code	
2	I.13. Place of Loadi	ing					I.14. Date and	time of de	eparture		
	Name										
	Address										
	Approval Number	r									
	Country			ISO	Code						
	I.15. Means of Trai	nsport					I.16 Entry Poi	nt			
	Mode	Internatio	nal	Identificati	on						
		transport document									
							-				
	I.18. Transport cor						I.17. Accompa	nying doc	uments		
	Controlled Ambient 🗆 Frozen 🗆 Chilled 🗆						Commercial	Commercial document Date of issue			
	I I I I I I I						reference		Date t	JI 1350C	
						Country		Place issue	of		
ľ	I.19. Container No / Seal No						I				
ŀ	I.20. Certified as										
	Pharmaceutical us	se 🗆		Other 🗌			Technical use				
ŀ	I.21. For transit thi	rough a thi	rd count				1.22 For trans	it through	Member State(s)		
	Country	rough a thi		ISO Code			1.22. FOT traits	n nirougn	Member State(s)		
	EU Exit			BCP code							
	Authority EU Entry						Country		ISO Co	ode	
ļ	Authority			BCP code							
	I.23. Total number	of package	es		1.25. Tota	l net weight			I.25. Total gross we	eight	
	I.28. Description of	f consignm	ent								
	1. 02 MEAT AND E										
	0206 Edible offa				eep, goats,	horses, asses, 1	nules or hinnie	s, fresh, cł	nilled or frozen		
	020610 Of bo 02061098 c			i or chilled	skirt						
	Commodity		Species		SKILL	Slaughterhou		Manufac	turing plant	Package coun	+
	Commonly		species	5		Slaughterhou	56	Mailulac		Package could	l
	Net weight						Batch number			1	
1											
1											

	II. Health infor	rmation									
	I, the undersigned, official veterinarian(8) of the (insert name of competent veterinary authority of the Member State of the EU), after due inquiry and to the best of my knowledge, do hereby certify that the animal by-products described above:										
	II.1.	Consists exclusively of products taken from animals slaughtered within the EU member state(s) of: (insert name of country (ies) here)									
t II: Certificati	II.2.	legally imp as free of th from which	Are all derived from animals o born and raised in (insert country name here) or o egally imported from (insert country name here), which is/are recognized by (2) Canada as free of the following diseases of concern (listed in notes by susceptible species) OR o that the animals from which the products are derived were present in the country of slaughter (insert country name here) without restrictions (outside of quarantine) for the (3)time required by species of prigin;								
	II.3.		nales", or th					icinal purposes" or "À des as pharmaceutiques", as the			
	II.4.	The product was manufactured in an establishment that was operating under a Hazard Analysis Critical Control Point (HACCP) principles based system determined by the Canadian Food Inspection Agency (CFIA) to be equivalent to the Food Safety Enhancement Program (FSEP) established by the Agency and from a country and an establishment approved to export both edible and inedible meat products to Canada by CFIA;									
II.5. Has been prepared exclusively with the following animal by-products (na type):							oducts (nam	e species of origin and tissue			
	II.6.	None of the animals from which the products/by-products are derived were suspected or confirmed of the following disease(s), nor any other reportable disease as defined by Canada (either through confirmatory negative testing, or "suspect" testing): anthrax, foot-and-mouth disease, rinderpest or Bovine Spongiform Encephalopathy and none of the animals from which the products/by-products are derived were under any official restrictions by the competent veterinary authority for any serious epizootic (4)disease to which the species from which the by-product was derived is susceptible and that can be transmitted by the by-product (see Notes II.1 and II. below);									
	□ [(5)II.7.)II.7. Additional certification for either swine or lagomorph origin products:									
		□ [(5)II.7.1.	For products derived from swine (or containing a mixture of products that include swine origin material), the animals from which the products were derived:								
			II.7.1.1.	Showed no	o clinical sigr	n of Aujeszky Dise	ase (AD) on	the day of slaughter;			
			II.7.1.2.			ict with animals fi ransport to and at		hments not considered free r;			
			II.7.1.3.	Have not b	een vaccina	ted against AD; ar	ıd				
		_	designa Annex I three ki		ad no contact with any swine or swine products that were in an area not esignated free from AD in accordance with Commission Decision 2008/185/EC nnex I as amended, or the premises of origin and all farms within a radius of aree kilometres were free from any clinical or epidemiological evidence of AD r a period of 12 months prior to collection of the products/by-products.]						
		□ [(5)II.7.2.	For products derived from lagomorphs, the animals:								
			II.7.2.1.	Showed no clinical sign of myxomatosis on the day of shipment to the approved abattoir; and							
			II.7.2.2.	-	ent where n	or for the six mon o case of myxoma	-	slaughter, in an fficially reported during that			
			(1)either	○ [7.2.2.1.	haemorrha disease has vaccinatior	not been present has been carried	, shown, by for at least l out in the p	free from rabbit serological testing, that the one year, and that no previous 12 months and such by the competent veterinary			

						manado	are for export to carlada				
	II. Health info	rmation									
			(1)or	。 [II.7.2.2.2.	during the II.4.); and s	in an establishment where no o 60 days prior to transport to th howed no lesions of RHD at pos nt does not include any pelts fr	e approved abattoir (as per st-mortem inspections; and				
	II.8.	All the anim	nals by-pro	ducts which	are covered	l by the present certificate:					
Part II: Certification		II.8.1.	inspection, veterinaria	, both carrie an(8) or an c	ed out by an official veter	ortem inspection and were sul inspector under the supervisio inarian(8) of the competent vet Canada by CFIA;	n of an official				
ບຶ ::		II.8.2.	were all:								
ヨピ		(1)either	○ [II.8.2.1	stunned (h	umanely rei	ndered unconscious) before sla	ughter;]				
Par		(1)or	○ [II.8.2. 2	 [II.8.2. 2 (In the case of animals slaughtered in conformance with ritual slaughter procedures) The products within this shipment and covered by this certifica are derived from animals that received (Kosher, Halal) slaughter, as based documentation provided by religious authorities or by (Kosher, Halal) label declaration and the poultry products within this shipment and covered by t certificate are derived from birds slaughtered by rapid decapitation withou prior electrical (6)stunning;] 							
	II.9.	The contain proof; and	ner in whicl	n the produ	cts/by-produ	cts are being shipped is comple	etely enclosed and leak-				
	(1)either	○ [II.10.	The produ	cts contain 1	no ruminant	origin products/by-products;]					
	(1)or	∘ [II.11.	were slaug slaughtere into the an of the anim	The product contains products/by-products of ruminant origin, and the ruminant animals were slaughtered as per BSE slaughter process. They were not subjected, before being slaughtered, to a stunning process in which a device is used to inject compressed air or gas into the animal's cranial cavity; nor to a pithing process involving laceration, after stunning of the animal, of the animal's central nervous tissue by means of an elongated rod-shaped instrument that is introduced into the animal's cranial cavity.]							
	AND										
	(1)either	○ [II.11.1.		s) name(s)) v		als born and raised for the first •) a negligible risk for BSE coun	-				
	(1)or	○ [II.11.2.	country(ies any of the	s) name(s)) a	and legally i	nt animals both born and raise mported into (insert country na bovine animals:					
		(1)either	。 [II.11.2.1.	root gangli ages (if pro for BSE cou	ia of cattle ag oduct contain untries as re	inal ganglia, eyes, palatine tons ged 30 months or older; and the ns any tissues from ruminant a cognized by Canada but no tiss BSE (7)countries);]	e distal ileum of cattle of all nimals from controlled risk				
		(1)or	○ [II.11.2.2.	cord and the transverse sacrum), frages of box	he vertebral processes of com bovine a vine animals	e skull, the brain, trigeminal ga column, (excluding the verteb f the thoracic and lumbar verte animals aged 12 months or olde (if product contains any tissue k for BSE countries as recogniz	rae of the tail, the ebrae, and the wings of the er and the distal ileum of all s from ruminant animals				
	Notes										
Part I											
	• Box reference I.6.: Indicate CFIA permit number if applicable (if for end use Technical or other)										
	· Box r approval n		1.: indicate	the shipping	g establishm	ent and the Member State com	petent veterinary authority				
		reference I.1	2.								
		Canada Esta				atory when the end use is phar	maceutical or cosmetic. If				

no Health Canada Establishment license number is provided then the certificate must not be issued.

(CA) Raw animal products and by-products for pharmaceutical manufacture for export to Canada

EUR	ODE	ΔΝ	TINI	ON
LON		ALV.	UINI	UIV.

	II. Health information								
	o The EL must be verified on the Health Canada web site Drug Establishment Licences Listing. Searches may be done either by a) Licence number (Site optional), b) Company name, c) Activity and/or Province. http://webprod5.hc-sc.gc.ca/el-le/prepare-search-recherche-del-leppp.do?lang=eng								
Certif	• Box reference I.22.: Commodities certified for must identify the end use. Note that "Pharmaceutical use" includes also cosmetic manufacture, "Technical use" applies to commodities not intended for human or animal consumption, and " Other" is intended for purposes not listed elsewhere in this classification. Where "Technical use . or "Other" is chosen, a CFIA Import Permit is required. The Import Permit number must be clearly indicated in Box Reference I. 6								
	• Box reference I.25 Identification of the commodities. This is where the Approval number of establishments eligible (approved by CFIA Meat Programs) for export to Canada of edible meat products should be listed. Based on Annex A certification requirements as negotiated between CFIA and EU SANTE. For more details visit: http://www.inspection.gc.ca/food/meat-and-poultry-products/manual-of-procedures/chapter-10/annex-a/european-union/eng/1336803459318/1336803636873								
	Custom code and title: Use the appropriate Harmonised Sy	vstem (HS) code.							
	Part II								
	• (1) Delete as appropriate.								
	• (2) Under the EU Canada Veterinary agreement, for t country or zone freedom, CFIA recognizes EU disease erad directives. When a disease incursion occurs in a previousl full freedom is only regained when the control zone meets	lication and control zones wh y free area, and Canada has r	en they are published in EU ecognized the control zones,						
	Diseases of concern for Canada for animal products & by-p	products covered by this certi	ficate are:						
	o For poultry: Notifiable Avian Influenza and Newcastle	disease							
	o For ruminants: Contagious bovine pleuropneumonia; I des petits ruminants; Rift valley fever and Sheep pox and g								
	o For swine: African swine fever; Classical swine fever (I disease; Vesicular Stomatitis	Hog cholera); Foot and Mouth	Disease; Swine vesicular						
	o For Horses &other equids: African horse sickness and V	Vesicular Stomatitis							
	o For lagomorphs (commercially reared): rabbit viral ha	emorrhagic disease and myxa	amotisi						
	o Vesicular Stomatitis - EU zoning is recognized by Canad CCVA and that zone recognized by Canada (once the EC zo an area or zone with an active outbreak of the disease, no declared outbreak, control or monitoring zone for vesicula	ning decision is published) ar r have been in direct contact	nd the animals are not from						
	o Rabbit viral haemorrhagic disease and myxamotosis. A establishment (farm) freedom is acceptable provided that								
	List of Countries which Canada has recognized as being fr	ee from the certain diseases:							
	Terrestrial Animal Health Status By Disease: (note after di- recognition will be indicated by removal of the zoning not	-	free zones, full freedom						
	http://www.inspection.gc.ca/animals/terrestrial-animals/di disease/eng/1306649804251/1306649991822	iseases/status-by-							
	Terrestrial Animal Health Status by Country: (note after di recognition will be indicated by removal of the zoning not		free zones, full freedom						
	http://www.inspection.gc.ca/animals/terrestrial-animals/di country/eng/1306648587424/1306649135327	iseases/status-by-							
	(3) Time requirements for an animal to be considere with animals of the importing country without restriction		ock (imported and housed						
	For avian (poultry & ratite, or other): 21 days;								
	For ruminants: 90 days;								
	For swine: 90 days;								
	For horses & other equids: 60 days;								
	For lagomorphs (commercially reared): 60 days.								

II. Health information

• (4) Note this section refers only to those diseases to which the species of animals from which the products/byproducts are derived, is susceptible (as identified above).

(5) Keep if appropriate.

• (6) The option for rapid decapitation without prior stunning is to be applied ONLY to poultry and not ratites (strike out inapplicable.

• (7) Canada publishes a list of BSE Categorization of countries on the Bovine Spongiform Encephalopathy Import Policy for Bovine Animals and Their Products and By-Products TAHD-DSAT-IE-2005-9-5 http://www.inspection.gc.ca/animals/terrestrial-animals/imports/policies/general/2005-

9/eng/1321066760292/1426255335689 The CFIA BSE import policy for bovine animals and animal products, animal by-products, germplasm, animal food, meat, meat by-products and veterinary biologics, of bovine origin, adheres

closely to the recommendations of the World Organisation for Animal Health (OIE) and the list of negligible risk and controlled risk for BSE countries is updated regularly. The updates are initiated following the updating of lists of negligible and controlled risk for BSE countries by the OIE at the annual general council.

• (8) The official veterinarian who signs this certificate must meet the Canadian definition of "official veterinarian"; which is defined under the Health of Animals Regulations as a veterinarian employed by the government of that country (vétérinaire officiel).

Further clarifications for Part I

CFIA Import Permit:

The Import Permit number in box I.6 is required if end use anything other than pharmaceutical or cosmetic as indicated in I.22.

Establishment Approval Number:

Approval number in box I.11 is the EU Member State Central Competent Authority approval number of the establishment from which the product is being exported and must be verified on the applicable CFIA database.

Health Canada Establishment Licence Number:

Approval Number -2 means the Health Canada Establishment Licence Number (EL) and must be indicated in box I.12 The EL must be verified on the Health Canada web site Drug Establishment Licences Listing. Searches may be done either by a) Licence number (Site optional), b) Company name, c) Activity and/or Province. http://webprod5.hc-sc.gc.ca/el-le/prepare-search-recherche-del-leppp.do?lang=eng

• Description of container in box I.21 is required in detail only where a standard shipping container (large metal box of a standard design and size used for the transport of goods by road, rail, sea, or air) is not used and there is no seal number and no container number. Example – Products shipped in sealed impermeable carton with plastic liners with (insert company name "A", brand name or commodity name on label "Porcine Pancreas Insul-Z" and description of the tamper evident mechanism (such as tamper evident tape or labelling).

• Note that within the signature block below the term official inspector must be struck out & initialled by the signing official veterinarian and the name of competent authority of named of Member State must appear as well as the stamp.

Name (in capital letters)
Date of signature
Stamp

Certifying Officer

Qualification and title Signature