

Part I : Details of consignment	I.1. Consignor Name Address Country		ISO Code		I.2. IMSOC Reference Specimen not to be used for exports from EU		I.2.a. Local Reference	
	I.5. Consignee Name Address Country		ISO Code		I.3. Central competent authority		I.4. Local competent authority	
	I.7. Country of origin		ISO Code		I.9. Country of destination		ISO Code	
	I.8. Region of origin		Code		I.10. Region of destination		Code	
	I.11. Place of Dispatch Name Address Approval Number Country		ISO Code		I.12. Place of destination Name Address Approval Number Country		ISO Code	
	I.13. Place of Loading Name Address Approval Number Country		ISO Code		I.14. Date and time of departure			
	I.15. Means of Transport				I.16 Entry Point			
	Mode	International transport document	Identification					
	I.18. Transport conditions Ambient <input type="checkbox"/> Controlled temperature <input type="checkbox"/> Frozen <input type="checkbox"/> Chilled <input type="checkbox"/>				I.17. Accompanying documents Commercial document reference Date of issue Country Place of issue			
	I.19. Container No / Seal No							
I.20. Certified as Transhumance <input type="checkbox"/> Pollination <input type="checkbox"/> Animal Feedingstuff <input type="checkbox"/> Pharmaceutical use <input type="checkbox"/> Game Restocking <input type="checkbox"/> Fattening <input type="checkbox"/> Registered equidae <input type="checkbox"/> Quarantine <input type="checkbox"/> Artificial reproduction <input type="checkbox"/> Training <input type="checkbox"/> Further process <input type="checkbox"/> Racing <input type="checkbox"/> Production of petfood <input type="checkbox"/> Breeding and production <input type="checkbox"/> Consignments according to Regulation No 999/2001 <input type="checkbox"/> Competition <input type="checkbox"/> Storage <input type="checkbox"/> Breeding <input type="checkbox"/> Ornamental use/research <input type="checkbox"/> Laboratory <input type="checkbox"/> Rodent food <input type="checkbox"/> Pets <input type="checkbox"/> Other <input type="checkbox"/> Production <input type="checkbox"/> Technical use <input type="checkbox"/> Ornamental bird food <input type="checkbox"/> Relaying <input type="checkbox"/> Slaughter <input type="checkbox"/> Approved Bodies <input type="checkbox"/> Human consumption <input type="checkbox"/> Pet food <input type="checkbox"/> Sales <input type="checkbox"/> Organic fertilizers <input type="checkbox"/> Unregistered equidae <input type="checkbox"/> Circus exhibition <input type="checkbox"/> Category 3 fish oil/fish meal for detoxification according to Regulation 2015/786 <input type="checkbox"/>								
I.21. For transit through a third country <input type="checkbox"/> Country _____ ISO Code _____ EU Exit Authority _____ BCP code _____ EU Entry Authority _____ BCP code _____				I.22. For transit through Member State(s) <input type="checkbox"/> Country _____ ISO Code _____				
I.23. Total number of packages		I.24. Total quantity		I.25. Total net weight		I.25. Total gross weight		

Part I : Details of consignment	I.28. Description of consignment				
	1. 04 DAIRY PRODUCE; BIRDS' EGGS; NATURAL HONEY; EDIBLE PRODUCTS OF ANIMAL ORIGIN, NOT ELSEWHERE SPECIFIED OR INCLUDED				
	0401 Milk and cream, not concentrated nor containing added sugar or other sweetening matter				
	Commodity	Species	Quantity	Batch number	Manufacturing plant
Cold store	Cutting plant	Date of freezing	Date of production	Date of slaughter	
Net weight	Product Description	Package count	Identification mark		

SPECIMEN

Part II: Certification	II. Health information	
II.1.	<p>Animal Health Attestation</p> <p>I, the undersigned official veterinarian, declare that I am aware of the relevant provisions of Directive 2002/99/EC and of Regulation (EC) No 853/2004 and hereby certify that the raw milk described above has been obtained from animals:</p> <ul style="list-style-type: none"> (a) under the control of the official veterinary service, (b) which were in a country or part thereof that has been free of foot-and-mouth disease and of rinderpest for a period of at least 12 months prior to the date of this certificate, and where vaccination against foot-and-mouth disease has not been carried out during that period, (c) belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest, and (d) subject to regular veterinary inspections to ensure that they satisfy the animal health conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004 and in Directive 2002/99/EC; 	
II.2.	<p>Public Health attestation</p> <p>I, the undersigned official inspector, declare that I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the raw milk described above was produced in accordance with those provisions, in particular that:</p> <ul style="list-style-type: none"> (a) it comes from holdings registered in accordance with Regulation (EC) No 852/2004 and checked in accordance with Annex IV to Regulation (EC) No 854/2004, (b) it was produced, collected, cooled, stored and transported in accordance with the hygiene conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004, (c) it meets the plate and somatic cell count criteria laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004, (d) the guarantees on the residues status of raw milk provided by the monitoring plans for the detection of residues or substances submitted in accordance with Council Directive 96/23/EC, and in particular, Article 29 thereof, are fulfilled; (e) pursuant to testing for residues of antibacterial drugs carried out by the food business operator in accordance with the requirements of Annex III, Section IX, Chapter 1, Part III, point 4 of Regulation (EC) No 853/2004, it complies with the maximum residue limits for residues of antibacterial veterinary medicinal products laid down in the Annex to Regulation (EU) No 37/2010; (f) it has been produced under conditions guaranteeing compliance with the maximum residue levels for pesticides laid down in Regulation (EC) No 396/2005, and maximum levels for contaminants laid down in Regulation (EC) No 1881/2006. 	

Part II: Certification	II. Health information								
	<p>Notes</p> <p>(*) Those countries subject to the transitional import arrangements include: an EU member State; Iceland; Liechtenstein; Norway and Switzerland.</p> <p>References to Great Britain in this certificate include Channel Islands and Isle of Man.</p> <p>References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018).</p> <p>This certificate is intended for raw milk from third countries or parts thereof authorised in column A of Annex 1 to Regulation (EU) No 605/2010 intended for further processing in Great Britain before being used for human consumption.</p> <p>Part I:</p> <p>Box reference I.7: Provide name and ISO code of the country or part thereof as appearing in Annex I to Regulation (EU) No 605/2010.</p> <p>Box reference I.11: Name, address and approval number of the establishment of dispatch.</p> <p>Box reference I.15: Registration number (railway wagons or container and road vehicle), flight number (aircraft) or name (ship). In case of unloading and reloading, the consignor must inform the border control post of introduction into Great Britain.</p> <p>Box reference I.16: Do not use this box until the end of the transitional staging period.</p> <p>Box reference I.19: Use the appropriate Harmonised System (HS) code under the following headings: 04.01; 04.02 or 04.03.</p> <p>Box reference I.20: Indicate total gross weight and total net weight.</p> <p>Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.</p> <p>Box reference I.28: Manufacturing plant: introduce the approval number of the production holding(s), collection centre or standardization centre approved for exportation to Great Britain.</p> <p>Part II:</p> <p>The colour of the signature shall be different to that of the printing. The same rule applies to stamps other than those embossed or watermark.</p>								
	<p>Certifying Officer</p> <table border="0"> <tr> <td>Name (in capital letters)</td> <td>Qualification and title</td> </tr> <tr> <td>Date of signature</td> <td>Signature</td> </tr> <tr> <td>Stamp</td> <td></td> </tr> </table>				Name (in capital letters)	Qualification and title	Date of signature	Signature	Stamp
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Date of signature	Signature								
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Část I	I.1. Odesílatel Název Adresa Země Kód ISO		I.2. Referenční číslo IMSOC Specimen not to be used for exports from EU I.2.a. Local Reference																
	I.5. Příjemce Název Adresa Země Kód ISO		I.3. Ústřední příslušný orgán I.4. Local competent authority																
	I.7. Země původu Kód ISO		I.9. Country of destination Kód ISO																
	I.8. Region of origin Kód		I.10. Region určení Kód																
	I.11. Place of Dispatch Název Adresa Číslo schválení Země Kód ISO		I.12. Místo určení Název Adresa Číslo schválení Země Kód ISO																
	I.13. Místo nakládky Název Adresa Číslo schválení Země Kód ISO		I.14. Date and time of departure																
	I.15. Dopravní prostředky		I.16. Entry Point																
	<table border="1"> <thead> <tr> <th>Typ</th> <th>Doklad</th> <th>Identifikace</th> </tr> </thead> <tbody> <tr><td> </td><td> </td><td> </td></tr> <tr><td> </td><td> </td><td> </td></tr> <tr><td> </td><td> </td><td> </td></tr> <tr><td> </td><td> </td><td> </td></tr> </tbody> </table>		Typ	Doklad	Identifikace														
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I.18. Transport conditions Okolní <input type="checkbox"/> Controlled temperature <input type="checkbox"/> Zmrazené <input type="checkbox"/> Chlazený <input type="checkbox"/>		I.17. Průvodní doklady Referenční číslo obchodního do k ladu Datum vydání Země Místo vydání																	
I.19. Č. kontejneru / č. plomby																			
I.20. Certified as Transhumance <input type="checkbox"/> Pollination <input type="checkbox"/> Krmivo pro zvířata <input type="checkbox"/> Použití pro farmaceutické účely <input type="checkbox"/> Game Restocking <input type="checkbox"/> Výkrmová <input type="checkbox"/> Evidování koňovité <input type="checkbox"/> Karanténa <input type="checkbox"/> Umělé rozmnožování <input type="checkbox"/> Training <input type="checkbox"/> Další postup <input type="checkbox"/> Racing <input type="checkbox"/> Production of petfood <input type="checkbox"/> Breeding and production <input type="checkbox"/> Consignments according to Regulation No 999/2001 <input type="checkbox"/> Competition <input type="checkbox"/> Storage <input type="checkbox"/> Breeding <input type="checkbox"/> Ornamental use/research <input type="checkbox"/> Laboratory <input type="checkbox"/> Rodent food <input type="checkbox"/> Zvířata v zájmovém chovu <input type="checkbox"/> Jiné <input type="checkbox"/> Production <input type="checkbox"/> Technické použití <input type="checkbox"/> Ornamental bird food <input type="checkbox"/> Sádkování <input type="checkbox"/> Porážka <input type="checkbox"/> Schválené orgány <input type="checkbox"/> Lidská spotřeba <input type="checkbox"/> Krmivo pro domácí zvířata <input type="checkbox"/> Sales <input type="checkbox"/> Organic fertilizers <input type="checkbox"/> Unregistered equidae <input type="checkbox"/> Cirkus/výstava <input type="checkbox"/> Category 3 fish oil/fish meal for detoxification according to Regulation 2015/786 <input type="checkbox"/>																			
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I.23. Celkový počet balení	I.24. Celkové množství	I.25. Celková čistá hmotnost	I.25. Celková hrubá hmotnost																

I.28. Description of consignment

1. 04 MLÉKO A MLÉČNÉ VÝROBKY; PTAČÍ VEJCE; PŘÍRODNÍ MED; JEDLÉ PRODUKTY ŽIVOČIŠNÉHO PŮVODU, JINDE NEUVEDENÉ ANI NEZAHRNUTÉ**0401** Mléko a smetana, nezahuštěné, neobsahující přidaný cukr ani jiná sladidla

Komodita	Druh	Množství	Číslo šarže	Výrobní zařízení
Chladírenské zařízení	Bourárna	Datum zmrazení	Datum výroby	Datum porážky
Čistá hmotnost	Product Description	Počet balení	Identifikační značka	

Část I

SPECIMEN

Part II: Certification	II. Informace týkající se zdraví		
	<p>II.1. Animal Health Attestation</p> <p>I, the undersigned official veterinarian, declare that I am aware of the relevant provisions of Directive 2002/99/EC and of Regulation (EC) No 853/2004 and hereby certify that the raw milk described above has been obtained from animals:</p> <ul style="list-style-type: none"> (a) under the control of the official veterinary service, (b) which were in a country or part thereof that has been free of foot-and-mouth disease and of rinderpest for a period of at least 12 months prior to the date of this certificate, and where vaccination against foot-and-mouth disease has not been carried out during that period, (c) belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest, and (d) subject to regular veterinary inspections to ensure that they satisfy the animal health conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004 and in Directive 2002/99/EC; <p>II.2. Public Health attestation</p> <p>I, the undersigned official inspector, declare that I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the raw milk described above was produced in accordance with those provisions, in particular that:</p> <ul style="list-style-type: none"> (a) it comes from holdings registered in accordance with Regulation (EC) No 852/2004 and checked in accordance with Annex IV to Regulation (EC) No 854/2004, (b) it was produced, collected, cooled, stored and transported in accordance with the hygiene conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004, (c) it meets the plate and somatic cell count criteria laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004, (d) the guarantees on the residues status of raw milk provided by the monitoring plans for the detection of residues or substances submitted in accordance with Council Directive 96/23/EC, and in particular, Article 29 thereof, are fulfilled; (e) pursuant to testing for residues of antibacterial drugs carried out by the food business operator in accordance with the requirements of Annex III, Section IX, Chapter 1, Part III, point 4 of Regulation (EC) No 853/2004, it complies with the maximum residue limits for residues of antibacterial veterinary medicinal products laid down in the Annex to Regulation (EU) No 37/2010; (f) it has been produced under conditions guaranteeing compliance with the maximum residue levels for pesticides laid down in Regulation (EC) No 396/2005, and maximum levels for contaminants laid down in Regulation (EC) No 1881/2006. 		

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