Health Certificate for the Export of Bovine Semen from Switzerland to Kosovo (D2) I.2. Certificate reference number Name Address Part I: Details of consignment presented I.3. Central Competent Authority Postal code / Region I.4. Local Competent Authority I.6. No.(s) of related original certificates No.(s) of accompanying documents .5. Consignee Postal code / Region I.7. Dealer Name Approval number I.8.Country of origin ISO code I.9. Region of origin Code I.10. Country of destination ISO code I.11. Region of destination Code I.12. Place of origin / Place of harvest I.13. Place of destination Holding Holding Assembly centre Dealer's premise Assembly centre Dealer's premise Approved body Semen centre Approved aquaculture holding Approved body Semen centre Approved aquaculture holding Embryo team Establishment Other Establishment Other Approval number Approval number Address Address Postal code / Region Postal code / Region I.14. Place of loading Postal code / Region I.16. Means of transport Aeroplane Ship Railway wagon Other Road vehicle Identification: Number(s): Member state I.18. Animal species/Product I.19. Commodity code (CN code) I.20. Number/Quantity I.21 Temperature of products I.22. Number of packages Chilled .24.Type of packaging I.25. Animals certified as/products certified for:: Approved bodies Breeding Slaughter Transhumance Fattening Artificial reproduction Pets Registered equidae Game restocking Animal feedingstuff Other Human consumption I.26. Transit through 3rd country I.27. Transit through Member states 3rd country ISO code ISO code Member state ISO code Member state BIP unit no. Entry point Member state ISO code I.28. Export I.29. Estimated journey time ISO code Exit point Code I.30. Route plan Yes No I.31. Identification of the animals Species; Scientific name; Identification mark; Breed; Quantity; Collect. Date Code; Collection date; Approval No of the centre of origin

II. Health information	II.a. Certificat reference number	II.b.Local reference number:	

## 1. ANIMAL HEALTH ATTESTATION

I, the undersigned official veterinarian, hereby certify that:

## 1.1 SWITZERLAND

(Name of exporting country) (3)

was free from rinderpest and foot-and-mouth disease during the 12 months immediately prior to collection of the semen for export and up until its date of dispatch and no vaccination against these diseases took place during the period.

- 1.2 The semen described above was collected before 31 December 2004 at a semen collection centre which:
  - 1.2.1 meets the conditions laid down in Chapter I of Annex A to Directive 88/407/EEC
  - 1.2.2 is operated and supervised in accordance with the conditions laid down in Chapter II of Annex A to Directive 88/407/EEC;
- 1.3 The centre at which the semen to be exported was collected was free from rabies, tuberculosis, brucellosis, anthrax and contagious bovine pleuropneumoniae during 30 days prior to the date of collection of the semen to be exported and the 30 days after collection (in the case of fresh semen, until the day of dispatch);
- 1.4 At the time the semen described above was collected, all bovine animals at the semen collection centre:
  - 1.4.1 came from herds and/or were born to dams which satisfy the conditions in paragraph 1 (b) and (c) of Chapter I of Annex B to Directive 88/407/EEC;
  - 1.4.2 have tested negative, within 30 days preceding the quarantine isolation period, to:
    - the tests referred to in points 1(d) (i), (ii) and (iii) of Chapter I of Annex B to Directive 88/407/EEC; and
    - a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis / infectious pustular vulvo-vaginitis, and
    - a virus isolation test (fluorescent antibody test or immunperoxidase test) for bovine viral diarrhoea, deferred until the animal reached the age of six months in the case of younger animals;
  - 1.4.3 had undergone the 30-day quarantine isolation period and had tested negative to the following health tests:
    - a serological test for brucellosis carried out in accordance with the procedure described in Annex C to Directive 64/432/EEC;
    - either an immunofluorescent antibody test or a culture test for campylobacter foetus infection on a sample of preputial material or artificial vagina washings, or, in the case of a female animal, a vaginal mucus agglutination test (1);
    - a microscopic examination and culture test for trichomonas foetus on a sample of preputial material or artificial vagina washings, or in case of a female animal, a vaginal mucus agglutination test (1);
  - 1.4.4 had tested negative, at least once a year, to the routine tests referred in points 1 (a), (b) and (c) of Chapter II of Annex B to Directive 88/407/EEC;
- 1.5 At the time the semen described above was collected,
  - 1.5.1 all female bovine animals in the centre had tested negative at least once a year to a vaginal mucus agglutination test for campylobacter foetus infection, and
  - 1.5.2 all bulls used for semen production had tested negative either to an immunofluorescent antibody test or to a culture test for campylobacter foetus infection on a sample of preputial material or artificial vagina washings carried out within 12 months prior to collection;
- 1.6 The semen to be exported was obtained from donor bulls which:
  - 1.6.1 satisfy the conditions laid down in Annex C to Directive 88/407/EEC;
  - 1.6.2 either were resident in the exporting country during the six months immediately prior to collection of the semen for export (1),
    - were imported from ......(3) after spending less than six months in the exporting country and at the time of import satisfied the animal health conditions applying to donors of the semen which is intended for export to the Community (1)
  - 1.6.3 stand in a semen collection centre at which:
    - (i) all bovine animals tested negative at least once a year to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis / invectious pustular vulvo-vaginitis (1), or
    - (ii) bovine animals not vaccinated against infectious bovine rhinotracheitis tested negative at least once a year to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis / infectious pustular vulvo-vaginitis and at which testing for infectious bovine rhinotracheitis was not carried out on bulls which had received their first vaccination against infectious bovine rhinotracheitis at the insemination centre after they had tested negative to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis / infectious pustular vulvo-vaginitis and which had been regularly revaccinated at intervals of not more than six months since the first vaccination (1);

eriand			and stored before 31. December 2004			
II. Health informati	ion	II.a. Certificat reference number	II.b.Local reference number:			
1.6.4	fulfil the import conditions for bovine semen laid down in OIE depending on the status of the country or zone of res	n the Bluetongue Chapter of the Terre	estrial Animal Health Code of the			
1.6.5	were resident in the country of export in which the follow; and tested negative on two occasion and a virus neutralisation test for all above listed serotyp taken prior to and not less than 21 days following collect	ns not more than 12 months apart to a es of EHD, carried out in an approved	n agar gel immunodiffusion test (4)			
1.6.6	were resident in the country of export in which the follow; and tested negative, prior to entry a to a virus neutralisation test for all above listed serotypes	and at six monthly intervals to an again	gel immunodiffusion test (4) and			
1.6.7	1.6.7 tested negative on two occasions not more than 12 months apart to a serum neutralisation test for Akabane virus carried out in an approved laboratory on samples of blood taken prior to and not less than 21 days following collection of the semen; *					
	emen to be exported was collected after the date on which thing country;	he centre was approved by the compe	tent national authorities of the			
	emen to be exported was processed, stored and transported to its amendment by Directive 2003/43/EEC;	under the conditions which satisfy the	terms of Directive 88/407/EEC			
1.9 Blueto	ongue Disease: The semen was collected in full compliance	e with Commission Regulation (EC) N	No 1266/2007 as amended;			
<u> </u>						
(1) Delete as no (2) [Box referer Identification Approval no (3) Countries li (4) Standards for Animals  **** To be used ****  To be used ***	nce No. 1.31 in Part 1]: on mark: corresponding to the identification of the donor an umber of the centre of origin: to be filled in if different from isted in Annex 1 to Decision 2004/639 or EHD virus diagnostic tests are described in the Bluetong only by Australia, Canada and the USA. only by Australia and the USA.	nimals and the date of collection n box reference No 1.12				
	only by Canada. only by Australia.					
territory; (b) Be made ou	ate must: p in at least one official language of the Member State of do at to a single consignee y the semen in the original	estination and of the Member State w	here the semen will enter Community			
The colour	of the stamp and signature must be different from that of th	e other particulars in the certificate.				
Official veterinar	rian or official inspector					
Name (in capital	letters):					
Date:						
Stamp		Qualification and title:				

Signatur

Attachement to		
----------------	--	--

## I.31 Identification of the animals / products

(page 1 of 1)

		of the animals	products			1		(page 1 01 1)
Species	Scientific name	Identification mark	ς.	Breed	Quantity	Collect. date Code	Collection date	Appr.No of the centre of origin
		Tatal				<u> </u>		
		Total						