

Department for Environment, Food and Rural Affairs

Export of treated blood products, excluding of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals intended for dispatch to or for transit through the European Union (EU) or Northern Ireland (NI)

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No: 8309NFG

For export of treated blood products, excluding of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals intended for dispatch to or for transit through the European Union or Northern Ireland.

NOTES FOR GUIDANCE (NFG) FOR THE CERTIFYING OFFICIAL VETERINARIAN (OV), CERTIFICATION SUPPORT OFFICER AND EXPORTER

1. APPLICABLE LEGISLATION

[Council Regulation \(EC\) No 1069/2009](#) and [Commission \(EU\) Regulation 142/2011](#) (as amended)

Any other EU legislation referenced in the certificate must be complied with and can be accessed on the following link:

<https://eur-lex.europa.eu/homepage.html>

IMPORTANT

These notes provide guidance to Certifying Officers and exporters (CO). The NFG should have been issued to you together with the relevant export certificate applicable for exports of treated blood products, excluding equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals intended for dispatch to or transit through the EU or Northern Ireland. The NFG should not be read as a standalone document but in conjunction with the veterinary certificate.

We strongly suggest that exporters obtain full details of the importing country's requirements from the veterinary authorities in the country concerned, or their representatives in GB, in advance of each consignment.

[Please note, policies are being reviewed. NFG will be further amended to provide specific guidance. Traders should look at NFGs regularly for any updates]

2. SCOPE OF THE CERTIFICATE

This Model 8309 veterinary certificate maybe used for the export of treated blood products, excluding equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals intended for dispatch to or transit through the EU or Northern Ireland, in accordance with the relevant requirements described in Regulation (EU) No 142/2011.

Blood products are defined in Annex I of Regulation (EU) No 142/2011 as meaning derived products from blood and fractions of blood, excluding blood meal; they include dried/ frozen/ liquid plasma, dried whole blood, dried/ frozen/ liquid red cells or fractions thereof and mixtures.

Only blood referred to in Article 8(c) and (d) and Article 10(a), (b), (d) and (h) of Regulation (EC) No 1069/2009 may be used.

3. CERTIFICATION BY AN OFFICIAL VETERINARIAN (OV)

In **England, Scotland and Wales**, this certificate must be signed by a Government Veterinary Officer or by an OV appointed by the Animal and Plant Health Agency (APHA) on behalf of Ministers in Defra, the Scottish Government or the Welsh Government and who hold the appropriate Official Controls Qualification (Veterinary) (OCQ (V)) authorisation.

OVs must sign and stamp, with the OV stamp, the health certificate in ink of a different colour to that of the printing of the Export Health Certificate (EHC). There is no requirement to sign and stamp in a specific colour.

The OV should keep a copy of the signed certificate and any supporting documents for at least three years after signature or receipt/dispatch of the consignment, whichever is later. These can be electronic copies.

EHC in foreign languages of the EU Member States (MSs).

EHC should be in English and the foreign languages of the Border Control Post (BCP) of entry in the EU, as well as in the language of the EU MS of destination if this a different country from the point of entry to the EU. The required EHC must accompany the consignment.

Listing of the EU MS BCPs can be found here:

https://ec.europa.eu/food/animals/vet-border-control/bip-contacts_en

The foreign language certificate as received from the APHA Centre for International Trade at Carlisle or via the Export Health Certificates on-line system (EHCO) and bearing the same unique reference number as the English certificate, should be considered an official and accurate translations of the English, as published in EU legislation.

The (sub-) paragraphs / options and how they are numbered and formatted is identical in the English and foreign language editions and to the legislation published by the EU Commission. Therefore, when the same phrases/sentences in the foreign language versionss as in the English version are struck through, both versions can and must be signed (as opposed to being initialled) by the OV as a genuine and properly authorised translation of the English.

This also applies to any instructions in the guidance notes to strike out certain paragraphs or to certify statements that the country is free of certain notifiable diseases etc.

Signing, stamping and pagination

The foreign language version/s and any schedules (if any) may be stapled to the English version but doing so and then fan stamping the multiple sheets is not enough to create one indivisible single document according to the EU Commission.

Therefore, each page (including schedules) should be individually signed and stamped and bear the reference number of the certificate. The pages comprising the complete document should be sequentially numbered so they are part of a finite sequence which covers the English, foreign language version/s and any schedule pages.

For example, if the certificate consists of four A4 pages printed back to back on two sheets of A4 paper with a schedule that is three A4 pages long, all 11 pages must be stamped and **signed** (as above) and numbered 1/11 to 11/11.

COs will have to make handwritten corrections to page numbering as may be required. E.g. 1/4 to 4/4 (if present) on the foreign language parts in the example given above will need to be crossed out and the 1/11 to 11/11 entered.

The EHC accompanying the consignment will then comprise the original English EHC and any required additional foreign language/s. These should be arranged in order with the English version on the top, followed by the foreign language/s version/s, and finally the pages of the schedule (if any) at the bottom.

As per general guidance for certifiers on APHA's [Official Veterinarian Training](#) , any handwritten corrections or permitted deletions to a certificate should be stamped and **initialled**. This includes the deletion of optional statements in Part II of the certificate and alterations to content in Part I. The same applies if a pre-populated text in a box in Part I of the EHC needs to be amended. (E.g. if Box I.7 which is pre-populated as 'United Kingdom' 'GB', needs to be amended for triangular trade where third country origin 'Products Of Animal Origin' are being certified in the original third country packaging with the original third country Identification Marks, in which case the country of origin will be the third country in question and not the United Kingdom). Please follow the guidance on corrections in the link below:

[Instruction: Official Veterinarian Training](#)

We advise that individual stamping and initialling of diagonal lines drawn through blank boxes in Part I is not necessary. This is to reduce excessive stamping on the certificate. However, we are aware that some BCPs advise otherwise and request stamping and initialling of manually crossed out blank boxes in Part I of the certificate. In such cases OV should conform to the BCPs request to facilitate the clearance of the goods.

You can find further information on Export Health Certificates (EHC) Online Guidance for Certifiers in the link below:

[Using export health certificate \(EHC\) online: certifier guidance - GOV.UK](#)

UK approved establishments will be uploaded to [Europa](#) website in due course, until the establishments are in Europa website you can find the list of UK approved establishments in the link below:

<https://www.gov.uk/government/publications/businesses-approved-to-export-to-the-eu>

Please check the guidance on completion of part I of the EHC at the bottom of the EHC and in the links provided in the NFG. For completion of Box I.8 - Region of Origin Code, if applicable; the territory code should be as listed in the relevant legislation that is provided

under the notes at the bottom of the EHC. This is only for species or products affected by regionalisation measures or by the setting up of approved zones in accordance with a European Community Decision. The approved regions or zones must be indicated as described in the Official Journal of the European Union

Part I: DETAILS OF THE CONSIGNMENT

Please complete all the boxes in Part I of the certificate.

The Harmonised System (HS) Code is a commodity classification system used as a basis for customs tariffs and for international trade statistics.

It is the exporter's responsibility to ensure that the HS code is entered correctly and accurately reflects the products being consigned.

Further information on HS Codes can be found online at:

<https://www.gov.uk/trade-tariff/sections> and <http://madb.europa.eu/madb/euTariffs.htm>

PART II: CERTIFICATION

Animal Health Attestation

II.1 The OV signing the certificate must have read and understood Regulations (EC) No 1069/2009, in particular Articles 8(c), 8(d) and 10, and Commission Regulation (EU) No 142/2011, in particular Chapter II of Annex XIV and must ensure that the blood products meet the requirements of the certificate. The starting material used must be Category 3 ABP or limited Cat 1 material as specified in the Scope Section 2.

The following specific guidance in conjunction with the RCVS Principles of Certification may be followed: **The OV must have familiarity with sourcing, procurement, segregation, processing, and handling and storage arrangements in place at the establishment and ensure that the consignment meets the conditions required in the certificate. Where the OV is required to certify conditions outside of their personal knowledge, they must request and be provided with appropriate supporting documentation from another veterinarian (if appropriate) and/or the exporter.**

II.2 This can be certified on the basis of a declaration from the establishment operator confirming that the blood products are not intended for human consumption.

II.3 Starting/source material

The OV must establish that the plant is approved and supervised by the competent authority for the production of derived products for uses outside the feed chain for farmed animals. This can be established through the process detailed in Section 7 - GB APPROVED ESTABLISHMENTS TO EXPORT TO THE EU. The OV can ask for a copy of the approval document.

The starting materials must be category 3 or limited category 1 materials as detailed in the Scope Section 2.

The correct subcategory or sub-categories of ABP under II.2 must be selected, and the other subcategories deleted as instructed. Familiarity with the sourcing arrangement of the raw material by the establishment is necessary as supported by physical inspection and by examination of relevant documentation or other records including commercial documents, veterinary statements and valid written exporter declarations to ensure the correct subcategory or sub-categories is/are selected.

II.4 This may be certified on the basis of familiarity with sourcing arrangements at the processing establishment and examination of relevant records including a written exporter declaration confirming the establishment at which the blood was collected. Approval of these establishments can be confirmed through the process detailed in Section 7 - GB APPROVED ESTABLISHMENTS TO EXPORT TO THE EU.

II.5, II.6, II.7 Processing methods in relation to species of origin of the source material.

The OV should develop due familiarity with sourcing, procurement, segregation, processing, handling and storage arrangements at the establishment. This should be supported by physical inspection and by examination of relevant documentation or other records including the approval document, commercial documentation, and valid written declaration from the plant operator confirming species of origin and treatment applied. Attestations not applicable to the source species should be deleted.

II.5 Relates to blood products derived from even toed ungulates (Artiodactyla), - eg bovine, ovine, caprine but excluding swine, odd toed ungulates (Perissodactyla), the elephant family (Proboscidea) and cross breeds. The blood products must have undergone at least one of four treatments. The appropriate treatments must be selected and the others deleted.

II.6 Relates to blood products derived from swine, poultry or other avians, - the blood products must have undergone at least one of three treatments. The appropriate treatments must be selected and the others deleted.

II.7 Relates to blood products derived from species other than those listed above at II.5 and II.6 but additionally excluding equines. The treatment applied should be entered in the available space.

II.8 and II. 9 Packaging, labelling, transport and storage.

The OV must establish these statements in relation to the storage of the products, hygiene of packaging or transport containers and labelling with “NOT FOR HUMAN OR ANIMAL CONSUMPTION” are met through knowledge of plant operations and physical inspection as supported by a declaration from the establishment.

Details of disinfectants approved for this purpose can be accessed via:

http://disinfectants.defra.gov.uk/DisinfectantsExternal/Default.aspx?Module=ApprovalsList_Sl (as amended)

II.10 Precautions to avoid contamination with pathogens

Procedures must be in place to ensure that all precautions have been taken to avoid pathogenic contamination after treatment.

The OV must establish that processes are in place to prevent contamination by pathogenic agents after treatment through familiarity with processing, handling and storage arrangements at the establishments and should obtain a relevant written declaration from the plant operator confirming these measures have been taken.

II.11 BSE risk status

NOTE: THIS ATTESTATION ONLY APPLIES WHERE ABP OR ABP PRODUCTS ARE DERIVED FROM RUMINANTS.

Species material other than ruminants.

This section should be deleted in its entirety.

The OV should obtain a written declaration from the exporter confirming the species of the source materials. The OV must be familiar with the plant sourcing procedures and complete a documentary check of the records.

Ruminant species material other than bovine, caprine or ovine only

The first “either” option should be signed for and all other subparagraphs should be deleted.

The OV must obtain a written declaration from the exporter confirming the species of the source materials and their BSE risk classification. The OV must be familiar with the plant sourcing procedures and complete a documentary check of the records.

Bovine, caprine or ovine species material only

GB sourced material

If GB sourced ABP material or ABP products are used only the second “or” subparagraphs (a) and/ or (c) can be signed for. All other subparagraphs should be deleted.

In relation to the second “or” subparagraph (a) this subparagraph may only be signed for if the EHC permits the use of such material. The OV should check that such source material is permitted to be used as stated in Section 2 - SCOPE OF THE CERTIFICATE above and

must obtain a written declaration from the exporter confirming the species of the source materials and their BSE risk classification. The OV must be familiar with the plant sourcing procedures and complete a documentary check of the records.

Imported material

If imported ABP material can be used then the OV should refer to Section 5 (CONSIGNMENTS OR PARTS OF THE CONSIGNMENT ORIGINATING FROM, NI, EU MEMBER STATES OR FROM THIRD COUNTRIES (TRIANGULAR TRADE)).

4. COLLECTION OF EVIDENCE

Personnel may be authorised to collect evidence which may be used to support veterinary certification. In GB, the Certification Support Officer (CSO) role has been developed by APHA.

CSOs can be utilised by OVs for gathering evidence relating to this certificate. The CSOs must be authorised by APHA and they must hold the appropriate Official Controls Qualification (Animal Health Professional) (OCQ (AHP)-CSO) qualification.

The OV must direct the CSO as to how and where any necessary evidence relevant to the requirements of the Export Health Certificate (EHC) should be obtained. CSOs may not carry out any functions that require the exercise of veterinary judgement and are restricted to the execution of administrative checks.

They may only carry out such inspections, factual verification and evidence collection as specified by the directing OV, who remains responsible for the certification of the product. CSOs are not authorised to sign an EHC in their own right or on behalf of an OV.

Any documentary evidence collected by the CSO must be stamped, signed and dated by the CSO, before being submitted by them as supporting evidence to the OV. It is required that the OV is familiar with the product process and evidence required to start with, before directing the CSO to provide future evidence on an ongoing basis.

Additional guidance and principles of implementation are provided in the OV Instructions [Export document](#) section of the APHA [Official Veterinarian Training](#).

5. CONSIGNMENTS OR PARTS OF THE CONSIGNMENT ORIGINATING FROM, NI, EU MEMBER STATES OR FROM THIRD COUNTRIES (TRIANGULAR TRADE)

This section of the guidance applies to exports to the EU and movements from GB to NI but does not apply to movements of retail products to Northern Ireland under the Northern Ireland Retail Movement Scheme (NIRMS).

An ABP consignment for export from GB can contain animal products that originate from NI, EU and third countries, only if those products have undergone further processing in GB.

Processing should be understood in the context of Commission Regulation (EU) No 142/2011 and is different than the definition that applies in the context of products of animal origin for human consumption.

ABP imported into GB, which is only unloaded, stored, and reloaded, or which is only rewrapped in GB, cannot be re-exported to the EU or moved to NI except under the NIRMS or under the customs transit procedure (see below). Guidance on triangular trade can be found here: [Triangular Trade Briefing Note](#)

To avoid the restriction applicable to triangular trade, businesses can make use of the customs transit procedure for goods from third countries landed in GB, to move through GB, directly to NI. Consignments being moved under the customs transit procedure are not subject to triangular trade rules. Guidance on the transit procedure can be found in the triangular trade briefing note above.

6. GB APPROVED ESTABLISHMENTS TO EXPORT TO THE EU

The exporting establishment must be authorised and listed by the GB as a 'GB approved establishment' for ABP not for human consumption (ABP). A list of approved establishments can be found on the European Commission's list of approved establishments' link below:

https://ec.europa.eu/food/safety/international_affairs/trade/non-eu-countries_en

Please note that the list is updated regularly and ONLY establishments on the list are approved to export to the EU and does not include establishments with pending applications for approval/registration.

If the final product contains animal products from other establishments, or products were previously processed in different establishments in the production chain, then these establishments should also be listed on the EU website as GB approved establishments.

For approved establishments in Northern Ireland the "EC" suffix which is present in the health/ID mark of approved food establishments, should not be included when referring to establishment approval numbers in the certificate. This may also be relevant to certain ABP consignments – e.g. where the ABP is generated at an approved slaughterhouse without separate ABP approval.

7. CERTIFIED COPIES OF EXPORT HEALTH CERTIFICATES

When completing export certification, the CO and, if applicable, FCCO must make photocopies of, or scan and save all documents they certify. OV's must retain copies of certification documents in accordance with RCVS Certification principles:

<https://www.rcvs.org.uk/setting-standards/advice-and-guidance/code-of-professional-conduct-for-veterinary-surgeons/supporting-guidance/certification/>

COs must retain copies of all export documentation for a period of two years. A certified copy of this EHC does not need to be returned to the APHA CITC. For the purposes of completing routine Quality Assurance checks on export certification, CITC may request certified copies of certification from COs.

Further details on Post Certifying Procedures, 'certified copies' of certification and the types of documents that should be retained by COs can be found on the OV Instruction Exports [document](#) of the APHA Official Veterinarian Training.

8. LEGAL STATEMENT

References in this guidance to "assimilated EU Regulation" should be interpreted as references to assimilated law, as defined under the European Union (Withdrawal) Act 2018.

9. DISCLAIMER

This certificate is provided on the basis of information available at the time and may not necessarily comply fully with the requirements of the importing country. It is the exporter's responsibility to check the certificate against any relevant import permit or any advice provided by the competent authority in the importing country. If these do not match, the exporter should contact the Animal and Plant Health Agency (APHA) in Carlisle, via the link below:

<https://www.gov.uk/government/organisations/animal-and-plant-health-agency>

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Section 5 - CONSIGNMENTS OR PARTS OF THE CONSIGNMENT ORIGINATING FROM, NI, EU MEMBER STATES OR FROM THIRD COUNTRIES (TRIANGULAR TRADE) is amended to align with the [Triangular Trade Briefing Note \(ABP\)](#)

References to Vet Gateway replaced by APHA's Official Veterinarian Training

Legal statement is updated.