

Department for Environment, Food and Rural Affairs

Notes for Guidance: Export Health Certificate for entry into the European Union or Northern Ireland of honey and other apiculture products intended for human consumption 8391

November 2025

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No: 8391 NFG

EHC for entry into the EU or NI of honey, and other apiculture products intended for human consumption.

NOTES FOR GUIDANCE (NFG) FOR THE CERTIFYING OFFICERS (COs) AND EXPORTERS

1. APPLICABLE LEGISLATION

[Regulations \(EC\) No. 178/2002](#)

[Regulation \(EC\) No. 852/2004](#)

[Regulation \(EC\) No. 853/2004](#)

[Regulation \(EC\) 396/2005](#)

[Commission Regulation \(EC\) 1881/2006](#)

[Regulation \(EU\) 2022/2292](#)

[Regulation \(EU\) 2017/625](#)

[Council Directive 2001/110/EC](#)

[Commission Implementing Regulation \(EU\) 2020/2235](#)

[Commission Implementing Regulation EU 2023/2744](#)

[Commission Delegated Regulation \(EU\) 2023/905](#)

Any EU legislation referenced in the EHC must be complied with and EU legislation can be accessed on the following link <https://eur-lex.europa.eu/homepage.html>

Please note that Official Control Regulations 2017/625 have repealed Regulation (EC) No 854/2004, 882/2004 and Directive No 96/23/EC.

Consolidated legislation

Consolidated texts, which integrate the basic instruments of Union legislation with their amendments and corrections in a single, non-official document, are available. Each consolidated text contains a list of all legal documents, taken into account for its construction.

You can search for consolidated texts by using the 'find results by document number' option on the European Commission website. Once you have selected the relevant legislation, click 'document information', and then scroll down to 'all consolidated versions' and select the most recent version.

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

Please note that the consolidated text may not contain the latest amendment to the legislation, as it takes several weeks for this to be updated.

Texts provided in this section are intended for information only. Please note that these texts have no legal value. For legal purposes please refer to the texts published in the 'Official Journal of the EU'.

IMPORTANT

These notes provide guidance to COs and exporters. The NFG should have been issued to you together with the relevant export certificate applicable for dispatch of honey, royal jelly and other apiculture products intended for human consumption to the EU or NI. 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 the UK, 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 certificate is for movements into the EU or NI of honey and other apiculture products intended for human consumption.

It can only be used for definitive imports into the EU or NI of these goods, it cannot be used for these products transiting the EU or NI to another third country.

3. CERTIFICATION BY AN OV

In **England, Scotland and Wales**, this certificate can be signed by a Government Veterinary Officer (e.g. APHA, FSA or FSS employed veterinary officers) or by an OV appointed by the 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.

In England, Scotland and Wales this certificate may also be signed by a Food Competent Certifying Officer (FCCO) who is authorised by APHA.

The FCCO or OV must sign and stamp, with their Official stamp, the health certificate in ink of a different colour to that of the printing of the EHC. There is no requirement to sign and stamp in a specific colour.

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

EHC in foreign language/s of the EU Member States (MSs):

EHC should be in English and the foreign language of the Border Control Post (BCP) of entry in the EU. The original copy of the required EHC must accompany the consignment to the BCP of entry.

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 Online 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 European Commission. Therefore, the same phrases/sentences in the foreign language versions as in the English version should be struck through and these deletions should be stamped and initialled in both versions. Both versions must also be signed (as opposed to being initialled) and stamped by the OV, the foreign language certificate is deemed to be a genuine and properly authorised translation of the English version.

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.

Additional information can be found in APHA: [Official Veterinarian Training](#)

SIGNING AND STAMPING

When signing a certificate, the CO should ensure that the certificate contains no deletions or alterations, other than those which are indicated on the certificate to be permissible and any corrections to permitted entries, subject to such changes being initialled and stamped (in the margin) by the CO. Permissible deletions are normally indicated in the 'Notes' section at the end of the certificate, with the instruction 'Keep as appropriate' or 'delete if not applicable'.

- Where the certificate contains optional or contextual statements, the statements which are not relevant shall be crossed out, individually initialled, and stamped by the CO, or completely removed from the certificate.
- Permitted paragraphs and sections may be crossed out by applying a 'Z' across the section or paragraph rather than crossing out line by line.
- There is no requirement for a date and time to accompany each stamp. The date is only entered at the required entry field in Part I of the certificate, and at the end where the CO signs, stamps and dates that action.
- We are aware of some BCPs demanding that all handwritten information in Part 1 of the EHC is initialled and stamped, including handwritten scoring out of otherwise blank boxes. There is no legal requirement in EU legislation that all the hand-written information entered in the certificate must be signed and stamped. It is only in the case of correction, in any part of the certificate, or in the case of statements to be crossed out, that the certifier must add signature (or initials) and stamp. This has been confirmed by the European Commission. The Commission noted however, in the

case of a hand-written certificate, it is expected that the same one person completes the document. If not, the BCP might suspect that empty boxes were completed by another person after the certificate has been signed by the official

You should consider checking with the specific BCP regarding their preference when it comes to the stamping and initialling of handwritten scoring out of otherwise blank boxes in Part I of the EHC.

- **Clarification from the European Commission means that all pages (as opposed to sheets of paper) are signed and stamped once individually in place of fan stamping and in addition to any permitted alterations. There is no requirement to fan stamp.**
- COs are reminded to consult the NFG prior to the certification of each EHC. NFG will be updated with this new information in due course.

Further Information COs should make sure they are familiar with all relevant guidance and other documents relating to EHCs and that they discuss requirements with exporters in advance.

See [Export-health-certificate-ehc-online-certifier-guidance](#)

You can also contact APHA's Centre for International Trade (CIT) on 03000 200 301.

PART I: DETAILS OF THE CONSIGNMENT

All boxes in Part I of the certificate must be completed. When a box is not applicable/optional, and not filled, please score it through.

Please use schedule to be attached to the certificate if there is not enough space to fill the information. See section 'Addition of Schedules' for further information.

Please complete all the boxes in Part I of the certificate in accordance with the guidance laid down in Chapter 4 of Annex I to Commission Implementing Regulation (EU) 2020/2235 that can be accessed via this link: [Commission Implementing Regulation EU 2023/2744](#)

<https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX:32020R2235>

I.11 Please refer to the bottom of the certificate for guidance on the registration number.

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

I.27: Please insert **HS Code** and **Combined Nomenclature (CN) title**

It is the exporter's responsibility to ensure that the HS code and CN title is entered correctly and accurately reflects the product(s) being consigned, as defined by the World Customs Organisation as referred to in Council Regulation (EEC) No 2658/87, please see link:

<https://eur-lex.europa.eu/legal-content/en/ALL/?uri=CELEX%3A31987R2658>

Further information on HS Codes for customs tariffs can be found online at:

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

I.27- Type of packaging: identify the type of packaging according to the definition given in Recommendation No 21 (10) of UN/CEFACT (United Nations Centre for Trade Facilitation and Electronic Business). Please see link:

https://www.unece.org/fileadmin/DAM/cefact/recommendations/rec21/rec21_Rev10e_Annex-V-VI_2019.xls

PART II: CERTIFICATION

II.1 Public Health Attestation

The OV signing the export veterinary certificate must ensure that the public health attestations set out in Part II of the veterinary certificate have been complied with.

They must ensure that they are aware of the provisions of Regulation (EC) Nos 178/2002, 852/2004, 853/2004, 2017/625. and Council Directive 96/23/EC and 2001/110/EC

To certify point **II.1**, the products described must come from a registered establishment under 852/2004. In practice this means that the establishment has to be registered by the local authority but it does not have to be listed on the EU website.

For guidance on certifying Residue Check Guarantees please refer to Section 4.

(a),(b) - This can be certified on the basis of the CO's own knowledge of the listed legislation (HACCP attestation). For honey and apicultural products derived from bees in the UK, this may be certified on the basis of the product(s) being produced in a registered and listed establishment which satisfies the requirements of Regulation (EC) 852/2004 and Article 127(3), point (e)(ii), of Regulation (EU) 2017/625.

(c) - For honey and apicultural products that are sourced from the UK this paragraph can be certified on the basis that the national surveillance scheme implements Council Directive 96/23/EC, which are transposed into national legislation by The Animals and Animal Products (Examination for Residues and Maximum Residue Limits) (England and Scotland) Regulations 2015 and parallel legislation in the devolved administrations.

Said provisions fulfil the guarantees covering live animals and products thereof provided by the residues plans submitted in accordance with Article 6(2) of Commission Delegated Regulation (EU) 2022/2292. The UK is listed in Annex -I to Commission Implementing Regulation 2021/405 for the concerned animals and products covered under this EHC.

See section 4 for further advice on residue check guarantees. The UK has a surveillance programme in place to monitor residues of authorised veterinary medicines, prohibited substances, and other contaminants in domestically produced foodstuffs of animal origin. The testing results of the level of pesticides and residue in food are published in an annual report. Annual reports can be found on gov.uk.

<https://www.gov.uk/government/publications/expert-committee-on-pesticide-residues-in-food-prif-annual-report>

d) This can be certified if the honey does not contain any added food ingredient, including food additives or extraneous sugars. Honey blends can still be certified as long as the product complies with the requirements of Annex I and II of [Council Directive 2001/110/EC](#), and all other requirements under this certificate.

II.1 (a) - This attestation on antimicrobial medicinal products is added which must be crossed out or deleted until 3 September 2026.

4. RESIDUE CHECK GUARANTEES

The UK has a surveillance programme in place to monitor for residues of authorised veterinary medicines, prohibited substances, and other contaminants in domestically produced foodstuffs of animal origin. Sample collection is conducted at the point of production i.e. at farm and slaughterhouse.

The domestic legislative basis for this monitoring is outlined in The Animal and Animal Products (Examination for Residues and Maximum Residue Limits) (England and Scotland) Regulations of 2015 and equivalent legislation in Wales ([2019](#)) and NI ([2016](#)). The monitoring conducted in GB is in accordance with the legislative requirements of Directive 96/23 (EC), 96/22 (EC), Decision 97/747 (EC) and 470/2009 (EC) concerning residue testing of products of animal origin. The residues tested in the programme are in accordance with Annex I and II of Directive No 96/23 (EC), specifically, and include veterinary medical products, banned substances and environmental contaminants. In practice, monitoring conducted in the UK meets the requirement of Regulation (EU) 2022/2292 for veterinary medicine and prohibited substances and Annex I of 2022/932, for contaminants.

With regards to maximum levels used to determine sample non-compliance, for authorised veterinary medicines GB work to the GB Maximum Residue Limits (MRLs) published [here](#); these MRLs are aligned to the EU veterinary MRLs published under Reg (EU) [37/2010](#). If a pesticidal compound has an MRL for food-producing species then this MRL is used as the respective non-compliance threshold, but if a pesticide does not have a foodstuff MRL then the MRLs as listed in Regulation (EC) 396/2005 are applied. For contaminants, such as heavy metals and mycotoxins, the limits as set out in Reg (EC) 1881/2006 are used to determine sample non-compliance.

The results of the statutory surveillance programme can be accessed on the link below:

<https://www.gov.uk/government/collections/residues-statutory-and-non-statutory-surveillance-results>

The EHC residue testing requirements can be certified based on evidence of compliance to the national surveillance programme, which complies with the relevant EU legislation.

The national monitoring programme for pesticide MRLs in food and feed in place under Regulation 396/2005 is underpinned by national legislation, The Pesticides (Maximum Residue Levels) Regulations (England and Wales) 2008 (as amended) and devolved administration equivalents. A national monitoring programme for Maximum Residue Levels is managed by the Health and Safety Executive. This involves testing a selection of produce that has already been placed on the market in Great Britain to provide assurance that only authorised pesticides, within permitted levels, are present. The results are published in an annual report. Annual reports can be found on gov.uk.

<https://www.gov.uk/government/publications/expert-committee-on-pesticide-residues-in-food-prif-annual-report>

Any EHC residue pesticide requirements can be certified based on evidence of compliance with the pesticide residue monitoring scheme.

<https://www.gov.uk/government/collections/pesticide-residues-in-food-results-of-monitoring-programme>.

5. COLLECTION OF EVIDENCE

In GB, the Certification Support Officer (CSO) role has been developed by APHA. CSOs can collect evidence, directed by an OV, which may be used to support OV certification of matters which do not require a clinical assessment or judgement e.g. for POAO and ABPs.

In **England, Scotland and Wales**, 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 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](#).

Groupage Export Facilitation Scheme (GEFS)

For groupage exports from Great Britain, where certain types of products are produced from a stable supply chain and are fully packaged for the final consumer, exporters who are GEFS members may use 30 day support attestations to provide information to OVs to facilitate completion of this certificate.

For further information including the definition of groupage exports, the template 30-day support attestation which must be used and requirements for exporters, suppliers and vets to use the scheme see: [Export groups of products using the Groupage Export Facilitation Scheme \(GEFS\) - GOV.UK](#)

You can check that exporters are GEFS members by emailing the exporter's name, GEFS membership number and the address of the exporting premises to GEFS@defra.gov.uk

6. CONSIGNMENTS OR PARTS OF THE CONSIGNMENT ORIGINATING FROM NI, EU MEMBER STATES OR FROM A THIRD COUNTRY (TRIANGULAR TRADE)

NI origin:

For NI origin raw materials which have then been processed into a final product in GB, or are presented in their original state and bearing a UK(NI) identification mark, the CO can certify certain matters relating to EU compliance at a national level.

Where the EHC refers to EU approval status of the premises of origin or manufacture in NI, this can be certified under the terms of the EU-UK Withdrawal Agreement and the Windsor Framework. The Windsor Framework applies EU SPS legislation in NI.

Approved premises in NI continue to implement the full requirements of Regulation (EC) Nos. 852/2004 and 853/2004 and Regulation (EU) No. 2017/625 and all relevant supporting EU legislation as set out in the Windsor Framework. This compliance is indicated by the presence of the EU oval health and identification marks applied to the products.

Some examples, but not a complete list, of how assurance can be established at national level are listed below.

Compliance with the microbiological criteria set out in Regulation (EC) No. 2073/2005 can be certified if the products originate in an EU approved premises in NI and bearing the EU oval ID mark.

Public health statements referring to compliance with EU requirements for testing for residues as set out in Regulation (EU) No_ 2017/625, Directive (EC) Nos 96/22 and 470/2009 can be certified by the CO on the basis of a national residue surveillance programme implemented in NI under The Animals and Animal Products (Examination for residues and maximum Residues Limits) Regulation (NI) 2016. This forms part of the UK national surveillance programme.

With regards to controls for Transmissible Spongiform Encephalopathies, guidance provided in this document relating to statements about the method of slaughter of animals in GB also applies to animals slaughtered in NI and can be certified by the CO on that basis.

Disease clearance for animals or products originating in NI can be completed using auto-clearance NDC found here:

<https://www.daera-ni.gov.uk/articles/notifiable-diseases-northern-ireland>

Where regional or local level disease clearance is required, this can be certified upon request on the basis of information from NI in the form of a declaration or a supporting health attestation.

Animal health statements which refer to the prohibition of certain vaccination programmes e.g. against FMD or CSF or ASF can be certified at a national level by the CO on the basis that NI also enforces a ban on such vaccinations in accordance with EU regulations.

Statements relating to implementation of a national system for identification and registration of bovine animals can be certified on the basis of the requirement to register all bovine animal births, moves and deaths on the DAERA database.

Animal welfare statements can be certified by the CO on the basis that relevant inspections, monitoring and controls are implemented in NI through The Welfare of Animals at the Time of Killing Regulations (NI) 2014 as amended, in compliance with Regulation (EC) No. 1099/2009.

Animal By-Products are handled in accordance with EU Regulation 1069/2009, which is implemented by the EU Implementing Regulation 142/2011, and ABP statements for materials originating in NI, can be certified on that basis.

When the certificate requires specific information to be included, such as the date of slaughter or the date of introduction into NI, the GB exporter/CO must request this information from the NI exporter. This NI exporter may forward the request to the relevant NI CO to provide this information. This supporting information must be in writing and kept by the GB CO. The GB CO is not required to attach it as a supporting document to the EHC, unless requested by the EU BCP or told otherwise.

EU origin:

For imported goods that need to be certified for export from GB, these are normally subject to import certification, or the availability of a Common Health Entry Document (CHED) issued by the BCP of entry to verify that they are compliant with GB import requirements and for placing on the GB market. COs including OV's may use these official documents to provide supporting evidence of compliance with relevant requirements for the re-export of products. In this context OV's may rely on the CHED issued by an Official Fish Inspector (a non-veterinarian) for Fishery Products and live bivalve molluscs, live echinoderms, live tunicates or live marine gastropods for human consumption, cleared via a GB BCP.

Where the CHED or accompanying import certificate are not available or do not provide sufficient supporting information, the COs should seek a supporting attestation from an 'authorised veterinarian' who has personal knowledge of the matters in question. This may be further supported by relevant commercial information or records. It is the responsibility of the GB exporter to obtain the necessary supporting information to enable the CO to verify compliance with export requirements.

For goods sourced in the EU and EFTA countries, especially those that are not accompanied by a veterinary certificate or CHED issued by a BCP - COs may rely on the oval ID mark applied at approved food establishments in the EU as evidence that the goods were produced compliant with EU food production requirements for placing on the EU market - but care must be taken not to extrapolate this to animal health requirements not covered by the obligations of a food approved establishment, i.e. matters that extend beyond the scope of Regulations 852/2004 and 853/2004.

Third country origin:

It is also possible that some consignments may contain POAO that have been imported to GB from non-EU countries and further processed in GB, which GB exporters intend to export to EU (known as Triangular Trade). In these cases, COs may obtain a copy of the EHC for the import of such commodity from the Third Country to the GB.

GB COs are not required to attach a copy of the Third Country EHC as a supporting document to the EHC, unless requested by the EU BCP or specifically instructed in the NFG.

It is the GB exporter's ultimate responsibility to obtain any necessary support documents (from the EU member state exporter/Third Country exporter), to enable GB COs to be able to certify the products in good time before the export to the EU.

7. ADDITION OF SCHEDULES

When the space in Part I or Part II of the certificate is insufficient to accommodate full details of the consignment a schedule may be used. In the relevant section of the certificate the CO should annotate the certificate 'see attached schedule'. A new schedule should be created (typed or clearly written) containing the same information as that required in the certificate. The schedule must include the certificate reference number on each page and must be signed, dated, and stamped by the CO in a colour other than the printed text on each page and under the last entry. The schedule forms part of the certificate. All pages of the certificate, including the schedule, must be sequentially numbered. Any blank spaces in the schedule or the certificate should be struck through with diagonal lines.

Further guidance is available here: [Official Veterinarian Training](#)

8. CERTIFIED COPIES OF EHCs

When completing export certification, the CO and, if applicable, FCCO must make photocopies of, or scan and save all documents they certify. OVs 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 in the OV instructions: [Official Veterinarian Training](#).

9. 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.

10. DISCLAIMER

This certificate and NFG are 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 APHA in Carlisle.

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PB 8391 NFG

Version History

EHC

Version Published XX November 2025

Part II:

II.1 (a) reference of legislation is added related to listing of honey and apiculture establishment.

II.1a: Reference to Delegated Regulation (EU) 2023/905 is replaced with the Commission Implementing Regulation (EU) 2024/2598.

Published 30 August 2024

Part II:

II.1 (a) - Attestation about the administration of antimicrobial medicinal products is added.

Notes - Footnote 3 is added.

Published 31 May 2024

Part II:

II.1 (c): Council Directive for residue plan 96/23 EC and Commission Implementing Regulation 2011/163 for listing of concerned animal and products, is replaced by Commission Delegated Regulation (EU) 2022/2292 for control plan and Commission Implementing Regulation (EU) 2021/405 for listing.

II.1 (d): Related to maximum residue level for pesticides and maximum level of contaminants is removed.

II.1 (e) is now point (d) and “with exception of honey” is removed from this attestation.

Notes:

Footnotes 1 and 2 are added for deleting if not applicable and for “applicable only to honey”.

NFG

Version 6: Published xx November 2025:

II.1: (a): Legislation reference is added.

Section 6: is updated replacing reference to Northern Ireland protocol with Windsor framework.

Links to APHA Gateway are replaced with APHA Official Veterinarian Training

Legal statement is updated.

Version 5: Published 30 August 2024

Applicable Legislation: Commission Delegated Regulation (EU) 2023/905 added

Part II: II.1 (a) - Guidance is added about the attestation related to antimicrobial medicinal products.

Version 4: Published 31 May 2024

Applicable Legislation is amended with addition of Regulation (EU) 2022/2292, 2023/2744.

Part I: Detail of the Consignment: Link to amended Regulation (EU) 2023/2744 is added for completing Part I of the EHC.

II.1. (c): Further clarity is added that the national surveillance scheme and mentioned provisions, fulfil the guarantees covering live animals and products provided by the residues plans submitted in accordance with Delegated Regulation (EU) 2022/2292.

II.1 (d): Guidance is removed as per EHC.

Section 4: Residue check guarantees: Further information is added - “In practice, monitoring conducted in the UK meets the requirement of Regulation (EU) 2022/2292 for veterinary medicine and prohibited substances.”

Version 3: Published 28 March 2023

Triangular trade section EU paragraph:

Amended to standardise the advice we provide on documentary evidence across POAO NFGs.