

Official health certificates: food-producing horses and horse products

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EU to propose revised official health certificates for food-producing horses and horse products

[Draft](#) Commission Implementing Regulation amending Annex III to Implementing Regulation (EU) 2020/2235 and Annex II to Implementing Regulation (EU) 2021/403 as regards model certificates for entry into the Union of consignments of certain products of domestic solipeds origin intended for human consumption and certain categories of equine animals

[Draft Annexes](#)

Update

The European Commission has informed the World Trade Organization Sanitary and Phytosanitary Measures (WTO SPS) Committee ([G/SPS/N/EU/891](#)) that it intends to amend the official certificates required to export horses that will be used in food, and horse products, to the European Union (EU). The certificates need to be amended to accurately reflect the EU's prohibition of the use of certain substances, such as thyrostatic (antithyroid) substances and stilbenes, which should apply not only to the exporting non-EU country, but also to the non-EU country where the horse was born and reared (if different).

Impacted products

Horses for use in food, horse products

What is changing?

The EU is proposing changes to the official certificates required for the export to the European Union of horses that will be used in food, and horse products. This affects the following model certificates:

- EQU, MP-PREP, MPNT, MPST in Regulation 2020/2235
- EQUI-X, EQUI-Y in Regulation 2021/403.

Details of the changes to these model certificates are given in Table 1.

Why?

The EU prohibits the use of certain substances, such as thyrostatic (antithyroid) substances and stilbenes, during the lifetime of a horse if the meat from that horse is intended to be consumed as food in the EU (Directive [96/22/EC](#)). Non-EU countries involved in the production of horse meat must be listed in Regulation [2021/405](#), Annex -I (see [Third country lists for public health – explained](#)).

Currently the wording of the official certificates that must accompany consignments of these products reflects these requirements only for the exporting country. It does not cover a non-EU country where the horse was born and reared, if different to the exporting country. This draft Regulation intends to align the wording of the certificates to the requirements of Directive [96/22/EC](#), Art. 3 (a), and Art. 11.

Timeline

The amended certificates are expected to apply from **30 October 2026**.

Recommended Actions

Where a non-EU country is exporting horses or their products that originated in another non-EU country, the official veterinarian signing the export certificate must check that this additional non-EU country fully complies with the EU rules (including non-use of prohibited substances), and is listed with an “X” for “equine” in Annex -I of Regulation 2021/405.

The WTO consultation closed on 30 December 2025.

Background

Council Directive [96/22/EC](#) bans the use of the following substances in food-producing horses [Art. 3 (a)], and bans imports into the EU of horses (and their products) that have been administered these substances (Art. 11):

- Aristolochia spp. and preparations, chloramphenicol, chlorpromazine, colchicine, dapsone, dimetridazole, metronidazole, nitrofurans, ronidazole (substances listed in Regulation 37/2010, Annex, Table 2)
- thyrostatic substances, stilbenes, stilbene derivatives, their salts and esters, oestradiol 17 β and its ester-like derivatives (Directive 96/22/EC, Annex II).

Other substances with action on hormones (oestrogenic, androgenic, or gestagenic action) and beta-agonists (used to enhance lean muscle gain) are also prohibited unless they are necessary for therapeutic treatments and/or animal husbandry (Directive [96/22/EC](#), Annex III).

Horse products can only be imported from countries listed with an “X” for “equine” in Annex -I of Regulation [2021/405](#), including the non-EU country where the horse was born and reared, if different from the exporting country. Countries that authorise the use of the above-mentioned substances in their legislation cannot be listed in this Annex (Directive [96/22/EC](#), Art. 11).

The rules establishing controls of residues of pharmacologically active substances were reviewed in 2022 (see [Official controls of veterinary drug residues in products of animal origin](#) and [Official controls on the use of pharmacologically active substances and their residues](#)).

For an overview of official certificates, see [EU official health certificates for exports to the EU – explained](#).

Resources

Council Directive [96/22/EC](#) concerning the prohibition on the use in stockfarming of certain substances having a hormonal or thyrostatic action and of beta-agonists

Commission Delegated Regulation (EU) [2022/1644](#) with specific requirements for the performance of official controls on the use of pharmacologically active substances authorised as veterinary medicinal products or as feed additives and of prohibited or unauthorised pharmacologically active substances and residues thereof

Commission Implementing Regulation (EU) [2022/1646](#) on uniform practical arrangements for the performance of official controls as regards the use of pharmacologically active substances authorised as veterinary medicinal products or as feed additives and of prohibited or unauthorised pharmacologically active substances and residues thereof, on specific content of multi-annual national control plans and specific arrangements for their preparation

European Commission (2025) [Guidelines on EU requirements for entry of animals and products of animal origin – Control plans for residues of veterinary medicines, pesticides and contaminants](#)

Sources

[Draft](#) Commission Implementing Regulation as regards model certificates for entry into the Union of consignments of certain products of domestic solipeds origin intended for human consumption and certain categories of equine animals

[Draft Annexes](#)

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Table & Figures

Table 1 Points to be modified in the model official certificates			
Regulation	Model certificate	Chapter	Point
2020/2235, Annex III	EQU	4	II.1.7
	MP-PREP	24	II.1.11
	MPNT	25	II.1.11
	MPST	26	II.1.11
2021/403, Annex II	EQUI-X	13	II.6
	EQUI-Y	14	II.6



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Source: [Draft Regulation](#) and [Draft Annexes](#)

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