Exporter Guidance:

Veterinary Certificate for the export of Products Containing Dairy and Egg from the United States to South Africa

The certificate, Veterinary Certificate for the export of Products Containing Dairy and Egg from the United States to South Africa may only be used for the export of protein drinks / supplements for human consumption containing both dairy and egg ingredients.

The Declarations in section IV are being endorsed based upon U.S. government regulatory controls, oversight and certification and the U.S. disease status. These declarations may not be endorsed based upon a notarized affidavit.

A. The following documents for the commodity (protein drink/ supplement) being exported and the animal origin ingredients must accompany the completed Veterinary Certificate for the export of Products Containing Dairy and Egg from the United States to South Africa (SA VC) when it is submitted to the Veterinary Services Area Office for endorsement

1. **Sanitary Certificate(s)** issued by a U.S. competent health authority (e.g. Agricultural Marketing Service, Food and Drug Administration, State Department of Health or State Dairy Board) that include the following information & statements:
   a. Manufacturer identification (name, address and approval number) – this information must be the same as the manufacturer (name, address) included in item I.A. on the SA VC;
   b. Commodity (protein drink / supplement) being exported – name of the commodity (protein drink or supplement) must be the same as the commodity (protein drink / supplement) identified for export in item I.B of the SA VC;
      i. The lot number of the processed egg product included in the protein drink or supplement must be included, **in parenthesis**, in the identification of the commodity being exported.
      NOTE: the inclusion of this information is for APHIS VS purposes only
   c. Processing / pasteurization statement – certification the dairy product(s) / ingredient(s) included in the commodity (protein drink/ supplement) for export received a pasteurization treatment or have been subjected to pasteurization;
      i. The exact pasteurization process parameters (time & temperature or pH) the dairy product(s) / ingredient(s) included in the commodity (protein drink/ supplement) for export were subjected to must be included **IF** a specific pasteurization / processing option (a. - c.) is selected on the SA VC.
   d. The following **exact** statements:
      i. *The product was manufactured in facilities inspected and approved by the competent authority and subjected to regular audits or inspections aimed at ensuring that the processing is properly and hygienically carried out, to produce a product that is fit for human consumption, AND, EITHER*
      ii. *To the best of our [my] knowledge the products contains no harmful levels of contaminants [additives];*
      OR
      iii. *The products do not, to the best of my [our] knowledge and belief, contain any harmful additive [contaminants];*

2. **FSIS Form PY-200, Egg Products Inspection and Grading Certificate** issued by the USDA Food Safety and Inspection Service, that includes the following information:
   a. Applicant – should identify the manufacturer / producer (name and address, including zip code)
   b. Name and Address of Receiver or Buyer – this information must be identified and be the same as the name and address of facility / manufacturer identified on Sanitary Certificate (described in item 1. above) manufacturing the commodity (protein drink/ supplement) for export
   c. Type of product – the type of processed egg product should be identified in this area
   d. Lot Number – must be identified


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i. This number must match the lot number of the processed egg product included in the protein drink or supplement being exported included in parenthesis in the identification of the commodity (protein drink or supplement) being exported.

NOTE: The identification and inclusion of this information is for APHIS VS purposes ONLY

e. Inspector (signature) – the signature of the FSIS inspector must be included

f. Date – the date the form was signed must be included

3. FDA export certificate (Certificate of Free Sale) issued by the Food & Drug Administration, that includes, the following information:

a. Product – the certificate must include the name of the commodity/ies (protein drink or supplement) being exported, and be the same as the name of the commodity (protein drink or supplement) included on the Sanitary Certificate (see A.1.b above) and commodity (protein drink/ supplement) name included in item I.B of the SA VC.

b. A valid copy of this certificate must be provided at least once a year to the VS Area Office where the SA VC are being submitted for endorsement.

c. VS requires the following conditions to be met in order for the certificate to be considered valid:

i. The certificate must have been issued by the FDA within the previous 12 months;

ii. The certificate must be within the period of validity, as designated by the FDA;

iii. The name of the commodity (protein drink / supplement) being offered for export must be identified on the certificate

NOTE: It is the exporter’s responsibility to insure the VS Area Office has a valid copy of the FDA export certificate (Certificate of Free Sale).

B. Special conditions for endorsements regarding processing of the animal origin ingredients

1. Pasteurization time and temperatures for dairy products

a. Identification of the particular pasteurization process (time & temperature or pH) to which the dairy product(s) were subjected is not required; item IV.B on the SA VC requires certification the dairy product was pasteurized.

b. If a particular pasteurization process is identified / selected (i.e. item IV.B.1. option a. b. or c.) on the SA VC, the Sanitary Certificate provided for the dairy product must also include the pasteurization processing parameters, (time & temperature or pH).

c. The manufacturer must contact the competent health authority with regulatory authority over their dairy processing plant to obtain information and make arrangements for the inclusion and certification of the pasteurization processing parameters (time & temperature or pH) the dairy products were subjected to on Sanitary Certificates.

NOTE: APHIS VS is not involved in this process.

2. Processing time and temperatures for egg product(s)

a. Section IV.C of the SA VC includes certification requirements for avian influenza, Newcastle disease and salmonella.

b. Avian influenza certification requirements – items IV.C options 1 or 2

i. Certification of country freedom from highly pathogenic notifiable avian influenza or the egg product(s) was/were treated to time and temperature suitable for the inactivation of highly pathogenic notifiable avian influenza virus in eggs and egg products;

ii. APHIS VS may certify the egg product has been treated to time and temperature suitable for the inactivation of highly pathogenic notifiable avian influenza virus in eggs and egg products (option #2) only when the FSIS Form PY-200, Egg Products Inspection and Grading Certificate includes the following information:

- type of [egg] product – this information must be consistent with the type of products identified in item IV.C.2 a through e, and

- processing time and temperatures – the processing parameters listed on the PY-200 must meet or exceed the time and temperature processing requirements listed on the SA VC for the type of [egg] product included in the commodity (protein drink or supplement) being exported.

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The manufacturer must work with FSIS to obtain information and make arrangements for the FSIS inspector to include and certify the processing time and temperatures the egg or egg products were subjected to on the PY-200.

NOTE: APHIS VS is not involved in this process.

c. **Newcastle disease** certification requirements – items IV.C options 3 or 4
   i. Certification of country or zone freedom from Newcastle disease or the egg product(s) was/were treated to a time and temperatures suitable for the inactivation of Newcastle disease virus, as defined by the OIE, in eggs and egg products;
   ii. APHIS VS may certify the egg product has been treated to a time and temperature suitable for the inactivation of Newcastle disease in eggs and egg products (option #4) only when the FSIS Form PY-200, *Egg Products Inspection and Grading Certificate* includes the following information:
      - type of [egg] product – this information must be consistent with the type of products identified in item IV.C.2 a through e, and
      - processing time and temperatures – the processing parameters listed on the PY-200 must meet or exceed the time and temperature processing requirements listed on the SA VC for the type of [egg] product included in the commodity (protein drink or supplement) being exported.

The manufacturer must work with FSIS to obtain information on what will be required in order for their inspectors to include and certify the specific processing time and temperatures the egg or egg products were subjected to on the PY-200.

NOTE: APHIS VS is not involved in this process.

d. **Salmonella** certification requirement – item IV.C.5
   i. This statement may be endorsed by APHIS VS based upon the certification statement included on the PY-200 indicating the egg product is compliant with 9 CFR 590.

C. Reminders & additional Information: VS - Export Products Policies etc.

   a. Letterhead Certificates may not be edited, altered, or amended, except as directed on the IREGS or in the Guidance documents.
   b. Exporters are responsible for confirming, **prior to shipment**, that they have all certification required for the consignment.
   c. Most countries will not accept certificates issued after consignments have shipped; exporter should obtain any required certificates prior to shipping.
   d. Exporters should contact the VS Area Office with jurisdiction in the area the production facility or exporter is located, with any questions or concerns.