

PA-OLVAC +I+E Vaccino inattivato in emulsione iniettabile per tacchini

Authorised

- Turkey haemorrhagic enteritis virus, Inactivated
- Newcastle disease virus, Inactivated
- Riemerella anatipestifer, serotype 1, Inactivated
- Riemerella anatipestifer, serotype 3, Inactivated
- Influenza A virus, subtype H9N2 (avian), Inactivated
- Influenza A virus, subtype H6N2 (avian), Inactivated

Product identification

Medicine name:

PA-OLVAC +I+E Vaccino inattivato in emulsione iniettabile per tacchini

Active substance:

Turkey haemorrhagic enteritis virus, Inactivated

Newcastle disease virus, Inactivated

Riemerella anatipestifer, serotype 1, Inactivated

Riemerella anatipestifer, serotype 3, Inactivated

Influenza A virus, subtype H9N2 (avian), Inactivated

Influenza A virus, subtype H6N2 (avian), Inactivated

Target species:

Turkey

Route of administration:

Subcutaneous use

Product details

Active substance and strength:

Turkey haemorrhagic enteritis virus, Inactivated

1733.00 enzyme-linked immunosorbent assay unit / 0.50 millilitre(s)

Newcastle disease virus, Inactivated

50.00 50% Protective Dose / 0.50 millilitre(s)

Riemerella anatipestifer, serotype 1, Inactivated

1.00 billion colony forming units / 0.50 millilitre(s)

Riemerella anatipestifer, serotype 3, Inactivated

1.00 billion colony forming units / 0.50 millilitre(s)

Influenza A virus, subtype H9N2 (avian), Inactivated

Influenza A virus, subtype H6N2 (avian), Inactivated

Pharmaceutical form:

Emulsion for injection

Withdrawal period by route of administration:**Subcutaneous use:**

-

Turkey

- Meat and offal. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QI01CL01

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Italy

Package description:

Available only in [Italian](#)

Available only in [Italian](#)

Available only in [Italian](#)

Available only in [Italian](#)

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Full application (Article 12(3) of Directive No 2001/82/EC)

Marketing authorisation holder:

Fatro S.p.A.

Marketing authorisation date:

9/03/2004

Manufacturing sites for batch release:

Fatro S.p.A.

Responsible authority:

Ministry Of Health

Authorisation number:

This information is not available for this product.

Date of authorisation status change:

9/03/2009

To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet

Documents

Combined File of all Documents

This document does not exist in this language (English). You can find it in another language below.