

# Improvac (--)- Solution for injection

Authorised

- Gonadotropin releasing factor analogue diphtheria toxoid conjugate

## Product identification

**Medicine name:**

Improvac (--)- Solution for injection

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**Active substance:**

Gonadotropin releasing factor analogue diphtheria toxoid conjugate

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**Target species:**

Pig

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**Route of administration:**

Subcutaneous use

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## Product details

**Active substance and strength:**

Gonadotropin releasing factor analogue diphtheria toxoid conjugate

Presentation\_strength:300 µg Reference:Hse Index:0

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**Pharmaceutical form:**

Solution for injection

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

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**Pig**

- Not applicable. 0 day Zero days

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**Anatomical therapeutic chemical veterinary (ATCvet) codes:**

QG03XA91

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**Legal status of supply:**

Veterinary medicinal product subject to veterinary prescription

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**Authorisation status:**

Valid

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**Authorised in:**

Austria , Belgium , Bulgaria , Croatia , Cyprus , Czechia , Denmark , Estonia , Finland , France , Germany , Greece , Hungary , Iceland , Ireland , Italy , Latvia , Liechtenstein , Lithuania , Luxembourg , Malta , Netherlands , Norway , Poland , Portugal , Romania , Slovakia , Slovenia , Spain , Sweden , United Kingdom (Northern Ireland)

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**Available in:**

Austria , Belgium , Czechia , Denmark , Finland , Greece , Hungary , Iceland , Italy , Latvia , Lithuania , Luxembourg , Netherlands , Poland , Romania , Slovakia , Spain , Sweden , United Kingdom (Northern Ireland)

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**Package description:**

Packaging:Polyethylene bottle with rubber closure and aluminium cap,

Package\_size:1 bottle, Content:250 ml

Packaging:Polyethylene bottle with rubber closure and aluminium cap,

Package\_size:1 bottle, Content:100 ml

Packaging:Polyethylene bottle with rubber closure and aluminium cap,

Package\_size:4 bottles, Content:250 ml

Packaging:Polyethylene bottle with rubber closure and aluminium cap,

Package\_size:10 bottles, Content:100 ml

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

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

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**Marketing authorisation holder:**

Zoetis Belgium

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**Marketing authorisation date:**

11/05/2009

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**Manufacturing sites for batch release:**

Zoetis Belgium

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**Responsible authority:**

European Commission

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**Authorisation number:**

This information is not available for this product.

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**Date of authorisation status change:**

14/10/2010

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To consult adverse reactions on veterinary medicinal products please go to [www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Combined File of all Documents

English (PDF)

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