

PARACOX ΠΟΣΙΜΟ ΕΝΑΙΩΡΗΜΑ

Not authorised

- Eimeria tenella, strain HP, Live
- Eimeria praecox, strain HP, Live
- Eimeria necatrix, strain HP, Live
- Eimeria mitis, strain HP, Live
- Eimeria maxima, strain CP, Live
- Eimeria maxima, strain MFP, Live
- Eimeria brunetti, strain HP, Live
- Eimeria acervulina, strain HP, Live

Product identification

Medicine name:

PARACOX ΠΟΣΙΜΟ ΕΝΑΙΩΡΗΜΑ

Active substance:

Eimeria tenella, strain HP, Live
Eimeria praecox, strain HP, Live
Eimeria necatrix, strain HP, Live
Eimeria mitis, strain HP, Live
Eimeria maxima, strain CP, Live
Eimeria maxima, strain MFP, Live
Eimeria brunetti, strain HP, Live
Eimeria acervulina, strain HP, Live

Target species:

Chicken

Route of administration:

Oral use

Product details

Active substance and strength:

Eimeria tenella, strain HP, Live

500.00 oocyst(s) / 0.10 millilitre(s)

Eimeria praecox, strain HP, Live

100.00 oocyst(s) / 0.10 millilitre(s)

Eimeria necatrix, strain HP, Live

500.00 oocyst(s) / 0.10 millilitre(s)

Eimeria mitis, strain HP, Live

1000.00 oocyst(s) / 0.10 millilitre(s)

Eimeria maxima, strain CP, Live

200.00 oocyst(s) / 0.10 millilitre(s)

Eimeria maxima, strain MFP, Live

100.00 oocyst(s) / 0.10 millilitre(s)

Eimeria brunetti, strain HP, Live

100.00 oocyst(s) / 0.10 millilitre(s)

Eimeria acervulina, strain HP, Live

500.00 oocyst(s) / 0.10 millilitre(s)

Pharmaceutical form:

Oral suspension

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

-

Chicken

- Meat and offal. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QI01AN01

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Surrendered

Authorised in:

Greece

Package description:

Available only in Greek

Available only in Greek

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Complete application (stand-alone)

Marketing authorisation holder:

Intervet Hellas A.E.

Marketing authorisation date:

17/01/1994

Manufacturing sites for batch release:

Merck Sharp & Dohme Animal Health S.L.

MSD Animal Health UK Limited

Responsible authority:

National Organization For Medicines

Authorisation number:

31896/93/18-01-1994/K-0085001

Date of authorisation status change:

15/07/2021

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