

OPTIVAR 500 MG BEE-HIVE STRIPS FOR HONEY BEES

Authorised

- Amitraz

Product identification

Medicine name:

OPTIVAR 500 MG BEE-HIVE STRIPS FOR HONEY BEES

Active substance:

Amitraz

Target species:

Honey bee

Route of administration:

In-hive use

Product details

Active substance and strength:

Amitraz

500.00 milligram(s) / 1.00 Strip

Pharmaceutical form:

Bee-hive strip

Withdrawal period by route of administration:**In-hive use:**

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Honey bee

- Honey. 0 day

Do not use during honey flow. Do not extract honey from the brood chamber. Do not harvest honey for human consumption while the treatment is in place. Brood combs should be replaced with new foundation at least every three years. Do not reuse brood frames as honey frames.

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP53AD01

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Cyprus

Package description:

Heat-sealed multi-layer sachet made of two films (PE/PA/Alu/PET and PE/Alu/oPA) containing 4 strips

Heat-sealed multi-layer sachet made of two films (PE/PA/Alu/PET and PE/Alu/oPA) containing 10 strips

Heat-sealed multi-layer sachet made of two films (PE/PA/Alu/PET and PE/Alu/oPA) containing 12 strips

Heat-sealed multi-layer sachet made of two films (PE/PA/Alu/PET and PE/Alu/oPA) containing 60 strips

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Bibliographic application (Article 22 of Regulation (EU) 2019/6)

Marketing authorisation holder:

Veto-Pharma

Marketing authorisation date:

23/06/2026

Manufacturing sites for batch release:

Veto-Pharma

Responsible authority:

Veterinary Services, Ministry Of Agriculture, Natural Resources And Environment

Authorisation number:

CY01038V

Date of authorisation status change:

9/04/2026

Reference member state:

France

Procedure number:

FR/V/0515/001

Concerned member states:

Bulgaria Croatia Cyprus Czechia Estonia Greece Hungary Ireland Italy
Latvia Lithuania Poland Portugal Romania Slovenia Spain
United Kingdom (Northern Ireland)

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

Documents

eu-puar-frv0515001-mr-rpe1052-en.pdf