Packaging ‘blue-box’ requirements and additional information on labelling/package leaflet for products authorised via national, mutual recognition, decentralised or centralised procedures

Additional information on labelling/package leaflet that may be required or permitted nationally in accordance with Articles 58 and 64 of Directive 2001/82/EC, as amended, is outlined below.

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Additional Requirements for the Labeling

Legal Status
The following are the specific requirements for the expression of the legal status:

- “Rezept- und apothekenpflichtig” = available only on prescription and only in pharmacies;
- “Apothekenpflichtig” = available only in pharmacies;
- If the supply is not restricted to pharmacies, this has to be declared appropriately.

Price
The price is not wanted on the label.

Identification
The Europäische Artikelnummerierung (EAN, barcode) is accepted on the label, but not required.

Additional Requirements for the Package Leaflet
For veterinary medicinal product containing prohibited substances in accordance with § 1 section 2 of the Austrian Anti-Doping Convention 2007, the following sentence has to be added:

“Die Anwendung des Arzneimittels [Bezeichnung des Arzneimittels einsetzen] kann bei Dopingkontrollen zu positiven Ergebnissen führen.” (= The administration of the medicinal product may result in positive doping controls).

Identification
The marketing authorisation number “Z.Nr.” is required in the package leaflet.
Belgium (BE)

Additional Requirements for the Labelling

Legal Status
For medicinal products restricted to special prescription (narcotics), a number code assigned by the Minister of Health and a double red line are mandatory. This double red line must be as large as the largest character on the label. The red lines should be parallel, 1 – 3 cm apart and in an angle of 45° starting from the left lower corner to the right upper corner.

Identification
A bar code is accepted on the label, but not required.
Additional Requirements for the Labelling and Package leaflet

The primary and outer packaging of the VMP containing narcotic substances shall be identified (marked) diagonally by two red lines (strips), while the psychotropic ones shall be marked by two blue strips.

In the cases where the marketing authorisation of VMP has been issued in accordance with the centralized procedure, each authorized or required by the National Veterinary Service (NVS) additional information shall be placed (written) within an area bordered by a frame that shall clearly outline it from all the other data.

The data placed on the primary and on the outer packaging and also in the instruction for use shall be written in Bulgarian language.

The labels of the homeopathic VMPs shall involve the note ‘Хомеопатичен ветеринарномедицински продукт’.

The note ‘Само за ветеринарномедицинска употреба’ shall be placed on each primary and outer packaging of any VMP.

The data in the instruction for use may be written in several languages, one of which must be Bulgaria, but if only the data written in all these languages are identical.

Where the VMP must be sold and used under veterinarian’s prescription, the note ‘По лекарско предписание’ shall be placed on each primary and outer packaging of the VMP concerned, excluding the homeopathic ones.

In the other cases the note ‘Без лекарско предписание’ shall be placed on each primary and outer packaging of the VMP.

All VMPs intended for food production animals and the VMPs, which are subject to special measures to be taken by the veterinarian in order to avoid any risk related to the animals treated or the persons applying the VMP or the environment, shall also be subject to the same requirements, i.e. mandatory identification by the note ‘По лекарско предписание’.
CROATIA (HR)

No Additional Requirements for the Labelling and Package Leaflet.
No Additional Requirements for the Labelling and Package Leaflet.

**Legal Status**

There is no requirement for the legal status to appear on the label.

**Identification**

Information for the identification and authenticity are not required on the label. Bar codes are accepted on the label but are not required.
Additional Requirements for the Labelling

**Legal Status**

The following words are specific for categories of prescription

- depending on category of veterinary medicinal products that are subject to prescription:

<Na předpis.> or <Veterinární léčivý přípravek je vydáván pouze na předpis.> (Where this information is not included in the approved text for packaging according to the current veterinary product template)

When the veterinary medicinal product is only intended for administration by a veterinary surgeon, the above mentioned sentence shall followed by the present statement:
<Poručen pro použití veterinárním lékařem.>

For specific veterinary medicinal product with restricted prescription (narcotic and psychotropic substances) according the national act No. 167/1998 Coll.

The picture shall be completed with blue strip if the substance is additionally classified in accordance with §13, section 1 of the above mention national act.

- depending on category of veterinary medicinal products that are not subject to prescription:

For veterinary medicinal products available in pharmacy:
<Bez předpisu.> or <Veterinární léčivý přípravek je vydáván bez předpisu.> (Where this information is not included in the approved text for packaging according to the current veterinary product template)

For specific veterinary medicinal products that are not subject to prescription and are available in authorised merchants – petshops:
<Vyhrazený veterinární léčivý přípravek.>

The following words (Indikační omezení = prudent use) are required for certain veterinary medicinal products containing antimicrobial substances, in accordance with national law. Concerning the veterinary medicinal products which are the subjects to prescription and which are intended to be reserved for the treatment of severe infections and clinical conditions which have responded poorly, or are expected to respond poorly (refers to very acute cases when treatment must be initiated without bacteriological diagnosis), to first line treatment. Their use should be whenever possible based on results of susceptibility testing of target pathogens and on detection of resistance to other classes of antibiotics.

<Indikační omezení>
The following words shall be used for homeopathic products:
< Veterinární homeopatický léčivý přípravek>
When the veterinary homeopathic product is approved by the simplified registration procedure, the use text shall be as follows:
< Veterinární homeopatický léčivý přípravek bez schválených léčebných indikací>

Identification
There is no requirement for the EAN1 bar codes to appear on the label. The EAN bar codes are accepted when they are put on the label.

Additional information
Recycling symbols are accepted.
DENMARK (DK)

Additional Requirements for the Labelling

Legal Status
There is no specific requirement in respect of the legal status.

Identification
The Nordic number is required on the outer labelling of all medicinal products, except radiopharmaceuticals, certain vitamins and mineral products, homeopathic, herbal and traditional herbal medicinal products. It may be written as “Vnr XX XX XX”.
A bar code is accepted but not required on the labelling.

Additional requirements (only for national, MR & DC products)
Other warnings to be included in the labelling are listed in section 29(1-3) and section 31(4-6) of the Danish executive order no 869 of 21 July 2011, as amended on labelling etc. of medicinal products (Danish title: Bekendtgørelse nr. 869 af 21. juli 2011, med senere ændringer, om mærkning m.m. af lægemidler).

Additional Requirements for the Package Leaflet (only for national, MR & DC products)

<table>
<thead>
<tr>
<th>Section / Explanation</th>
<th>English translation</th>
<th>Danish text required in the package leaflet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section 8, Dosage for each species, route(s) and method of administration</td>
<td>Please notice that your veterinarian may have prescribed the medicinal product for a different therapeutic indication and/or at a different dosage than stated in the package leaflet. Always follow the veterinarian's prescription and the instructions on the dosage label.</td>
<td>Vær opmærksom på, at dyrlægen kan have foreskrevet anden anvendelse eller dosering end angivet i denne information. Følg altid dyrlægens anvisning og oplysningerne på doseringsetiketten.</td>
</tr>
<tr>
<td>Section 6, Adverse reactions: - After the sentence &quot;If you notice any serious effects..&quot;</td>
<td>Side effects can thereby be reported to the Danish Health and Medicines Authority and the knowledge about side effects can be improved. The owner of the animal can also report side effects directly to the Danish Health and Medicines Authority. You can find guidance on the Danish Health and Medicines Authority's website (see Pharmacovigilance). <a href="http://www.sst.dk">www.sst.dk</a></td>
<td>Bivirkningerne kan dermed blive indberettet til Sundhedsstyrelsen, og viden om bivirkninger kan blive bedre. Dyrets ejer kan også indberette bivirkninger direkte til Sundhedsstyrelsen. De/du finder skema og vejledning under Bivirkninger på Sundhedsstyrelsens netsted. <a href="http://www.sst.dk">www.sst.dk</a></td>
</tr>
</tbody>
</table>

Additional information (only for centralised products):
Products containing inflammable material must bear the international warning symbol: 😡
ESTONIA (EE)

No Additional Requirements for the Labelling and Package Leaflet.
Additional Requirements for the Labelling

**Legal Status**
There is no requirement for the legal status to appear on the label.

**Identification and Authenticity**
The Nordic number is required on the label of all medicinal products, except immunological products, radiopharmaceuticals and herbal remedies. It is written as “Vnr XX XX XX”.
A bar code is accepted on the label but not required.

**Symbols and Pictograms**

• Products containing inflammable material must bear the international warning symbol:

![Flammable Symbol](image-url)
Legal Status

The information that the product is a prescription only medicine has to appear in dark ink on a red background rectangle as:

- “A NE DELIVRER QUE SUR ORDONNANCE » for products intended for non-food producing.
- “A NE DELIVRER QUE SUR ORDONNANCE DEVANT ETRE CONSERVEE PENDANT AU MOINS 5 ANS” for products intended for food producing animals.
- « VACCIN : DELIVRANCE SOUMISE A ORDONNANCE » for vaccines.
- « PRODUIT IMMUNOLOGIQUE : DELIVRANCE SOUMISE A ORDONNANCE » for immunological products other than vaccines (sera for instance).

The information that the product is a veterinary medicine has to be mentioned as “USAGE VETERINAIRE” written in dark ink in the same red background rectangle.

For medicinal products containing an active substance subject to a special regulation in France (narcotic, psychotropic or so called “substances vénéneuses”), it must be added:

- In the red background rectangle “ RESPECTER LES DOSES PRESCRITES” and, if the medicinal products is to be administered by a route different than nasal, oral, per lingual, sublingual, rectal, vaginal, urethral or by injection, “NE PAS FAIRE AVALER”.

Above the red background rectangle

- an empty (white) rectangle with a red border for List I substances
- or
- an empty (white) rectangle with a green border for List II substances

Identification

If appropriate, the French MA number has to appear on the label.
Additional Requirements for the Labelling

Legal Status
The legal status is required on the label:

- “apothekenpflichtig” = to appear in the case of medicinal products that are not subject to medical prescription but are only available in pharmacies or from veterinarians
- “verschreibungspflichtig; Betäubungsmittel” in case of narcotics (only for national, MR & DC products)
- No separate statement is necessary in the case of products, which are neither prescription only nor pharmacy only.

Identification and Authenticity

- In respect of sera, the animal species from which they were obtained, in respect of vaccines, particulars of the host system serving the multiplication process of the virus shall be given. (only for label & outer package)
- A barcode is accepted on the label. A distribution number (PZN, i.e. Pharmazentralnummer) is accepted on the label.

Additional information:

- In case of samples the indication “unverkäufliches Muster” (sample – not for sale) is required. (only for national, MR & DC products)
- A special symbol concerning the recycling of the packaging material is accepted such as the “Grüne Punkt”.

Additional Requirements for the Package leaflet (only for national, MR & DC products)

- Specific requirements in case of narcotics have to be stated.
**GREECE (EL)**

**Additional Requirements for the Labelling**

**Legal Status**

Veterinary medicinal products subject to a special prescription (narcotics) must have a letter/code assigned by the Ministry of Health and Welfare with a special colour (read/green) according to the classification and the following text must appear on the label:

1. Products belonging to list B must mention in red letters:
   «Β, χορηγείται με ειδική συνταγή Ναρκωτικών».

2. Products belonging to the exceptions of list B must mention in green letters:
   «ΒΣ, χορηγείται με απλή συνταγή Ναρκωτικών».

3. Products belonging to list Γ must mention in red letters:
   «Γ, χορηγείται με ειδική συνταγή Ναρκωτικών».

4. Products belonging to the exceptions of list Γ must mention in green letters:
   «ΤΣ, χορηγείται με απλή συνταγή Ναρκωτικών».

5. Products belonging to list Δ must mention in green letters:
   «Δ, χορηγείται με συνταγή του Ν. 1729/98». 
HUNGARY (HU)

No Additional Requirements for the Labelling and Package Leaflet.
ICELAND (IS)

Additional Requirements for the Labelling

**Identification**

The Nordic Article Number is required on the outer package of all veterinary medicinal products except radiopharmaceuticals, homeopathics and herbal remedies. It is written as “Vnr xx xx xx”.

A barcode is accepted on the package but not required.
IRELAND (IE)

The following requirements are in addition to those of Directive 2001/82/EC as amended and the QRD templates.

<table>
<thead>
<tr>
<th>Immediate packaging</th>
<th>Outer packaging</th>
<th>Package Leaflet</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Legal status</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abbreviation for route of sale and supply, as appropriate:</td>
<td>Abbreviation for route of sale and supply, as appropriate:</td>
<td>Abbreviation for route of sale and supply and explanatory phrase, as appropriate:</td>
</tr>
<tr>
<td>VPO</td>
<td>VPO</td>
<td>VPO Veterinary Practitioner Only</td>
</tr>
<tr>
<td>VPO-1</td>
<td>VPO-1</td>
<td>VPO-1 Veterinary Practitioner Only</td>
</tr>
<tr>
<td>POM</td>
<td>POM</td>
<td>POM Prescription Only Medicine</td>
</tr>
<tr>
<td>POM(E)</td>
<td>POM(E)</td>
<td>POM(E) Prescription Only Medicine (Exempt)</td>
</tr>
<tr>
<td>PS</td>
<td>PS</td>
<td>PS Pharmacy Only</td>
</tr>
<tr>
<td>LM</td>
<td>LM</td>
<td>LM Licensed Merchant</td>
</tr>
<tr>
<td>CAM</td>
<td>CAM</td>
<td>CAM Companion Animal Medicine</td>
</tr>
</tbody>
</table>

**Identification - nationally authorised products**

<table>
<thead>
<tr>
<th>Veterinary Product Authorisation (VPA) number</th>
<th>Veterinary Product Authorisation (VPA) number</th>
<th>Veterinary Product Authorisation (VPA) number</th>
</tr>
</thead>
</table>

**Other requirements**

For nationally authorised immunological products only: If a product is classified as LM the following warning is required: “Prior to first time use on a farm, it is strongly recommended that the advice of a veterinary practitioner is sought”.

For other products, the following warning is required: “Prior to first time use on a farm, it is strongly recommended that the advice of a veterinary practitioner is sought”.

For veterinary antisepsics and disinfectants, the following warning is required: “Prior to first time use on a farm, it is strongly recommended that the advice of a veterinary practitioner is sought”.

For veterinary immunological products, the following warning is required: “Prior to first time use on a farm, it is strongly recommended that the advice of a veterinary practitioner is sought”.

For veterinary pharmaceutical products, the following warning is required: “Prior to first time use on a farm, it is strongly recommended that the advice of a veterinary practitioner is sought”.
ITALY (IT)

Additional Requirements for the Labelling

Legal Status

For products “subject to prescription”:

• When the veterinary medicinal product contains an active substance of the pharmacological groups listed in article 2, letters c) and h) of the D.M. 28/07/2009:
  “Da vendersi soltanto dietro presentazione di ricetta medico-veterinaria in triplice copia non ripetibile” or “Medicinale veterinario soggetto a ricetta.... (veterinary prescription to be decided, on a case by case basis), secondo D.P.R. 309/90 e successive modifiche, tabella medicinali sezione ....” (with the correct letter specified by the Italian authority on a case by case basis according to "Decreto Presidente della Repubblica 9 ottobre 1990, n. 309” as amended).
followed by:
“La somministrazione e detenzione del medicinale deve essere effettuata esclusivamente dal medico veterinario.”

• When the veterinary medicinal product contains an active substance of the pharmacological groups listed in art. 2 of the D.M. 28/07/2009:
  “La somministrazione del medicinale deve essere effettuata esclusivamente dal medico veterinario.”

The veterinary prescription to be decided according to articles 75 and 76 of D. Lgs. n. 193, 06/04/2006.

• When the veterinary medicinal product is intended for food producing animals, according to D. Lgs. n. 193, 06/04/2006, art. 76:
  “Da vendersi dietro presentazione di ricetta medico-veterinaria in triplice copia non ripetibile”
  (to be sold only with three copies of a non-renewable vet. med. prescription)

• When the veterinary medicinal product is intended for food-producing animals or companion animals, according to D. Lgs. n. 193, 06/04/2006, art. 75:
  “Da vendersi dietro presentazione di ricetta medico-veterinaria non ripetibile in copia unica” (to be sold only with a non-renewable vet. med. prescription)

• or, only for a veterinary medicinal product intended for companion animals, according to D. Lgs. n. 193, 06/04/2006, art. 76, section 6:
  “Da vendersi dietro presentazione di ricetta medico-veterinaria ripetibile” (to be sold with a renewable vet. med. prescription)

• In the case of a veterinary medicinal product authorized without prescription.
  “Medicinale veterinario senza obbligo di ricetta medico veterinaria”
For veterinary medicinal product containing psychotropic substances, the following sentence has to be specified:

“Medicinale veterinario soggetto a ricetta.... (veterinary prescription to be decided, on a case by case basis), secondo D.P.R. 309/90 e successive modifiche, tabella medicinali sezione ....” (with the correct letter specified by the Italian authority on a case by case basis according to “Decreto Presidente della Repubblica 9 ottobre 1990, n. 309” as amended).

**IDENTIFICATION**

The marketing authorization identification number (A.I.C.) and the barcode/GTIN (DM 17/12/2007) are required in the label.

The national identification number (N.I.N.) substitutes the A.I.C. number (for centralised authorised products).

**OTHER INFORMATION**

Any other information according to D. Lgs. n. 193, 06/04/2006, articles 30 or 58, section 5, published in the “Supplemento ordinario alla Gazzetta Ufficiale” n. 121, 26/05/2006 is required, where applicable.

The manufacturer responsible for the batch release should be mentioned on the outer label.

**Additional Requirements for the package leaflet**

Information about the waste collection system should be included.
Additional Requirements for the Package leaflet

Restrictions regarding supply must be mentioned in Section 15.

*Only for centralised authorised products*

**Legal Status**

1. For immunologicals, general anesthetics, narcotic analgesics, psychotropics and for specific products which have special precautions for use or they can cause potential hazard to human or animal health:

   “Izplatīšanai tikai praktizējošam veterinārārstam.”

   (For supply only to veterinary practitioner).

2. For all veterinary medicinal products: restrictions regarding supply must be mentioned under Section 15 of Package leaflet.
LIECHTENSTEIN (LI)

No Additional Requirements for the Labelling and Package Leaflet.
LITHUANIA (LT)

No Additional Requirements for the Labelling and Package Leaflet.
No Additional Requirements for the Labelling and Package Leaflet.
MALTA (MT)

No Additional Requirements for the Labelling and Package Leaflet.
Additional Requirements for the Labelling

**Legal Status**

**UDD** – (Uitsluitend door dierenartsen toe te dienen – administration only by veterinarian), or

**UDA** – (Op recept van dierenarts, af te leveren door de dierenarts of apotheker – prescription by veterinarian, for supply by veterinarian or pharmacist), or

**URA** – (Op recept van dierenarts, af te leveren door de dierenarts, apotheker of vergunninghouder – prescription by veterinarian, for supply by veterinarian, pharmacist or licensed retailer), or

**VRIJ** – (Freely available without prescription in pharmacies, pet shops and by licensed retailers)

**Identification**

The national identification number is required on the label.
Additional Requirements for the Labelling

**Legal status:**
There are no additional requirements

**Identification**
The Nordic number is required on the outer label of all medicinal products except for radiopharmaceuticals, homeopathics and herbal remedies. It is written as "Vnr XX XX XX". A barcode is accepted on the label but not required.

**Additional warnings:**
Products containing inflammable material must bear the international warning symbol:

- Brannfarlig + symbol

- English translation: Inflammable + symbol
Additional Requirements for the Labelling

Legal Status
The following are the specific requirements for the expression of the legal status in the boxed area:

- **Legal status regarding categories of administration:**
  - Wydawany z przepisu lekarza - Rp = available on prescription only
  - Wydawany bez przepisu lekarza - OTC = available without prescription

- **Legal status regarding distribution/dispensing:**
  - Do podawania wyłącznie przez lekarza weterynarii = Administration by the veterinary surgeon only
  - Do podawania pod nadzorem lekarza weterynarii = Administration by the veterinary surgeon or under their direct responsibility
  - Do podawania przez właściciela lub opiekuna zwierzęcia = Administration by the owner or animal keeper

The description of the legal status must be exactly the same as in the Marketing License (= Pozwolenie na dopuszczenie do obrotu)

Identification and Authenticity
The EAN code is required on the label.
Items mandatory for both pharmacologicals and immunologicals

1. Legal status and withdrawal period are required in the blue box, if not mentioned elsewhere on the labels
2. For generic veterinary medicinal products “MVG” should be included in the outer packaging.
3. If applicable, specific statements, symbols or safety warnings concerning the handling/administration/storage/disposal of the veterinary medicinal product may be required on the label as, for example:

   A INJEÇÃO ACIDENTAL É PERIGOSA - ANTES DE UTILIZAR LEIA O FOLHETO INFORMATIVO. (accidental injection is dangerous – Read the package leaflet before use)

4. Identification

   “AIM nº” – the marketing authorisation number is required on the label for National Procedures, MRP and DCP. A barcode is accepted.
   “N° de Código Nacional “ - the national code of the veterinary medicinal product is required on the label for centralized procedures. A barcode is accepted.

5. “Manter fora do alcance e da vista das crianças” (Keep out of reach and sight of children) if not mentioned elsewhere on the label
6. The expression “Uso Veterinário” must be stated in an entirely green boxed area.
7. If not already mentioned, in the package leaflet should appear the description of the visual appearance of the veterinary medicinal product’s pharmaceutical form as marketed e.g. shape, texture, colour, imprint: In case of veterinary medicinal products intended for reconstitution, the appearance of the veterinary medicinal product before and after reconstitution.

Specific items for pharmacologicals to be also included

1. If applicable, specific statements concerning the administration and/or availability of the veterinary medicinal product may be required on the label as one of the following:
   • “Só pode ser administrado pelo médico-veterinário” (administered by a veterinarian only)
   • “Só pode ser administrado sob controlo do médico veterinário” (to be administered under the responsability of a veterinarian)
2. Products for external use should state "Uso externo“ in a entirely red boxed area on the label.
3. Medicated premixes: “Só pode ser vendido a unidades de fabrico de alimentos compostos para animais”

Specific items for IVMP – Immunological Veterinary Medicinal Products to be also included

1. The following sentences are mandatory unless authorised otherwise:
   “Só pode ser administrado pelo médico veterinário” (to be administered by the veterinarian only)
   or
“Só pode ser administrado sob controlo do médico veterinário” (to be administered under the responsibility of a veterinarian)

2. Name/address of the local representative/distributor in labels and package leaflet

USO EXTERNO

USO VETERINÁRIO

Nº de Código Nacional (Nº Cód. Nac.)
or
Nº de AIM (depending if it is a Community or a national authorisation)
Additional Requirements for the Labelling

**Price**
There is no requirements for the price to appear on the label and package leaflet.

**Reimbursement**
There is no requirement for the reimbursement conditions to appear on the label.

**Legal Status**
The legal status is required to be expressed on the label for prescription-only products.
The following mentions must appear in the boxed area:
For medicinal products supplied in pharmacy based on veterinary prescriptions:
- Se elibereaza pe baza de prescriptie medicala : **PRF** = available with prescription
- Se elibereaza pe baza de prescriptie medicala speciala cu timbru sec: **PTS** = available with special prescription / stamp
- Se elibereaza fara prescriptie medicala : **OTC** = available without prescription

**Symbols or pictograms**
Medicinal products containing inflammable material must bear the international warning symbol:

![Warning Symbol]

**If veterinary medicinal product contains a narcotic or psychotropic substance indicate a follows:**
- **Produsul medicinal veterinar contine o substanta narcotica** (Veterinary product contains a narcotic substance.)
- **Produsul medicinal veterinar contine o substanta psihotropa** (Veterinary product contains a psychotropic substance.)

**Identification**
Veterinary Product Authorisation (VPA) number on the label.
In justified cases VPA may be omitted
Barcodes are accepted on the label, but are not required.
Additional Requirements for the Labelling

**Legal status**
For specific veterinary medicinal products with restricted prescription (narcotic and psychotropic substances) according the national act No. 362/2011 Coll:

<VETERINÁRNÝ LIEK OBSAHUJE OMAMNÚ LÁTKU.>
or
<VETERINÁRNÝ LIEK OBSAHUJE PSYCHOTROPNÚ LÁTKU.>

For **homeopathic** veterinary products shall be used the following words:
<HOMEOPATICKÝ VETERINÁRNÝ LIEK.>

When the homeopathic veterinary product is approved by the simplified authorisation procedure shall be used the following words:
<HOMEOPATICKÝ VETERINÁRNÝ LIEK BEZ SCHVÁLENÝCH TERAPEUTICKÝCH INDIKÁCIÍ.>

When the homeopathic veterinary product was not pharmacologically, toxicological and clinical tested shall be used the following words:
<VETERINÁRNÝ LIEK NEBOL KLINICKÝ SKÚŠANÝ.>

For the **radioactive veterinary** medicinal product shall be used the international symbol for radioactivity and the quantity of radioactivity.

**Identification**
The GTIN bar codes are accepted when they are put on the label

**Additional information**
Recycling symbols are accepted

**Package leaflet**
Where veterinary medicinal product is subject to prescription:

<Na veterinárne použitie - Výdaj len na veterinárny lekársky predpis>
SLOVENIA (SI)

No Additional Requirements for the Labelling and Package Leaflet.
Additional Requirements for the Labelling

The following requirements for outer packaging of veterinary medicines are according to Real Decreto 1246/2008, de 18 de julio, por el que se regula el procedimiento de autorización, registro y farmacovigilancia de los medicamentos veterinarios fabricados industrialmente.

**Legal status:**
The medicinal product may only be supplied in accordance with a prescription (legends and pictograms), including controlled drugs (psicotropics according to Annex I or II of RD 2829/1977):

- Supplied with a prescription: Dispensación con receta veterinaria
- Supplied with a prescription for narcotics: Dispensación con receta de estupefacientes
- Supplied with a prescription for psicotropics: Dispensación con receta de psicótropos anexo I del RD 2829/1977
- Supplied with a prescription for psicotropics: Dispensación con receta de psicótropos anexo II del RD 2829/1977

**Requirements of preservation:**

- Preservation in a fridge: Conservación en frigorífico
- Preservation in a freezer Conservación en congelador

**Further information**

- The medicinal product may only be administered by a surgeon: Administración exclusiva por el veterinario AV
- In all medicinal products for veterinary use: “USO VETERINARIO”.
- In the particular case of medicinal premixes for feed: “Premezclas medicamentosas para piensos”.
- National code number (Código Nacional) is granted for every authorised format of all medicinal products and required on the outer packaging. It is written as C.N. XXXXXX.X in the main face on the upper right corner.
- In cases requiring a prescription symbol or “AV” should be included on a dedicated place of the outer package, upper right corner, outside the blue-box.
Identification and authenticity

The Nordic number is required on the label of all medicinal products, except radiopharmaceuticals and herbal remedies. It is written as “Vnr XX XX XX”. A bar code is accepted on the label but not required.

Additional requirements regarding OTC for Labels and the Package Leaflet

As the Legal status may differ between MS there is a need for the possibility to make additional amendments as adequate for the legal status.

Additional information:

• Products containing inflammable material must bear the international warning symbol:-

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UNITED KINGDOM (UK)

Additional Requirements for the Labeling

Legal Status

1. The medicinal product may only be supplied in accordance with a prescription:

   **POM-V**
   Medicines may only be prescribed by a registered veterinary surgeon for an animal under his care. The prescription may be dispensed by any registered veterinary surgeon or registered pharmacist.

   **POM-VPS**
   Medicines which can be prescribed and supplied by a Veterinarian Surgeon, Pharmacist or a registered Suitably Qualified Person (SQP) or it may be supplied separately by one of the above in accordance with a written prescription from that person.

2. The medicinal product may be sold or supplied without a prescription:

   **NFA-VPS**
   Medicines which can be supplied without a prescription by a Veterinary Surgeon, Pharmacist or a Suitably Qualified Person (SQP).

   **AVM-GSL**
   Medicines which may be supplied by any retailer. Products which do not require specific advice concerning their method of use, and pose minimal safety risks.

3. Controlled Drug (CD):
   Medicinal products considered to be dangerous and likely to be subject to abuse. Additional precautions in respect of storage and supply are required. These products are also **POM-V**.

\[\text{CD} \text{, followed by Sch 2 or Sch 3 as appropriate}\]

Identification

Information for the identification and authenticity are not required on the label. Barcodes are accepted on the label, but are not required.

Additional Information

‘Keep out of reach of children’ – if not included elsewhere on product literature

‘Keep the container in the outer carton’