DRAFT BLISTER LABEL TEXT

1. NAME OF THE VETERINARY MEDICINAL PRODUCT		
NOROCLAV 500 mg PALATABLE TABLETS FOR DOGS		
2. NAME OF THE MARKETING AUTHORISATION HOLDER		
Norbrook Laboratories Limited		
3. EXPIRY DATE		
XX/XX/XXXX		
4. BATCH NUMBER		
XXXX-XX		
5. THE WORDS "FOR ANIMAL TREATMENT ONLY"		
FOR ANIMAL TREATMENT ONLY		

Do not store above 25°C

DRAFT CARTON TEXT

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Noroclav 500mg Palatable Tablets for Dogs

2. STATEMENT OF ACTIVE AND OTHER SUBSTANCES

Noroclav 500mg Palatable Tablets for Dogs are for oral administration to dogs: each tablet contains Amoxicillin 400 mg (as Amoxicillin Trihydrate), Clavulanic Acid 100 mg (as Potassium Clavulanate) and Carmosine Lake (E122) 2.45mg

3. PHARMACEUTICAL FORM

Tablets.

The tablets can be divided into equal halves.

4. PACKAGE SIZE

10 Tablets

20 Tablets

25 Tablets

100 Tablets

5. TARGET SPECIES

Dogs

6. INDICATION(S)

For the treatment of the following infections caused by beta-lactamase producing strains of bacteria sensitive to amoxicillin in combination with clavulanic acid:

- Skin infections (including superficial and deep pyodermas) caused by susceptible *Staphylococci*.
- Urinary tract infections caused by susceptible *Staphylococci* or *Escherichia* coli.
- Respiratory infections caused by susceptible Staphylococci.
- Enteritis caused by susceptible Escherichia coli.

It is recommended to carry out suitable tests for sensitivity when initiating the treatment. The treatment should only proceed if sensitivity is proven to the combination.

7. METHOD AND ROUTE(S) OF ADMINISTRATION

To ensure a correct dosage, bodyweight should be determined as accurately as possible to avoid underdosing.

Administration is via the oral route. The dosage rate is 12.5 mg combined actives/kg bodyweight twice daily.

The following table is intended as a guide to dispensing Noroclav Palatable Tablets at the standard dose rate of 12.5 mg/kg twice daily. The tablets may be crushed and added to a little food.

Bodyweight (kg)	Number of tablets per dose twice daily
20 kg	1/2
40 kg	1
60 kg	1½
80 kg	2

Duration of therapy:

Routine cases involving all indications: The majority of cases respond to between 5 and 7 days therapy.

Chronic or refractory cases: In these cases where there is considerable tissue damage, a longer course of therapy may be required in that it allows sufficient time for damaged tissue to repair.

8. WITHDRAWAL PERIOD

Not Applicable.

9. SPECIAL WARNING(S)

Read the package leaflet before use

Do not use in animals with known cases of hypersensitivity to penicillins or other substances of the beta-lactam group.

Do not use in rabbits, guinea pigs, hamsters or gerbils.

Do not use in animals with serious dysfunction of kidneys accompanied by anuria or oliguria.

Do not use where resistance to the combination is known to occur.

Do not administer to horses and ruminating animals.

Operator Warnings:

Penicillins and cephalosporins may cause hypersensitivity (allergy) following injection, inhalation, ingestion or skin contact. Hypersensitivity to penicillins may lead to cross-reactions to cephalosporins and vice versa. Allergic reactions to these substances may occasionally be serious.

Handle this product with great care to avoid exposure, taking all recommended precautions.

If you develop symptoms following exposure such as a skin rash, you should seek medical advice and show the package leaflet or the label to the physician.. Swelling of the face, lips or eyes or difficulty with breathing are more serious symptoms and require urgent medical attention.

Wash hands after use.

10. EXPIRY DATE

XX/XXX

11. SPECIAL STORAGE CONDITIONS

Do not store above 25°C. Store in a dry place. Return any halved tablet to the open blister pack. Any divided tablet portion remaining after 24 hours should be discarded. Keep the blister in the outer carton.

12. SPECIAL PRECAUTIONS FOR THE DISPOSAL OF UNUSED PRODUCTS OR WASTE MATERIALS

Dispose of any unused product or empty containers in accordance with guidance from your local waste regulation authority

13. THE WORDS "FOR ANIMAL TREATMENT ONLY" AND CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE, if applicable

FOR ANIMAL TREATMENT ONLY

14. THE WORDS "KEEP OUT OF THE REACH AND SIGHT OF CHILDREN"

KEEP OUT OF THE REACH AND SIGHT OF CHILDREN

15. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Marketing Authorisation Holder:

(EU)

Norbrook Laboratories (Ireland) Limited Rossmore Industrial Estate Monaghan Ireland

(UK)

Norbrook Laboratories Limited Station Works Camlough Road Newry Co. Down BT35 6JP Northern Ireland

DISTRIBUTED IN THE UK BY:

Norbrook Laboratories Limited Carnbane Industrial Estate Newry Co. Down BT35 6QQ Northern Ireland

DISTRIBUTED IN IE BY:

Norbrook Laboratories (Ireland) Limited Rossmore Industrial Estate MONAGHAN

16. MARKETING AUTHORISATION NUMBER(S)

ManA 2000 Vm: 02000/4259 VPA: 10999/118/001

17. MANUFACTURER'S BATCH NUMBER

XXXX-XX

LEGAL CATEGORY:

UK- POM -V

To be supplied only on veterinary prescription.

IE - POM Prescription Only Medicine

FURTHER INFORMATION

Amoxicillin is a broad-spectrum antibiotic active against a wide range of Grampositive and Gram-negative bacteria. However, many clinically important bacteria produce beta-lactamase enzymes which destroy this antibiotic. Clavulanic acid inactivates these enzymes, rendering the organisms susceptible to the amoxicillin. Studies in laboratory animals have not produced any evidence of teratogenic effects. Use only according to the risk/benefit assessment by the responsible veterinarian.

DRAFT INSERT TEXT

1. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER AND OF THE MANUFACTURING AUTHORISATION HOLDER RESPONSIBLE FOR BATCH RELEASE

Marketing Authorisation Holder:

(UK)

Norbrook Laboratories Limited

Station Works

Camlough Road

Newry

Co. Down

BT35 6JP

Northern Ireland

(EU)

Norbrook Laboratories (Ireland) Limited Rossmore Industrial Estate

Monaghan

Ireland

Manufacturer Responsible for Batch Release:

Norbrook Manufacturing Limited

Rossmore Industrial Estate

Monaghan

Ireland

Norbrook Laboratories Limited

Station Works

Newry

Co. Down, BT35 6JP

2. NAME OF THE VETERINARY MEDICINAL PRODUCT

Noroclav 500mg Palatable Tablets for Dogs.

3. STATEMENT OF THE ACTIVE SUBSTANCE(S) AND OTHER INGREDIENTS

Noroclav 500mg Palatable Tablets for Dogs are for oral administration to dogs. Each Noroclav 500 mg Palatable Tablet for Dogs contains Amoxicillin 400 mg (as Amoxicillin Trihydrate.), Clavulanic Acid 100 mg (as Potassium Clavulanate) and Carmosine Lake (E122) 2.45mg The tablets can be divided into equal halves.

4. INDICATION(S)

For the treatment of the following infections caused by beta-lactamase producing strains of bacteria sensitive to amoxicillin in combination with clavulanic acid:

- Skin infections (including superficial and deep pyodermas) caused by susceptible *Staphylococci*.
- Urinary tract infections caused by susceptible *Staphylococci* or *Escherichia coli*.
- Respiratory infections caused by susceptible Staphylococci.
- Enteritis caused by susceptible Escherichia coli.

It is recommended to carry out suitable tests for sensitivity when initiating the treatment. The treatment should only proceed if sensitivity is proven to the combination.

5. CONTRAINDICATIONS

Do not use in animals with known cases of hypersensitivity to penicillins or other substances of the beta-lactam group.

Do not use in rabbits, guinea pigs, hamsters or gerbils.

Do not use in animals with serious dysfunction of kidneys accompanied by anuria or oliquria.

Do not use where resistance to the combination is known to occur.

Do not administer to horses and ruminating animals.

6. ADVERSE REACTIONS

Hypersensitivity unrelated to dose can occur with these agents.

Gastrointestinal symptoms (diarrhoea, vomiting) may occur after administration of the product.

Allergic reactions (e.g. skin reactions, anaphylaxia) may occasionally occur.

In case of occurrence of allergic reaction, the treatment should be withdrawn.

If you notice any serious effects or other effects not mentioned in this leaflet, please inform your veterinary surgeon.

7. TARGET SPECIES

Dogs

8. DOSAGE FOR EACH SPECIES, ROUTE(S) AND METHOD OF ADMINISTRATION

To ensure a correct dosage, bodyweight should be determined as accurately as possible to avoid underdosing.

Administration is via the oral route. The dosage rate is 12.5 mg combined actives/kg bodyweight twice daily.

The tablets may be crushed and added to a little food.

The following table is intended as a guide to dispensing Noroclav Palatable Tablets at the standard dose rate of 12.5 mg/kg twice daily.

Bodyweight (kg)	Number of tablets (500 mg) per dose twice
	daily
20 kg	1/2
40 kg	1
60 kg	1½
80 kg	2

Duration of therapy:

Routine cases involving all indications: The majority of cases respond to between 5 and 7 days therapy.

Chronic or refractory cases: In these cases where there is considerable tissue damage, a longer course of therapy may be required in that it allows sufficient time for damaged tissue to repair.

9. ADVICE ON CORRECT ADMINISTRATION

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10. WITHDRAWAL PERIOD

Not Applicable

11. SPECIAL STORAGE PRECAUTIONS

Do not store above 25°C; store in a dry place.

Return any halved tablet to the open blister pack. Any divided tablet portion remaining after 24 hours should be discarded.

Keep out of the reach and sight of children.

12. SPECIAL WARNINGS

Inappropriate use of the product may increase the prevalence of bacteria resistant to amoxicillin/clavulanic acid.

In animals with hepatic or renal failure, the dosing regimen should be carefully evaluated.

Use of the product should be based on susceptibility testing and take into account local antimicrobial policies. Narrow spectrum antibacterial therapy should be used for first line treatment where susceptibility testing suggests likely efficacy of this approach.

Studies in laboratory animals have not produced any evidence of teratogenic effects. Use only according to the benefit/risk assessment by the responsible veterinarian.

The potential for allergic cross-reactivity with other penicillins should be considered. Penicillins may increase the effect of aminoglycosides. Caution is advised in the use in small herbivores other than those reported in contraindications.

Chloramphenicol, macrolides, sulphonamides and tetracyclines may inhibit the antibacterial effect of penicillins because of the rapid onset of bacteriostatic action.

Operator Warnings:

Penicillins and cephalosporins may cause hypersensitivity (allergy) following injection, inhalation, ingestion or skin contact. Hypersensitivity to penicillins may lead to cross reactions to cephalosporins and vice versa. Allergic reactions to these substances may occasionally be serious.

Handle this product with great care to avoid exposure, taking all recommended precautions.

If you develop symptoms following exposure such as a skin rash, you should seek medical advice and show the package leaflet or the label to the physician. Swelling of the face, lips or eyes or difficulty with breathing are more serious symptoms and require urgent medical attention. Wash hands after use

13. SPECIAL PRECAUTIONS FOR THE DISPOSAL OF UNUSED PRODUCT OR WASTE MATERIALS

Dispose of any unused product and empty containers in accordance with guidance from your local waste regulation authority.

14. DATE ON WHICH THE PACKAGE LEAFLET WAS LAST APPROVED

October 2022

15. OTHER INFORMATION

LEGAL CATEGORY:

UK - POM-V

To be supplied only on veterinary prescription.

IE - POM

Prescription Only Medicine

PACKAGE QUANTITIES:

Packs of 2, 4, 5 and 20 blister strips containing 5 tablets per strip.

Not all pack sizes may be marketed.

FURTHER INFORMATION

Amoxicillin is a broad-spectrum antibiotic active against a wide range of Gram-positive and Gram-negative bacteria. However, many clinically important bacteria produce beta-lactamase enzymes which destroy this antibiotic. Clavulanic acid inactivates these enzymes, rendering the organisms susceptible to the amoxicillin. Noroclav Palatable Tablets are safe for use during pregnancy and lactation. Studies in laboratory animals have not produced any evidence of teratogenic effects. Use only according to the risk/benefit assessment by the responsible veterinarian.

ManA 2000

UK - Vm: 02000/4259

IE – VPA No : 10999/118/001

FOR ANIMAL TREATMENT ONLY.

Approved: 28 October 2022