FREEDOM OF INFORMATION SUMMARY

ORIGINAL ABBREVIATED NEW ANIMAL DRUG APPLICATION

ANADA 200-681
Carprofen Tablets
Flavored Tablets
Dogs

For the relief of pain and inflammation associated with osteoarthritis and for the control of postoperative pain associated with soft tissue and orthopedic surgeries in dogs.

Sponsored by:
Dechra Veterinary Products LLC
Executive Summary

Carprofen Tablets flavored tablets are approved to relieve pain and inflammation in dogs with osteoarthritis and to control postoperative pain associated with both soft tissue and orthopedic surgeries in dogs. Carprofen Tablets are a generic version of Rimadyl® caplets.

<table>
<thead>
<tr>
<th>Generic Animal Drug</th>
<th>Proprietary Name</th>
<th>Established Name</th>
<th>Application Type and Number</th>
<th>Sponsor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carprofen Tablets</td>
<td>carprofen tablets</td>
<td>ANADA 200-681</td>
<td>Dechra Veterinary Products LLC</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Brand Name Animal Drug, also called the Reference Listed New Animal Drug (RLNAD)</th>
<th>Proprietary Name</th>
<th>Established Name</th>
<th>Application Type and Number</th>
<th>Sponsor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rimadyl® caplets</td>
<td>carprofen tablets</td>
<td>NADA 141-053</td>
<td>Zoetis Inc.</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviated New Animal Drug Application for a generic animal drug.

New Animal Drug Application for a brand name animal drug.

Bioequivalence

The Federal Food, Drug, and Cosmetic (FD&C) Act allows an animal drug sponsor to submit an abbreviated new animal drug application (ANADA) for a generic version of an approved brand name animal drug (also called the reference listed new animal drug or RLNAD). This law typically requires the sponsor to show that the generic drug is bioequivalent to the approved RLNAD. Broadly, bioequivalence means the generic drug is absorbed by and performs the same way in the animal’s body as the RLNAD, which has already been shown to be safe and effective when used according to the label. The FD&C Act doesn’t require the sponsor to submit new effectiveness or target animal safety data in the ANADA for a generic animal drug.

For this approval, the sponsor conducted one in vivo blood-level study to show that the 25 mg tablet of Carprofen Tablets was bioequivalent to the 25 mg tablet of Rimadyl® caplets. No serious adverse events were reported during the study. The sponsor also conducted comparative in vitro dissolution studies to show that the dissolution profiles for the 75 mg and 100 mg strength Carprofen Tablets were similar to the dissolution profile for the 25 mg strength Carprofen Tablets. Because all strengths had rapid dissolution, the 75 mg and 100 mg tablets qualified for a waiver from the requirement to perform separate in vivo bioequivalence studies (a biowaiver). Therefore, FDA granted a biowaiver for these strengths.

Conclusions

Based on the data submitted by the sponsor for the approval of Carprofen Tablets, FDA determined that the drug is safe and effective when used according to the label.
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I. GENERAL INFORMATION

A. File Number
   ANADA 200-681

B. Sponsor
   Dechra Veterinary Products LLC
   7015 College Blvd.
   Suite 525
   Overland Park, KS 66211
   Drug Labeler Code: 017033

C. Proprietary Name
   Carprofen Tablets

D. Drug Product Established Name
   carprofen tablets

E. Pharmacological Category
   Non-steroidal anti-inflammatory drug (NSAID)

F. Dosage Form
   Flavored Tablet

G. Amount of Active Ingredient
   25 mg, 75 mg, or 100 mg of carprofen per tablet

H. How Supplied
   Each tablet size is scored and packaged in bottles containing 30, 60, or 180 tablets.

I. Dispensing Status
   Prescription (Rx)

J. Dosage Regimen
   The recommended dosage for oral administration to dogs is 2 mg/lb (4.4 mg/kg) of body weight daily. The total daily dose may be administered as 2 mg/lb of body weight once daily or divided and administered as 1 mg/lb (2.2 mg/kg) twice daily. For the control of postoperative pain, administer approximately 2 hours before the procedure. Tablets are scored and dosage should be calculated in half-tablet increments.

K. Route of Administration
   Oral
L. **Species/Class**
   Dogs

M. **Indications**
   Carprofen Tablets are indicated for the relief of pain and inflammation associated with osteoarthritis and for the control of postoperative pain associated with soft tissue and orthopedic surgeries in dogs.

N. **Reference Listed New Animal Drug (RLNAD)**
   Rimadyl®; carprofen tablets; NADA 141-053; Zoetis Inc.

II. **BIOEQUIVALENCE**
   The Federal Food, Drug, and Cosmetic Act (FD&C Act), as amended by the Generic Animal Drug and Patent Term Restoration Act (GADPTRA) of 1988, allows for an abbreviated new animal drug application (ANADA) to be submitted for a generic version of an approved new animal drug (RLNAD). The ANADA sponsor is required to show that the generic product is bioequivalent to the RLNAD, which has been shown to be safe and effective. Effectiveness, target animal safety and human food safety data (other than tissue residue data) are not required for approval of an ANADA. If bioequivalence is demonstrated through a clinical endpoint study in a food-producing animal, then a tissue residue study to establish the withdrawal period for the generic product is also required.

   For this ANADA, one *in vivo* blood-level study was conducted to demonstrate product bioequivalence using the generic and RLNAD carprofen 25 mg tablets. The RLNAD is available in 25, 75, and 100 mg tablet sizes. The *in vivo* blood-level study was conducted in 24 healthy, fasted beagle dogs. The pivotal parameters to evaluate bioequivalence are the observed maximum plasma drug concentration ($C_{MAX}$) and area under the concentration-time curve (AUC) from time 0 to the last sampling time before the first unquantifiable concentration after $C_{MAX}$. Bioequivalence was demonstrated between the 25 mg Rimadyl® (carprofen tablets) caplets and the 25 mg Carprofen Tablets flavored tablets by the average bioequivalence approach as described in the Statistical Methods section below. A waiver from the requirement to demonstrate *in vivo* bioequivalence (biowaiver) for the generic 75 mg and 100 mg flavored tablets was requested. Dissolution data was used to demonstrate that the generic 75 mg and 100 mg carprofen flavored tablets are comparable to the generic 25 mg flavored tablet strength used in the *in vivo* blood-level bioequivalence study. Therefore, a biowaiver for the generic 75 mg and 100 mg carprofen flavored tablets was granted. The study information is summarized below.

A. **Blood-level Bioequivalence Study in Dogs**
   One blood-level bioequivalence study was conducted to determine the comparative bioavailability of the generic and RLNAD formulations of carprofen tablets (25 mg).

   **Title**: Pivotal Two-Way Oral Bioequivalence Study of Carprofen (Flavored Tablets) in Beagles (Study No. 017-01604).

   **Study Dates**: March 7, 2018 to November 14, 2018
Study Locations:

In-life phase: Las Cruces, NM

Bioanalytical testing: Colorado Springs, CO

Study Design:

Objective: The objective of this study was to determine the comparative in vivo blood-level bioequivalence data for the generic 25 mg Carprofen Tablets and the RLNAD 25 mg Rimadyl® (carprofen tablets) caplets in fasted dogs. The study was conducted according to Good Laboratory Practices (GLP) regulations.

Study Animals: 24 intact (non-pregnant) female beagle dogs between 2 and 8 years of age and weighing from 9 to 14 kg on the day of initial dose administration (study day 0).

Experimental Design: A randomized, masked, two-period, two-sequence, single-dose crossover study.

Drug Administration: Each animal received 25 mg of either the generic or RLNAD carprofen tablets according to their randomized treatment sequence (generic/RLNAD or RLNAD/generic).

Measurements and Observations: The plasma concentrations of carprofen were measured using a validated bioanalytical method. Pharmacokinetic parameters were determined for each animal individually in each period. Animal observations were made throughout the study for assessment of general health and adverse events.

Statistical Methods: The laboratory study was conducted as a randomized, masked, two-period, two-sequence, single-dose crossover design using 24 dogs with a 14-day washout between periods. Appropriate randomization of animals to sequence and pen/treatment order was performed. Primary variables evaluated were $C_{MAX}$ and AUC. Time to maximum concentration ($T_{MAX}$) was summarized and evaluated clinically.

A mixed-effect model was used to evaluate bioequivalence. The model included fixed effects of treatment, sequence and period, and random effects of subject nested within sequence. Prior to analysis, $C_{MAX}$ and AUC were natural logarithm transformed. Bioequivalence is established because the back-transformed estimated upper and lower bounds of the 90% confidence interval for geometric mean ratios (generic:RLNAD) of both $C_{MAX}$ and AUC are contained within the acceptance limits of 0.80 to 1.25.

Results: As seen in the table below, both $C_{MAX}$ and AUC fall within the prescribed bounds (Table II.1). The mean values of $T_{MAX}$ obtained for the generic article and RLNAD were summarized.
Table II.1. Bioequivalence Evaluation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Generic Mean</th>
<th>RLNAD Mean</th>
<th>Ratio°</th>
<th>Lower 90% CI</th>
<th>Upper 90% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC (ng/mL)*hour</td>
<td>371155†</td>
<td>360234†</td>
<td>1.03</td>
<td>0.98</td>
<td>1.09</td>
</tr>
<tr>
<td>(C_{\text{MAX}}) (ng/mL)</td>
<td>45492†</td>
<td>45510†</td>
<td>1.00</td>
<td>0.92</td>
<td>1.09</td>
</tr>
<tr>
<td>(T_{\text{MAX}}) (hour) (SD‡)</td>
<td>1.32 (0.83)†</td>
<td>1.57 (1.21)†</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
</tr>
</tbody>
</table>

† Geometric mean  
‡ Arithmetic mean and standard deviation (SD)  
° Ratio = Generic:RLNAD  
CI = confidence interval  
NE = not estimated

**Adverse Reactions:** There were no serious adverse events reported during the study.

**Conclusion:** The *in vivo* bioequivalence study demonstrated that the generic 25 mg Carprofen Tablets and the RLNAD 25 mg Rimadyl® (carprofen tablets) caplets are bioequivalent in dogs.

**B. Bioequivalence Waiver**

A pivotal *in vivo* blood bioequivalence study was conducted using the 25 mg carprofen flavored tablets strength. A waiver from the requirement to perform *in vivo* bioequivalence studies (biowaiver) for the generic 75 mg and 100 mg flavored tablets was requested. To qualify for a biowaiver for each of these product strengths, comparative *in vitro* dissolution studies were conducted to determine the dissolution profiles of generic 75 mg and 100 mg carprofen flavored tablets. The similarity factor (\(f_2\)) calculation was used to evaluate dissolution profile comparisons. Comparisons were made between the following tablets:

- Generic 25 mg and generic 75 mg tablets  
- Generic 25 mg and generic 100 mg tablets

The objective was to satisfy \(f_2\) criteria between the generic 25 mg flavored tablet strength and the generic 75 mg and 100 mg flavored tablet strengths.

Test conditions were as follows:

- Dissolution apparatus: USP Apparatus 2  
- Dissolution medium: 0.05 M Phosphate buffer, pH 7.5  
- Dissolution medium volume: 900 mL  
- Temperature: 37 °C ± 0.5°C  
- Paddle speed: 50 rpm  
- Number of vessels: 12  
- Data points: 10, 15, and 30 minutes

The generic drug lot number used in the *in vivo* bioequivalence study was the same lot used to support the *in vitro* profile comparisons. Analytical method
validation was required to ensure that the quantification of drug concentrations in all samples was accurate and precise.

To allow use of mean data, the percent coefficient of variation at the earlier time points (e.g., 15 minutes) should not be more than 20%, and at other time points should not be more than 10%. The percent coefficient of variation for all generic product profiles was within acceptable limits. Only one measurement should be considered after 85% dissolution of both products. The $f_2$ should be greater than 50 to ensure sameness or equivalence of two profiles.

Study results demonstrate similar dissolution profiles for all comparisons. However, because of rapid dissolving characteristics (>85% in 15 minutes) in all strengths, a dissolution profile comparison using the $f_2$ test is unnecessary. When comparative profiles between tablets do not require an $f_2$ test because of rapid dissolution or when the $f_2$ value is ≥50, the product strengths used in the comparison qualify for a biowaiver. Therefore, a biowaiver for the generic 75 mg and 100 mg carprofen flavored tablets is granted.

III. HUMAN FOOD SAFETY

This drug is intended for use in dogs. Because this new animal drug is not intended for use in food-producing animals, CVM did not require data pertaining to drug residues in food (i.e., human food safety) for approval of this ANADA.

IV. USER SAFETY

The product labeling contains the following information regarding safety to humans handling, administering, or exposed to Carprofen Tablets:

Keep out of reach of children. Not for human use. Consult a physician in cases of accidental ingestion by humans.

V. AGENCY CONCLUSIONS

This information submitted in support of this ANADA satisfies the requirements of section 512(c)(2) of the FD&C Act. The data demonstrate that Carprofen Tablets, when used according to the label, is safe and effective.