

Carprovet Chewable 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.

Therapeutically equivalent to the pioneer drug so you can expect the same safety and efficacy at a substantially lower price.

Backed by Dechra's Veterinary Technical and Sales Support Teams.

Available in 25 mg, 75 mg, and 100 mg beef-flavored scored tablets in 30, 60 and 180 count bottles.

STRENGTH	BOTTLE COUNT	ACTUAL TABLET SIZE
25 mg	30, 60, 180	2.5 Mg
75 mg	30, 60, 180	7.5 MG
100 mg	30, 60, 180	10.0 MG

To order, please contact your Dechra or distributor representative or call (866) 683-0660. For Full Prescribing Information please visit www.dechra-us.com.

24-hour Veterinary Technical Support available (866) 933-2472. Nonurgent Technical Support available via email support@dechra.com.



Important Safety Information: As with other NSAIDs, signs of carprofen intolerance may include appetite loss, vomiting and diarrhea, which could indicate side effects involving the digestive tract, liver or kidneys. **Some of these side effects, in rare situations, may be serious, resulting in hospitalization or even death. Pet owners should be advised to discontinue treatment if side effects occur and contact their veterinarian.** Concomitant use of carprofen with other anti-inflammatory drugs, such as other NSAIDs or corticosteroids, should be avoided because of the potential increase of adverse reactions. Refer to the prescribing information and "Dog Owner Information Sheet" for complete details or visit www.dechra-us.com

Carprovet®

(carprofen) Chewable Tablets Non-steroidal anti-inflammatory drug For oral use in dogs only

CAUTION: Federal law restricts this drug to use by or on the order of a licensed veterinarian.

DESCRIPTION: Carprovet (carprofien) is a non-steroidal anti-inflammatory drug (NSAID) of the propionic acid class that includes ibuprofen, naproxen, and ketporofen. Carprofen is the nonpropientary designation for a substituted carbazole, 6-chloro-o-methyl-9H-carbazole-2-acetic acid. The empirical formula is 1,541-f2(INO) and the molecular weight 273.72.

The chemical structure of carprofen is:

Carprofen is a white, crystalline compound. It is freely soluble in thand, but practically insoluble in water at 25°C.

CLINICAL PHARMACOLOFY: Carprofen is a non-narcotic, non-steroidal anti-inflammatory agent with characteristic analgesic and antipyretic activity approximately equipotent to indomethacin in animal models. The mechanism of action of carprofen, like that of other NSAIDs, is believed to be associated with the inhibition of peoloxyegenase action. White inhibition of coloxyegenases, CDX-1, synthesizes prostaglandins necessary for normal gastrointestinal and renal function. The inducible ocycloxyegenases, CDX-2, generates prostaglandins necessary for normal gastrointestinal and renal function. The inducible ocycloxyegenases, CDX-2 generates prostaglandins necessary for normal gastrointestinal and renal function. The inducible ocycloxyegenases, CDX-2 generates prostaglandins necessary for normal gastrointestinal and renal function. The prostaglandins includes anti-inflammatory activity. The specificity of a particular NSAID for CDX-2 versus CDX-1 may ray from species to species. In an in vitro study using canine cell cultures, carprofine demonstrated selective inhibition of CDX-2 versus CDX-1 discinct relevance of these data has not been shown. Carprofen has also been shown to inhibit the release of several prostaglandins in two inflammatory cell systems: rat polymorphonuclear leukocytes PMNI) and human memanistid synovial cell systems in fallmamatory reactions: Several studies have demonstrated that carprofen has modulatory effects on both humoral and cellular immune responses. So that also indicate that carprofen inhibits the production of selectosak-activating factor (DAP). PECF, and PSV bis hibitory effect in prostaglandin hibosynthesis. Based upon comparison with data obtained from intravenus administration, carprofen is rapidly and nearly completely absorbed (more than 90%) bis hibitory of the prostaglandin problem of the select glucuronides of 2 phenolic metabolists. 7-hydroxy carprofen and the

commonly used include cardiac, anticonvulsant and behavioral medications. It has been suggested that treatment with carproten may reduce the level of inhalant nasehticis needed; 13 ff additional pain medication is warranted after administration of the total daily dose of Carprovet, alternative analgesia should be considered. The use of another NSAID is not recommended. Consider appropriate washout times when switching from one NSAID to another or when switching from corticosteroids use to NSAID use. Store out of reach of dogs in a secured location. Severe adverse reactions may occur if large quantities of tablets are ingested if you suspect your dog has consumed Carprovet chewable tablets above the labeled dose, please call your veterinarian for immediate assistance and notify Dechra at (866) 932-2472.

INFORMATION FOR DOG OWREENSE Carprovet, like other drugs of its class, is not free from adverse reactions. Downers should be advised of the potential for adverse reactions and be informed of the clinical signs associated with drug intolerance. Adverse reactions may include decreased appetite, vomiting, diarrhea, dark or tarry stobs, increased variet or consumption, increased variet rous increased variety or consumption, increased variety or cons

Percentage of Dogs with Abnormal Health Observations Reported in Clinical Field Study (2 mg/lb once daily)			
Observation	Carprofen (n=129)	Placebo (n=132)	
Inappetence	1.6	1.5	
Vomiting	3.1	3.8	
Diarrhea/Soft stool	3.1	4.5	
Behavior change	0.8	0.8	
Dermatitis	0.8	0.8	
PU/PD	0.8	-	
SAP increase	7.8	8.3	
ALT increase	5.4	4.5	
AST increase	2.3	0.8	
BUN increase	3.1	1.5	
Bilirubinuria	16.3	12.1	
Ketonuria	14.7	9.1	

Clinical pathology parameters listed represent reports of increases from pre-treatment values; medical judgment is necessary to determine clinical relevance. During investigational studies of surgical pain for the caplet formulation, no clinically significant adverse reactions were reported. The product vehicle served as control.

Percentage of Dogs with Abnormal Health Observations Reported in Surgical Pain Field Studies with Caplets (2 mg/lb once daily)			
Observation*	Carprofen (n=148)	Placebo (n=149)	
Vomiting	10.1	13.4	
Diarrhea/Soft stool	6.1	6.0	
Ocular disease	2.7	0	
Inappetence	1.4	0	
Dermatitis/Skin lesion	2.0	1.3	
Dysrhythmia	0.7	0	
Apnea	1.4	0	
Oral/Periodontal disease	1.4	0	
Pyrexia	0.7	1.3	
Urinary tract disease	1.4	1.3	
Wound drainage	1.4	0	

"A single dog may have experienced more than one occurrence of an event. During investigational studies for the chewable tablet formulation, gastrointestinal signs were observed in some dogs. These signs included vomiting and soft stools.

Post-Approval Experience:

Post-Approval Experience:

Although not all adverse reactions are reported, the following adverse reactions are based on voluntary post-approval adverse drug experience reporting. The categories of adverse reactions are listed in decreasing order of frequency by body system.

Castrointestinal: Voniting, diarrhea, constipation, inappetence, melena, hematemesis, gastrointestinal ulceration, gastrointestinal bleeding, pancreatitis. Hepatic: happetence, vomiting, jaundice, acute hepatic toxicity, hepatic enzyme elevation, abnormal liver function test(s), hyperbilirubinemia, biblirubiniria, hypocalbuminemia. Approximately one-fourth of hepatic reports were in Labrador Retrievers.

Neurologic: Akaia, paresis, paralysis, sezures, vestibulari signs, disorientation.

Urinary: Hematuria, polyuria, polydipsia, urinary incontinence, urinary tract infection, azotemia, acute renal failure, tubular abnormalities including acute tubular necrosis, renal bubular acidosiss, gluozosmia.

Behavioral: Sedation, lethargy, hyperacibrity, restlessness, aggressiveness.

Hematologic: Immune-mediated hemohylic amenia, immune-mediated thrombocytopenia, blood loss anemia, epistaxis.

Dermatologic: Pruritus, increased shedding, alopecia, pyotraumatic moist dermatitis (hot spots), necrotizing pannicultis/vasculitis, ventral ecchymosis.

Immunologic or hypersensitivity: Facial swelling, hives, erythema.

In rare situations, death has been associated with some of the adverse reactions listed above.

To report suspected adverse events, for technical assistance or to obtain a copy of the safety data sheet (SDS), contact Dechra at (866) 933-2472.

For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS, or www.fda.gov/reportanimalae

DOSAGE AND ADMINISTRATION: Always provide Client Information Sheet with prescription. Carefully consider the potential benefits and risk of Carprovet and other treatment options before decling to use Carprovet sevential experience the potential benefits and ri

ANIMAL SAFETY: Laboratory Studies in unanestiretized days and unincent into studies here desired the color of administration.

In target animal safety studies, carprofen was administered orally to healthy Beagle dogs at 1, 3, and 5 mg/lb twice daily (1, 3 and 5 times the recommended total daily dose) for 42 consecutive days with no significant adverse reactions. Serum albumin for a single female dog receiving 5 mg/lb twice daily decreased to 2.1 gl/d, after 2 weeks of treatment, returned to the pre-treatment value (2, 6 gl/d) after 4 weeks of treatment, and was 2.3 gl/d, at the final 6-week evaluation. Over the 6-week treatment period, black or bloody stools were observed in 1 dog (1 incident) treated with 1 mg/lb twice daily and in 1 dog (2 incidents) treated with 3 mg/lb twice daily. Redness of the coloric mucosa was observed in 1 male that received 3 mg/lb

yold. at the final 6-week evaluation. Over the 6-week treatment period, black or bloody stools were observed in 1 dog (1 incident) treated with 1 mg/lb twice daily. The old of a dogs receiving 10 mg/lb or ally twice daily. The old of a dogs receiving 10 mg/lb or ally twice daily. The old of a dogs receiving 10 mg/lb or ally twice daily. The old of a dogs receiving 10 mg/lb or ally twice daily. The old of a dogs receiving 10 mg/lb or ally twice daily. The old of a dogs receiving the dogs receiving this dose was lower (2.38 g/dl) than each of 2 placebo control groups (2.88 and 2.93 g/dl, respectively). Three incidents of black or bloody stool were observed in 1 dog, 1 mg/lb or dogs receiving the laminary propria in 2 of the 5 dogs. In separate safety studies lasting 13 and 52 weeks, respectively, dogs were administered orally up to 11 mg/lb/day (5.7 times the recommended total daily dose of 2 mg/lb) of cargoride. In both studies, the drug was well toterated clinically by all of the animals. No gross or histologic changes were seen in any of the treated animals. In both studies, dogs receiving the highest doses had average increases in serum L-alanine aminotransferse (ALT) of approximately 20 IU.

In the 52-week study, minor dematologic changes occurred in dogs in each of the treatment groups but not in the control dogs. The changes were described as slight redness or reash and were diagnosed as non-specific dermachitis. The possibility exists that these mild lesions were treatment related, but no dose relationship was observed.

Chical field studies were conducted with 549 dogs of different breeds at the recommended oral doses for 14 days (297 dogs were included in a subdy evaluating 1 mg/lb wice daily and 252 dogs were included in a separate study evaluating 2 mg/lb once daily), in both studies the quive accordance of the controlled controlled to the incidence of clinical adverse reactions for carporden-reated animals was no higher than placebo-treated animals placebo contained inactive ingredients found in carp

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