Can Humans Get Cancer from Their Pet?

William C. Eward, MD, DVM
Duke Medical Center
Durham, North Carolina

Jolle Kirpensteijn, DVM, PhD, DACVS, DECVS
Hill’s Pet Nutrition
Topeka, Kansas

Cancer is a complex biological process through which a normal cell acquires capabilities that cause its transformation into a tumorigenic and eventually malignant cell. If the cell is not recognized and/or eliminated by the immune system, a tumor will form.

Cancer in pets is naturally occurring and is as common in dogs and cats as it is in humans. It is a leading cause of death in dogs and cats. The similarities between pets and humans— with respect to anatomy, physiology, and tumor onset and progression — make canine tumor models a valuable tool for identifying new cancer-associated genes and for enhancing understanding of tumor molecular biology in humans. Because dogs and cats often share the same environments as humans, one might wonder whether there is risk for tumor development caused by pets, especially in immunocompromised humans. Although no human cancer is known to spread naturally from human to human (or animal), the existence of such diseases in dogs and Tasmanian devils raises the question of whether humans could be at risk.

Few tumors can pass from one living host to another. These tumors, called naturally transmissible tumors, emerge when a tumor cell gains the ability to pass infectious material between individuals. There are 2 known naturally occurring contagious cancers:

- Canine transmissible venereal tumors are nonfatal, affect the orogenital area of dogs, and are transmitted through direct contact between individual dogs via sexual contact or bites/licks.

- Devil facial tumor disease spreads among Tasmanian devils through fighting and biting and has close to 100% mortality.

Neither disease is infectious to humans, and there is no scientific evidence that humans, even if immunocompromised, are susceptible to these or any other cancers by direct transmission.

Pet ownership may actually decrease the incidence of cancer in humans.
Metacam® (meloxicam)

1.5 mg/mL Oral Suspension (equivalent to 0.05 mg per drop)
0.5 mg/mL Oral Suspension (equivalent to 0.02 mg per drop)

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.

Warning: Repeated use of meloxicam in cats has been associated with acute renal failure and death. Do not administer additional doses of oral or oral-metacam to cats. See Contraindications, Warnings, and Precautions for detailed information.

Description: Meloxicam is a non-steroidal anti-inflammatory drug (NSAID) of the oxicam class. Each milliliter of Metacam Oral Suspension contains meloxicam equivalent to 0.5 mg/mL or 1.5 mg/mL and sodium benzoate (1.5 milligrams) as a preservative. The chemical name for Meloxicam is 4-hydroxy-2-(methyl-N-(5-methyl-2-thiazolyl)-2H-1,2-benzothiazine-3-carboxamide-1, 1-dioxide. The formulation is a yellowish viscous suspension with the odor of honey.

Indications: Metacam Oral Suspension is indicated for the control of pain and inflammation associated with osteoarthritis in dogs.

Contraindications: Dogs with known hypersensitivity to meloxicam should not receive Metacam Oral Suspension. Do not use Metacam Oral Suspension in cats. Acute renal failure and death have been associated with the use of meloxicam in cats.

Warnings: Not for use in humans. Keep this and all medications out of reach of children. Consult a veterinarian in case of accidental ingestion by humans. For oral use in dogs only. As with any NSAID, all dogs should undergo a thorough history and physical examination before the initiation of NSAID therapy. Appropriate laboratory testing to establish hematological and serum biochemical baseline data is recommended prior to and periodically during administration. Owner should be advised to observe their dog for signs of potential drug toxicity and should be given a client information sheet about Metacam.

Precautions: The safe use of Metacam Oral Suspension in dogs younger than 6 months of age, dogs used for breeding, or in pregnant or lactating dogs has not been evaluated. Meloxicam is not recommended for use in dogs with renal, hepatic, or cardiac disease or with bleeding disorders, as safety has not been established in these dogs with these disorders. As a class, cyclo-oxygenase inhibitory NSAIDs may be associated with gastrointestinal, renal, and hepatic toxicity. Sensitivity to drug-associated adverse events varies with the individual patient. Dogs that have experienced adverse reactions from one NSAID may experience adverse reactions from another NSAID. Patients at greatest risk for renal toxicity are those that are dehydrated, on concomitant diuretic therapy, or those with existing renal, cardiovascular, and/or hepatic dysfunction. Concurrent administration of potentially nephrotoxic drugs should be carefully approached. NSAIDs may inhibit the prostaglandins that maintain normal hemostatic function. Such anti-prostaglandin effects may result in clinically significant disease in patients with underlying or pre-existing disease that has not been previously diagnosed. Since NSAIDs possess the potential to induce gastrointestinal ulcerations and/or perforations, concomitant use with other anti-inflammatory drugs, such as NSAIDs or corticosteroids, should be avoided. If additional pain medication is needed after administration of the total daily dose of Metacam Oral Suspension, a non-NSAID or non-corticosteroid class of analgesia should be considered. The use of another NSAID is not recommended. Consider appropriate washout times when switching from corticosteroid use or from one NSAID to another in dogs. The use of concomitantly protein-bound drugs with Metacam Oral Suspension has not been studied in dogs. Commonly used protein-bound drugs include digoxin, anticonvulsant and behavioral medications. The influence of concomitant drugs that may inhibit metabolism of Metacam Oral Suspension has not been evaluated. Drug compatibility should be monitored in patients requiring adjunctive therapy.

Adverse Reactions: Field safety was evaluated in 306 dogs.1 Based on the results of two studies, GI abnormalities (vomiting, soft stools, diarrhea, and inappetance) were the most common adverse reactions associated with the administration of meloxicam. The following adverse events are based on post-approval adverse drug experience reporting. Not all adverse reactions are reported to FDA. It is not always possible to reliably estimate the actual frequency or establish a causal relationship to product exposure using these data. The following adverse events are listed in decreasing order of frequency by body system.

Gastrointestinal: vomiting, anorexia, diarrhea, melena, gastrointestinal ulceration
Urinary: azotemia, elevated creatinine, renal failure
Neurological/Behavioral: lethargy, depression
Hepatic: elevated liver enzymes
Dermatologic: pruritus

Death has been reported as an outcome of the adverse events listed above. Acute renal failure and death have been associated with use of meloxicam in cats.


Warning and in rare situations result in death (see Adverse Reactions). Owners should be advised of the importance of periodic follow up for all dogs during administration of any NSAID.

Effectiveness: The effectiveness of meloxicam was demonstrated in two field studies involving a total of 277 dogs representing various breeds, between six months and sixteen years of age, all diagnosed with osteoarthritis. Both of the placebo-controlled, masked studies were conducted for 14 days. All dogs received 0.2 mg/kg on day 1. All dogs were maintained on 0.1 mg/kg oral meloxicam from days 2 through 14 of both studies. Fluorochromes were evaluated by veterinarians included lameness, weight-bearing, pain on palpation, and overall improvement. Parameters assessed by owners included mobility, ability to rise, limping, and overall improvement. In the first field study (n=109), dogs showed clinical improvement with statistical significance after 14 days of meloxicam therapy. In the second field study (n=48), dogs receiving meloxicam showed a clinical improvement after 14 days of therapy for all parameters; however, statistical significance was demonstrated only for the overall investigator evaluation on day 7, and for the owner evaluation on day 14.1

Reference: 1. FOI for NADA 141-213 (Metacam® (meloxicam) 0.5 milligram/mL and 1.5 milligram/mL Oral Suspension).

Manufactured for: Boehringer Ingelheim Vetmedica, Inc.
St. Joseph, MO 64066 U.S.A.
US Patent No. 5,646,220
Metacam is a registered trademark of Boehringer Ingelheim Vetmedica GmbH, licensee to Boehringer Ingelheim Vetmedica, Inc.
6015161, 601516, 601519-1007
Code 061511, 061521, 061531, 061571

In conclusion, no virus transmission—including retroviruses such as human papillomavirus, which can be spread between humans—between pets and humans has been shown to cause cancer in humans.

In fact, reports have shown that pet ownership may actually decrease the incidence of cancer in humans. A population-based case-control study in the San Francisco Bay area showed that pet owners had a reduced risk for non-Hodgkin’s lymphoma as compared with those who never owned a pet.3 However, this research remains to be confirmed.

References


Clinical View

References


