Clinical pharmacology of tramadol and tapentadol, and their therapeutic efficacy in different models of acute and chronic pain in dogs and cats

Adriana Domínguez-Oliva, Alejandro Casas-Alvarado, Agatha Elisa Miranda-Cortés, Ismael Hernández-Avalos

ABSTRACT

Opioids are considered the gold standard to manage acute or chronic or mild to severe pain. Tramadol is a widely prescribed analgesic drug for dogs and cats; it has a synthetic partial agonism on μ-opioid receptors and inhibits the reuptake of norepinephrine and serotonin. However, the biorotation and resultant metabolites differ between species and depend on cytochrome P450 interactions. Dogs mainly produce the inactive N-desmethyl tramadol metabolite, whereas cats exhibit an improved antinociceptive effect owing to rapid active O-desmethyltramadol metabolite production and a longer elimination half-life. Tapentadol, a novel opioid with dual action on μ-receptors and noradrenaline reuptake inhibitory activity, is a promising option in dogs, as it is less reliant on metabolic activation and is unaffected by cytochrome polymorphisms. Although scientific evidence on the analgesic activity of tapentadol in both species remains limited, experimental studies indicate potential benefits in animals. This review summarizes and compares the pharmacology, pharmacokinetics, and therapeutic efficacy of tramadol and tapentadol in dogs and cats with different pain conditions. According to the available data, tramadol seems a more suitable therapeutic option for cats and should preferably be used as a component of multimodal analgesia in both species, particularly dogs. Tapentadol might possess a superior analgesic profile in small animals, but additional studies are required to comprehensively evaluate the activity of this opioid to manage pain in dogs and cats.

Introduction

Analgesia is an essential aspect of veterinary patient care that prevents pathophysiological complications owing to pain, reduces the stress response during surgical procedures [1], and improves overall animal welfare [2]. To select an appropriate analgesic therapy, the etiology, type, severity, and chronicity of the process must be considered to prevent central and peripheral sensitization in response to a noxious stimulus [3–5]. Currently, multimodal analgesia is considered the most appropriate strategy for pain management [6,7], and commonly used therapies include non-steroidal anti-inflammatory drugs (NSAIDs) [8], non-opioid NSAIDs such as paracetamol [9], local anesthetics [10], alpha-2 adrenergic agonists [11], N-methyl-D-aspartate (NMDA) antagonists [12], or opioids like tramadol and tapentadol [13].

In general, opioids are considered the gold standard to manage mild to intense, short, or long-term pain and represent a perioperative protocol to improve recovery times and decrease post-surgical pain scores [14]. They act on central and peripheral opioid receptors (primarily, G protein-coupled receptors), cause hyperpolarization of the neurons, and lessen the secretion of excitatory neurotransmitters to interrupt the transduction, transmission, modulation, and perception of a noxious stimulus [13].

Tramadol and tapentadol are central non-traditional opioids with dual effects [14]. Tramadol is a synthetic...
opioid agonist at μ-opioid receptors (MOR) that suppress serotonin (5-HT) and norepinephrine (NE) reuptake [15]. It acts primarily on the descending inhibitory pathway but requires frequent dosage intervals and concomitant administration with other drugs to maintain adequate therapeutic efficacy [14]. Conversely, tapentadol is a recently developed drug with the earliest evidence documented in 2010 [16]. This opioid is considered the first MOR – norepinephrine reuptake inhibitor (NRI) agent – given its combined agonist action on MORs and the potent inhibition of NE reuptake [17]. The analgesic efficacy and therapeutic safety of these drugs depend on the delivery method, gene diversity between species, and their association with other analgesics [18]. In addition, their use in small animals remains controversial due to discrepancies between clinical and experimental outcomes [13].

In the present review, we aim to summarize the pharmacokinetics, pharmacodynamics, main adverse effects, contraindications, and therapeutic efficacy of tramadol and tapentadol in dogs and cats during different acute and chronic pain models. Through Web of Science, PubMed, Google Scholar, Scopus, and Elsevier, relevant keywords regarding the administration of tramadol and tapentadol to manage pain in dogs and cats were used to identify and select publications associated with these opioids. The chosen publications covered the following inclusion criteria: 1) articles after 2000 where tramadol or tapentadol pharmacokinetics was first reported; 2) reports from 2010 to 2021 where tramadol and tapentadol were used in a clinical setting as analgesic therapy for different surgical procedures or painful conditions; and 3) studies where their antinociceptive properties were tested in dogs or cats, or current animal models.

Clinical Pharmacology of Tramadol

Tramadol is an atypical opioid with a weak action on μ-receptors and modulates the descending noradrenergic and serotonergic pathways by inhibiting the reuptake of the monoamines (5-HT and NE), as well as 5-HT release [19]. In addition, this opioid is known to possess mechanisms such as an agonist activity at TRPV1 receptors, inhibition of G protein-coupled receptors (alpha-2 adrenoceptors, neurokinin receptor 1, and muscarinic receptors), inhibitory effects on nicotinic acetylcholine receptors, and NMDA receptor antagonism [13,20]. These characteristics mediate the opioid and non-opioid mechanisms of action [21]. Given its rapid onset of action (10–20 min), it is frequently employed in small animals to manage moderate to severe perioperative pain [22] and osteoarthritic diseases [20].

Surveys conducted in Colombia [23] and Brazil [24] have revealed that tramadol is the most commonly prescribed opioid in small animals (more than 80%) [23], corresponding to 58% and 50% of preoperative analgesic management in dogs and cats, respectively, and 62% and 53% in terms of post-surgical therapy. On the contrary, in Canada, the United States, and Europe, butorphanol and buprenorphine were reported as the most commonly used drugs. These distinct utilization patterns can be attributed to the availability of tramadol as an over-the-counter drug in Brazil and Colombia [24].

According to the available literature, tramadol offers a better therapeutic range to control pain when co-administered with other drugs rather than monotherapy, and its pharmacokinetic properties considerably differ between dogs and cats and even within species. These differences are compared and presented in Table 1 [25–32] when administered by different routes and distinct doses. Table 2 [25–32] shows a comparison between half-life, peak concentration, and time peak of the main active O-desmethytramadol (M1) metabolite in dogs and cats. Although data on cats and dogs has been previously reported, the pharmacokinetics of tramadol in other species such as koala has been recently determined and compared with those in dogs and cats following a subcutaneous (SC) dose of tramadol (2 and 4 mg/kg); the results revealed a half-life of 2 ± 0.3 h, which was similar to that in cats (3–4 h) [33].

Tramadol undergoes biotransformation by demethylation, oxidation, and hepatic conjugation by the uridine diphosphate (UDP) UDP-glucuronyltransferase (UGT) enzymes in phase II metabolism, forming at least 30 different compounds [18]. However, only three of these metabolites have analgesic activity: M1, N,O-didesmethytramadol (M5) (a result of the metabolism of M1), and tramadol [34]. M1 is the principal active metabolite and the most relevant component that mediates the analgesic effects due to its high affinity to MOR (up to 200 to 300-fold greater than the parent compound) [16,19]. N-desmethytramadol (M2) is another metabolite considered inactive and formed by the liver microsomes, mainly in dogs [35].

Tramadol is a mixture of two trans-enantiomers: (+) (1R, 2R)-tramadol and (−) (1S, 2S)-tramadol, which confers antinociceptive effects [36]. (+)-Tramadol has a greater affinity to MOR and is responsible for inhibiting 5-HT reuptake and its extracellular increase. Contrarily, (−)-tramadol is a potent inhibitor of NE reuptake [37]. (+)-M1 is an active metabolite with agonism to μ-receptor, while (−)-M1 inhibits NE reuptake [35]. Additionally, a greater association to MOR is observed on M5, compared to (−)-M1 and tramadol [18]. Accordingly, the binding of the (+)-tramadol stereoisomer and (+)-M1 to μ-receptors are pivotal for producing the analgesic effect [37]. Some reports have suggested that MOR agonism is predominantly effective during acute pain management. On
Table 1. Comparison of mean ± SD values for some tramadol pharmacokinetic variables in dogs and cats.

| Species | Route and dose | Parameters (units) | Mean ± SD values | References |
|---------|----------------|--------------------|------------------|------------|
| Dogs    | IV (4 mg/kg)   | F%                 | ND               | [25]       |
|         |                | Vd (l/kg)          | 3.01 ± 0.45      |            |
|         |                | Cl (ml/min/kg)     | 54.63 ± 8.19     |            |
|         |                | T_{max} (h)        | ND               |            |
|         |                | C_{max} (ng/ml)    | ND               |            |
|         | IV (4 mg/kg)   | F%                 | ND               | [26]       |
|         |                | Vd (ml/kg)         | 1,003 ± 472      |            |
|         |                | Cl (ml/h/kg)       | 923 ± 460        |            |
|         |                | T_{max} (h)        | ND               |            |
|         |                | C_{max} (µg/ml)    | ND               |            |
|         | IV (4 mg/kg)   |                     | 4.77 ± 1.07      | [27]       |
|         | 2 years        | Cl (ml/min/kg)     | 29.9 ± 7.3       |            |
|         | IV (4 mg/kg)   |                     | 4.73 ± 1.43      | [27]       |
|         | 8–10 years     | Cl (ml/min/kg)     | 23.7 ± 5.4       |            |
|         | IV (2 mg/kg)   | F%                 | ND               | [28]       |
|         |                | Vd (ml/kg)         | 1,995.89 ± 1,165.24 |          |
|         |                | Cl (ml/h/kg)       | 1,748.99 ± 1,239.6 |          |
|         |                | T_{max} (h)        | ND               |            |
|         |                | C_{max} (µg/ml)    | ND               |            |
|         | EPI (2 mg/kg)  | F%                 | 82               | [28]       |
|         |                | Vd (ml/kg)         | ND               |            |
|         |                | Cl (ml/h/kg)       | ND               |            |
|         |                | T_{max} (h)        | 1.15 ± 0.31      |            |
|         |                | C_{max} (µg/ml)    | 0.18 ± 0.12      |            |
|         | IM (4 mg/kg)   | F%                 | 92 ± 9           | [29]       |
|         |                | Vd (ml/kg)         | 293 ± 151        |            |
|         |                | Cl (ml/h/kg)       | 1,131 ± 146      |            |
|         |                | T_{max} (h)        | 0.34 ± 0.05      |            |
|         |                | C_{max} (µg/ml)    | 2.52 ± 0.43      |            |
|         | Oral (11.2 ± 2.0 mg/kg) | F% | 65 ± 38 | [25] | Vd (l/kg) | ND | |
|         |                | Cl (ml/min/kg)     | ND               |            |
|         |                | T_{max} (h)        | 1.04 ± 0.51      |            |
|         |                | C_{max} (ng/ml)    | 1,402.75 ± 695.52 |          |
|         | Oral (5 to 7 mg/kg) | F% | ND | [30] | Vd (l/kg) | 5.4 | Cl (ml/min/kg) | 30 | T_{max} (h) | 3.54 | C_{max} (ng/ml) | 195 |
| Cats    | IV (2 mg/kg)   | F%                 | ND               | [31]       |
|         |                | Vd (l/kg)          | 3.0 ± 0.1        |            |
|         |                | Cl (ml/min/kg)     | 20.8 ± 3.2       |            |
|         |                | T_{max} (min)      | ND               |            |
|         |                | C_{max} (ng/ml)    | 1,323 ± 92       |            |
|         | IV (2 mg/kg)   | F%                 | ND               | [32]       |
|         |                | Vd (ml/kg)         | 1,953.65 ± 418.68 |          |
|         |                | Cl (ml/h/kg)       | 895 ± 366.62     |            |
|         |                | T_{max} (h)        | ND               |            |
|         |                | C_{max} (µg/ml)    | ND               |            |
|         | Oral (5.2 mg/kg) | F% | 93 ± 7 | [31] | Vd (l/kg) | 5.1 ± 0.3 | Cl (ml/min/kg) | 18.6 ± 3.2 | T_{max} (min) | 25 | C_{max} (ng/ml) | 914 ± 232 |

\( \text{F\%} = \) bioavailability, \( \text{ND} = \) not determined, \( \text{Vd} = \) apparent volume of distribution at steady-state, \( \text{Cl} = \) clearance, \( \text{T_{max}} = \) time of maximum plasma concentration, \( \text{C_{max}} = \) maximum plasma concentration.
the contrary, the inhibition of 5-HT and NE recapturing is appropriate during chronic pain states [18]. Moreover, compared to the inhibition of 5-HT reuptake, NE has a greater role during pain modulation [16].

In dogs and cats, the considerable variations in metabolite formation can be attributed to the cytochrome P450 (CYP) (CYP2D15 in dogs) inhibition and genetic polymorphisms [38] (Fig. 1). The CYPs constitute the main hepatic enzymes for drug biotransformation in human and non-human animals, and genetic variations reportedly influence clinical outcomes, given their impact on therapeutic effects of metabolism-dependent analgesics tramadol. The complete loss of a single protein that encodes a CYP or a single nucleotide polymorphism can alter the enzyme conformation, potentially reducing its enzymatic capacity. CYP2D is solely responsible for M1 production in dogs, and mutant alleles have been linked to differences between species and breeds [38]. For instance, in Bullmastiffs, Border collies, Rottweilers, and English cocker spaniels, van Hagen et al. [39] reported that all breeds demonstrate differences in coding sequences or exons 4, 5, and 6, but differences in exon 2 were exclusively detected in the Border collies. In other CYPs, such as the CYP2C41 gene, the site of gene deletion was consistently absent in Bearded Collies, Boxers, Bernese Mountains, Briards, French bulldogs, and Irish Wolfhounds. On the contrary, the enzyme was present in breeds such as Chinese shar-pei, Siberian husky, Schapendoes, and Kangals. Mutations in CYP oxidoreductase (POR) are reportedly prevalent in breeds like Greyhounds, which can metabolize CYP2B11 substrates but not CYP2D15; this could be translated into an altered metabolism in all breeds related to Greyhounds [40]. Although in vivo and clinical trials to determine the pharmacokinetics of tramadol within breeds are limited, these genetic modifications are associated with poor antinociceptive effects in Beagles, as mentioned by Kögel et al. [21] and Schütter et al. [41]. However, not all polymorphisms result in enzymatic deficits, indicating that the capacity of the mutant CYP is not always reduced [39]. In the case of dogs, this implies that, based on the available data and

| Table 2. Comparison of mean ± SD values for some M1 pharmacokinetic variables in dogs and cats. | Route and dose | Parameters (units) | Mean ± SD values | References |
|---|---|---|---|---|
| **Species** | **Parameters** | **Mean ± SD values** | **References** |
| **Dogs** | | | | |
| IV (4.4 mg/kg) | $\lambda_z$ (1/h) | 0.44 ± 0.12 | [25] |
| | $T_{max}$ (h) | 0.43 ± 0.20 | |
| | $C_{max}$ (ng/ml) | 146 ± 40.51 | |
| IV (4 mg/kg) | $\lambda_z$ (1/h) | 0.45 ± 0.39 | [29] |
| | $T_{max}$ (h) | 0.94 ± 0.52 | |
| | $C_{max}$ (ng/ml) | 0.02 ± 0.01 | |
| EPI (2 mg/kg) | $\lambda_z$ (1/h) | 3.77 ± 1.74 | [28] |
| | $T_{max}$ (h) | 1.14 ± 0.72 | |
| | $C_{max}$ (ng/ml) | 0.20 ± 0.08 | |
| IM (4 mg/kg) | $\lambda_z$ (1/h) | 0.34 ± 0.06 | [29] |
| | $T_{max}$ (h) | 0.88 ± 0.18 | |
| | $C_{max}$ (ng/ml) | 0.6 ± 0.01 | |
| Oral (11.2 ± 2.0 mg/kg) | $\lambda_z$ (1/h) | 0.33 ± 0.07 | [25] |
| | $T_{max}$ (h) | 0.50 ± 0.02 | |
| | $C_{max}$ (ng/ml) | 449.13 ± 210.10 | |
| Oral (5 to 7 mg/kg) | $\lambda_z$ (1/h) | 4.67 | [30] |
| | $T_{max}$ (h) | 2.82 | |
| | $C_{max}$ (ng/ml) | 4.60 | |
| Rectal (4 mg/kg) | $\lambda_z$ (1/h) | 0.45 ± 0.39 | [26] |
| | $T_{max}$ (h) | 0.94 ± 0.52 | |
| | $C_{max}$ (ng/ml) | 20 ± 12 | |
| IV (2 mg/kg) | $\lambda_z$ (min) | 261 ± 28 | [31] |
| | $T_{max}$ (min) | 55 ± 17 | |
| | $C_{max}$ (ng/ml) | 366 ± 31 | |
| Cats | IV (2 mg/kg) | $\lambda_z$ (h) | 3.54 ± 1.17 | [32] |
| | $T_{max}$ (h) | 0.25 ± 0.0 | |
| | $C_{max}$ (ng/ml) | 0.81 ± 0.23 | |
| Oral (5.2 mg/kg) | $\lambda_z$ (min) | 289 ± 19 | [31] |
| | $T_{max}$ (min) | 53 ± 13 | |
| | $C_{max}$ (ng/ml) | 655 ± 77 | |

$\lambda_z$ = plasma half-life, $T_{max}$ = time of maximum plasma concentration, $C_{max}$ = maximum plasma concentration.
inconsistencies between breeds, therapeutic dosing can be less efficacious in some dogs due to high enzymatic bio-transformation, increased clearance, and minimal plasma concentrations [42], or could be attributed to a metabolically less active enzyme that produces fewer active metabolites [39].

Conversely, cats are known to alter the metabolism of medications, given their enzymatic deficiency in functional UGT (UGT1A6 and UGT1A9), N-acetyltransferase and thiopurine S-methyltransferase. This loss reduces or completely suppresses the CYP catalytic activity, resulting in slow clearance, a higher plasma concentration of some compounds, and an enhanced risk of developing more adverse effects and toxicity [42]. To date, it remains unknown which feline CYP enzyme is responsible for the conversion of tramadol to M1 [43]; however, an in vitro report by Izes et al. [44] revealed that the rate of depletion of M1 in dogs exhibited an intrinsic clearance (Cl\text{\textsubscript{m}}) of 22.8 μl/min/mg, whereas in feline microsomes, the concentration of M1 was not depleted owing to the lack of CYP2B-like metabolism in this species. Accordingly, cats demonstrate reduced glucuronidation activity in phase II due to the lack of CYP2B6, the enzyme participating in the metabolism of M1 and is present in humans and dogs, with CYP2B11 as an analog in canines. Therefore, the lack of CYP2B6 in cats and the complete absence of phase II glucuronidation in feline microsomes, along with the minimal depletion of M1 in phase I, may clarify why cats present higher M1 concentrations in the bloodstream. On the contrary, dogs not only produce less M1 but also demonstrate rapid metabolite conjugation owing to the activity of the CYP2B11 enzyme, a pathway unavailable to cats. Recently, Ono et al. [45] reported that the CYP2C subfamily revealed some pseudogenes in cats, and its hepatic and intestinal presence was negligible, limiting the biotransformation; in addition, fewer than one-third CYPs were found to be available for phase I metabolism in cats when compared with dogs [46]. However, although these routes are deficient in cats, other pathways such as sulfation, acetylation, and methylation can also contribute to the metabolism and elimination of tramadol [44].

Notably, the metabolism of tramadol relies on hepatic enzymes, and this influences the amount of M1 produced [35], particularly in dogs, where the generation of M1 is 3.9-fold slower than in cats, but the production of the inactive M2 is faster (4.8-fold) [38]. To enhance the bioavailability of tramadol in dogs, particularly in Beagles, a soft capsule containing acetaminophen and tramadol was developed and demonstrated better bioavailability in dogs; however, additional studies and its application in a clinical setting as an analgesic adjuvant are crucial to evaluate the therapeutic potential of this new formulation [47]. The metabolism via CYPP450s in the intestinal mucosa can also influence the high amounts of the inactive compound. Likewise, the oxidation of M1–M5, which has a lower potency for MOR and low central nervous system (CNS) penetration ability, is faster in dogs than in cats [38]. The main route for drug excretion (M1, M2, and M5) is via urine (90%), and a small amount undergoes biliary and fecal elimination (10%) [16].

Tramadol is commonly administered via oral and intravenous (IV) routes. IV administration is typically administered at 2–4 mg/kg, as a sole drug or as multimodal analgesia, while oral therapy ranges between 4 and 10 mg/kg/6h in dogs and 1–4 mg/kg in cats [48]. In both dogs and cats, the route of administration reportedly influences certain pharmacokinetic parameters and the bioavailability of tramadol. For example, Di Salvo et al. [49] determined that intranasal administration in dogs (4 mg/kg) presented a half-life of 1.03 ± 0.53 h, a T\text{\textsubscript{\text{max}}} 0.67 ± 0.22 h, and a C\text{\textsubscript{\text{\text{\textsubscript{\text{max}}}}}} 123.17 ± 46.33 ng/ml, with an F\% between 3.26% to 20.6%. Interestingly, despite the observed bioavailability, bitches undergoing elective surgery exhibited similar analgesia. On the contrary, extradural (ED) administration reached a bioavailability of 82%, a longer half-life (2.66 ± 0.50 h) and higher drug concentration of 0.18 ± 0.12 μg/ml [28]. Tables 1 and 2 summarize the values observed following administration via these different routes.

Regarding differences between species, tramadol presented a shorter half-life in dogs (1–2 h) following oral administration with 10 mg/kg [50]. KuKanich [51] suggests that canines require higher doses (15 mg/kg/6–8 h, orally); however, therapeutic concentrations remain difficult to attain. Furthermore, repeated dosing decreased absorption and plasma concentration [50]. Typically, high doses of tramadol are well tolerated in dogs; however, nausea, salivation, anorexia, and sedation are common side effects. These adverse reactions may result from 5-HT and NE reuptake rather than opioid-linked mechanisms [41]. Some reports have indicated that administration of 40 mg/kg/day for 1 year resulted in adverse effects such as mydriasis, reduced body weight, restlessness, difficulty walking, salivation, vomiting, tremors, and seizures [50].

Pharmacokinetic data suggest that, given the higher bioavailability and slower clearance, the half-life of tramadol and M1 is longer in cats (3–4 h for tramadol; 4 to 6 h for the metabolite) [51,52]. In addition, M1 concentrations following oral administration of 5.2 mg/kg are higher than those reported in humans; hence, oral doses of 1–2 mg/kg/12 h or 5–10 mg per cat every 12 h are considered appropriate [51]. Following IV injection, higher M1 formation has been detected due to its slow conjugation, and there are no reports of other metabolites found in cats [38].

Although tramadol has a broad safety in animals, cats can exhibit excitement, dysphoria, euphoria, and mydriasis,
and accordingly, a lower dosage is recommended to avoid these manifestations [53]. In addition, the co-administration of drugs like famotidine or omeprazole may decrease the risk of gastrointestinal issues when tramadol is used with NSAIDs [50].

The effects of tramadol can be partially antagonized by naloxone, alpha-2 antagonists (yohimbine), and serotonin antagonists (ketanserin and ondansetron). Furthermore, acetylcholine could prevent the inhibitory action of M1 on muscarinic receptors [51]. The opioid, along with selective reuptake inhibitors, seems to increase its effectiveness; however, it should be noted that coadministration with selective 5-HT reuptake inhibitors such as fluoxetine and trazodone, monoamine oxidase inhibitors (e.g., selegiline), and tricyclic antidepressants (e.g., amitriptyline, domipramine) can increase the risk of developing 5-HT toxicity, known as “serotonin syndrome” [18,48].

As highlighted, the variability of M1 concentrations between species indicates that tramadol is more suitable for cats than dogs. In addition, the extended elimination half-life and greater amounts of plasma M1 can be attributed to lower glucuronidation activity observed in cats [22,37], in contrast to dogs, who have reported plasma M1 levels below therapeutic ranges even when compared with humans [4].

**Figure 1.** Pharmacokinetic differences of tramadol in dogs and cats [29,38].

Therapeutic Efficacy of Tramadol in Dogs

In dogs, the subtherapeutic concentration of M1, given the abundance and type of liver enzymes necessary for its metabolization [48], implies that tramadol monotherapy is unlikely to have meaningful benefits for pain control in this species [51]. A recent meta-analysis published by Donati et al. [54] documented a substantially low certainty of the evidence in terms of its analgesic action for managing
postoperative pain. However, the results of multiple studies using the surgical model of ovariohysterectomy (OVH) in bitches, a procedure where tramadol is commonly administered during the perioperative period or as rescue analgesia [55], revealed divergent conclusions.

For example, in one of the earliest reports where authors compared tramadol IV (2 mg/kg) to another well-known opioid (i.e., morphine at 0.2 mg/kg), they did not detect any differences in analgesia levels in bitches undergoing OVH [56]. In addition, the concentrations of catecholamines, cortisol, and glucose remained unaltered, and the postoperative pain scores were low in both groups. Although two animals treated with tramadol and only one treated with morphine required rescue analgesia, Mastrocinque and Fantoni [56] concluded that tramadol has a similar analgesic effect as morphine to control post-surgical pain. On the contrary, according to the Glasgow Composite Measure Pain Scale, nefopam IV at 2 mg/kg conferred superior analgesia than the same dose of tramadol, whereas patients receiving tramadol required less rescue analgesia (1 mg/kg IV of nefopam) [57]. Furthermore, Meunier et al. [58] reported that the preoperative IV application of 0.2 mg/kg meloxicam and 4 mg/kg tramadol as a multimodal analgesic regimen for free-roaming dogs subjected to OVH provided adequate analgesia but no clinical benefit over the use of meloxicam alone. Interestingly, the need for rescue analgesia was approximately four times greater in animals administered meloxicam than in the opioid group.

The SC route has been used in bitches both pre- and post-surgery (4 mg/kg) every 12 h to evaluate hematological and biochemical changes in pain stress after OVH [59], as well as following surgical treatment of pyometra (3 mg/kg every 6 h), as a multimodal approach along with SC meloxicam (0.2 mg/kg) [60]. Rossetti et al. [61] compared the analgesic effect of tramadol SC (3 mg/kg) and maropitant (1 mg/kg) during the post-surgical period in bitches. The authors reported that 5 out of 10 animals in the tramadol group and 6 out of 10 animals in the maropitant group required rescue analgesia with morphine, indicating that none of the drugs are appropriate as monotherapy to control immediate surgical pain. Conversely, epidural (EPI) opioids offer excellent long-term analgesia at a lower dose than systemic administration, as the opioid directly binds to receptors in the spinal cord, inhibiting the nociceptive signal and modulating the release of epinephrine and NE in descending pathways [62]. Mastrocinque et al. [62] compared EPI tramadol with intramuscular (IM) tramadol administration in elective surgery. The authors revealed that EPI was safe and afforded better neuroendocrine modulation of nociception with good cardiorespiratory stability, although it did not offer a more beneficial analgesic effect than systemic medication. In addition, preemptive EPI analgesia using tramadol at 1 and 4 mg/kg of lidocaine was compared with a morphine + lidocaine protocol during orchitectomy and OVH. According to the University of Melbourne’s pain scores, both protocols were similar and had an efficient analgesic effect within 24 h post-surgery. Nevertheless, the synergic effects of the premedication (ketamine + xylazine) might have enhanced the analgesic properties of the drugs [63]. Intraperitoneal (IP) tramadol at 4 mg/kg, combined with lidocaine (8.8 mg/kg IP) following OVH, provided better analgesia, based on scoring methods and lower cortisol and glucose levels at 6 and 3 h post-surgery, respectively; however, the differences were not statistically relevant when compared with animals in other groups [64]. Giorgi et al. [26] investigated rectal tramadol administration and observed that the plasma concentrations were below the effective therapeutic dose, undergoing rapid metabolism to its inactive compounds M2 and M5. Intranasal administration of tramadol (4 mg/kg) after canine OVH presented a similar effect to IV administration of both tramadol and 0.22 mg/kg of methadone, with no considerable alterations in pain scores or cardiorespiratory parameters [49].

Constant rate infusion (CRI) of tramadol has also been used during OVH at a dosage of 22 µg/kg/min (loading dose: 3 mg/kg IV) in combination with dissociative anesthesia. The drug combination decreased anesthetic requirements, shortened recovery times without adverse effects, and provided cardiorespiratory stability [65]. Furthermore, the coadministration of tramadol (4 mg/kg IV) and ketamine and lidocaine by CRI as preemptive analgesia reportedly improved the antinociceptive effect of the opioid following postoperative elective sterilization [66]. In healthy subjects, the combination of a tramadol–lidocaine infusion decreased the minimum alveolar concentration of sevoflurane, in comparison to the use of the opioid alone (2.2% ± 0.3% and 1.7% ± 0.3%, respectively) [67]. Likewise, CRI using tramadol (0.5, 1.0, and 2.0 mg/kg/h) and 5 mg/kg of ketoprofen SC at 50 mg/ml in dogs undergoing laparotomy reduced glucose levels and pain scores during 1 h post-surgery without cardiorespiratory alterations observed on following days [14].

The effect of tramadol on thermal and mechanical acute nociceptive pain has revealed differences in several investigations. In Beagles exposed to thermal stimuli using the tail-flick test, high doses of IV tramadol (6.81 and 10 mg/kg), compared with tapentadol (2.15, 4.64, 6.81 mg/kg IV) and morphine (0.464, 0.681, 1.0 mg/kg), did not induce antinociception and side effects such as ataxia and a short seizure [21]. The dose-dependent antinociception and better analgesic performance of morphine and tapentadol were attributed to their direct action without relying on metabolic activation. Schütter et al. [41] also described the effect of tramadol IV (1 and 4 mg/kg) as questionable to treat acute nociception in
Beagles. Although a slight increase in the mechanical threshold was reported, no clinically relevant antino-
ciception was noted at the thermal threshold, and M1 concentrations were below the previously reported con-
centration [41].

Some authors have suggested the accumulative effect of tramadol after repeated oral dosing as the underlying reason for its effectiveness during chronic or osteoar-thritic pain [41]; however, Budsberg et al. [68] reported that dogs orally administered 5 mg/kg/8 h failed to
demonstrate a clinical benefit for elbow or stifle joint osteoarticular disease when compared with carprofen
(2.2 mg/kg/12 h) and a placebo. Conversely, Benitez et al. [69] achieved similar analgesia to a hydrocodone–acetaminophen protocol following oral administration of tramadol (5–7 mg/kg/8 h) after tibial plateau leveling osteotomy. Piras et al. [70] determined that cimicoxib and tramadol have similar analgesic potency for long-term analgesia for the same procedure. However, treatment with a selective COX-2 inhibitor conferred superior func-
tional improvements. Therefore, tramadol may be unsuit-
able as monotherapy for osteoarthritic pain. Accordingly, it has been established that tramadol (at 3–5 mg/kg
every 8 h for 4 weeks), in addition to NSAIDs, increases the peak vertical force in 50% of the subjects, a simi-
lar percentage to dogs treated with gabapentin (61%); hence, both drugs can be recommended in this clinical condition and as a part of multimodal analgesia [71]. In a controlled trial conducted by Malek et al. [72], animals with naturally occurring hip osteoarthritis received tram-
adol at 4 mg/kg/8 h orally for 2 weeks. The animals showed improved physical parameters, pain, and mobility in all treatment groups. The authors revealed two significant findings regarding tramadol: i) the improved pain interference was superior to the improvement in pain severity; and ii) decreasing plasma tramadol con-
centrations following 2 weeks due to the impaired bio-
transformation in domestic canines. Therefore, tramadol treatment for chronic pain management is still question-
able [72]. The prevalence of low M1 and high M2 con-
centrations was reported by Giudice et al. [19] in dogs with neuropathic pain due to degenerative lumbosacral stenosis, where 50% of patients did not show a reduc-
tion in pain scores after 1 week of oral dosage (3 mg/
kg/8 h) together with prednisolone; accordingly, the treat-
ment protocol was replaced with oral gabapentin. In a recent work by the same author [73], buprenorphine
(0.02 mg/kg IM), a partial mu-opioid receptor agonist, offered better postoperative analgesia in dogs subjected
to hemilaminectomy when compared with 3 mg/kg IM tramadol; the study also revealed that tramadol might be a better option for follow-up treatment. Based on these findings, tramadol seems appropriate for reducing lumbosacral pain to some degree, but some dogs might fail to respond despite high doses and a shortened dosing time. Ripplinger et al. [74] supported this statement after evaluating 180 dogs undergoing vertebral surgery, where 15% of 46 dogs receiving tramadol (3 to 8 mg/kg SC) revealed a higher percentage of persistent pain than other opioids (morphine and methadone) (15%, 4.8%, and 3.3%, respectively). The authors attributed this finding to pharmacokinetics and suggested that trama-
dol should not be employed as monotherapy or first-line therapy for long-term orthopedic pain management.

In other procedures such as surgical resection of cutaneous tumors, the effects of preemptive tramadol administration (3 mg/kg every 8 h), orally for 48 h before surgery, were compared with carprofen (2.2 mg/kg/12 h) and no preemptive analgesia, in combination with pre-
operative hydromorphone [75]. According to pain scores, there was no difference between animals that received and did not receive preemptive analgesia, and this could be due to the premedication time, surgical procedure, and the surgeon itself. In mastectomies and OVH, tramadol (5
mg/kg IM) did not offer superior analgesia when com-
pared with methadone (0.5 mg/kg IM), another opioid. The group medicated with methadone required less com-
plementary analgesia and showed lower drug consump-
tion with better postoperative pain scores [76]. Using the same surgical procedure, Reis et al. [77] determined that the addition of tramadol (2 and 4 mg/kg EPI) to an anesthetic protocol of tramadol + propofol + levobupiva-
caine potentiated and reduced the propofol requirement while providing adequate analgesia, with vascular stabil-
ity but without differences when compared with propo-
fol + levobupivacaine alone. Furthermore, the analgesia provided by oral tramadol (12 mg/kg/24 h) (combined with an anesthetic protocol of methadone, bupivacaine, and carprofen) for lateral thoracotomy exhibited higher pedometric activity than patients receiving fentanyl transdermal patches [78].

Accordingly, available scientific evidence indicates that tramadol may be an acceptable alternative for dogs only when administered with other analgesic drugs [48]. In canines, its efficacy depends on hepatic activation and is influenced by genetic or pharmacological factors. Moreover, the administration of CYP substrates or inhibi-
tors can increase opioid bioavailability, thereby prolonging and intensifying analgesic and adverse effects [38]. For example, ketoconazole and cimetidine are inhibi-
tors of CYP3A, an enzyme that produces M2. Their use with tramadol can reduce first-pass metabolism and the formation of M2 and M5 from M1 metabolite [79]. On the contrary, phenobarbital has the opposite effect and reduces the analgesic efficacy of this opioid by altering drug metabolism and M1 concentrations [38].
Therapeutic Efficacy of Tramadol in Cats

In cats, opioids are not always considered first-choice drugs owing to the reduced glucuronidation activity that can increase the risk of toxicity when doses and intervals are not administered appropriately [80]. The main side effects following high doses of tramadol are sympathetic responses such as mydriasis, hypertension, tachycardia, and an increase in adrenal hormone concentration (e.g., NE, epinephrine, dopamine, met-enkephalin). Dysphoria, excitement, anxiety, and vocalizations have also been reported [81]. Nevertheless, tramadol is a potential alternative to NSAIDs, drugs not well-tolerated or contraindicated in some cases, and for treating certain pre-existing diseases [7]. The bitter taste of tramadol is also considered a limiting factor in cats [51].

The effect of tramadol in surgical settings as perioperative analgesia has been examined in various studies in recent years. Teixeira et al. [22] evaluated 2 mg/kg/8 h SC tramadol alone and in combination with IV dipyridam at different doses (25 mg/kg every 8, 12, or 24 h) in patients undergoing OVH. Biochemical, hematological, and behavioral parameters were assessed. The authors revealed that cats receiving dipyridam + tramadol required less complementary analgesic; however, it did not offer a superior analgesic combination than the use of the opioid alone [22]. Likewise, IV administration of 3 mg/kg tramadol during OVH and orchiectomy, along with meloxicam (0.05 mg/kg SC), revealed that 89.4% of the animals presented only mild pain [82]. In a recent work from Bovo et al. [83], the IM administration of tramadol (2 mg/kg) produced a similar analgesic efficacy to morphine (0.5 mg/kg), with 40% of patients requiring rescue analgesia in both groups; however, tramadol induced a lower degree of sedation than the pure agonist.

A comparative study between IM tramadol at 2 and 4 mg/kg, and IM pethidine (6 mg/kg) determined the dose-dependent analgesic effect of tramadol [53]. At the highest dose, serum cortisol values and pain levels were lower, and the animal did not require additional analgesia. Associations of tramadol (2 mg/kg IM) and midazolam (0.2 mg/kg IM) as pre-anesthetic protocol for OVH were compared with dexametomidine. The tramadol group had fewer secondary effects (nausea and vomits), a faster recovery time, and better cardiovascular stability than the alpha-2 adrenergic agonist [84]. Furthermore, given the absence of cardiorespiratory changes and physiological parameters within normal ranges, Evangelista et al. [53] suggested that tramadol is a reliable perioperative drug for cats undergoing OVH at the recommended doses. In addition, Li et al. [85] investigated a mixture of tiletamine/zolazepam–xylazine–tramadol during sterilization. This protocol enhanced the induction of the anesthesia and the antinociceptive effect of the drugs without cardiopulmonary side effects. A study by Nascimento et al. [86] mentioned the addition of tramadol at 2 mg/kg IM as rescue analgesia in patients subjected to laser acupuncture and electroacupuncture therapies before OVH. Despite its analgesic effect, 3 out of 30 cats required additional meloxicam when the opioid failed to improve postoperative pain scores. On the contrary, Martins et al. [87] determined that perioperative tramadol (2 mg/kg IM) along with acepromazine and tiletamine/zolazepam conferred a higher analgesic effect, with lower heart rate and a stable respiratory rate in prepubertal cats undergoing OVH.

Experimentally, tramadol (1 mg/kg) and acepromazine (0.1 mg/kg), both SC, were evaluated on pressure and thermal thresholds. Tramadol monotherapy offered limited effects on nociceptive thermal and pressure stimuli, but improvements were observed when administered with acepromazine [80]. In many cases, the neuroleptoanalgesia afforded by acepromazine increases the weak analgesic effect of the opioid. In particular, this pre-anesthetic association has also been considered a suitable option for ophthalmic surgery in cats [88], with benefits such as unaltered intraocular pressure and sustained mydriasis when co-administered with tramadol (3 mg/kg IM), distinct from acepromazine alone (0.05 mg/kg IM) [89]. The analgesic potency of 2 mg/kg IM tramadol or EPI tramadol co-administered with lidocaine (3 mg/kg EPI) was also evaluated in terms of painful mechanical stimuli following pressure application from hemostatic forceps in the skin of several body regions. Both administration routes demonstrated a short anesthetic onset and similar motor blockade duration without cardiorespiratory changes. Regarding analgesia, EPI tramadol with lidocaine produced more prolonged analgesia than IM administration (120 ± 31 min vs. 71 ± 17 min, respectively) [37].

In osteoarticular diseases associated with chronic pain, nociception, inflammation, central and peripheral sensitization, hyperalgesia, allodynia, and a reduction in mobility, tramadol is recommended despite necessitating prolonged treatment periods [20]. Clinical trials in geriatric cats with osteoarthritis medicated with different dosages (1–4 mg/kg) of oral tramadol indicate that 2 mg/kg/12 h for 5 days increased the level of activity and the global quality of life; however, adverse effects such as euphoria, dysphoria, sedation, hyporexia, and diarrhea were present at high doses (4 mg/kg) [43]. Monteiro et al. [20] observed a favorable response in mobility, biomechanical aspects, and hypersensitivity in patients medicated with oral tramadol at 3 mg/kg/12 h for 19 days and three-non-treatment months during the therapy. Conversely, another study evaluated the analgesic efficacy of oral tramadol (3 mg/kg/12 h) with oral transmucosal meloxicam (approximately 0.05 mg/kg/24 h) for 25 days. Animals administered tramadol did...
not show a clinical benefit over meloxicam alone, except in the case of central hypersensitivity [34]. In addition, some cats showed mydriasis, depression, hyporexia, hypersalivation, and vomiting, although these reactions could not be attributed to a specific treatment.

Tramadol has also been used for treating polytraumatized cats [90], and it is known to significantly diminish central sensitization in neuropathic pain. Additionally, transdermal patches of tramadol exist, but there is no literature on its use in domestic species [91].

Clinical Pharmacology of Tapentadol

Tapentadol was formulated based on morphine, tramadol, and its active M1 metabolite building, but exists as a single enantiomer commercially available for oral administration [92,93]. It is a central acting atypical opioid with agonism toward MOR while simultaneously inhibiting neuronal reuptake of NE without any clinical evidence of serotonergic activity, unlike tramadol [21,94] (Fig. 2). It blocks modulation and perception of the nociceptive pathway, acting on the ascending tracks via MOR agonism [93]. The affinity of tapentadol for MOR, compared to morphine, is approximately 50-fold below but 50-fold to 120-fold greater than tramadol. Thus, it has an analgesic potency almost identical to morphine, with adequate oral absorption, minimal physiological effects, and a similar plasma concentration in animals and humans [94,95].

The pharmacokinetic profile of tapentadol makes it an attractive analgesic option when compared with tramadol [96], especially in those species where active M1 concentrations are negligible, such as dogs. Following IV (20 mg) and oral (200 mg single dose) administration in dogs, measurable levels of plasma concentration were detectable for up to 6 h and 15–240 min, respectively [97], with a terminal half-life of 0.5–1 h [98]. In cats, the parenteral route achieved a bioavailability above 90% with a short elimination half-life (2–3 h) [91]. On the contrary, oral administration revealed a decreased bioavailability due to reduced glucuronidation (Table 3) [95,97,99].

The advantage of tapentadol administration as an active compound is the safety and efficacy that does not warrant CYP450 liver enzyme-dependent metabolic activation; this reduces the inter-individual analgesic variabilities due to genetic polymorphisms and decreases the possibility of adverse interactions when administered with other

![Figure 2](http://bdvets.org/javar/)

**Figure 2.** Mechanism and sites of action of the opioids tramadol and tapentadol. $\alpha_2R$ = alpha-2 receptors, GLU = glutamate, SP = substance P.
Moreover, it is considered a good option for patients with mild or moderate hepatic or renal impairment without affecting its analgesic effect [92]. The metabolism is mainly through phase II glucuronidation, with sulfation playing a minor role and minimal contribution from CYP enzymes [18,95]. Apparently, the metabolites produced do not provide analgesia, resulting in fewer side effects (e.g., sialorrhea or sedation) [97]; however, some studies have reported an accumulative effect after repeated dosing [95].

The antinociceptive effect of the marked NE reuptake inhibition on terminal endings and the descending pain pathway suppresses the noxious signaling by stimulating alpha-2 adrenergic receptors in the nociceptive fibers. The inhibitory mechanism of NE is reportedly predominant during the modulation of chronic and neuropathic pain [17,93]. Likewise, because tapentadol does not use serotonergic pathways, there are no reports of serotonin syndrome in animals, and the possibility of developing this toxicity is reduced [18]. However, a study has documented serotonin-related toxicity when administered in association with tramadol, duloxetine, venlafaxine, amitriptyline, sertraline, desvenlafaxine, and escitalopram, but a clear correlation cannot be established in dogs and cats [100].

| Table 3. Comparison of mean ± SD values for some tapentadol pharmacokinetic variables in dogs and cats. |
| Species | Route and dose | Parameters (units) | Mean ± SD values | References |
|---------|---------------|-------------------|-----------------|------------|
| Dogs    | IV (50 mg/dog) | F% | ND | [97] |
|         |               | Vd (ml/kg) | 5,917–7,236 | |
|         |               | Cl (ml/min/kg) | 59.8–116.6 | |
|         |               | T\text{\textsubscript{max}} (min) | ND | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | ND | |
|         | IV (30 mg/kg) | F% | ND | [95] |
|         |               | Vd (ml/kg) | ND | |
|         |               | Cl (ml/min/kg) | ND | |
|         |               | T\text{\textsubscript{max}} (h) | 3.5 ± 1.2 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 31.0 ± 7.5 | |
|         | IV (20 mg/kg) | F% | ND | [95] |
|         |               | Vd (ml/kg) | ND | |
|         |               | Cl (ml/min/kg) | ND | |
|         |               | T\text{\textsubscript{max}} (h) | 2.4 ± 1.2 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 19.7 ± 5.5 | |
|         | IV (10 mg/kg) | F% | ND | [95] |
|         |               | Vd (ml/kg) | ND | |
|         |               | Cl (ml/min/kg) | ND | |
|         |               | T\text{\textsubscript{max}} (h) | 2.7 ± 0.9 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 10.2 ± 1.8 | |
|         | Oral (200 mg/dog) | F% | 4.4 ± 2.3 | [97] |
|         |               | Vd (ml/kg) | 7,024–55,358 | |
|         |               | Cl (ml/min/kg) | 1,138–2,384 | |
|         |               | T\text{\textsubscript{max}} (min) | 47.5 ± 6.1 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 240–3,640 | |
| Cats    | IV (5 mg/kg)  | F% | ND | [99] |
|         |               | Vd (ml/kg) | 8.79 ± 1.97 | |
|         |               | Cl (ml/min/kg) | 35.60 ± 7.05 | |
|         |               | T\text{\textsubscript{max}} (h) | ND | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | ND | |
|         | IM (5 mg/kg)  | F% | 93.93 ± 9.91 | [99] |
|         |               | Vd (ml/kg) | 7.53 ± 2.95 | |
|         |               | Cl (ml/min/kg) | 37.85 ± 5.68 | |
|         |               | T\text{\textsubscript{max}} (h) | 0.25 ± 0.26 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 1,406 ± 779 | |
|         | SC (5 mg/kg)  | F% | 90.01 ± 6.52 | [99] |
|         |               | Vd (ml/kg) | 7.06 ± 2.10 | |
|         |               | Cl (ml/min/kg) | 40.13 ± 9.97 | |
|         |               | T\text{\textsubscript{max}} (h) | 0.63 ± 0.31 | |
|         |               | C\text{\textsubscript{max}} (ng/ml) | 906 ± 356 | |

F% = bioavailability, ND = not determined, Vd = apparent volume of distribution at steady-state, Cl = clearance, T\text{\textsubscript{max}} = time of maximum plasma concentration, C\text{\textsubscript{max}} = maximum plasma concentration.
Conversely, unlike tramadol, opioid antagonists like naloxone cannot suppress the effects of tapentadol, corroborating the dual mechanism of this drug [17].

It is generally well tolerated, but it may cause excitability in cats and depression, panting, and salivation in dogs. Disadvantages include antimuscarinic activity and low oral bioavailability, although in animals with deficient glucuronidation, such as cats, tapentadol bioavailability may be enhanced [97]. Similar to tramadol, tapentadol is bitter and can be limiting in this species [91].

It can treat mild to severe pain when animals need prolonged analgesia [99]. In chronic pain, tapentadol is administered as part of multimodal protocols combined with NSAIDs. To date, there are no recommended dosages in companion animals, but clinical doses mentioned by Gaynor and Muir [98] range between 5 and 10 mg/kg, every 12 or 8 h, with a half-life of 0.5–1 h in dogs.

Studies evaluating the analgesic effect of tapentadol in dogs and cats are limited [101]. Moreover, recent experimental animal models have shown good antinociception associated with mechanical stimulation and acute orofacial pain in adult rats medicated with tapentadol at 2 and 5 mg/kg [102]. In another study, tapentadol demonstrated a dose-dependent effect and inhibited the electrophysiological neuronal activity in the locus coeruleus in a rat model of diabetic polyneuropathy [103]. In addition, tapentadol enhances the inhibitory activity of NE descending pathways and prevents nociceptive signaling at the spinal cord of rats with osteoarthritis [104]. These authors also mentioned that alpha-adrenergic drugs such as yohimbine and atipamezole could block tapentadol action. Tapentadol has also exhibited a dose-dependent antinociceptive effect (approximately 99%) in rodents without causing gastric alterations [105], whereas a synergistic effect was observed when co-administered with NSAIDs (e.g., diclofenac and ketorolac) in rats assessed using the writhing test [106].

**Experimental Research and Future Expectations for the Use of Tapentadol in Dogs and Cats**

As stated earlier, investigations assessing tapentadol and its analgesic effects in small animals are scarce, but some studies have elucidated its potential in veterinary medicine. In dogs, Howard et al. [95] reported the pharmacokinetics and pharmacodynamics of oral tapentadol (at 10, 20, and 30 mg/kg). The highest plasma concentration was detected approximately 2.7–3.5 h after administration, with rapid absorption and no side effects [95]. On the contrary, high doses (6.8 mg/kg IV), produced sedation, ataxia, salorhea, and diarrhea in Beagle dogs [21]. In a similar work, Kögel et al. [21] used the tail-flick model test to assess acute nociceptive pain, comparing the analgesic potential of IV tramadol, morphine, and tapentadol (2.15, 4.64, and 6.8 mg/kg, respectively). Unlike tramadol, morphine and tapentadol induced antinociception through direct action on MOR without the need for metabolic activation, indicating their potential for reliable analgesia in clinical settings.

In dogs with orthopedic pain due to naturally occurring cranial cruciate ligament rupture, tapentadol (30 mg/kg orally and single dose) lowered subjective lameness scores 4 h after administration [94]; however, no significant improvement in the gait analysis was observed, which was evaluated by a pressure-sensitive gait system. As the plasma level required to achieve adequate analgesia in dogs remains unknown, Kieves et al. [94] concluded that the ranges obtained in this study (9–49 ng/ml) might need to be higher, along with repeated doses, to achieve the therapeutic efficacy with long-term treatment; however, further studies are needed.

Concerning tapentadol in cats, Doodnaught et al. [92] evaluated the thermal antinociception of oral dosages of 5.7 and 11.4 mg/kg in healthy cats. The thermal threshold and skin temperature rise for up to 2 h after administering the highest dose. The duration of thermal antinociception was similar between tapentadol at 11.4 and 0.02 mg/kg of buprenorphine IM, with salivation as the only documented side effect [92]. This observation is consistent with the results by Lee et al. [99], where 5 mg/kg IV caused salivation, agitation, and panting. In this same study, the authors determined the bioavailability of parenteral IM and SC tapentadol (94% and 90%, respectively).

Some authors have carried out pharmacokinetic and toxicological investigations in other species, including recent studies in Amazon parrots [107] and laying hens [108]. In parrots, the short half-life (24.8 min) following one dose of 30 mg/kg per oral (PO) indicated its limited use in this species; in hens, only IV administration (1 mg/kg) exhibited minimum therapeutic efficacy when compared with PO administration (5 mg/kg), showing a half-life of 0.9 h and a low bioavailability. Additional studies have been carried out in red-eared slider turtles [109], yellow-bellied slider turtles [110], Wistar rats [111], and goats [112]. Antinociceptive effects have been evaluated in rabbits [113], rats, and mice [105], all of which demonstrated adequate analgesia at 5 mg/kg IV dosages.

As previously reported, analgesic assessments of tapentadol in companion animals are insufficient to reach a definitive conclusion before recommending its use in veterinary practice. Additionally, some limitations in this study and the selected papers should be noted. Given the inclusion criteria and previously mentioned publications, all works that employed tramadol and tapentadol as an analgesic or antinociceptive therapy in different pain-related settings in dogs and cats were included. The sample size of animals...
used in selected articles was not deemed an exclusion/inclusion criterion in the present review; however, some reported data included clinical trials with minimal subjects, leading to a false absence of statistical differences between groups or subjects. Nonetheless, these findings contribute to critical and scientific knowledge that needs to consider these factors. Regarding tapentadol, the limited available data for an analgesic protocol in veterinary medicine is, per se, a limitation to objectively propose or reject the application of this drug; hence, animal models and nociception tests can help veterinarians to broaden their view on this analgesic alternative.

To summarize the characteristics of tramadol and tapentadol, the doses and mechanisms of action are presented in Table 4 [81,91,114]. Table 5 [16-18,20,25,26,31,35,93,95,99] presents a brief comparison of relevant pharmacological aspects and differences between tramadol and tapentadol in dogs and cats.

### Serotonin Syndrome Induced by Tramadol and Serotonin Reuptake Inhibitors

It has been established that tramadol possesses a non-opioid mechanism of action that promotes the inhibition of 5-HT reuptake, increasing its extracellular levels in the brain [115]. 5-HT is stored primarily in the presynaptic nerve terminals, in enterochromaffin cells, and within platelets [116]. It is an inhibitory neurotransmitter used in

| Table 4. Dosage and pharmacological characteristics of tramadol and tapentadol [81,91,114]. |
|---|---|---|---|---|
| **Drug** | **Species** | **Doses** | **Route** | **Mechanism of action** | **Adverse effects** |
| Tramadol | Dogs | 4–10 mg/kg/6–8 h | IV, IM, PO | Agonism to μ-receptors | Tremors, myoclonus, tachycardia, agitation, excitement, hypertension, and seizures (signs of serotonin toxicity) |
| | Cats | 1–4 mg/kg/12–24 h | IM, IV, PO | Inhibits monoamine reuptake (NE and 5-HT) |
| Tapentadol* | Dogs | 2–30 mg/kg | IV, PO | μ-Opioid receptor agonist | Salivation, agitation, panting, mydriasis, euphoria, and sedation |
| | Cats | 5–10 mg/kg q 12 h | IV, IM, PO, SC | NRI |

PO = oral.
* doses taken from the available literature.

| Table 5. Summary of some pharmacological differences between tramadol and tapentadol in dogs and cats [16–18,20,25,26,31,35,93,95,99]. |
|---|---|---|---|---|
| **Tramadol** | **Tapentadol** |
| **Dogs** | **Cats** | **Dogs** | **Cats** |
| Administration routes | IV, IM, PO, SC | IV, PO |
| Absorption | Intestinal | Intestinal |
| Oral bioavailability | 60%–83% | 60%–90% | 4.4% | Unknown |
| Plasma concentration (ng/ml) (M1) | 146–449 | 366–850 | 240–3640 | 906–1,406 |
| Metabolism | Hepatic | Hepatic |
| Main metabolites | M2 | M1 and M5 | Tapentadol-O-glucuronide |
| Active metabolite | M1 | Active parent compound |
| Terminal half-life (M1) | 1–2 h | 4–6 h | 2–5 h | 1–3 h |
| Excretion | Urine and feces | Urine and feces |
| Interactions | Combination with a SSRI can cause serotonin syndrome | The antagonist atipamezole, and yohimbine could potentially block the effect of tapentadol |
| Differences between species | CYP polymorphism affects its analgesic efficacy in dogs | It does not rely on metabolism to produce its therapeutic effects, reducing the inter-species differences. |

SSRI = Selective serotonin reuptake inhibitor.
animals to treat separation anxiety, canine cognitive dysfunction, and compulsive disorders, as well as an adjuvant for pain control [116].

The concomitant administration of drugs that can increase the concentration of 5-HT, such as tramadol and other medication listed in Table 6 [51,117], potentiates the risk of developing serotonin syndrome [115].

This syndrome affects dogs and cats, typically occurs within the first 30 min to 2 h post-ingestion and demonstrates multiple clinical signs in patients (Table 7) [115,117].

Indrawirawan and McAlees [118] published a case of accidental tramadol overdose (80 mg/kg orally) in a cat. The animal exhibited agitation, hypersalivation, hypertension, tachycardia, abdominal pain, head tremors, myoclonus, hyperreflexia, paresis, disorientation, and altered mental status. In cats, deficient hepatic methylation and glucuronidation enhance the predisposition to severe adverse effects [50]. Toxicity has also been recorded in dogs, with similar signs mentioned in the case report [50].

The initial treatment requires the administration of serotonergic antagonists such as cyproheptadine (1.1 mg/kg q 1–2 h PO) or chlorpromazine (3 mg/kg) [119]. Supportive therapy includes phenobarbital or propofol in case of seizures (benzodiazepines can increase arousal), beta-blockers (propranolol), muscle relaxants (methocarbamol), antihypertensives and fluid therapy (crystalloids), physical methods to reduce hyperthermia, tracheal intubation if neuromuscular paralysis occurs, and cardiorenal, electrolyte and glucose monitorization [117,119].

The main complications include rhabdomyolysis, myoglobinuria, intravascular coagulation, renal insufficiency, respiratory and CNS depression, and even death [115]. For preventing serotonin syndrome, the combination of serotonergic drugs should be avoided and administered doses need to be regulated, especially in individuals with reduced liver metabolism [120].

Table 6. Drugs that can contribute to serotonin syndrome, in addition to another serotonergic drug, or overdosing [51,117].

| Type or drug | Examples |
|-------------|----------|
| Analgesic   | Tramadol |
| Selective 5-HT reuptake inhibitors | Citalopram, escitalopram, fluoxetine, fluvoxamine, nefazodone, paroxetine, sertraline, trazodone. |
| Tricyclic antidepressant | Amitriptyline, clomipramine, desipramine, dosulepin, doxepin, imipramine, nor-triptyline, opipamol, trimipramine. |
| NRI and 5-HT reuptake inhibitors | Desvenlafaxine, duloxetine, milnacipran, sibutramine, venlafaxine. |
| Atypical antipsychotics | Aripiprazole, asenapine, bupropion, clozapine, lurasidone, olanzapine, quetiapine, risperidone, ziprasidone. |
| Selective MAO inhibitors | Brofaromine, metralindole, minaprine, moclonemide, pirlindol, toloxatone, rasagline, selegiline, amitraz. |
| Non-selective MAO inhibitors | Hydralazine, isocarboxazid, isoniazid, phenelzine, phenoxyprazine, safrazine, carazazone, furazolidone, linezolid. |
| Other | 5-HT, chlorpheneramine, dextrometorphan, lithium, metoclopramide, ondansetron, tryptophan. |

MAO = monoamine oxidase.

Table 7. Clinical signs of serotonin syndrome in companion animals [115,117].

| Body system          | Clinical signs                                                                 |
|----------------------|-------------------------------------------------------------------------------|
| Cardiovascular       | Tachycardia, bradycardia, arrhythmia, hypertension                            |
| Respiratory          | Tachypnea                                                                     |
| Nervous system       | Sedation, lethargy, hyperexcitability, temporary blindness, nystagmus         |
| Mental status        | Confusion, agitation, excitement, vocalizations, aggression, coma              |
| Neuromuscular         | Hyperreflexia, muscle spasms and hyperthermia, trembling, rigidity, convulsions, recumbency, weakness, hyperesthesia |
| Gastrointestinal     | Hypersalivation, nausea, vomiting, diarrhea, abdominal pain                  |
| Metabolic            | Fever or hyperthermia with diaphoresis                                        |
Conclusion
Tramadol and tapentadol are analgesics administered to manage pain in dogs and cats. These drugs have opioid and non-opioid mechanisms that influence their analgesic effect and efficacy in companion animals. Tramadol, a pain reliever widely employed in animals, is dependent on liver metabolism to produce the principal active M1 metabolite. Compared with cats, dogs differ in their capacity to metabolize M1 due to polymorphisms in hepatic enzymes, which affect the analgesia induced by tramadol. On the contrary, the novel opioid tapentadol has potential advantages over tramadol as it does not depend on hepatic biotransformation, acts through NE reuptake inhibition, and demonstrates stronger MOR agonism; however, to date, the investigations assessing the analgesic efficacy of tapentadol remain limited in veterinary medicine. Given these points and available evidence on both drugs, tramadol appears to be a more suitable therapeutic option for cats and, preferably, should be administered as a multimodal analgesia protocol in both species, particularly dogs. Tapentadol might have a superior analgesic profile in animals, but the effectiveness of this opioid needs to be further clarified before recommending its use for managing acute and chronic pain in dogs and cats.

List of Abbreviations
CNS: central nervous system; COX-2: cyclooxygenase 2; CRt: constant rate infusion; CYP: cytochrome P450; EPI: epidural; h: hours; 5-HT: serotonin; IM: intramuscular; IV: intravenous; M1: O-desmethytramadol; M2: N-desmethytramadol; M5: N,O-didesmethytramadol; MOR: μ-opioid receptor; NE: norepinephrine; NMDA: N-methyl-D-aspartate; NRI: norepinephrine reuptake inhibitor; NSAIDs: non-steroidal anti-inflammatory drugs; POR: CYP oxidoreductase; SC: subcutaneous; TRPV1: transient receptor potential vanilloid 1; UDP: uridine diphosphate; UGT: UDP-glucuronon transferase; OVH: ovariohysterectomy.

Acknowledgment
The authors did not receive any funds for this study.

Conflict of interests
The authors declare that they do not have any competing interests.

Authors’ contributions
IHA conceptualized, drafted, and supervised the final version. IHA, ADO, AEMC, and ACA contributed to the original draft, data curation, investigation, writing, review, and editing of the manuscript. ADO and AEMC developed the methodology, wrote, and edited the review. ACA and ADO retrieved articles, wrote, reviewed, and edited the manuscript. All authors have read and approved the final manuscript.

References
[1] Hernández-Avalos I, Flores-Gasca E, Mota-Rojas D, Casas-Alvarado A, Miranda-Cortés AE, Domínguez-Oliva A. Neurobiology of anesthetic-surgical stress and induced behavioral changes in dogs and cats: a review. Vet World 2021; 14(2):393–404; www.doi.org/10.14202/vetworld.2021.393-404
[2] Nazifi S, Shojaee TA, Mohammadi S, Erjaae H, Mirzaie A. The effect of tramadol and meloxicam, alone and in combination on oxidative stress status in dogs. Comp Clin Pathol 2019; 28(4):1055–60; https://doi.org/10.1007/s00580-019-02927-w
[3] Grubb T. Following the pathway for effective patient analgesia. Hellenic J Companion Anim Med 2021; 10(1):23–31.
[4] Simon BT, Scallan EM, Von Pfeil DJF, Boruta DT, Nibbrett BM, Beauchamp G. Perception and opinions of pet owners in the United States about surgery, pain management, and anesthesia in dogs and cats. Vet Surg 2017; 47(2):277–84; https://doi.org/10.1111/vsu.12753
[5] Truchetti G, Otis C, Briville AC, Beauchamp G, Pang D, Troncy E. Management of veterinary anaesthesia in small animals: a survey of current practice in Quebec. PLoS One 2020; 15(1):e0227204; https://doi.org/10.1371/journal.pone.0227204
[6] Grubb T, Sager J, Gaynor JS, Montgomery E, Parker JA, Shafford H, et al. 2020 AAHA anesthesia and monitoring guidelines for dogs and cats. J Am Anim Hosp Assoc 2020; 56(2):59–82; https://doi.org/10.5326/JAAHA-MS-7055
[7] Bradbrook CA, Clark L. State of the art analgesia-recent developments in pharmacological approaches to acute pain management in dogs and cats. Vet J 2018; 238:76–82; https://doi.org/10.1016/j.tvjl.2018.06.003
[8] Tomsić K, Rakinić K, Sokolov C, Selikár A. A survey study on the recognition and treatment of pain in dogs and cats by Slovenian veterinarians. Vet Anaesth Analg 2021; 48(3):334–43; https://doi.org/10.1111/vaa.2020.11.007
[9] Hernández-Avalos I, Valverde A, Iancovich-Camarillo JA, Sánchez-Aparicio P, Recillas-Morales S, Osorio-Avalos J, et al. Clinical evaluation of postoperative analgesia and serum biochemical changes of paracetamol compared to meloxicam and carprofen in bitches undergoing ovariohysterectomy. PLoS One 2020; 15(2):e0223697; https://dx.doi.org/10.1371%2Fjournal.pone.0223697
[10] Grubb T, Lorbispre H. Local and regional anaesthesia in dogs and cats: overview of concepts and drugs (Part 1). Vet Med Sci 2020; 6(2):209–17; https://doi.org/10.1002/vms3.219
[11] Valverde A, Skelding AM. Alternatives to opioid analgesia in small animal anesthesia: alpha-2 agonists. Vet Clin North Am Small Anim Pract 2019; 49(6):1013–27; https://doi.org/10.1016/j.cvsm.2019.07.010
[12] Miranda-Cortés E, Ruiz García A, Olivera-Ayub AE, Garza-Malacara G, Ruiz-Cervantes JG, Toscano-Zapien JA, et al. Cardiorespiratory effects of epidurally administered ketamine or lidocaine in dogs undergoing ovariohysterectomy surgery: a comparative study. Iran J Vet Res 2020; 21(2):92–6; https://doi.org/10.22099/ijvr.2019.34065.5039
[13] Simon BT, Steagall PV. The present and future of opioid analgesics in small animal practice. J Vet Pharmacol Ther 2016; 40(4):315–26; https://doi.org/10.1111/jvpt.12377
[14] Ugwu N, Eze CA, Udegbufun R. Assessment of the analgesic potency of constant rate infusion of tramadol hydrochloride and
as an adjunct to ketoprofen in laparotomized bitches. Sophia J Vet Sci 2017; 15:80–9; https://doi.org/10.4314/sjkys.v15i4.12

[15] Yadav V, Chang D, Holander EM, Bordelon CJ, Kai A, Kaye AD, et al. Ketrorolac, oxymorphone, tapentadolan, and tramadol: A comprehensive review. Anesthesiol Clin 2017; 35(2):e1–20; https://doi.org/10.1016/j.anclin.2017.01.001

[16] Giorgi M. Tramadol vs tapentadol: a new horizon in pain treatment? Am J Anim Vet Sci 2012; 7:7–11; https://doi.org/10.3844/ajavs.2012.7.11

[17] Polati E, Canonic PL, Schweiger V, Collino M. Tapentadol: an overview of the safety profile. J Pain Res 2019; 12:1569–76; https://doi.org/10.2147/jpr.s191545

[18] Barbosa J, Farja J, Querós O, Moreira B, Carvalho F, Diniz-Oliveira RJ. Comparative metabolism of tramadol and tapentadol: a toxicological perspective. Drug Metab Rev 2016; 48(4):577–92; https://doi.org/10.1080/03602532.2016.1229788

[19] Giudice E, Crinó C, Giuseppe B, Barillaro G, Crupi R, Macrì E, et al. Clinical findings in degenerative lumbar spinal stenosis (DLSS) in ten dogs—a pilot study on the analgesic activity of tramadol and gabapentin. J Vet Behav 2019; 33:7–15; https://doi.org/10.1016/j.jveb.2019.05.004

[20] Monteiro BP, Klinkck MP, Moreau M, Guillot M, Steagall PV, Pelleiter JP, et al. Analgesic efficacy of tramadol in cats with naturally occurring osteoarthritis. PLoS One 2017; 12(4):e0175565; https://doi.org/10.1371/journal.pone.0175565

[21] Kögel B, Terlinden R, Schneider J. Characterisation of tramadol, morphine and tapentadol in an acute pain model in Beagles. Vet Anaesth Analg 2014; 41(3):297–304; https://doi.org/10.1111/ava.12140

[22] Teixeira LG, Martins LR, Schimites PL, Dorneles GL, Aiello G, Oliveira JS, et al. Evaluation of postoperative pain and toxicological aspects of the use of dipyramine and tramadol in cats. J Feline Med Surg 2020; 22(6):647–75; https://dx.doi.org/10.1111/jfms.13906

[23] Lorena SE, Luna SP, Lascelles BD, Corrente JE. Current attitudes regarding the use of perioperative analgesics in dogs and cats by Brazilian veterinarians. Vet Anaesth Analg 2014; 1:83–9; https://doi.org/10.1111/vaa.12104

[24] Morales-Vallecilla C, Ramírez N, Villar D, Díaz MC, Bustamante S, Teixeira LG, Martins LR, Schimites PI, Dornelles GL, Aiello G, Oliveira JS, et al. Limited expression of functional cytochrome P450 2C subtypes as chiral selectors and experimental design method optimization. Chem Pap 2019; 25(4):278–86; https://dx.doi.org/10.1007/s11696-019-00789-8

[25] Kögel B, Terlinden R, Schneider J. Pharmacokinetics of tramadol, morphine and tapentadol in an acute pain model in Beagles. Vet Anaesth Analg 2014; 41(3):297–304; https://doi.org/10.1111/ava.12140

[26] Giorgi M. Tramadol vs tapentadol: a new horizon in pain treatment? Am J Anim Vet Sci 2012; 7:7–11; https://doi.org/10.3844/ajavs.2012.7.11

[27] Teixeira LG, Martins LR, Schimites PL, Dorneles GL, Aiello G, Oliveira JS, et al. Evaluation of postoperative pain and toxicological aspects of the use of dipyramine and tramadol in cats. J Feline Med Surg 2020; 22(6):647–75; https://dx.doi.org/10.1111/jfms.13906

[28] Monteiro BP, Klinkck MP, Moreau M, Guillot M, Steagall PV, Edge DK, et al. Analgesic efficacy of an oral transmucosal spray formulation of meloxicam alone or in combination with tramadol in cats with naturally occurring osteoarthritis. Vet Anaesth Analg 2016; 43(6):643–51; https://doi.org/10.1016/j.vaa.12360

[29] Pérez JTE, Mealey KL, Grubb TL, Greene SA, Court MH. Tramadol metabolism to O-desmethyl tramadol (M1) and N-desmethyl tramadol (M2) by dog liver microsomes: species comparison and identification of responsible canine cytochrome P450s. Drug Metab Dispos 2016; 44(12):1963–72; https://doi.org/10.1124/dmd.116.1171902

[30] Kögel B, Terlinden R, Schneider J. Characterisation of tramadol, morphine and tapentadol in an acute pain model in Beagles. Vet Anaesth Analg 2014; 41(3):297–304; https://doi.org/10.1111/ava.12140

[31] Pérez JTE, Mealey KL, Schneider D, Grubb TL, Greene SA, Court MH. Identification of canine cytochrome p450s (CYPs) metabolizing the tramadol (+)-M1 and (+)-M2 metabolites to the tramadol (+)-M5 metabolite in dog liver microsomes. J Vet Pharmacol Ther 2018; 41(6):815–24; https://doi.org/10.1111/jvp.12706

[32] van Hagen MA, Schipper L, Oosterveer-van der Doelen MA, Voos-Loohuis M, Gehring R, Leegwater PA. Analysis of polymorphisms of canine cytochrome P450 CYP2D15. J Vet Pharmacol Ther 2020; 43(6):602–7; https://doi.org/10.1111/jvp.12890

[33] Martinez SE, Pandey AV, Court MH. Isoform-dependent effects of cytochrome p450 oxidoreductase polymorphisms on drug metabolism by cytochrome P450 enzymes in dogs. FASEB J 2019; 33(S1):506–9; https://doi.org/10.1096/fasebj.2019.33.1_supplement.5069

[34] Schüttler AF, Tümsenj J, Kästner SBR. Influence of tramadol on acute thermal and mechanical cutaneous nociception in dogs. Vet Anaesth Analg 2017; 44(2):309–16; https://doi.org/10.1016/j.vaa.2016.02.003

[35] Karakus E, Prinzinger C, Leiting S, Geyer J. Sequencing of the canine cytochrome p450 CYP2C41 gene and genotyping of its polymorphic occurrence in 36 dog breeds. Front Vet Sci 2021; 8:663175; https://doi.org/10.3389/fvets.2021.663175

[36] Overall KL. Behavioral pharmacotherapeutics. Pharmacotherapeutics for veterinary dispensing. John Willey & Sons Inc., Hoboken, NJ, pp 377–401, 2019.

[37] Iñiguez AM, Kimble B, Govendr M. Intrinsic clearance rate of O-desmethyl tramadol (M1) by glucuronide conjugation and phase I metabolism by feline, canine and common brush-tailed possum microsomes. Xenobiotica 2020; 50(7):776–82; https://doi.org/10.1080/00498254.2019.1672718

[38] Ono Y, Sugiyama S, Matushita M, Kitazawa T, Amano T, Uno Y, et al. Limited expression of functional cytochrome P450 2C subtypes in dogs.
in the liver and small intestine of domestic cats. Xenobiotica 2019; 49(6):627–35; https://doi.org/10.1080/00498254.2018.1493543

[46] Lautz LS, Jedli MZ, Girotti MF, Nevela C, Dorne JLM. Metabolism and pharmacokinetics of pharmaceuticals in cats (Felis sylyvestris catus) and implications for the risk assessment of feed additives and contaminants. Toxicol Lett 2021; 338:114–27; https://doi.org/10.1016/j.toxlet.2020.11.014

[47] Cho JH, Choi HG. Acetaminophen and tramadol hydrochloride-loaded soft gelatin capsule: preparation, dissolution and pharmacokinetics in beagle dogs. Pharm Dev Technol 2021; 26(5):576–81; https://doi.org/10.1089/pdt.2021.19030

[48] Ruel HLM, Steagall PV. Adjuvant analgesic in acute pain management. Vet Clin North Am Small Anim Pract 2019; 49(6):1127–41; https://doi.org/10.1016/j.cvsm.2019.07.005

[49] Di Salvo A, Contin MB, Nannarone S, Bufalari A, Giorgi M, Moretti G, et al. Pharmacokinetics and analgesic efficacy of intranasal administration of tramadol in dogs after ovariohysterectomy. Vet Anaesth Analg 2020; 47(4):557–66; https://doi.org/10.1111/j.1467-2995.2020.00121

[50] KuKanich B. Outpatient oral analgesics in dogs and cats beyond nonsteroidal antiinflammatory drugs: an evidence-based approach. Vet Clin North Am Small Anim Pract 2013; 43:1190–2; https://doi.org/10.1016/j.cvsm.2013.04.007

[51] KuKanich B. Pain management in veterinary species. Pharmacotherapeutics for veterinary dispensing. Wiley Blackwell, Hoboken, NJ, pp 173–88, 2019.

[52] Guedes AGP, Meadows JM, Pympendop BH, Johnson EG. Evaluation of barbed suture for celiorrhaphy and subcutaneous closure in bitches with pyometra submitted to ovariohysterectomy. Acta Gr Bras 2021; 36(5):e360502; https://dx.doi.org/10.1590%2FABCB360502

[53] Rossetti RB, Mastrocinque S, Macedo J. Tratamento da dor pós-operatória persistente após ovariohysterectomia em cadelas: estudo comparativo entre tramadol ou maropitant. Revista Interdiscip Saúde Educ 2020; 1(2):109–21. (In Portuguese).

[54] Mastrocinque S, Almeida TF, Tatarunas AC, Imagawa VH, Otsuki DA, Materia JM, et al. Comparison of epidural and systemic tramadol for analgesia following ovariohysterectomy. J Am Anim Hosp Assoc 2012; 48(5):310–9; https://doi.org/10.5326/jaaha-ms-5795

[55] Maksimovic A, Lutvikalad E. Efficiency of epidurally injected lidocaíne, lidocaíne and morphine or lidocaíne and tramadol for postoperative analgesia in dogs following orchietomy and ovariohysterectomy. Int J Vet Sci 2021; 10(1):13–8; http://dx.doi.org/10.47278/journal.ijvs.2020.015

[56] Farokhzad B, Sabiza S, Jalali MR, Baniadam A. Intraperitoneal administration of lidocaíne or tramadol alone or in combination on postoperative pain after ovariohysterectomy in dogs. Vet Med Sci 2021; 7(3):634–41; https://doi.org/10.1016/j.vms.2021.03.034

[57] Ospina-Arquéllas DA, Ramírez CA, Burke E, Echeverry B. Analgesic infusions with tramadol or lidocaíne in bitches undergone to lateral ovariohysterectomy under dissociative anesthesia protocol. Rev Electron Vet 2017; 18(5):1–13. (In Spanish).

[58] Kaka U, Rahman NA, Abubakar AA, Goh YM, Fakurazi S, Omar MA, et al. Pre-emptive multimodal analgesia with tramadol and ketamine-lidocaíne infusion for suppression of central sensitization in a dog model of ovariohysterectomy. J Pain Res 2018; 11:743–52; https://doi.org/10.2147%2FJPR.S152475

[59] Thengchaiasi N, Mahidol C. Evaluating the effects of continuous intravenous infusions of tramadol and tramadol-lidocaíne on sevoflurane minimum alveolar concentration (MAC) and entropy values in dogs. J Vet Med Sci 2019; 81(5):682–8; https://doi.org/10.1292%2Fjvms.19-0448

[60] Budberg SC, Torres BT, Kleine SA, Sandberg GS, Berjeski AK. Lack of effectiveness of tramadol hydrochloride for the treatment of pain and joint dysfunction in dogs with chronic osteoarthritis. J Am Vet Med Assoc 2018; 252(4):427–32; https://doi.org/10.1136%2Fjvet.252.4.427

[61] Benitez ME, Rouch JK, McMurphy R, KuKanich B, Legallet C. Clinical efficacy of hydrocode-acetaminophen and tramadol for control of postoperative pain in dogs following tibial plateau leveling osteotomy. Am J Vet Res 2015; 76(9):755–62; https://doi.org/10.2460/ajvr.76.9.755

[62] Póras LA, Mancusi D, Olimpo M, Gastaldi L, Rosso V, Panaro E, et al. Postoperative analgesia following TPLO surgery: a comparison between cimicoxib and tramadol. Res Vet Sci 2021; 136:351–9; https://doi.org/10.1016/j.rvsc.2021.03.010

[63] Mikes J, Bajjesen J, Christensen P, Andersen-Ranberg E, Vítgy A, Poulsen HJ, et al. Tramadol and gabapentin improve peak vertical force in osteoarthritic dogs already receiving non-steroidal anti-inflammatory drugs. In BSAVA Congress Proceedings 2020. BSAVA Library, p 535, 2020; http://dx.doi.org/10.22233/9781910443774.87.3

[64] Malek S, Sample SJ, Schwartz Z, Nemke B, Jacobson PB, Cozzi EM, et al. Comparison of oral and intravenous pain management with tramadol in client-owned dogs with hip osteoarthritis. BMC Vet Res 2012; 8(1):1–17; https://doi.org/10.1186%2F1746-6148-8-185

[65] Śliwka P, Martyniuk A, Maciak D, Piotrowska O, Piotrowski R. Postoperative pain in dogs undergoing hemilaminectomy: comparison of the analgesic activity of buprenorphine and tramadol. J Vet Behav 2017; 19:45–9; http://dx.doi.org/10.1016/j.jveb.2017.02.003

[66] Ripplinger A, Aiello G, Chaves RO, Andrades AO, Beckmann DV, Polidoro D, et al. Efeitos adversos da morfina, metadona e tramadol no pós-operatório de cães submetidos à cãrumia da...
columna vertebral: 180 casos (2011–2016). Pesqui Vet Bras 2018; 38:1431–7; https://doi.org/10.1590/1678-5150-PVB-5307 (in Portuguese).

[75] Karrasch NM, Lerce P, Aarnes TK, Gardner HL, London CA. The effects of preoperative oral administration of carprofen or tramadol on postoperative analgesia in dogs undergoing cutaneous tumor removal. Can Vet J 2015; 56(8):817–22.

[76] Uscategi RAR, Tiosso C, Moro JV, Mostachio GQ, Padilha-Nakaghi LC, Feliciano MAR, et al. Pre-emptive methadone or tramadol analgesia for mastectomy and ovariomyectomy in bitches. Rev Colomb Genc Pecu 2017; 30:39–47. (in Spanish).

[77] Reis HA, Mangabeira RO, Coelho APG, Costa RB, Barbosa VF. Influência do tramadol associado à levobupivacaina epidural sobre a taxa de procoletiva variáveis fisiológicas, em cadelas submetidas à mastectomia e à ovari-histerectomia. Arq Bras Med Vet Zootec 2020; 72(5):1639–45; https://doi.org/10.1590/1678-4162-11199 (in Portuguese).

[78] Read K, Khuton M, Murphy H. Comparison of transdermal fentanyl and oral tramadol for lateral thoracotomy in dogs: cardiovascular and behavioural data. Vet Anaesth Analg 2019; 46(1):116–25; https://doi.org/10.1111/eva.12641.

[79] Jiménez TEP, Kukanich B, Joo H, Mealey KL, Grubb TL, Greene SA. Oral coadministration of fluconazole with tramadol markedly increases plasma and urine concentrations of tramadol and the O-desmethyltramadol metabolite in healthy dogs. Drug Metab Dispos 2019; 47(1):15–25; https://doi.org/10.1124/dmd.118.083444.

[80] Simon BT, Steagall PV. Feline procedural sedation and analgesia: when, why and how. J Feline Med Surg 2020; 22(11):1029–45; https://doi.org/10.1080/10986118.2019.1685880.

[81] Steagall PV. What makes cats different/challenging and what is critical for cats? Clin North Am Small Anim Pract 2020; 35(4):749–67; https://doi.org/10.1016/j.cvsm.2020.02.002.

[82] Muñoz-Rodríguez L, Santisteban-Arena R, Rios-Torres M, Rios-Ceballos V. Evaluación del dolor postoperatorio en felinos sometidos a columna vertebral: 180 casos (2011–2016). Pesqui Vet Bras 2018; 38(3):1–6; https://doi.org/10.1590/1678-5150-PVB-5307 (in Portuguese).

[83] Li L, Dong J, He J, Cui J, Yu X, Tan D, et al. Anaesthetic effects of tiletamine-zolazepam-xylazine-tramadol combination in cats under going surgical sterilization. Acta Vet Brno 2015; 4:181–5; https://doi.org/10.1292/jvms.2015.8018.

[84] Aarnes TK, Doodnaught GM, Evangelista MC, Steagall PV. Pain. Eur J Pain 2019; 23(6):1185–95; https://doi.org/10.1002/ejp.1386.

[85] Fricová J, Lainczová H, Nedvídek J, Rokyta R. Effect of tapentadol on postoperative analgesia in dogs undergoing cutaneous tumor removal. Can Vet J 2020; 61(3):289–93; https://doi.org/10.1177/0008412119855809.

[86] Kieves NR, Howard J, Lerche P, Lakritz J, Aarnes TK. Effectiveness of tapentadol hydrochloride for treatment of orthopedic pain in dogs: a pilot study. Can Vet J 2020; 61(3):289–93; https://doi.org/10.1177/0008412119855809.

[87] Torres-Sanchez S, Borges GDS, Mico JA, Berrocoso E. Opioid and noradrenergic contributions of tapentadol to the inhibition of locus coeruleus. Physiol Res 2021; 69(Suppl 3):533–7; https://doi.org/10.1007/s10105-021-01515-z.

[88] Lee HK, Lebrowska-Wieruszewska B, Kim TW, Kowsaki CJ, Giorgi M, Meieker A, Mills PC. Pharmacokinetics of the novel atypical opioid tapentadol following oral and intravenous administration of tapentadol hydrochloride in dogs. Am J Vet Res 2018; 79(4):367–75; https://doi.org/10.1177/0361122217737101.

[89] Bouvier LI, Gutiérrez DPF, Kemper BK, Pereira LI, Kemper DAG. Comparação da ação analgésica do emprego sistêmico de tramadol e morfina em gatas submetidas à ovarião-histerectomia. In Congresso Medvet Internacional de Medicina Felina, pp 41–3. 2020.

[90] Lemos VCC, Sande JQ, Barbosa VF, Costa Neto JM, Martins Filho IF, Iwassa CHD. Avaliação da dexmedetomidina e do tramadol, associados ao midazolam, em gatas anestesiadas com isoflurano e isovolárico-histerectomia. Arq Bras Med Vet Zootec 2017; 69(6):1521–8; https://doi.org/10.1590/1678-4162-9426.

[91] Nascimento FF, Marques VI, Crociolli GC, Nicácio GM, Nicácio PAG, Lainczová H, et al. Effect of tapentadol on postoperative analgesia in dogs undergoing mastectomy and ovariohysterectomy. Rev Bras Med Vet Zootec 2018; 39(3):404–22; https://doi.org/10.1590/2427-1792.

[92] Fricová J, Lainczová H, Nedvídek J, Rokyta R. Effect of tapentadol on postoperative analgesia in dogs undergoing mastectomy and ovariohysterectomy. J Adv Vet Anim Res. 2020; 8(3): 404–422, September 2021.
[106] Zapata-Morales JR, Alonso-Castro AI, Granados-Soto V, Sánchez-Enriquez S, Isiordia-Espinoza MA. Assessment of the antinociceptive and ulcerogenic activity of the tapentadol-diclofenac combination in rodents. Drug Dev Res 2018; 79(1):38–44; https://doi.org/10.1002/ddr.21420

[107] Duvall A, Tully Jr TN, Carpenter JW, KuKanich B, Beaufrière H, Magnin GC. Pilot study of a single dose of orally administered tapentadol suspension in Hispaniolan Amazon parrots (Amazona ventralis). J Avian Med Surg 2021; 35(1):45–50; https://doi.org/10.1647/1082-6742-35.1.45

[108] De Vito V, Owen H, Marzoni M, Kim TW, Poapolathep A, Giorgi M. Pharmacokinetics of tapentadol in laying hens and its residues in eggs after multiple oral dose administration. Br Poult Sci 2017; 59(1):128–33; https://doi.org/10.1080/00071668.2017.1401705

[109] Giorgi M, De Vito V, Owen H, Demontis MP, Varoni MV. PK/PD evaluations of the novel atypical opioid tapentadol in red-eared slider turtles. Med Weter 2014; 70(9):530–5.

[110] Giorgi M, Salvadori M, De Vito V, Owen H, Demontis MP, Varoni MV. Pharmacokinetic/pharmacodynamic assessments of 10 mg/kg tramadol intramuscular injection in yellow-bellied slider turtles (Trachemys scripta scripta). J Vet Pharmacol Ther 2015; 38(5):488–96; https://doi.org/10.1111/jvp.12206

[111] Faria J, Barbosa J, Leal S, Afonso LP, Lobo J, Moreira R, et al. Effective analgesic doses of tramadol or tapentadol induce brain, lung and heart toxicity in Wistar rats. Toxicology 2017; 385:38–47; https://doi.org/10.1016/j.tox.2017.05.003

[112] Lavy E, Lee HK, Mahbesh Sf, Sebastian C, Baker Y, Giorgi M. Use of the novel atypical opioid tapentadol in goats (Capra hircus) pharmacokinetics after intravenous, and intramuscular administration. J Vet Pharmacol Ther 2014; 37(5):518–21; https://doi.org/10.1111/jvp.12123

[113] Giorgi M, Mills PC, Tayari H, Rota S, Bregghi G, Briganti A. Plasma concentrations of tapentadol and clinical evaluations of a combination of tapentadol plus sevoflurane for surgical anaesthesia and analgesia in rabbits (Oryctolagus cuniculus) undergoing orchietomy. Isr J Vet Med 2013; 68(3):141–8.

[114] Berry SH. Analgesia in the perioperative period. Vet Clin North Am Small Anim Pract 2015; 45(5):1013–27; https://doi.org/10.1016/j.cvsm.2015.04.007

[115] Jardim MPB, Farias LF, Cid GC, Souza, HJM. Poisoning in domestic cats in Brazil: toxicants, clinical signs, and therapeutic approaches. Arq Bras Med Vet Zootec 2021; 73(01):99–107; https://doi.org/10.1590/1678-4162-11856

[116] Tinson E, Cook S. Supporting the intoxicated patient: toxicants affecting the neurological and cardiovascular systems. In Pract 2020; 42(1):27–38; http://dx.doi.org/10.1136/inp.7080

[117] Barakat, A. Revisiting tramadol: a multimodal agent for pain management. CNS Drugs 2019; 33:81–90; https://doi.org/10.1007/s40263-019-00623-5

[118] Indrawirawan Y, McAlees T. Tramadol toxicity in a cat: case report and literature review of serotonin syndrome. J Feline Med Surg 2014; 16(7):572–8; https://doi.org/10.1177/1098612x14539088

[119] James NK, Wismer TA, Diniz PPVP. Duloxetine ingestion in 364 dogs. J Am Vet Med Assoc 2019; 255(10):1161–6; https://doi.org/10.2460/javma.255.10.1161

[120] Hassamal S, Miotto K, Dale W, Danovitch I. Tramadol: understanding the risk of serotonin syndrome and seizure. Am J Med 2018; 131(11):1382.e1–6; https://doi.org/10.1016/j.amjmed.2018.04.025