In dogs with metaldehyde intoxication, are benzodiazepines more effective than methocarbamol in relaxing muscles and reducing tremors?

A Knowledge Summary by

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KNOWLEDGE SUMMARY

The term tremor used throughout the paper has been more recently replaced by the term twitches which is acknowledged by the author (Lowrie & Garosi, 2016).

PICO question
In dogs with metaldehyde intoxication, are benzodiazepines (e.g. diazepam, midazolam) more effective than methocarbamol in relaxing muscles and reducing tremors?

Clinical bottom line

Category of research question
Treatment

The number and type of study designs reviewed
Five papers were critically reviewed. There were five retrospective case series

Strength of evidence
Weak

Outcomes reported
Currently, five retrospective case series exist in the literature which discuss metaldehyde intoxication cases treated mainly with benzodiazepines, a few of which had methocarbamol. There is not really any study to compare directly benzodiazepines with methocarbamol. In addition to that, factors such as commercial (e.g. the low availability of methocarbamol in the UK market compared to the US market), administrational (e.g. multiple administration routes of benzodiazepines) and pharmacological (e.g. lack of anticonvulsant function of methocarbamol), have played an important role in the treatment choice. Several case reports exist as well

Conclusion
Currently, there is insufficient evidence to determine whether benzodiazepines are more effective than methocarbamol in relaxing muscles and reducing occurrence of muscle tremors

How to apply this evidence in practice
The application of evidence into practice should take into account multiple factors, not limited to: individual clinical expertise, patient’s circumstances and owners’ values, country, location or clinic where you work, the individual case in front of you, the availability of therapies and resources.

Knowledge Summaries are a resource to help reinforce or inform decision-making. They do not override the responsibility or judgement of the practitioner to do what is best for the animal in their care.
**Clinical Scenario**
A 4-year-old male neutered dog is presented to you as an emergency due to acute ongoing generalised muscle tremors. Prior to the episode, the dog was out for a walk at the neighborhood. Physical examination reveals hyperthermia (40.5°C), neurological examination reveals generalised muscle tremors, however the dog is bright, alert and responsive. Based on generalised muscle tremors and absence of other neurological findings, you suspect that the hyperthermia is secondary to the tremors, and you neurolocalise forebrain, cerebellum, meninges (pyrexia), peripheral nerve or multifocal, as it is difficult to clinically establish the origin of generalised muscle tremors. You observe some watery discharge of blue/green colour from the anus, compatible with the colour of the commercial form of slug bait, and thus you suspect metaldehyde intoxication. Would you choose methocarbamol or benzodiazepines to relax the muscles and reduce the tremors of the dog?

**The evidence**
Five studies of indirect relevance to the PICO were reviewed, all of them being retrospective in nature. Due to a lack of prospective or retrospective studies with direct correlation of benzodiazepines and methocarbamol treatment without administration of other medications (e.g. antiepileptic drugs), the strength of the evidence is extremely low.

**Summary of the evidence**

| Firth (1992) |
|-------------|
| **Population:** | Dogs with snail bait poisoning (metaldehyde or methiocarb) and follow-up. This study was conducted in Australia |
| **Sample size:** | 56 dogs |
| **Intervention details:** | • 26/56 dogs were intoxicated by metaldehyde (30/56 dogs were intoxicated by methiocarb)  
• There is no information whether the intoxicated dogs manifested epileptic seizures, tremors or both  
• There is no information whether the dogs were amenable or not to administration of oral medications upon presentation  
• Treatment was achieved with sedatives, general anaesthetics and/or muscle relaxants including:  
  a. diazepam premedication (9/26)  
  b. diazepam/ketamine general anaesthesia (GA) (12/26)  
  c. diazepam/ketamine/lidocaine GA (7/26)  
  d. lidocaine/ketamine GA (1/26)  
  e. methocarbamol (post-GA) (6/26)  
• Methocarbamol was given only after performing GA with either one of the above mentioned GA protocols, but not as sole medication  
• Two dogs were lost to follow-up |
| **Study design:** | Retrospective, single centre, case series |
| **Outcome studied:** | • Different management protocols (including the use of premedication, general anaesthesia and post-general anaesthesia relaxants)  
• The correlation between the treatment modality and patients’ response and outcome |
**Main findings: (relevant to PICO question):**
There was full recovery of 100% (24/24) metaldehyde intoxication cases using multimodal treatment, most of which included diazepam. However, there was no comparison between the usage of benzodiazepines and methocarbamol.

**Limitations:**
- This is a retrospective, single centre, case series study with a low level of evidence.
- The usage of methocarbamol was additional to a GA protocol that might or might not include diazepam.
- The outcome was not correlated with the specific treatment that each dog had.

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**Yas-Natan et al. (2007)**

**Population:**
Dogs diagnosed with metaldehyde. This study was conducted in Israel (School of Veterinary Medicine, The Hebrew University of Jerusalem).

**Sample size:**
18 dogs

**Intervention details:**
- All cases were intoxicated by metaldehyde and presented with a variety of clinical signs.
- 16/18 dogs presented with epileptic seizures and 10/18 dogs presented with muscle tremors amongst other clinical signs.
- Only 2/18 dogs were amenable to administration of oral medications upon presentation as they were not presented with *status epilepticus* or altered mentation. Nevertheless, only injectable medications were administered within the study.
- Monotherapy (6/18 dogs) or multimodal treatment was administered (12/18 dogs) including one, or more than one, of the medications below:
  a. diazepam (17/18)
  b. phenobarbital (7/18)
  c. pentobarbital (6/18)
  d. isoflurane (9/18)
- Diazepam was administered in a dose of 0.28–6.3 mg/kg IV q24h.

**Study design:**
Retrospective, single centre, case series

**Outcome studied:**
- Clinical signs and clinicopathological findings.
- Different management protocols.
- The correlation between the treatment modality and patients’ response and outcome.

**Main findings: (relevant to PICO question):**
Dogs with metaldehyde intoxication which were mostly treated with multimodal treatment including diazepam had overall a good outcome (survival rate of 83%).
### Limitations:
- This is a retrospective, single center, case-series study with a low level of evidence.
- This is a study which does not concentrate on the treatment. No information provided for the type of treatment in conjunction with the outcome.
- The treatment of the dogs with metaldehyde intoxication is multimodal, and the majority of dogs were treated with barbiturates. As the barbiturates are successful antiepileptic drugs, this makes the conclusion of diazepam efficacy unreliable.

### Zimmermann et al. (2010)

| Population: | Dogs with *status epilepticus* due to acute intoxications. This study was conducted in Germany (Ludwig Maximilian University of Munich) |
| Sample size: | 14 dogs (three of them diagnosed with metaldehyde intoxication) |
| Intervention details: | • 3/3 cases were intoxicated by metaldehyde and presented with *status epilepticus*  
• 0/3 presented with muscle tremors  
• None of the dogs (0/3) were amenable to administration of oral medications as per *status epilepticus* upon presentation  
• All metaldehyde intoxicated dogs were administered phenobarbital alone or with other medications, whilst two of them were treated with diazepam. Specifically:  
  a. dog 1: phenobarbital, pentobarbital  
  b. dog 2: diazepam, acepromazine, atropine, phenobarbital, pentobarbital  
  c. dog 3: lidocaine, diazepam, propofol, pentobarbital  
• For dog 2, diazepam was administered initially 0.5–1.0 mg/kg IM and then IV, whilst for dog 3 only IV |
| Study design: | Retrospective, single centre, case series |
| Outcome studied: | • Describe intoxication aetiology for dogs with *status epilepticus*  
• Describe clinical presentation  
• Different management protocols  
• The correlation between the treatment modality and patients’ response and outcome |
| Main findings: (relevant to PICO question): | • Dogs with metaldehyde intoxication which were treated in hospital with phenobarbital (2/3), pentobarbital (3/3), diazepam (2/3) and other medications survived  
• All dogs were discharged with oral antiepileptic treatment (phenobarbital), which was tapered gradually until discontinuation for a total period of 4 weeks post-discharge |
Telephone follow-up was done for all dogs in the study (median follow-up time 2.6 years); the three dogs with metaldehyde intoxication were alive with no further seizures

**Limitations:**
- This is a retrospective, single centre, case series study with a low level of evidence.
- This study is a general intoxication study, which does not concentrate specifically on metaldehyde intoxication. Thus, the case number is very low (n=3), where only 2/3 have been treated with diazepam
- The treatment of the dogs with metaldehyde intoxication is multimodal, and particularly all dogs are treated with pentobarbital or phenobarbital. As phenobarbital and pentobarbital are of the barbiturate family, they are very successful antiepileptic drugs, the conclusion of diazepam efficacy is unreliable
- The whole study does not provide detailed findings for every individual case (follow-up time, outcome, etc.) making the strength of evidence even lower

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**Jull et al. (2011)**

**Population:** Dogs with *status epilepticus* due to acute intoxications. This study was conducted in the UK (Royal Veterinary College, Animal Health Trust)

**Sample size:** 20 dogs (17 cases were intoxicated by metaldehyde and presented with *status epilepticus*)

**Intervention details:**
- All dogs (17/17) had epileptic seizures (*status epilepticus*), however, it is not stated whether some of these dogs had muscle tremors as well
- None of the dogs (0/17) were amenable to administration of oral medications as per *status epilepticus* upon presentation
- Metaldehyde intoxicated dogs received monotherapy or multimodal therapy. More specifically:
  a. 8/17: diazepam (monotherapy)
  b. 3/17: diazepam + phenobarbital
  c. 1/17: diazepam + propofol
  d. 1/17: midazolam (monotherapy)
  e. 2/17: midazolam + propofol
  f. 2/17: phenobarbital + propofol

**Study design:** Retrospective, multi-centre (two centres), case series

**Outcome studied:** Whether prolonged *status epilepticus*, secondary to a chemoconvulsant, can induce spontaneous recurrent seizures in dogs
Main findings: (relevant to PICO question):

- Dogs with metaldehyde intoxication which were treated with benzodiazepine monotherapy (diazepam or midazolam) survived and did not manifest any post intoxication seizures
- 3/17 dogs of the metaldehyde intoxication group were discharged with oral antiepileptic treatment (phenobarbital) which was tapered gradually until discontinuation
- Median follow-up time for the 20 dogs was 757 days. The 17 dogs with metaldehyde intoxication survived and none of them manifested any post intoxication seizures
- All metaldehyde intoxicated dogs survived

Limitations:

- This is a retrospective, multi-centre, case series study with a low level of evidence
- The multi-center nature of the study increases the possibility of non-standardised protocols between the centres, and thus the strength of the study
- There was no case treated with methocarbamol
- Due to the nature of the study, no direct comparison between the different treatments can be reliably assumed

Bates et al. (2012)

Population: Dogs with suspected metaldehyde intoxication (slug bait poisoning) with follow-up; cases reported to the Veterinary Poisons Information Service (VPIS) through phone calls by veterinary practices. This study was conducted in the UK (1985–2010)

Sample size: 772 dogs

Intervention details:

- 597/772 dogs were symptomatic
- Only 528/597 dogs developed increased muscular activity, such as tremor, twitching, muscle spasms or fasciculation, epileptic seizures or opisthotonos
- 290/597 dogs were presented with convulsions and 136/597 with tremors, whilst the rest had a variety of other neurological or extraneural signs
- There is no information whether the dogs were amenable or not to administration of oral medications upon presentation
- Treatment, on either symptomatic or asymptomatic intoxicated dogs was achieved with one, or more than one, of the medications below:
  - 392/772 benzodiazepines
  - 227/772 barbiturates
  - 90/772 propofol
  - 70/772 acepromazine
  - 4/772 isoflurane
  - 2/772 methocarbamol
  - 1/772 ketamine
| Study design: | Retrospective, multi-centre, case series |
|--------------|----------------------------------------|
| Outcome studied: | • To analyse retrospectively telephone enquiries of referring veterinarians who confront dogs poisoned by metaldehyde  
• Describe clinical features  
• The correlation between the treatment modality and patients’ response and outcome |
| Main findings: (relevant to PICO question): | • Benzodiazepines remain a major option for metaldehyde intoxication treatment for the 50% of the cases, however there is evidence that refractory cases require further medications/anaesthetics  
• The use of barbiturates and benzodiazepines remained fairly constant over the period examined  
• Benzodiazepines were given to half the dogs in this cases series  
• Methocarbamol was used only in two cases most likely as a result of the decreased availability in the UK |
| Limitations: | • This is a retrospective, multi-centre, case series study with a low level of evidence  
• The multi-centre and questionnaire based nature of the study increases the possibility of non-standardised protocols between the centres, and thus the strength of the study  
• There were only two cases treated with methocarbamol, with no detailed reference as to the outcome  
• The outcome was not correlated with the specific treatment that each dog had, making any evaluation of the benzodiazepine or methocarbamol efficacy to each case impossible |

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### Appraisal, application and reflection

Metaldehyde intoxication is a common intoxication in dogs attributed to ingestion of slug bait, which consists of the carbamate named metaldehyde. Clinical signs include generalised muscle tremors and/or epileptic seizures, as well as a variety of other signs (Dolder, 2003). Among others (e.g. metabolic acidosis), one of the major causes of death in such cases is the hyperthermia secondary to the excessive generalised muscle tremors (Dolder, 2003). Consequently, one of the major therapeutic goals of the general practitioner is to decrease the muscle tremors, avoiding hyperthermia. As mentioned already, there are no prospective or retrospective studies in the literature to compare different treatments for metaldehyde intoxication in dogs focusing on benzodiazepines and methocarbamol.
Benzodiazepines bind to γ-aminobutyric acid (GABA) receptors of the brain resulting in increased GABA activity, which is the main neurotransmitter of the brain. Consequently, they are used as centrally acting skeletal muscle relaxants, but also as anxiolytics, sedatives, hypnotics and anticonvulsants (Podell, 1995; and Van Tulder et al., 2003). Benzodiazepines are quite beneficial as most of them can be administered through different routes (intravenous [IV], intramuscular [IM], per os [PO], intranasal [IN], intrarectal [IR]) (Podell, 1995; and Charalambous et al., 2017). Diazepam reaches therapeutic plasma levels within 10 minutes when administered IN or IV (Musulin et al., 2011) or IR (Papich & Alcorn, 2007) or 30 minutes to 2 hours when administered PO (Plumb, 2008). Diazepam IM has a slower and incomplete absorption (Plumb, 2008). The serum half-life of diazepam in dogs is 2.5–3.2 hours (Plumb, 2008). Diazepam’s major drawbacks include: (a) possible cause of contradictory response (central nervous system excitement) (Plumb, 2008); (b) sedative inefficacy (Plumb, 2008); (c) tolerance to its anticonvulsant effect in dogs (Frey et al., 1984); and (d) inability to administer as a constant-rate infusion (CRI) solution as its availability might be reduced within the plastic syringe (Cloyd et al., 1980). Midazolam’s unique solubility characteristics (water soluble injection but with high lipophilicity at body pH) give it a very rapid onset of action after injection (Plumb, 2008). Although midazolam IV provides the quickest onset of action (Plumb, 2008), IN route provides superiority when the time needed to place an IV catheter is taken into account and same efficacy (Charalambous et al., 2019). Midazolam IM is rapidly and completely absorbed, in contrast with diazepam IM. Midazolam PO is not commercially available, whilst midazolam IR is not clinically useful due to very low rectal bioavailability. Compared to diazepam, midazolam is nearly 3 times as potent, and has a faster onset of action (in humans 30–97 seconds), but a shorter duration of effect. Midazolam can also provide sedation if used with opioids, in contrast to diazepam (Plumb, 2008). As a take-home medication, recently, midazolam IN revealed to be superior to diazepam IR for status epilepticus (Charalambous et al., 2017). Midazolam’s major drawbacks include: (a) dose-dependence on plasma protein concentrations (as it is protein binding); (b) shorter serum half-life (within almost an hour) compared to diazepam, and therefore necessity for a CRI; and (c) respiratory depression when used with other narcotics (e.g. opioids) (Plumb, 2008).

Methocarbamol is a centrally acting muscle relaxant that selectively blocks polysynaptic reflex pathways in the spinal cord without any effect on monosynaptic pathways, whilst it has no direct effect on the contractile mechanism of the striated muscle, the nerve fibre or the motor end plate (Van Tulder et al., 2003; and Nielsen et al., 2005). It has been used in veterinary medicine in traumatic myopathies or intoxications (including tetanus) (Nielsen et al., 2005). Oral tablets are the only commercially available form of methocarbamol, although it can be prepared in an off-label enema in hospital. Methocarbamol has an onset of action of about 30 minutes after oral administration. Its peak levels in humans occur approximately 2 hours after dosing, and its serum half-life is about 1–2 hours (Plumb, 2008). In the US, methocarbamol IV is available as well, and successful management of tremors has been reported with methocarbamol CRI in cats (Draper et al., 2013). Methocarbamol’s major drawbacks include: (a) limited routes of administration in combination with availability limited to the oral form in Europe; (b) delayed onset of action compared to benzodiazepines IV; and (c) central nervous system depressant effects as a carbamate (sedation, salivation, lethargy, weakness, ataxia) (Plumb, 2008).

Most of the above mentioned retrospective studies include benzodiazepines and particularly diazepam as one of the most common first-line drugs for the treatment of metaldehyde intoxication. Firth (1992) reported metaldehyde intoxicated dogs treated with diazepam or methocarbamol. Both canine groups were treated with diazepam or methocarbamol as a part of a multimodal treatment which included additionally a general anaesthetic. All dogs recovered, but no comparison between the groups can be made for the efficacy of either diazepam or methocarbamol. Yas-Natan et al. (2007) described cases of metaldehyde intoxication treated with benzodiazepines, most of which were accompanied by barbiturates (phenobarbital or pentobarbital). Due to the administration of the above mentioned antiepileptic drugs, no conclusion can be made about the diazepam only efficacy to these patients, whilst no case with methocarbamol treatment is described. Zimmerman et al. (2010) treated all three metaldehyde intoxication cases with diazepam followed by barbiturates amongst other medications, with an aim to control the status epilepticus. All cases recovered,
however due to the multimodal nature of treatment, no conclusion could be made for the diazepam only efficacy. Jull et al. (2011) described similar therapeutic protocols.

Bates et al. (2012) described that general practitioners preference to use of benzodiazepines (392/772 cases) among other treatment protocols as either monotherapy or multimodal therapy, with barbiturates being used frequently (227/772 cases). Only 2/772 cases were reported to have used methocarbamol in the therapeutic protocol.

In practice, the vast majority of dogs suspected to be intoxicated by metaldehyde are presented with epileptic seizures (e.g. status epilepticus) and/or generalised muscle tremors. At the time of presentation, the general practitioner is not able to distinguish the origin of the clinical signs, and given the emergency nature of these cases, injectable benzodiazepines (and specifically diazepam) are the first choice. Injectable benzodiazepines offer rapid onset of action and have both antiepileptic and muscle relaxant properties. Additionally, both generalised muscle tremors and epileptic seizures usually include motor activity of the facial and masticatory muscles and thus jaw movements, which makes any oral administration unsafe for the veterinary surgeon. Therefore, these reasons, as well as the restricted administration routes of methocarbamol, could probably explain why the vast majority of studies include primarily benzodiazepines rather than methocarbamol.

Conclusions

In conclusion, there is not enough evidence to define whether benzodiazepines (e.g. diazepam, midazolam) or methocarbamol is better for the control of muscle tremors during metaldehyde intoxication, thus the answer of the current PICO remains open. Although it is reported that the availability of methocarbamol is limited in the UK (Bates et al., 2012) and there are no prospective studies describing its efficacy on tremors, it is suggested that methocarbamol is very successful in reducing muscle tremors during this intoxication (Dolder, 2003). Due to possible manifestation of epileptic seizures concurrently with the generalised muscle tremors and in the light of their anticonvulsant activity, their broader availability, their multiple administration routes and their rapid action when given IV, benzodiazepines are preferred for the initiation of the treatment in cases of metaldehyde intoxication by many vets; and they carry on with an antiepileptic drug (e.g. phenobarbital) or general anaesthesia (Firth, 1992; Yas-Natan et al., 2007; Zimmermann et al., 2010; Jull et al., 2011; and Bates et al., 2012). It is important to note that in the decision-making process, apart from the pharmacological features of each medication, all points of care should be taken into consideration such as: (a) best practice: each patient should be treated with the best practice that would be to treat the dog immediately with the faster acting drug; (b) the patient stress factor: that is no oral medications should be administered in a patient with risk of regurgitation or distress; and (c) safety of the staff: that is risks that could arise from administration of oral medications in a dog with generalised muscle tremors (including the jaw). Further studies are necessary to provide information on the efficacy of benzodiazepines or methocarbamol in patients with metaldehyde intoxication.
## Methodology Section

### Search Strategy

| Databases searched and dates covered: | CAB Abstracts, 1973 to 2019 week 25  
                                      | PubMed, 1966 to current               |
|--------------------------------------|-------------------------------------|
| Search terms:                        | **CAB Abstracts:**                  |
|                                      | 1. (dog OR dogs OR canine OR canines OR canis OR bitch OR bitches OR puppy OR puppies OR pup OR pups) OR exp dogs/ OR exp bitches/ OR exp puppies/ OR exp canidae/ OR exp canis/  
|                                      | 2. metaldehyde.mp. OR exp metaldehyde/ OR 'slug bait'.mp. OR tremorgenic  
|                                      | 3. (intoxication OR toxic* OR poison* OR toxicosis) OR exp toxicity/ OR exp poisoning/             |
|                                      | 4. 1 AND 2 AND 3                     |
|                                      | **PubMed:**                         |
|                                      | 1. dog OR dogs OR canine OR canines OR canis OR bitch OR bitches OR puppy OR puppies OR pup OR pups  
|                                      | 2. metaldehyde OR slug bait OR tremorgenic  
|                                      | 3. intoxication OR toxic* OR poison* OR toxicosis                                                    |
|                                      | 4. 1 AND 2 AND 3                     |
| Dates searches performed:            | 2 July 2019                          |

### Exclusion / Inclusion Criteria

| Exclusion: | Articles not available in English, articles which were not relevant to the PICO question. book chapters, literature reviews, single case reports, conference proceedings |
| Inclusion: | Original peer-reviewed articles in English language with more than one dog intoxicated by metaldehyde and treated with benzodiazepines and/or methocarbamol |
| Database   | Number of results | Excluded – non-English language | Excluded – not relevant with the PICO | Excluded – book chapters | Excluded – reviews | Excluded – case reports | Excluded – conference proceedings | Total relevant papers |
|------------|-------------------|---------------------------------|--------------------------------------|--------------------------|-------------------|------------------------|-------------------------------|----------------------|
| CAB Abs    | 113               | 16                              | 71                                   | 2                        | 11                | 5                      | 3                            | 5                    |
| PubMed     | 52                | 0                               | 36                                   | 0                        | 10                | 1                      | 0                            | 5                    |

Total relevant papers when duplicates removed 5

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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