ACEPROMAZINE MALEATE- acepromazine maleate injection Covetrus North America

Acepromazine maleate Injection

Approved by FDA under NADA # 015-030

Caution:

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

Description:

Acepromazine maleate injection, a potent neuroleptic agent with a low order of toxicity, is of particular value in the tranquilization of dogs, cats and horses. Its rapid action and lack of hypnotic effect are added advantages. According to Baker, the scope of possible applications for this compound in veterinary practice is only limited by the imagination of the practitioner.

Each mL contains: Acepromazine maleate 10 mg, sodium citrate 0.36%, citric acid 0.075%, benzyl alcohol 1% and water for injection.

Acepromazine [10-[3-(dimethylamino) propyl] phenothiazin-2-yl-methyl ketone] Maleate, USP has the following chemical structure:

Acepromazine maleate injection has a depressant effect on the central nervous system and, therefore, causes sedation, muscular relaxation and a reduction in spontaneous activity. It acts rapidly, exerting a prompt and pronounced calming effect.

Indications:

Dogs and cats: Acepromazine maleate injection can be used as an aid in controlling intractable animals during examination, treatment, grooming, x-ray and minor surgical procedures; to alleviate itching as a result of skin irritation; as an antiemetic to control vomiting associated with motion sickness.

Acepromazine maleate injection is particularly useful as a preanesthetic agent (1) to enhance and prolong the effects of barbiturates, thus reducing the requirements for general anesthesia; (2) as an adjunct to surgery under local anesthesia.

Horses: Acepromazine maleate injection can be used as an aid in controlling fractious animals during examination, treatment, loading and transportation. Particularly useful

when used in conjunction with local anesthesia for firing, castration, neurectomy, removal of skin tumors, ocular surgery and applying casts.

Dosage and administration:

The dosage should be individualized, depending upon the degree of tranquilization required. As a general rule, the dosage requirement in mg/lb of body weight decreases as the weight of the animal increases.

Acepromazine maleate injection may be given intravenously, intramuscularly or subcutaneously. The following schedule may be used as a guide to IV, IM or SC injections.

Dogs: 0.25–0.5 mg/lb of body weight **Cats:** 0.5–1 mg/lb of body weight

Horses: 2-4 mg/100 lb of body weight

IV doses should be administered slowly, and a period of at least 15 minutes should be allowed for the drug to take full effect.

Contraindications:

Phenothiazines may potentiate the toxicity of organophosphates and the activity of procaine hydrochloride. Therefore, do not use Acepromazine maleate injection to control tremors associated with organic phosphate poisoning. Do not use in conjunction with organophosphorus vermifuges or ectoparasiticides, including flea collars. Do not use with procaine hydrochloride.

Warning:

Do not use in horses intended for human consumption.

Precautions:

Tranquilizers are potent central nervous system depressants and they can cause marked sedation with suppression of the sympathetic nervous system. Tranquilizers can produce prolonged depression or motor restlessness when given in excessive amounts or when given to sensitive animals.

Tranquilizers are additive in action to the actions of other depressants and will potentiate general anesthesia. Tranquilizers should be administered in smaller doses and with greater care during general anesthesia and also to animals exhibiting symptoms of stress, debilitation, cardiac disease, sympathetic blockade, hypovolemia or shock. Acepromazine maleate injection, like other phenothiazine derivatives, is detoxified in the liver; therefore, it should be used with caution in animals with a previous history of liver dysfunction or leukopenia.

Hypotension can occur after rapid intravenous injection causing cardiovascular collapse.

Epinephrine is contraindicated for treatment of acute hypotension produced by phenothiazine-derivative tranquilizers since further depression of blood pressure can

occur. Other pressor amines, such as norepinephrine or phenylephrine, are the drugs of choice.

In horses, paralysis of the retractor penis muscle has been associated with the use of phenothiazine-derivative tranquilizers. Such cases have occurred following the use of Acepromazine maleate injection.

This risk should be duly considered prior to the administration of Acepromazine maleate injection to male horses (castrated and uncastrated). When given, the dosage should be carefully limited to the minimum necessary for the desired effect. At the time of tranquilization, it is not possible to differentiate between reversible protrusion of the penis (a normal clinical sign of narcosis) and the irreversible paralysis of the retractor muscle. The cause of this side reaction has not been determined. It has been postulated that such paralysis may occur when a tranquilizer is used in conjunction with testosterone (or in stallions).

Accidental intracarotid injection in horses can produce clinical signs ranging from disorientation to convulsive seizures and death.

Adverse reactions:

A few rare but serious occurrences of idiosyncratic reactions to acepromazine may occur in dogs following oral or parenteral administration. These potentially serious adverse reactions include behavioral disorders in dogs such as aggression, biting/chewing, and nervousness.

To report suspected adverse reactions, to obtain a Safety Data Sheet or for technical assistance, call 1-855-724-3461. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS, or www.fda.gov/reportanimalae.

Animal safety:

Acute and chronic toxicity studies have shown a very low order of toxicity.

Acute toxicity: The LD₅₀ dose of Acepromazine maleate injection in mice was determined by means of a probit transformation with the following results: 2

Intravenous route — 61.37 mg/kg Subcutaneous route — 130.5 mg/kg Oral route — 256.8 mg/kg

Chronic toxicity: Tests³ in rats revealed no deleterious effects on renal or hepatic function or on hemopoietic activity. In several groups of two male and two female beagle hounds treated for six months with daily oral doses of 20 to 40 mg/kg, no untoward effects were encountered. Hematologic studies and urinalysis gave values within normal limits. Another group of four dogs, given gradually increasing oral doses up to a level of 220 mg/kg daily and reaching a total daily dose of 2.2 g per dog, showed some signs of pulmonary edema and hyperemia of the internal organs, but no animal died.

When administered intramuscularly, Acepromazine maleate injection causes a brief sensation of stinging comparable with that observed with other phenothiazine

tranquilizers.

Effectiveness:

Controlled clinical studies in the United States and Canada have demonstrated the effectiveness and safety of Acepromazine maleate injection as a tranquilizer.

Good to excellent results were reported 1,4,5 in dogs, cats and horses given Acepromazine maleate injection for restraint during examination, treatment and minor surgery and for preanesthetic sedation. In dogs, the drug reportedly 4 helps control convulsions associated with distemper.

In both dogs and cats, good to excellent results were obtained⁴ when Acepromazine maleate tablets were used to control nervousness, excessive vocalization, neurotic and excitable behavior, vomiting associated with motion sickness, coughing and itching caused by dermatitis.

In horses, Bauman⁶ had good results using the drug as an aid in the control of painful spasm due to colic.

Other practitioners^{7,8} found the drug useful as a preanesthetic sedative for nervous or aggressive horses, but it had to be administered while the animals were quiet and not in an excited state. In a trial⁹ on more than 200 horses with a wide variety of disorders, Acepromazine maleate injection proved to be both effective and safe.

Storage:

Store at 20° to 25°C (68° to 77°F), excursions permitted between 15° and 30°C (between 59° and 86°F). When used as labeled, there is no limit on the number of punctures throughout the full expiry period.

How supplied:

Acepromazine maleate injection is supplied in 50 mL multiple dose vials.

NDC 11695-0095-5 - 10 mg/mL

50 mL vial

References:

- 1. Baker, J.M.: Paper presented at the Ontario Veterinary Association meeting, held in Toronto, Canada, 1958.
- 2. Pharmacology Reports, ClinByla Laboratories, Paris, France.
- 3. Stegen, M.G.: Pharmacology Report, Ayerst Laboratories, 1958.
- 4. Veterinary Medical Records, Ayerst Laboratories.
- 5. Foley, J.T.: Clinical Reports to Ayerst Laboratories, 1963.
- 6. Bauman, W.G.: Clinical Reports to Ayerst Laboratories, 1963.
- 7. Ford, R.W.: in Equine Panel Report, Mod. Vet. Pract. 40:45 (Nov. 1) 1959.
- 8. Baldwin, R.: in Equine Panel Report, Mod. Vet. Pract. 40:46 (Nov. 1) 1959.
- 9. Dunkin, T.E.: Clinical Reports to Ayerst Laboratories, 1963.

Distributed by:

Covetrus North America

400 Metro Place North

Dublin, OH 43017

covetrus.com

AH-Acepromazine Injection-03

Rev: 0919

382703-05

87018822

Principal Display Panel - 50 mL Container Label

Covetrus

NDC: 11695-0095-5

Acepromazine maleate

Injection

10 mg/mL

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50 mL

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A sterile solution for intravenous, intramuscular or subcutaneous injection.

Each mL contains: Acepromazine maleate 10 mg, sodium citrate 0.36%, citric acid 0.075%, benzyl alcohol 1% and water for injection.

Refer to folded package insert attached to this container for complete product information.

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Questions? (855) 724-3461 Reorder #003845 REV: 0919 AH-003845-03 382703-05 87018822

ACEPROMAZINE MALEATE

acepromazine maleate injection

Product Information			
Product Type	PRESCRIPTION ANIMAL DRUG	Item Code (Source)	NDC:11695- 0095
Route of Administration	INTRAVENOUS, INTRAMUSCULAR, SUBCUTANEOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ACEPROMAZINE MALEATE (UNII: 37862HP2OM) (ACEPROMAZINE - UNII:54EJ303F0R)	ACEPROMAZINE MALEATE	10 mg in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
BENZYL ALCOHOL (UNII: LKG8494WBH)	10 mg in 1 mL		
TRISODIUM CITRATE DIHYDRATE (UNII: B22547B95K)	3.6 mg in 1 mL		
ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)	0.75 mg in 1 mL		
WATER (UNII: 059QF0KO0R)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:11695-0095-5	50 mL in 1 VIAL, MULTI-DOSE		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
NADA	NADA015030	11/25/2020			

Labeler - Covetrus North America (603750329)

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