Amitraz poisoning: Rare but not uncommon - a veterinary compound causing high morbidity

Case Report

Amitraz Poisoning: Rare but not Uncommon - a Veterinary Compound Causing high Morbidity

Sapna Gupta¹, Dhaiwat Shukla²

¹Assistant Professor dept of emergency, medicine, Smt, NHL MMC,VSGH
²2nd year resident, dept of, medicine, Smt, NHL MMC,VSGH

INTRODUCTION
A 18 years old male has taken some unknown substance & after 30 mins called his friends that he has taken some poison & he is having nausea & vomiting .When friends reach for rescue patient was found unconscious in pool of vomitus with an empty bottle of Amitraz(12.5%w/v 25 ml).

Friends took him to local hospital where positive findings were as follows.
On general examination
Pulse was not palpable with cold extremities
• BP – 78 systolic on auscultation
• Pupils – miosis
Patient was given RT wash by saline & drip started immediately. Patient was referred to VS general hospital in gasping state.
Urgent intubation done. And was put on volume A/C mode of mechanical ventilator with FiO₂ 70%.
Complete toxicology screen was sent to rule out mixed poisoning.

Examination Finding

*Corresponding Author
Dr Sapna Gupta,
Asst Professor room,
Dept. of Emergency Medicine,
1st, floor, trauma centre, V.S general hospital, Ellisbridge,
Ahmedabad – 380007
Email: sapna_gupta76@yahoo.com

• BP – 90 Systolic.
• Right side middle zone air entry was reduced.

Patients investigation revealed.
• Hb – 14.6gm%
• TC – 14,500 / cumm
• APC – 1.67 lac / cumm
• Glucose –>600 mg%
• Creatinine–1.06 mg%
• Potassium –5.2
• Sodium – 141
• Billirubin – 0.4 mg%
• Serum Cholinesterase–4722(2900-5800) (NIOH sample) 8.2 (4.62 – 11.5) (Private Lab)
• Urine for toxicology screening was normal for morphine, cocaine, amphetamine, barbiturates, benzodiazepenes.

• ABGA
pH –7.275
PCO₂ – 37.4
PO₂ – 60.0
HCO₃ – (-) 17.0
O₂ Sat – 91.7%
Patient was given supportive treatment in form of higher antibiotics, inotropic support, antacids & atropine on as and when required basis
He was ventilated on assisted mode of ventilator for 48 hours because he was in drowsy state and required inotropic support.
On the 3rd day patient became alert and vitals were normal. Patients was conscious oriented following verbal command. So, extubated on 3rd dayand kept under vigilant observation for 24 hours. Next day he was sent to ward from ICU and further recovery was uneventful.
Amitraz poisoning: Rare but not uncommon - a veterinary compound causing high morbidity

**DISCUSSION**

Our patient was a cattle man (herder) in this case there was no diagnostic dialema as patient was found with bottle of Amitraz and with typical features of the same. But in case of doubt organophosphorous poisoning, clonidine poisoning and oral contraceptive poisoning should be considered as differential diagnosis.

**Compound Introduction**

Use: Amitraz is particularly effective against acarids, but it is used as a pesticide in many different fields. Therefore amitraz is available in many different forms, as in a wettable powder, emulsifiable concentrate, soluble concentrate / liquid, and impregnated collar (for dogs). It is characterized as an insect repellent, insecticide and pesticide synergist. These are the properties which make it especially useful as a pesticide.

The repellent effect causes insects to turn away from their target as this is treated with amitraz. It acts as an insecticide, which means that it can be used to control insects that are directly or indirectly harmful to men.

As a pesticide synergist, also increases the effect of some other pesticides if they are combined with amitraz. These can be traced back to the mechanisms of action, which lead to a wide field of effects, including direct lethality, excitant-repellent behavioral effects, and chemosterilization for the target species. In addition it generally causes low damage to non-target species, which is one of the advantages of amitraz. Furthermore amitraz is especially effective against insects as spider mites and ticks in their juvenile and resistant forms. For agricultural purposes amitraz is primarily used to control the pear psylla (Cacopsylla pyricola) on Oregon pear crops and whiteflies and mites on cotton or pear crops. It’s also applied to pome fruit, citrus fruit, cotton, stone fruit, bush fruit, strawberries, hops, cucurbits, aubergines, capiscums, tomatoes and ornamental plants to control all stages of tetranychid and eriophyid mites, pear suckers, scale insects, mealybugs, whiteflies, aphids and eggs and first instar larvae of lepidoptera. To apply amitraz, various techniques can be used as for example an airblast and concentrate spray to pears or by ground boom and aircraft to cotton. Territorial differences in amitraz use depend on the species of mites that infest the crops/trees/etc., the local practice and the number and size of the pear trees. An infestation e.g. by Tetranychus requires higher rates of amitraz. Taking those factors into consideration the application volumes of amitraz have been standardized in terms of maximum spray concentration and in the rate of amitraz per hectare.

Besides its application as pesticide on plants, amitraz is also used as an animal ectoparasiticide on cattle, goats, sheep, pigs and dogs. Thereby it’s exclusively applied externally. It achieves special efficiency against mites (first of all Demodex canis), but it also works against lice, flies and all development stages of ticks. In combination with additional agents it can be used against flea-infestation as well. For the treatment of dogs amitraz is available as a collar or as a spray- or wash solution and has an immediate effect against tick-infestation as well as a preventative effect. In some countries amitraz-emulsions are also applied to treat Demodicosis of cats or dogs, an exceeding infestation of mites of the family Demodicidae. For the treatment of cattle, sheep, goats and pigs amitraz is available as spray or wash solution, to treat or prevent infestations by mites, lice, flies and ticks. Thereby pigs and cattle should be sprayed and sheep and goats bathed. Other animal species as for example horses or Chihuahuas should not be treated with amitraz, because adverse effects may occur.

**Metabolism**

Since amitraz most common use is as a pesticide, it is important to consider that between animals and plants often different pathways for biotransformation occur. Most animal species, including humans can metabolize amitraz rapidly to form six metabolites during biotransformation, N-methyl-N′-(2,4-xylyl) formamide, Form-2′xylidine, 4-N-Methyl-formidoyl) amino-meta-toluix acid, 4-Formamido-meta-toluic acid, 4- Acetamido-meta-
Amitraz poisoning: Rare but not uncommon - a veterinarian compound causing high morbidity

toluic acid and 4-Amino-meta-toluic acid.\textsuperscript{13,14,15}

Amitraz Metabolism in Animals
In rats the metabolic pathway (figure 3) has been examined after oral administration of \textsuperscript{14}C-labelled amitraz, which was found to be effectively metabolized, degraded and excreted to four of the metabolites in urine and six in faeces.\textsuperscript{14} The metabolic pathway or rate did not differ between the sexes. Hornish and Nappier (1983)[full citation needed] detected that the metabolic pathway after dermal administration follows the same route of degradation as after oral uptake, because the parent compound, N-methyl-N'(2,4-xylyl) formamidine and form-2',4'-xylidine were found in urine and blood also after dermal administration.\textsuperscript{14} In humans, N-methyl-N'(2,4-xylyl) formamidine, form-2',4'-xylidine, 4-amino-meta-toluic acid, 4-acetamido-meta-toluic and 4-formamido-meta-toluic acids were recognized in the urine as well which indicates for the same or a similar metabolic pathway.\textsuperscript{15}

As illustrated in figure 3 the first step is a hydrolysis reaction to N-methyl-N'(2,4-xylyl)-formamidine, which already can be excreted in the urine but is still pharmacological active.\textsuperscript{14,15} Depending on the dose, the quantity of this metabolite in the urine can vary from 4\% at low doses to 23\%-38\% at high doses (e.g. in case of rats: 1–100 mg per kg body weight).\textsuperscript{14} As it isn't excreted it also can be oxidized to 4-N-Methyl-formidoylamin-meta-toluic acid, which can be further oxidized to 4-formamido-meta-toluic acid.\textsuperscript{14} Form-2,4-xylidine is formed directly by hydrolysis from amitraz or arises from N-methyl-N'(2,4-xylyl) formamidine.\textsuperscript{15} During this early stage of biotransformation N-methyl-N'(2,4-xylyl) formamidine and Form-2,4-xylidine may already form conjugates.\textsuperscript{[14]} But the major route followed after the formation of Form-2,4-xylidine is the oxidation to 4-formamido-meta-toluic acid, which is further metabolized to its acetyl conjugate, 4-acetamido-meta-toluic acid or 4-amino-meta-toluic acid.\textsuperscript{14,15} 4-formamido-meta-toluic acid and 4-acetamido-meta-toluic acid make 32\% of the metabolites found in urine and are detected at any administered dose. Therefore they are considered as two of the major metabolites in the amitraz pathway.\textsuperscript{14} Form-2',4'-xylidine and 4-amino-meta-toluic acid account only for 2\% of the total excretion.\textsuperscript{14} In insects different metabolites are formed. N-methyl-N'(2,4-xylyl)formamidine, Form-2,4-xylidine and 4-Amino-meta-toluic acid occur, but in addition several unidentified metabolites were detected, too.\textsuperscript{15}

Amitraz Metabolism in Plants
In plants the biotransformation of amitraz proceeds very rapidly. The predominant metabolites detected are N-(2,4-dimethylphenyl)-N'-methylformamidine (BST 27 271) and 2,4-dimethylformanilide (BST 27 919).\textsuperscript{8} N-(2,4-dimethylphenyl)-N'-methylformamidine (BST 27 271), 2,4-dimethylformanilide (BST 27 919) and N,N'-bis-dimethylphenyl formamidine (BTS 28 037) result from hydrolysis of amitraz. Thereby N-(2,4-dimethylphenyl)-N'-methylformamidine (BST 27 271) occurs in higher amounts than 2,4-dimethylformanilide (BST 27 919). N-(2,4-dimethylphenyl)-N'-methylformamidine (BST 27 271) can be further metabolized to 2,4-dimethylformanilide (BST 27 919) or 2,4-dimethylaniline (BTS 24 868).\textsuperscript{8} N,N'-bis-dimethylphenyl formamidine (BTS 28 037) can be transformed to 2,4-dimethylformanilide (BST 27 919) or directly react to 2,4-dimethylaniline (BTS 24 868), but the exact mechanisms of these biotransformations are not known yet.\textsuperscript{8} However, of 2,4-dimethylaniline (BTS 24 868) and N,N'-bis-dimethylphenyl formamidine(BTS 28 037) less than 1\% has been accounted, which makes them minor metabolites compared to N-(2,4-dimethylphenyl)-N'-methylformamidine (BST 27 271) and 2,4-dimethylformanilide (BST 27 919).\textsuperscript{8} Figure the suggested amitraz' metabolic pathway in plants.\textsuperscript{8}

Kinetics
The hydrolysis reactions of amitraz strongly depend on the environmental pH. Even though amitraz undergoes hydrolysis reactions at any pH, spectrophotometry, HPLC and GC-MS studies revealed that pH-depending differences occur, affecting both the sort of
Amitraz poisoning: Rare but not uncommon - a veterinary compound causing high morbidity

reaction-products and the reaction rate.\textsuperscript{[1]}\textsuperscript{[16]} Under basic conditions (pH>6) amitraz is metabolized to 2,4-dimethylphenylformamide. Followed by hydrolysis to 2,4-dimethylaniline, which also benefits from a basic pH.\textsuperscript{1,16} At very acidic pH (pH<3) 2,4-dimethylaniline has been observed as the main degradation product. Under less acidic conditions (pH 3–6) mainly N-(2,4-dimethylphenyl)-N'-methylformamidine and already amounts of 2,4-dimethylphenylformamide occur.\textsuperscript{1}

Mechanism of action
Amitraz is used as a pesticide and in the industry. Therefore amitraz exposure to humans is rare and occurs mainly through inhalation or dermal contact with the compound during its use or production.\textsuperscript{17} The toxic effects to humans following on amitraz-uptake include loss of consciousness, vomiting, respiratory failure, miosis, hypothermia, bradycardia, hyperglycemia and central nervous system depression.\textsuperscript{4}

The pharmacological activity of amitraz includes different mechanisms of action leading to toxic effects in humans as well as in animals. Many of these effects and most of the effects on humans are caused by its alpha-adrenergic agonist activity.\textsuperscript{4} Furthermore amitraz inhibits prostaglandin synthesis, interacts with the octopamine receptors of the central nervous system and inhibits monoamine oxidases.\textsuperscript{4}

Animal studies revealed that damages due to amitraz poisoning can be recovered even after exposure to a potentially lethal dose. This could mean that amitraz' effects are reversible or at least are recoverable.\textsuperscript{18} When an amitraz poisoning is lethal, death results from respiratory depression.\textsuperscript{18}

Alpha-adrenergic agonist activity
Amitraz is a central alpha-adrenoceptor agonist.\textsuperscript{17} That means that it selectively stimulates alpha adrenergic receptors, which are metabotropic G-protein-coupled receptors, that are usually targeted by catecholamines. Stimulating these receptors is in great extent the reason for the neurotoxic and preconvulsant effects of amitraz.\textsuperscript{19} Xylenes present in amitraz formulations additionally induces central nervous system depression.\textsuperscript{4} Adrenergic receptors can be divided into two subclasses, alpha1- and alpha2-adrenergic receptors. To determine whether amitraz interacts with subclass 1 oder subclass 2, subcutaneous injections of amitraz (0.3–3.0 mg/kg) were given to mice.\textsuperscript{20} Consequently a dose-dependent delay of gastrointestinal transit in conscious mice occurs. This effect could be antagonized by alpha2-adrenergic blocking agents, but administration of other antagonists did not reduce the depressant effect on the gastrointestinal transit.\textsuperscript{20} So it is suggested that amitraz-induced delay of gastrointestinal transit is mediated by postjunctional alpha2-adrenergic receptors and appears not to involve the activation of β-adrenergic, dopaminergic, serotonergic, histaminergic, cholinergic, GABAergic, or opioid receptors.\textsuperscript{20} Besides the neurotoxic effects other clinical effects observed in amitraz poisoning are related to alpha2-adrenergic agonistic activity.\textsuperscript{3} Adrenergic receptors are present in many different cells. The activation of these receptors by an agonist as amitraz generally induces a sympathetic response. This leads to an increased heart rate, dilation of the pupils, elevation of blood pressure and blood and energy supply focus on skeletal muscles.\textsuperscript{17}

Interaction with the octopamine receptor
It's thought that the mode of action of amitraz involves the interaction with the neuromodulator octopamine.\textsuperscript{21} This interaction is probably the reason for increased nervous activity of ticks as a response on amitraz.\textsuperscript{21,22} usual activation of the receptors may lead to changes in the concentration of intracellular second messengers such as cyclic nucleotides cyclic AMP (cAMP) and cyclic GMP, inositol-1,4,5-trisphosphate and Ca2+.\textsuperscript{23} Influencing this signal transduction system can lead to various events depending on the celltype.\textsuperscript{23} Since it has been discovered that the octopamine receptor coding gene is expressed on very high rates in the somata of the honeybee brain, it is suggested that it is involved in the processing of sensory inputs, antennal motor outputs and higher-order brain functions. The amitraz-octopamine

receptor- interaction restrains these normal functions of the octopamine receptor. Therefore it is efficient as an insect-pesticide. Still, resistance against amitraz can occur. A mutation can lead to a working version of the octopamine receptor but with an altered pesticide target side. This is probably the case for a very resistant Brazilian and Mexican tick strain, which have two nucleotide substitutions on the octopamine receptor coding gene compared with the Australian strains. A closer understanding of these resistance mechanisms would help to develop more rapid and accurate diagnostic tools for detecting resistance and steer development of alternative acaricides.

Inhibition of monoamine oxidases
In vitro a monoamine oxidase-inhibiting effect of amitraz has been found. Monoamine oxidases catalyze the oxidative deamination of monoamines and thereby form flavoproteins and inactivatet neurotransmitters. However, in vivo it has been observed that only at high doses of amitraz or its main metabolite N-2,4-dimethylphenyl-N-methyl-formamide monoamine oxidase inhibition occurs. In dogs it has been observed that after administration of such a dose an increase in plasmaglucose and suppression of insulin occurs.

Inhibition of prostaglandin synthesis
Like other formamidines amitraz inhibits the synthesis of prostaglandin E2 from arachidonic acid by bovine seminal vesicle microsomes. In a dose of 5 to 80 mg/kg body weight, given intraperitoneally to rats, amitraz reduces yeast-onduces fever and antagonizes the caragenin-induced swelling of the hind paw. Some of the physiological effects of amitraz probably go back to this aspirin-like activity and occur due to inhibition of prostaglandin synthesis.

Adverse effects
Adverse effects in mammals are caused by amitraz' alpha-adrenergic agonist activity. Symptoms can include low blood pressure and pulse, hypothermia, lethargy, absence of appetite, vomiting, increased blood sugar and digestive problems. Furthermore, skin- or mucosa-irritations may occur in dogs as a response to an amitraz containing collar. This can lead to itching, eczema, alopecia or conjunctivitis.

Treatment
In case of an amitraz overdose in humans atipamezole or yohimbine, which act as a2-antagonists, can be used as antidote. Initially it is important to remove the patient from the amitraz contaminated area. When amitraz has been inhaled the patient should first get respiratory protection. Additionally the patient should be supplied with 4 L oxygen per minute. In case of an intoxication via skin-contact, contaminated clothes should be removed first. Affected areas need to be washed with water. If eyes have been exposed to amitraz, anesthesia should be administered and the eyes carefully washed. After the oral intake of amitraz it is important to make the patient drink ca. 0.3 L water to reduce amitraz' irritating effect on the gullet. Furthermore, it is important to prevent the patient as much as possible from vomiting, to reduce the risk of further aspiration of amitraz. Subsequently, the patient need to be observed for at least 24 hours to ensure that the symptoms do not recur.

Toxicity
Human toxicity
In 2006 the United States Environmental Protection Agency (USEPA) re-assessed the classification for amitraz to a non-quantifiable “Suggestive Evidence of Carcinogenicity” descriptor, and in 2013 determined that quantification of risk using a non-linear approach for amitraz will adequately account for all chronic toxicity, including carcinogenicity, that could result from exposure to amitraz and its metabolites. Accidental exposure of men to greater amounts of amitraz can lead to death due to respiratory failure, mainly after oral uptake or inhalation. In Turkey during 1989, 41 cases of deadly amitraz intoxications have been detected. The observed toxic dose in about 50% of these patients has been 0.3 g to 1.25 g of 12.5% amitraz formulations and 0.5 to 2 g of 20% formulations. The remaining patients took doses up to 10 g. Other frequently occurring symptoms after massive amitraz intoxication are CNS depression, respiratory depression, miosis,
Amoriz poisoning: Rare but not uncommon - a veterinarian compound causing high morbidity

hypothesis, hyperglycemia, loss of consciousness, vomiting and bradycardia.3

CONCLUSION

Amitraz poisoning is an emerging health hazard as it is freely available to herders as a over the counter medicine and it can cause high mortality if not managed in time but life can be saved with very good supportive care.

REFERENCES


4. Jump up to: a b c d e f g h i Bonsall, J. L., & Turnbull, G. J. (1983). Extrapolation from safety data to management of poisoning with reference to amitraz (a formamidine pesticide) and xylene. Human Toxicology

5. Jump up to: a b c d e f g h Brown, P. M. (1977). Toxicolical problems associated with the manufacture of triazapentadienes. Proceedings of the Royal Society of Medicine, 70(1), 41-43


7. Jump up to: a b c d e f g h i Hollingworth, R. M. (1976). Chemistry, biological activity, and uses of formamidine pesticides. Environmental Health Perspectives, 14(April), 57-69


Amitraz poisoning: Rare but not uncommon - a veterinary compound causing high morbidity


