



CASE REPORT

Crotalus oreganus concolor: Envenomation Case with Venom Analysis and a Diagnostic Conundrum of Myoneurologic Symptoms

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A case of midget-faded rattlesnake (*Crotalus oreganus concolor*) envenomation of an adult male professional herpetologist occurred in a rural setting and resulted in an array of venom induced myoneurologic symptoms. The patient experienced blurry vision, total body paresthesia, dyspnea, chest tightness, and waves of spastic muscle movements of the hands and feet that resembled tetany. It was not apparent whether these symptoms were potentially venom induced or were related to stress-induced physiologic responses. Local envenomation effects were minimal, and coagulation parameters remained within normal limits. Antivenom was not administered per patient concerns related to a history of acute allergic reactions to antivenom. Venom was collected from the *Crotalus oreganus concolor* responsible for the bite, and analysis revealed the presence of high levels of myotoxins (SR calcium pump antagonists) and concolor toxin, a presynaptic neurotoxin that can have myotoxic effects and cause respiratory paralysis; several serine proteinases associated with coagulopathies were also present in the venom profile.

Keywords: rattlesnake, snakebite, hyperventilation, carpopedal, alkalemia, hypocalcemia

Introduction

The midget-faded rattlesnake (*Crotalus oreganus concolor*) is an uncommon, small (50–65 cm total body length) subspecies originally assigned to the *Crotalus viridis* complex and currently recognized as a subspecies in the *Crotalus oreganus* clade.^{1,2} It is indigenous to a restricted geographic range of the Colorado Plateau that includes west central Colorado, the Colorado and Green River basins, eastern Utah, southwestern Wyoming, and extreme northern Arizona.³⁻⁵

Crotalus oreganus concolor venom studies have revealed the presence of a potent lethal toxin, antigenically similar to Mojave toxin.^{6,7} It has been determined to be one of the most lethal crotaline venoms, nearly equal in toxicity to *Crotalus durissus terrificus* and *Crotalus scutulatus scutulatus* venoms based on murine lethality experiments.⁸ In addition, *Crotalus oreganus concolor* venom lethality is reported to range from 10- to 30-fold greater than multiple other crotaline species.^{8,9} A presynaptic phospholipase A₂-β-neurotoxin (concolor toxin) and nonenzymatic peptide myotoxins have been identified as major venom components.⁹⁻¹²

Envenomation of humans by *Crotalus oreganus concolor* is rare, and the case reported here illustrates what clinically appeared to be possible venom-induced myoneurologic symptoms that were confounded in interpretation by physiologic stresses. Analysis of venom from the *Crotalus oreganus concolor* responsible in this case suggested a possible correlation between specific venom components and several observed clinical symptoms.

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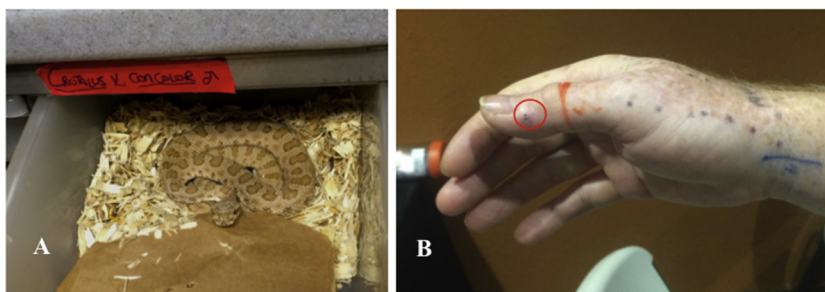


Figure 1. A, The midget faded rattlesnake (*Crotalus oreganus concolor*) responsible for the bite, in its housing drawer enclosure from which it struck, imparting a single fang puncture to the patient. B, Single fang puncture into the distal phalangeal region of the right thumb.

Case Presentation

A healthy 61-y-old male professional herpetologist sustained a bite from a captive (wild caught) adult male *Crotalus oreganus concolor* (Figure 1A). Patient history included prior crotaline envenomation from *Crotalus horridus*, *Crotalus horridus atricaudatus* (subspecies recognized at the time of the bite), and *Crotalus durissus ruruima*. Treatments involved Wyeth anticrotalic antivenoms and Instituto Butantan anticrotalico, with anaphylactic reactions having occurred. Patient medication history included migraine headache prescriptions (rizatriptan, atenolol, and amitriptyline).

The incident took place in a remote facility, and the bite occurred as the snake was being removed from a drawer-type housing unit, resulting in a single fang puncture into the right thumb (Figure 1B). The patient immediately experienced generalized “pins and needles” sensations in the bitten thumb that rapidly progressed to pain, followed by tingling of the lips and a feeling of chest tightness. Within 15 min the patient reported total body tingling. The first responder found the patient resting supine on the floor and reported the patient did not appear anxious or agitated, with no obvious signs of hyperventilation (20 breaths \cdot min $^{-1}$). The patient described feeling like he was “wearing a clay mask and a hat with hat band one size too small constricting around my forehead.” The affected limb was splinted for immobilization during a 1-h ground ambulance transport to the helispot and subsequent 45-min helicopter flight to a rural hospital. During ground transport, the patient reported breathing difficulty, and supplemental oxygen was provided via nasal cannula. Tightening of the tongue, blurred vision, and difficulty speaking were also experienced. Three distinct waves of transient neuromuscular spasms associated with inspirational weakness ensued. Spastic simultaneous contractures of first the left hand and then to a lesser degree the right (bitten) hand and foot/toes occurred, followed by a third wave that involved the left forearm, lower leg, and foot. The first responder described these waves as severe, tetany-like,

spastic muscle movement (Figure 2A). The patient was alert and oriented; blood pressure was normal by sphygmomanometer/cuff (113/72 mm Hg), and pulse was normal by palpitation (69 beats \cdot min $^{-1}$) (Figure 2B).

Approximately 2.5 h postbite the patient arrived to the emergency department, and the myoneurologic symptoms had abated. Examination revealed a single fang puncture in the distal phalangeal region of the right thumb and swelling that extended proximally to the thenar eminence. Chemistry panel, complete blood count, electrolytes, coagulation, and hematologic parameters were within normal limits, excepting a slightly elevated D-dimer (566 ng \cdot mL $^{-1}$ FEU; reference laboratory range 190–490 ng \cdot mL $^{-1}$ FEU) and elevated creatine kinase (CK) 3394 U \cdot L $^{-1}$ (reference laboratory range 22–198 U \cdot L $^{-1}$).

Over the following 24 h, there was a small decline in total serum calcium from 8.8 mg \cdot dL $^{-1}$ (3 h) to 7.7 mg \cdot dL $^{-1}$ (22 h) (reference laboratory range 8.5–10.1 mg \cdot dL $^{-1}$) and a corresponding slight decline in serum albumin from 4.2 g \cdot dL $^{-1}$ (3 h) to 3.5 g \cdot dL $^{-1}$ (22 h) (reference laboratory range 3.4–5.0 g \cdot dL $^{-1}$). There was no evidence of bleeding at any time. Swelling spread proximally to mid-forearm over the first 3 h, but local pain had lessened. Medical staff observed weak patient total body strength with poor coordination, requiring walking frame assistance. There was no evidence of compartment syndrome. Despite indications for antivenom, its administration was declined by the patient because of anaphylactic reactions from previous antivenom treatments.

The patient was observed for another 6 h and discharged at 25 h postbite (22 h postadmission) in stable condition. At 24 h after discharge, the patient remained weak but could walk without assistance. Myalgia in the right arm and generalized weakness persisted for several days. At 3-wk follow-up, the bitten thumb skin had sloughed and thumb numbness persisted, but other local symptoms had resolved completely and all laboratory studies were unremarkable. Venom from the responsible *Crotalus oreganus concolor* was collected for assay and analysis of venom components and their potential correlation to observed symptoms.

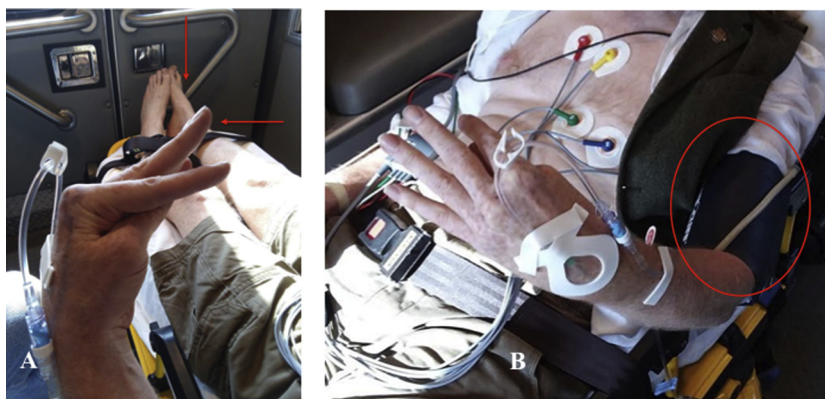


Figure 2. A, Tetany-like spasms showing flexion of the hands and extension of the feet (carpopedal spasms) that began 30 min after the bite. Three waves of approximately 10 min each, 20 min apart, were observed. B, Blood pressure and pulse were normal to low as measured by palpitation and the sphygmomanometer/cuff placed on the contralateral arm.

METHODS

Venom was extracted manually from the *Crotalus oreganus concolor* responsible for the bite approximately 4 d after the bite and centrifuged to pellet the cell debris, and the supernatant was lyophilized (fluid yield 120 μL ; ≈ 27 mg dry venom). Lyophilized venom was solubilized in phosphate-buffered saline ($8.0 \text{ mg} \cdot \text{mL}^{-1}$), and 200 μL (1.6 mg) was subjected to reversed-phase high-performance liquid chromatography (RP-HPLC) as reported previously; 1-min fractions were collected and lyophilized.¹³ Toxins were identified from the known elution profiles of purified toxins and quantified via peak area integration (percentage total area).

HPLC fractions were also subjected to polyacrylamide gel electrophoresis (SDS-PAGE) under reducing conditions to identify chromatogram peaks.¹³ Crude venom was assayed for protein content via Pierce BCA assay and for 6 enzyme activities common to rattlesnake venoms.^{10,14} These data were used in conjunction with HPLC chromatogram data to identify the protein families present and their relative abundance in the venom.

DISCUSSION

Our patient's symptoms illustrate a potential diagnostic conundrum clinicians may face in cases of snakebite envenomation: What symptoms might be venom-induced effects, and what symptoms might be related to stress-induced physiologic responses? Our patient experienced generalized paresthesia, blurred vision, waves of spastic, tetany-like symptoms of the hands and feet (carpopedal spasms), breathing difficulties, numbness, and generalized weakness that persisted for days, all symptoms consistent with venom-induced effects reported in other cases of North American rattlesnake envenomation, including by *Crotalus oreganus concolor*.^{9,15-17} However, in our patient, the tetany-like carpopedal spasms may not have been venom induced, but instead from patient

anxiety-induced hyperventilation or sphygmomanometer cuff inflation.^{18,19} *Crotalus oreganus concolor* venom contains myotoxic and neurotoxic components capable of eliciting neurotoxic symptoms.^{9,10,12} Purified *Crotalus oreganus concolor* venom toxins injected into mice elicit myotoxic effects of rapid ataxia, hind limb extension, and respiratory distress.^{7,20} Collectively, symptoms in the present case combined with the interpretation of the venom analysis for the inflicting snake provide for interesting discussion.

Venomous snakebite can trigger significant anxiety, leading to an autonomic response of hyperventilation-induced alkalemia with consequent precipitation of hypocalcemia, resulting in symptoms of breathlessness and paresthesia of the face and hands, although blood pressure remains normal.¹⁸ Tetany-like symptoms of painful sharp involuntary flexion of the wrist and ankle joints (carpopedal spasms), cramps, and muscle twitching may be clinically misleading. These physiologic responses are due to decreased extracellular ionized calcium, resulting in hyperexcitability of muscles and nerves.¹⁸ Ionized serum calcium was not measured in our patient; however, the patient was not observed to be tachypnic, although hyperventilation may occur via slow deep breathing. Hypocalcemic tetany of the lower extremities has been reported in a case of *Crotalus scutulatus* envenomation with minimal coagulopathy, but this was thought to have been subsequent to rhabdomyolysis.²¹ Our patient had normal blood pressure and may have deeply hyperventilated to the point of respiratory alkalosis followed by hypocalcemic tetany. The carpopedal symptoms fit the profile of stress-related physiologic responses, which were considered a contributing root cause.

Blood pressure measurement via sphygmomanometer with an inflatable cuff was used to monitor our patient's blood pressure during ambulance transport. Cuff inflation, in conjunction with hyperventilation-induced hypocalcemia, likely triggered a classic Trousseau sign.¹⁹ Cuff

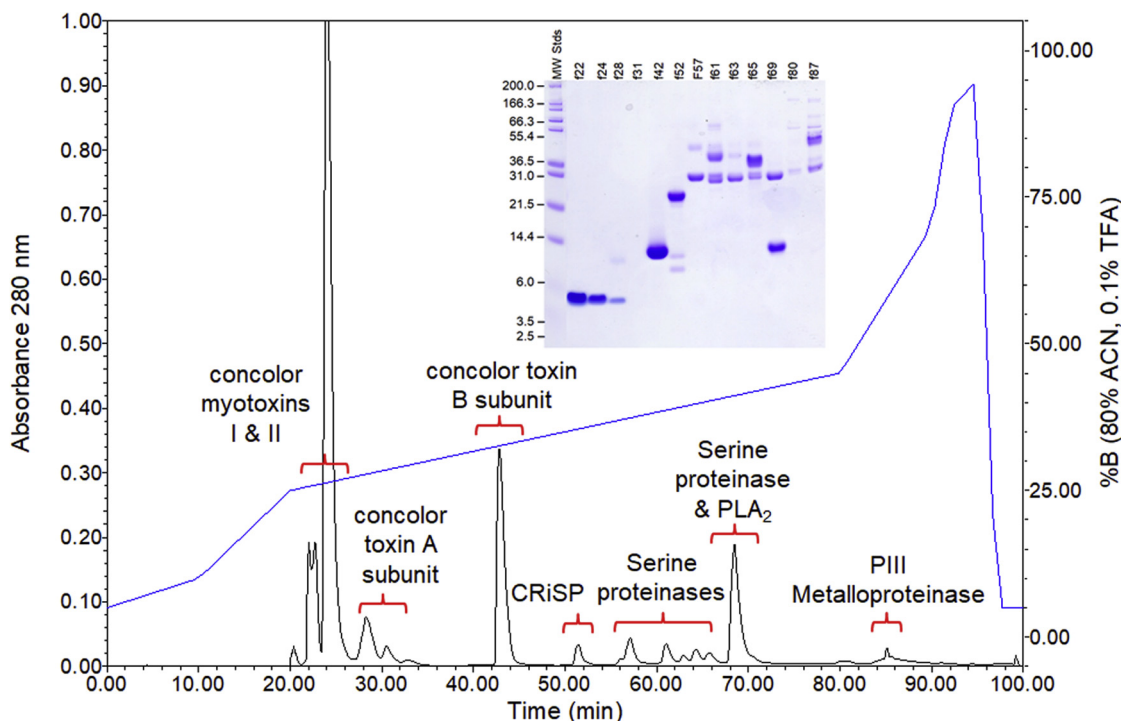


Figure 3. Reversed-phase high-performance liquid chromatography chromatogram of *Crotalus oreganus concolor* venom, 1.6 mg in 200 μ L phosphate-buffered saline. Identification of peaks are based on SDS-PAGE and enzyme activity assays. Inset: reducing SDS-PAGE of indicated high-performance liquid chromatography fractions. CRISP, cysteine-rich secretory protein.

inflation to greater than the mean arterial pressure can result in the hand adopting a posture of metacarpophalangeal joint flexion with the interphalangeal joints of the fingers and thumb extended so that the thumb is in an opposing posture, as observed in our patient (Figure 2B). This phenomenon has been reported in 1 to 4% of healthy individuals but has not been reported to be associated with cases of snake-bite envenomation.¹⁹ Although *Crotalus oreganus concolor* myotoxic venom components can induce muscle contractile activity, causing rapid tetany-like hind limb hyperextension in mice (a symptom somewhat similar to the extension of the ankles, feet, and toes observed in our patient), this symptom was likely a stress-induced response rather than venom induced^{20,22}

Our patient's other myoneurologic symptoms of paresthesia, respiratory distress, and total body weakness may have been related to the high myotoxin content and concolor toxin in the *Crotalus oreganus concolor*'s venom (Figure 3, Table 1). Studies in mice injected with *Crotalus oreganus concolor* venom showed they died from rapid respiratory failure, suggesting that venom may have contributed to our patient's respiratory difficulty, as has been reported to occur after *C. scutulatus*, *C. cerastes*, *C. oreganus helleri*, and *C. horridus* envenomation.^{16,17,23,24}

Venom was delivered to our patient via a single fang puncture, and it is reasonable to assume it was not a

maximum dose; this is a potential reason for the reduced severity and duration of symptoms. Neurotoxic effect duration after snake envenomation is highly variable.²⁵ The short duration of myoneurologic actions in our patient may have been related to dose-dependent pharmacokinetics or pharmacokinetic metabolic actions on a responsible toxin.

Myokymia has been reported after envenomation by *Crotalus oreganus concolor* and other North American rattlesnakes.^{15,26,27} In our patient, the periodic episodes of tetany-like carpedal spasms were distinctly different from myokymia or fasciculations.^{15,27}

Crotaline envenomation can result in hematologic effects ranging from mild to severe coagulopathy and

Table 1. Composition of *Crotalus oreganus concolor* venom based on percent area of reversed-phase high-performance liquid chromatography peaks and SDS-PAGE

Protein family	Percent
Myotoxin I and II	60
Concolor toxin subunit B	15
Snake venom serine proteinases	9
Concolor toxin subunit A	6
Phospholipase A2 (acidic)	6
Unidentified	3
Snake venom metalloproteinase (P-III)	1

Note that myotoxins I and II comprise the majority of venom proteins.

Table 2. Assays of *Crotalus oreganus concolor* venom for enzyme activities common to rattlesnake venoms

Enzyme assayed	Specific activity	Relative activity
Phospholipase A ₂ (nmol product·min ⁻¹ ·mg ⁻¹)	25.4	↓
Azocasein metalloproteinase (Abs 342 nm·min ⁻¹ ·mg ⁻¹)	0.01	↓↓
Kallikrein-like (nmol product·min ⁻¹ ·mg ⁻¹)	970.9	↑↑
Thrombin-like (nmol product·min ⁻¹ ·mg ⁻¹)	1388.8	↑↑
Phosphodiesterase (Abs 400 nm·min ⁻¹ ·mg ⁻¹)	1	↑
L-amino acid oxidase (nmol product·min ⁻¹ ·mg ⁻¹)	9.7	↓

↓, venom activity lower than average; ↑, venom activity higher than average

Activity levels in the responsible venom relative to averages of 22 taxa of rattlesnakes (Mackessy³³) are also indicated.

neurologic effects ranging from paresthesia to seizures and coma. Consequently, patients can exhibit multiple combinations of clinical manifestations varying in severity.²⁸ Thus, the absence of hematologic effects in our patient, in the presence of neurologic symptoms, was not uncharacteristic of rattlesnake envenomation. Neurologic symptoms after *C cerastes* and *C concolor* envenomation, without hematologic effects, have been reported.^{9,15-17}

Local wound complications in our patient were modest, including ecchymosis, swelling, pain, and skin sloughing. Elevated CK, a routinely used marker for tissue damage and myonecrosis, has been reported in 2 cases of *Crotalus oreganus concolor* envenomation (range 1800–17,000 IU·L⁻¹) and in the absence of coagulopathic complications.^{15,29}

Our patient's elevated CK value (3394 U·L⁻¹) at 22 h after envenomation was indicative of venom-induced local tissue/muscle damage to the bitten thumb.

Medications may influence snake envenomation symptoms, and our patient's medications (atenolol, rizatriptan, and amitriptyline) have reported adverse effects of blurred vision, numbness, chest tightness, paresthesia, and weakness similar to our patient's symptoms.^{30,31} Our patient reported having no adverse effects to these medications before the bite, suggesting drug/venom interactions were noncontributory to observed symptoms.

Venom analyses of the responsible snake revealed abundant PLA₂-based β neurotoxin (concolor toxin) and nonenzymatic peptide myotoxins.^{9,12,32} Enzyme analyses revealed the presence of 6 enzymes common to rattlesnake venoms (Table 2); however, their levels varied significantly from averaged values based on 22 taxa of rattlesnakes.³³ Notably, snake venom metalloproteinase activity (azocasein metalloproteinase) was barely detectable, and this likely was responsible for the lack of hemorrhage, rhabdomyolysis, or inflammation. These very low levels of snake venom metalloproteinase are characteristic of type II venoms, including that of *Crotalus oreganus concolor*.¹⁰ Thrombin-like and kallikrein-like serine proteinase activities in the venom were quite high, and thrombin-like activity was nearly twice the average value of many rattlesnake

venoms. Because of these high activity levels, coagulopathies including hypofibrinogenemia would have been expected; however, laboratory blood panels did not indicate any form of coagulopathy. Phospholipase A₂ levels were moderate but not noteworthy compared to other species; PLA₂ activity is sometimes associated with severe inflammation, myotoxicity, and occasionally renal damage/failure. The lack of these symptoms indicates that this acidic PLA₂ was not particularly toxic, consistent with similar enzymatic PLA₂s from other species.

RP-HPLC analysis of the venom (Figure 3) indicated the presence of myotoxins I and II, small peptide components common to the venoms of numerous species of rattlesnakes.³¹ In humans, they may be responsible for fasciculations (possibly muscle weakness, both acute and prolonged).²⁵ The myotoxin content of this venom was extremely high and comprised >59% of the total venom proteins; concolor toxin made up >21% of the venom total protein content (Figure 3, Table 1). The combined actions of these 2 protein families, comprising >80% of venom proteins, possibly contributed to the rapid paraesthesias and other tetany-like symptoms observed in the patient.

Conclusions

The case reported here is an interesting academic case in that certain symptoms, similar to previously reported cases of *Crotalus oreganus concolor* envenomation, may have been a hybridization of anxiety- and stress-induced symptoms combined with venom-induced symptoms. Analyses of ionized calcium and arterial blood gases and a formal neurologic evaluation would have provided for a more comprehensive assessment to confirm or rule out the source of some symptoms. Although the venom profile of the snake involved showed a high myotoxin content, this is unlikely to have played a role in the myoneurologic (carpopedal spasms) symptoms. Thus, the presented case illustrates a diagnostic conundrum, and medical personnel should be cognizant of the potential for confounding symptoms associated with rattlesnake envenomation and interpretation as to their cause.

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