Caseous Stomatitis Caused by *Pseudomonas aeruginosa* in *Boa constrictor amarali*

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**ABSTRACT**

**Background:** *Pseudomonas aeruginosa* is a bacterium that belongs to the microbiota of snakes, but it may also be an opportunistic pathogen and contaminate humans through fecal contact, bites, and injuries. In snakes, this microorganism may present high pathogenicity at certain conditions and have been associated with high morbidity and mortality. Reports of infection of *Boa constrictor* by this pathogen are rare. Thus, this study aimed to describe the *P. aeruginosa* oral infection in a snake specimen (*Boa constrictor amarali*), approaching the isolation and identification of the infectious agents involved, the antimicrobial sensitivity and resistance, and the therapeutic protocol adopted.

**Case:** A free-living adult female specimen of *Boa constrictor amarali* (Amaral’s boa), with no described previous history was rescued in an urban area by the Environmental Police. Clinical evaluations showed structures of caseous aspect in the oral cavity, with hyperemia spots in the mucosa. Samples of these lesions were sent for mycological examination, and fungal forms were not found. Samples were collected for isolation and culture. The antimicrobial susceptibility of the isolated microorganisms was determined by the modified Kirby-Bauer disk diffusion method. *P. aeruginosa* was isolated and showed susceptibility to amikacin, gentamicin, and polymyxin-B; intermediate susceptibility to azithromycin, and ciprofloxacin; and resistance to cephalixin, cefitiofur, chloramphenicol, and enrofloxacin. The treatment consisted of cleaning of the oral cavity, local infiltration of lidocaine for debridement of the caseous area that were later cauterized with iodine. Systemic antibiotic therapy was used, with intramuscular administration of amikacin (5 mg/kg) for the first dose and (2.5 mg/kg) for the other doses with intervals of 72 h, and oral administration of metronidazole (20 mg/kg) with intervals of 48 h, both during 21 days. Daily subcutaneous fluid therapy was performed as support treatment, using Lactated Ringer’s solution (25 mg/kg) and Vitamin C (10 mg/kg) with intervals of 24 h, being the cure observed at the end of treatment.

**Discussion:** This paper presents the pathological findings of the *Pseudomonas aeruginosa* oral infection in a *B. constrictor amarali*. This bacterium is an opportunistic pathogen that is commonly found in snakes, thus, humans in contact with these animals may be contaminated with this pathogen. However, oral cavity lesions associated with *P. aeruginosa* had not yet been related to *Boa constrictor amarali*, which is a non-venomous species. Few bacteria associated with reptile diseases are primary causative agents. Clinical bacterial infections generally tend to be secondary to viral infections. Predisposing factors for the development of bacterial diseases in these reptiles include immunodepression, malnutrition, poor adaptation to captivity, and the maintenance of these animals at temperatures and humidities outside their thermal comfort range. In the present study, the *P. aeruginosa* behaved as an opportunistic pathogen, resulting in clinical manifestations with caseous lesions in the oral cavity, probably due to an imbalance of the microbiota caused by stress or immunodepression. The antibiogram allowed the adoption of a correct therapeutic protocol based on the susceptibility of the pathogen, resulting in remission of lesions and clinical signs after 21 days of treatment.

**Keywords:** abscesses, bacteriosis, infectious diseases, snakes.
INTRODUCTION

_Boa constrictor amarali_ is a non-venomous snake of the Squamata order (Ophidia suborder) that is widely distributed in the Brazilian territory [9], especially in the Northeast, Southeast, and South regions [7].

Infectious diseases are the main causes of mortality of snakes, mainly by Gram negative microorganisms. The predominance of these bacteria in infectious processes is directly connected to the opportunistic character of the normal bacterial microbiota of the host, especially in infections caused by _Pseudomonas_ sp., _Aeromonas hydrophila_, _Proteus_ sp., _Salmonella_ sp., _Citrobacter_ sp., _Escherichia coli_, _Providencia_ sp. and _Xanthomonas maltophilia_ [9].

Reptiles host important zoonotic microorganisms, such as the bacterium _Pseudomonas aeruginosa_, therefore, humans can be contaminated with these microorganisms through fecal contact, bites, and injuries [4]. _Pseudomonas_ spp. may present high pathogenicity at certain conditions and have been associated with high morbidity and mortality in infected animals due to their ability to develop resistance to antibiotics and express multiple virulence factors [14].

_P. aeruginosa_ can be isolated from samples of diseased and healthy reptiles. The first cases of cross contamination between snakes and their healthy owners have been reported recently [1]. _P. aeruginosa_ is a microorganism that belongs to the microbiota of reptiles, especially in their oral cavity and intestinal tract; however, it can be an opportunistic pathogen with several clinical manifestations in susceptible animals [2].

CASE

A free-living adult female specimen of _Boa constrictor amarali_ (Amaral’s boa), with no described previous history was rescued in an urban area by the Environmental Police and taken to the Laboratory of Teaching and Research in Wild Animals of the Veterinary Hospital of the Federal University of Uberlândia (VH-UFU), Minas Gerais, Brazil. The animal was weighed in a digital scale and presented 3.2 kg.

The clinical evaluation showed dehydration signs, and presence of caseous structures in the oral cavity, in the upper-left, upper-right, and lower-right tooth regions, with hyperemia spots in the mucosa (Figure 1A). Samples of these lesions were collected and sent to the Veterinary Clinical Laboratory of the VH-UFU for mycological examination. This material was placed in slides and analyzed in an optical light microscope (10x and 40x objective) using low drops of 10% potassium hydroxide (Anidrol®) to clear the sample, but fungal forms were not found.

Samples were collected by passing a sterilized alginate cotton tip swab through the entire lesion. The sample was stored in a plastic tube containing a semi-solid transport medium Stuart (Absorve®) and sent to the Laboratory of Infectious Diseases of UFU. The sample was then transferred to a tube containing Thioglycollate broth (Himedia®) - a highly nutritive medium that helps the growth of microorganisms and incubated in a bacteriological oven at 37°C for 24 h [12].

Samples were plated for bacterial isolation on one Petri dish containing blood agar and other containing MacConkey agar (TM Media®), using the streaking technique with a platinum inoculating loop. These dishes were placed in a bacteriological oven at 37°C for 24 h for incubation [16]. Gram staining was carried out to identify gram-positive and gram-negative bacteria in the blood agar (Kasvi®) colonies. Colonies grown on MacConkey agar were identified through biochemical tests in the mediums phenol red, lysine, phenylalanine, citrate, urea, and SIM (sulfide, indole, motility), screening colonies that grow on selective medium for gram-negative bacteria of the Enterobacteriaceae family. A medium was used for each different colony grown on MacConkey agar to identify each bacterial genus or species. The oxidation-fermentation test was included for colonies of gram-negative bacteria that were not of the Enterobacteriaceae family.

![Figure 1. Oral cavity of a specimen of _Boa constrictor amarali_ with caseous stomatitis. A- Presence of caseous structures in the upper-left (white arrow), upper-right (red arrow), and lower-right (black arrow) tooth regions, with hyperemia spots in the mucosa (circles). B- Absence of caseous lesions and hyperemia spots in the mucosa after 21 days of treatment.](image-url)
Colonies of *Pseudomonas* spp. grew on MacConkey agar medium; these bacteria were identified by their colony morphology and typical odor. Biochemical tests confirmed this genus and identified the bacterium species *Pseudomonas aeruginosa*.

The antimicrobial susceptibility of the isolated microorganisms was determined by the modified Kirby-Bauer disk diffusion method [12]. The isolates were inoculated in Mueller Hinton (Kasvi®)® broth and incubated at 37°C until showing turbidity of 0.5 according to the Mac Farland scale. The isolates were streaked on Petri dishes containing Muller Hinton agar medium, using a swab. Subsequently, the antimicrobial drugs enrofloxacin, ciprofloxacin, amikacin, gentamicin, chloramphenicol, azithromycin, polymyxin-B, and cephalexin (Laborclin®)® were applied to the impregnated discs [9]. They were then incubated in an oven at 37°C for 24 h and the formed halos were evaluated to determine the susceptibility profile of the isolated pathogens.

*P. aeruginosa* presented susceptibility to amikacin, gentamicin, and polymyxin-B; intermediate susceptibility to azithromycin, and ciprofloxacin; and resistance to cephalexin, cephalothin, chloramphenicol, and enrofloxacin.

The treatment consisted of cleaning of the oral cavity with 0.9% saline solution (0.9% Saline) and oral 2.0% chlorhexidine (Riohex®), local infiltration of 2.0% lidocaine (Xylestesin®) for debridement of the caseous lesions, and cautery of the lesions with iodine (Riodeine®). Local cleaning and cautery with iodine were performed daily at lesion sites.

Based on the antimicrobial susceptibility testing (AST) and according to Klaphake [8], amikacin (Amicilon®)® was administered intramuscularly (5 mg/kg) for the first dose and 2.5 mg/kg for the other doses with intervals of 72 h, combined with oral administration of metronidazole [Benzoilmeteronidazol®]® at dose of 20 mg/kg with intervals of 48 h, both during 21 days. Metronidazole® can be given simultaneously with amikacin® for a larger spectrum.

Daily subcutaneous fluid therapy was performed as support treatment, using Lactated Ringer’s® solution (25 mg/kg) and Vitamin C [Monovin C®]® - 10 mg/kg] with intervals of 24 h [8]. Vitamin C can be a coadjuvant in cases of stomatitis and cutaneous conditions, and a supportive therapy for bacterial infections. Remission of lesions and clinical signs was observed after 21 days of treatment (Figure 1B).

The animal remained under controlled temperature of 30°C throughout the treatment for better metabolism of the drugs.

**DISCUSSION**

Infectious agents can be transmitted to snakes through wounds caused by trauma, and contact with contaminated environments, handle, and utensils. However, immunodepression, malnutrition, poor adaptation to captivity, and the maintenance of these animals at temperatures and humidities outside their thermal comfort range are considered predisposing factors for the development of bacterial diseases in these reptiles [15]. However, few bacteria associated with reptile diseases are primary causative agents. Clinical bacterial infections generally tend to be secondary to viral infections [8].

*Pseudomonas* spp. belong to the microbiota of snakes, especially in their oral cavity [4,6,17]. However, in conditions of high pathogenicity, clinical manifestations and lesions can develop in their oral cavity and even in other organs [4,11,18]. In the present study, the *P. aeruginosa* behaved as an opportunistic pathogen, resulting in clinical manifestations with caseous lesions in the oral cavity, probably due to an imbalance of the microbiota caused by stress or immunodepression.

*P. aeruginosa* has been isolated from skin vesicles of a coral snake (*Micrurus corallinus*), scales with caseous content of a sucuri snake (*Eunectes marinus*) [5], and subcutaneous nodules of a boa snake (*Boa constrictor amarali*) [11]. However, oral cavity lesions associated with *P. aeruginosa* had not yet been related to *Boa constrictor amarali*, which is a non-venomous species.

*P. aeruginosa* is an opportunistic pathogen for animals and humans, associated with high morbidity and, sometimes, high mortality. Studies have shown that captive reptiles have been carriers of *P. aeruginosa*, thus, they are sources of contaminations for humans who interact with these animals [3,4].

*P. aeruginosa* from skin lesions of snakes was isolated in previous studies and was observed resistance to all antibiotics tested in the AST; even with the establishment of systemic and topical antibiotic therapy, 90.9% of the animals died, denoting the multiresistance of the isolated agents [5]. Cases
of multiresistant bacteria isolated from reptiles has been reported [11,13,19], showing the limitation of the therapeutic protocols adopted [5] and the need of AST to determine the susceptibility or resistance profile of these agents to antimicrobials.

In the present study, the pathogen evaluated was not resistant to all antimicrobials tested, but it was resistant to cephalaxin, ceftiofur, chloramphenicol and enrofloxacin; moderately resistant to azithromycin, and ciprofloxacin; and susceptible to amikacin, gentamicin, and polymyxin-B. The AST allowed the adoption of a correct therapeutic protocol, based on the susceptibility of the microorganism, resulting in remission of lesions and clinical signs after 21 days of treatment.

REFERENCES


