**Devriesea agamarum** associated cheilitis in a North African spiny-tailed lizard (*Uromastyx acanthinura*) in Spain

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**Abstract**

An 11-year-old, male North African spiny-tailed lizard (*Uromastyx acanthinura*) was presented with hyperkeratotic cheilitis. Based on clinical examination, histology and microbiological testing, *Devriesea agamarum* was identified as the causative agent of the dermatological disorder. Moreover, an abundant culture of *D. agamarum* was obtained following sampling of the oral cavity. Cheilitis was present for three years in the spiny-tailed lizard and during the first year of that period the lizard was housed together with an ocellated lizard (*Timon lepidus*), an Algerian orange-tailed skink (*Eumeces algeriensis*) and a female *U. acanthinura*. The latter showed signs of chronic dermatitis and had deceased approximately 2.5 years prior to initial presentation of the male dab lizard because of renal failure. The other cohabiting lizards showed no dermal lesions and *D. agamarum* could not be demonstrated following intestinal, cloacal and oral sampling. Recurrence of the bacterial skin infection was observed following a first antimicrobial treatment. This was considered to result from failure to eliminate the bacterium from the treated dab lizard or re-infection from the environment or asymptomatic carriers. A second treatment, including disinfection of the enclosure with household bleach 0.1%, weekly mechanical debridement of the crusty lesion and treatment with ceftazidime at 20 mg/kg intramuscularly every 72 hours for 57 days resulted in resolution of the skin lesion, elimination of *D. agamarum* from the oral cavity based on repetitive microbiological sampling and no recurrence of the lesions during a 31 month follow up period. The present case is the first report of devrieseasis in Spain and highlights the importance of a multidirectional diagnostic and therapeutic approach towards controlling devrieseasis in captive lizard collections. Several disease aspects such as persistency are discussed in the light of the contemporary available literature.

**Keywords:** Cheilitis, Dermatitis, *Devriesea agamarum*, *Uromastyx acanthinura*.

**Introduction**

*Devriesea agamarum* has been isolated previously in captive lizards from Belgium, Austria, Switzerland, Croatia and from free-ranging lizards species from the French Caribbean island of Saint Barthélemy (Martel et al., 2008; Lukac et al., 2013; Schmidt-Ukaj et al., 2014; Rossier et al., 2016; Hellebuyck et al., 2017). The bacterium is considered worldwide as an important agent of dermal disease and/or septicaemia that has eradicated entire captive collections of lizards. Moreover, the bacterium was recently demonstrated to affect free-ranging endangered Antillean iguanas (*Iguana delicatissima*) and should be considered as one of the first bacterial diseases that might pose a conservation threat to free-ranging squamates (Hellebuyck et al., 2017).

This is the first reported case of devrieseasis in Spain; moreover, several disease aspects such as persistency are discussed in the light of the contemporary available literature.

**Case Details**

An 11-year-old male North African spiny-tailed lizard (*Uromastyx acanthinura*) weighing 800g, purchased at a reptile pet shop in Madrid eight years before initial presentation, was presented with a history of progressive crusty cheilitis since three years. Since two months, a 10% povidone-iodine solution was topically applied on a daily basis by the owner and resulted in a mild improvement of the skin lesions. The spiny-tailed lizard was housed with a female during a one-year-period that died 2.5 years prior to initial presentation of the male at the age of 24 years. The female had been raised in captivity since birth and was born out of wild caught animals originating from Morocco. Besides showing cheilitis and dermatitis with an identical clinical appearance as observed in the male, severe renal and visceral gout were determined as the eventual cause of death in the female spiny-tailed lizard based on gross pathological and histopathological examination.

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An adult male Algerian orange-tailed skink (*Eumececs algeriensis*) and a female ocellated lizard (*Timon lepidus*) of unknown age were introduced to the same enclosure as the male *Uromastyx* a couple of months after the female had died but remained clinically unaffected.

At the time of initial presentation, four full thickness biopsies were collected from the affected area of skin in the male lizard for microbiological and histopathological examination. Anesthesia was performed with intravenous propofol into the ventral tail vein (Lipuro®, B. Braun Spain, 08191 Barcelona, Spain; 10 mg/kg). The biopsies were fixed in formalin and embedded in paraffin followed by hematoxylin and eosin (HE) staining as well as gram staining. Histological evaluation of the sections of the skin biopsies revealed severe multifocal epidermal hyperplasia with marked hyperkeratosis, mild multifocal infiltration of lymphocytes and heterophils in the dermis with heterophilic exocytosis, intercellular edema and intralesional, predominantly filamentous gram positive bacteria with a beaded appearance and occasional branching within the keratin layers. Abundant growth of a gram negative bacillus belonging to the *Enterobacteriaceae* that could not be identified using conventional biochemical testing was observed following culturing of the skin lesion. The latter isolate was considered as a contaminant or normal part of the skin microbiota taking into account the presence of exclusively gram positive beaded and branching filamentous bacteria in the histological sections of the skin biopsies.

Treatment with ceftazidime (Ceftazidima Normon injectable 1g, Laboratorios Normon, 28760 Madrid, Spain; 15mg/kg intramuscularly every 72 hours) was initiated during three weeks and resulted in complete resolution of the skin lesions. During the next months, however, cheilitis gradually reappeared and the severity of the skin lesions was identical to those observed during initial presentation approximately eighteen months later (Figure 1).

At that moment, swabs were collected from the oral cavity and the scaly lip lesions as well as crusts that were obtained following debridement of the crusty lesions for microbiological examination. All samples were cultured during 24–48 h on colistin nalidixic acid (CNA, Oxoid GmbH, Wesel, Germany) agar at 37 °C and 5% CO₂. Pure cultures and abundant growth of small, whitish colonies of gram positive bacteria with a narrow zone of hemolysis were observed following culturing of the skin and oral samples respectively. Based on morphological aspects and the results of API Coryne, API 20 STREP, API 50 CH (bioMérieux, Marcy l’etoile, France) testing, the isolates were identified as *D. agamarum*.

Following 16S rRNA gene sequencing as previously described (Martel *et al*., 2008), 100 % similarity with the type strain of *D. agamarum* (IMP2 = LMG 24257T) was revealed for all isolates.

In addition, a radiological study of the head and whole body, as well as ultrasound were performed in the spiny-tailed lizard but did not reveal bone lesions, masses or other abnormalities, that could indicate disseminated bacterial infection.

Blood was collected by venipuncture of the ventral coccygeal vein in a heparinized tube for a serum biochemistry profile and hematologic evaluation at the time of radiological examination. A microcapillar filled with heparinized blood was centrifuged at 15,800 rpm for 2 minutes (StatSpin® VT, Iris Sample Processing, Massachusetts 02090-1825, USA) for hematocrit determination. The total white blood cell count by the Natt and Herrick method was obtained with heparinized blood (Campbell and Ellis, 2007).

The rest of the blood was centrifuged at 15,800 rpm for 90 seconds in the same centrifuge and the plasma obtained was used to obtain a basic biochemical profile (Bird/Reptile Rotor Profile for VetScan® VS2, Abaxis, California 94587, USA). Blood biochemistry and hematology results and reference values are presented in Table 1. To date, reference values for North African spiny-tailed lizards are not available in the literature, so values from closely related species (*Uromastyx acanthinura* and *Uromastyx leptieni*) were used for guidance (Naldo *et al*., 2009). Uric acid, albumin and aspartate aminotransferase were mildly raised (Table 1). Dehydration and mild tissue damage due to loss of appetite and chronic dermal inflammation were considered to most likely have caused these biochemical alterations (Campbell, 2014). Samples from the oral cavity and skin obtained from the orange-tailed skink and the ocellated lizard that cohabited with the dab lizard were submitted to microbiological examination as described above but cultured negative for *D. agamarum*.
Table 1. Hematology and blood chemistry of a North African spiny-tailed lizard (Uromastyx acanthinura) with devrieseasis.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Uromastyx acanthinura with devrieseasis</th>
<th>Reference values (U. aegyptia microlepis and U. leptieni)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCV (%)</td>
<td>31</td>
<td>4.9 - 44.5</td>
</tr>
<tr>
<td>WBC</td>
<td>5.2</td>
<td>1 - 8.1</td>
</tr>
<tr>
<td>Heterophils (x1000)</td>
<td>3.64</td>
<td>0.59 - 5.36</td>
</tr>
<tr>
<td>Lymphocytes (x1000)</td>
<td>1.3</td>
<td>0.27 - 4.05</td>
</tr>
<tr>
<td>Monocytes (x1000)</td>
<td>0.26</td>
<td>0 - 0.5</td>
</tr>
<tr>
<td>Total protein (g/dL)</td>
<td>5.2</td>
<td>2.6 - 7.4</td>
</tr>
<tr>
<td>Albumin (g/dL)</td>
<td>3.3**</td>
<td>1.2 - 3.1</td>
</tr>
<tr>
<td>Globulins (g/dL)</td>
<td>1.9</td>
<td>-</td>
</tr>
<tr>
<td>Uric acid (mg/dL)</td>
<td>8.78**</td>
<td>0.3 - 7.3</td>
</tr>
<tr>
<td>Glucose (mg/dL)</td>
<td>188</td>
<td>67.7 - 355.6</td>
</tr>
<tr>
<td>Calcium (mg/dL)</td>
<td>11.8</td>
<td>7.2 - 13.2</td>
</tr>
<tr>
<td>Phosphorus (mg/dL)</td>
<td>4.8</td>
<td>-</td>
</tr>
<tr>
<td>Sodium (mEq/L)</td>
<td>161</td>
<td>-</td>
</tr>
<tr>
<td>Potassium (mEq/L)</td>
<td>4.1</td>
<td>-</td>
</tr>
<tr>
<td>AST (UI/L)</td>
<td>183**</td>
<td>28.5 - 172</td>
</tr>
<tr>
<td>Bile Acids (umol/L)</td>
<td>37</td>
<td>-</td>
</tr>
</tbody>
</table>

* = (Naldo et al., 2009); ** = Values above the reference value.

The owner was advised to perform thorough cleaning and disinfection of the terrarium with bleach 0.1% and ceftazidime was administered at 20 mg/kg intramuscularly every 72 hours for 57 days. In addition, the crusty skin lesion was debrided mechanically every 7 days until the affected skin site regained its normal appearance. Cheilitis resolved after 50 days of antimicrobial treatment, local debridement and the passing of two shedding cycles (Figure 2). The previously affected area of skin as well as the oral cavity were resampled for microbiological evaluation and tested negative for the presence of D. agamarum.

Discussion

Currently, Devriesea agamarum is the only member of a newly described genus within the class Actinobacteria. This bacterium is a non-motile, non-spore-forming and non-acid-fast gram positive small rod that occur singly, in pairs or in short chains and that grow on sheep blood agar as small hemolytic mucoid and whitish colonies after 24 h of incubation at 25-42 °C under aerobic, microaerophilic or anaerobic conditions (Martel et al., 2008). Partial ribosomal RNA gene sequences and the full genome sequence of the D. agamarum strain IMP2 and a partial 16S ribosomal RNA gene sequence from a bearded dragon with concurrent Chrysosporium guaari infection have been deposited in the European Nucleotide Archive (Schmidt-Ukaj et al., 2014; Haesendonck et al., 2015).

While disease associated with this bacterium initially seemed restricted to dry land lizards, recent reports have documented devrieseasis in several lizard species that originate from (sub)tropical regions (Bauwens et al., 2014; Rossier et al., 2016; Hellebuyck et al., 2017). Various lizard species can act as asymptomatic carriers in which D. agamarum may be part of the microbiota of the oral cavity or the cloaca (Hellebuyck et al., 2009a; Devloo et al., 2011; Bauwens et al., 2014; Rossier et al., 2016).

The presence of asymptomatic carriers, considered as one of the factors that promote persistency in captive lizard collections, has also been put forward as a possible origin of devrieseasis in free-ranging iguanas (Hellebuyck et al., 2017). Besides the presence of asymptomatic carriers, long term environmental survival has been demonstrated to be another major contributing factor that leads to persistency of devrieseasis (Hellebuyck et al., 2010).

It is possible that persistency of the bacterium in the enclosure of the dab lizard from the present report combined with the presence of D. agamarum in the oral cavity, might explain therapeutic failure following the initial antimicrobial treatment of the dab lizard and recurrence of the skin lesions (Pasmans et al., 2008; Devloo et al., 2011). Although D. agamarum could not be isolated from the healthy ocellated lizard and the Algerian orange-tailed skink that cohabited with the North African spiny-tailed lizard, screening of healthy lizard should preferably be based on repetitive sampling of oral skin and cloacal swabs as D. agamarum is a relatively slow growing coryneform bacterium. Consequently, cultures derived from samples obtained from skin lesions or oral cavities collected in lizards with dermatitis are easily overgrown by other bacteria that might be part of the saurian microbiota.

Unless pure D. agamarum cultures are obtained, the diagnosis of devrieseasis as a cause of dermatological disease in lizards may easily be missed, especially if analyses are performed by microbiological laboratories that are not familiar with the bacterium.

Fig. 2. Lips in shedding after ceftazidime 20mg/kg every 48 hours for 57 days.
Following morphological identification, 16S rRNA sequencing is necessary to confirm the identity of the obtained isolate as *D. agamarum* (Martel et al., 2008). Skin disease in lizards and in other reptiles is primarily the result of inappropriate management in captivity, leading to secondary infection (Hellebuyck et al., 2012). A few pathogens such as *D. agamarum*, however, may act as primary etiological agents. In the case of *D. agamarum*, a breach of skin integrity seems necessary to allow the bacterium to establish experimental clinical infection (Hellebuyck et al., 2008; Hellebuyck et al., 2017). In naturally *D. agamarum* infected lizards, infection also seems to be correlated with traumatic lesions resulting from territorial or reproductive behavior (Bauwens et al., 2014; Hellebuyck et al., 2017). The deceased female spiny-tailed lizard that showed identical skin lesions and was housed together with the male during a one-year-period is presumed as the most likely source of infection. Although infection may have occurred through the environment following contamination by the female lizard, direct contact or infection of biting lesions that are often witnessed during territorial or reproductive behavior in dab lizards, may also have promoted transmission and the onset of clinical infection in the male dab lizard. Indirect factors that impaired skin integrity in the affected male *Uromastyx* could not be identified.

The most characteristic clinical finding presented in lizards suffering from *D. agamarum* associated disease is the chronic, crusted dermatitis. Some species, especially *Uromastyx* spp. and *Crotaphytus collaris*, show chronic skin lesions associated with high morbidity and low mortality. If the infection persists for a long time, however, the dermal lesions may interfere with normal feeding and foraging behavior and eventually become fatal (Martel et al., 2008; Hellebuyck et al., 2009a; Devloo et al. 2011; Lukac et al., 2013; Bauwens et al., 2014).

In other species such as *Agama imperialis*, *Sauromalus obesus* and *Physignathus cocincinus*, however, high morbidity and mortality is frequently observed resulting from the development of septicemia after showing minimal skin disease in most of the cases (Martel et al., 2008; Pasmans et al., 2008; Rossier et al., 2016).

*Devriesea agamarum* infection in the male spiny-tailed lizard presented in this case report was limited to the skin and no radiological, hematological or biochemical abnormalities indicative for systemic infection could be demonstrated. Other agents that might cause identical skin disease as seen for *D. agamarum* should be properly excluded (Hellebuyck et al., 2012; Schmidt-Ukaj et al., 2014).

Especially, dermatophytosis caused by *Nannizziosis* spp. and *Paranannizziosis* spp. (formerly known as CANV) in lizards should be differentiated as the clinical presentation as well as the predilection sites of affected skin are highly similar (Paré and Sigler, 2016). In the present case a tentative diagnosis of *D. agamarum* associated cheilitis was based on clinical examination taking into account the species predisposition, but the definitive diagnosis was made by bacteriological examination of the skin lesions and 16S rRNA sequencing of the obtained isolate.

Initial treatment with ceftazidime at 15 mg/kg intramuscularly every 72 hours was sufficient to resolve the skin lesions in the present case, but cheilitis recurred. Accordingly, it was decided to increase the dose of ceftazidime to 20 mg/kg every 72 hours, clean the enclosure with bleach and debride the lesions once a week. After 57 days of antimicrobial treatment, the resolution of skin lesions and negative cultures as well as the non-recurrence of lesions during a follow-up period of 31 months confirmed the efficacy of the applied treatment and the importance of a multidirectional approach towards the control of *D. agamarum* associated disease. In a reported case series, *D. agamarum* infected dab lizards that were treated for 15 days with ceftazidime at 10 mg/kg every 72 hours in conjunction with rinsing the lesions with an antiseptic and surgical debridement in one of the cases resulted in successful outcome in all treated lizards (Lukac et al., 2013).

For the present case, the empirical treatment was presumed to be ineffective due to inadequate circulating levels of the used antimicrobial or insufficient penetration of the poorly vascularized affected skin sites. Because in turtles and snakes the administration of ceftazidime at a dose of 20-22 mg/kg every 72 hours proved to achieve therapeutic plasma concentrations (Gibbons, 2014), the dose used in the second treatment applied in the *Uromastyx* in this case report was increased to 20 mg/kg every 72 hours. Unfortunately, there are no pharmacokinetic / pharmacodynamic studies in relation to the use of ceftazidime in *Uromastyx*. In the present report, a case of cheilitis caused by *Devriesea agamarum* is described for the first time in Spain. Based on the available literature and the widespread occurrence of asymptomatic carriers, devriesesiasis may be severely underdiagnosed. Mainly difficulties to correctly sample and interpret cultures derived from *D. agamarum* associated skin lesions may largely contribute to this.

The recurrence of the dermal lesions in the present case is a typical feature that is associated with devriesesiasis in captive lizard collections and besides the use of appropriate treatment protocols, a multidirectional control strategy is mandatory to resolve environmental survival of the bacterium and to identify asymptomatic carriers, two major factors that promote persistency of the disease.
Natural or acquired resistance against antimicrobial agents has not been documented so far (Hellebuyck et al., 2009b, 2010), but the importance of susceptibility testing of any *D. agamarum* isolate cannot be overly emphasized.

**Conflict of interest**
The Authors declare that there is no conflict of interest.

**References**


