Antimicrobial susceptibility and molecular characterization of Salmonella serovar Ndolo isolated from outbreaks in cattle and horses

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INTRODUCTION

The genus Salmonella has been associated with enteric and systemic infections in animals and humans, and is one of the most important zoonoses worldwide (CONRAD et al., 2017). Clinical manifestations of salmonellosis vary depending upon the serovar involved and the host’s susceptibility (BARROW et al., 2010; JUFFO et al., 2016). The most common Salmonella serovars responsible for outbreaks in calves are Typhimurium and Dublin (COURA et al., 2015; MOHLER et al., 2009); whereas, Typhimurium, Anatum, Newport, and Agona are serovars commonly reported in foals (CUMMINGS et al., 2016; OLIVO et al., 2016). Interestingly, some Salmonella serovars, such as Dublin in calves or Choleraesuis in pigs, are classified as host-adapted strains and zoonotic. Previously, this serovar was identified only in human infections. The presence of relevant virulence genes in all Salmonella Ndolo isolates and the detection of antimicrobial multi-resistant strains highlighted the importance of monitoring serovars associated with salmonellosis in domestic animals.

Key words: salmonellosis, zoonoses, antimicrobial resistance, ERIC–PCR, Salmonella Ndolo.

ABSTRACT: The present study aimed to describe and characterize, for the first time, two outbreaks of salmonellosis caused by Salmonella Ndolo in foals and calves in Brazil and compare the isolated strains with S. Ndolo previously identified in asymptomatic reptiles. The affected calves and foals presented fever, lethargy, and profuse diarrhea. Isolated strains were subjected to antimicrobial susceptibility testing, characterized according to virulence genes, and fingerprinted by ERIC-PCR. Salmonella Ndolo was identified in fecal samples from two foals and four calves. One isolate from a calf was resistant to amoxicillin/clavulanic acid, trimethoprim/sulfamethoxazole, and florfenicol. Strains from two other calves were resistant to oxytetracycline. All virulence genes tested were present in the isolates, and two major clusters of closely related strains were identified by ERIC-PCR, each per outbreak. This is the first report of Salmonella Ndolo infection in domestic and symptomatic animals. Previously, this serovar had been identified only in human infections. The presence of relevant virulence genes in all Salmonella Ndolo isolates and the detection of antimicrobial multi-resistant strains highlighted the importance of monitoring serovars associated with salmonellosis in domestic animals.

Susceptibilidade antimicrobiana e caracterização molecular de isolados de Salmonella serovar Ndolo de bovinos e equinos

RESUMO: O objetivo do presente estudo foi descrever e caracterizar, pela primeira vez, dois surtos de salmonelose causados por Salmonella Ndolo em potros e bezerros do Brasil e comparar esses isolados com Salmonella Ndolo previamente identificada em répteis assintomáticos. Os animais infectados apresentaram febre, letargia e diarreia profusa. Os isolados foram submetidos a testes de susceptibilidade a antimicrobianos e foram caracterizados conforme a presença de genes de virulência e diversidade genética, utilizando-se o ERIC-PCR. Salmonella Ndolo foi identificado em amostras fecais de dois potros e quatro bezerros. Um isolado de bezerro foi resistente a amoxicilina/acido clavulânico, trimetoprima/sulfametoxazol e florfenicol. Estírpes de dois outros bezerros foram resistentes a otxettracyclina. Todos os genes de virulência testados foram identificados nos isolados e dois grandes grupos de estírpes geneticamente relacionadas foram identificados pelo ERIC-PCR, um para cada surto. Esse é o primeiro relato de Salmonella Ndolo em animais domésticos e sintomáticos. Previamente, este serovar foi identificado apenas em infecções humanas. A presença de fatores de virulência relevantes em todos os isolados e a detecção de estírpes multirresistentes a antimicrobianos destaca a importância do monitoramento de sorovares associados a salmonelose em animais domésticos.

Palavras-chave: salmonelose, zoonose, resistência antimicrobiana, ERIC–PCR, Salmonella Ndolo.

INTRODUCTION

The genus Salmonella has been associated with enteric and systemic infections in animals and humans, and is one of the most important zoonoses worldwide (CONRAD et al., 2017). Clinical manifestations of salmonellosis vary depending upon the serovar involved and the host’s susceptibility (BARROW et al., 2010; JUFFO et al., 2016). The most common Salmonella serovars responsible for outbreaks in calves are Typhimurium and Dublin (COURA et al., 2015; MOHLER et al., 2009); whereas, Typhimurium, Anatum, Newport, and Agona are serovars commonly reported in foals (CUMMINGS et al., 2016; OLIVO et al., 2016). Interestingly, some Salmonella serovars, such as Dublin in calves or Choleraesuis in pigs, are classified as host-adapted strains and infect a limited number of species (BARROW et al., 2010). However, other Salmonella serovars, such as Typhimurium, are known to infect a broad range of host species and are considered important zoonotic agents (VRBOVA et al., 2018).
Although, some *Salmonella* serovars have been well-characterized as infectious agents in humans and animals, the potential pathogenic role of many other *Salmonella* serotypes is unknown. In this context, previous reports of *Salmonella* Ndolo infection are limited to humans from Brazil and several European countries (BERTRAND et al., 2013; KAUFFMANN et al., 1950; LEAL et al., 1987). The present research aimed to described and characterized, for the first time, two outbreaks of salmonellosis caused by *Salmonella* Ndolo in foals and calves in Brazil. Isolates obtained during these two outbreaks were evaluated for antimicrobial susceptibility, presence of virulence factors associated with *Salmonella* spp., and genetic diversity using ERIC-PCR.

**MATERIALS AND METHODS**

**First outbreak**

The first *Salmonella* Ndolo outbreak occurred in 2014 on a horse farm located in the metropolitan area of Belo Horizonte (Minas Gerais, Brazil). The owner reported the presence of diarrhea in three foals between 1 and 4 months old. Despite treatment with trimethoprim/sulfamethoxazole and flunixin meglumine for three days, one foal died and the other two exhibited no clinical improvement. Upon veterinary examination, foals were febrile with profuse, watery, malodorous diarrhea. Fecal samples were obtained rectally after digital stimulation from both diarrheic foals. Additionally, fecal samples were collected from all other horses (n=14) on the affected farm, including two other healthy foals and twelve healthy, adult horses. The samples were stored at 4°C until processing was performed at the Bacteriology and Research Laboratory at the School of Veterinary, Universidade Federal de Minas Gerais.

**Second outbreak**

The second *Salmonella* Ndolo outbreak occurred in 2017 on a dairy farm with approximately 600 lactating cows housed in free-stall barns, located in the Cristalina municipality (Goiás, Brazil). The owner reported an increased occurrence of diarrhea and weight loss in calves between 45 and 90 days old. They were housed in six groups of ten to twelve calves. Calves that developed diarrhea were promptly treated by the owner with amoxicillin, enrofloxacin, and flunixin meglumine. Despite treatment, nine calves died. Upon veterinary examination, several calves from this age group were lethargic, febrile, dehydrated, and passing diarrhea that contained fibrin fragments and undigested blood. Fecal samples from six affected calves were collected rectally and stored at 4°C until processed at the same Laboratory of the first outbreak.

**Isolation and characterization of Salmonella spp.**

Isolation of *Salmonella* spp. from the fecal samples was performed as previously described (RAMOS et al., 2018). Briefly, fecal samples were incubated in tetrathionate broth (Oxoid, USA) followed by plating on Hektoen enteric agar (BD, Germany). Identification of the *Salmonella* genus, species, and subspecies was performed as previously described (KWANG et al., 1996; LE MINOR & POPOFF, 1987). Characterization of the specific *Salmonella* serovar was determined according to antigenic characterization (GRIMONT & WEILL, 2007) at the Brazilian National Reference Laboratory of Enterobacteriaceae of the Oswaldo Cruz Foundation (FIOCRUZ - Brazilian Ministry of Health). Other analyses were performed on the fecal samples using previously described methods, including: detection of *Lawsonia intracellularis* by PCR (JONES et al., 1993), isolation and genotyping of *Clostridium perfringens* (DINIZ et al., 2017), isolation of *Escherichia coli* using MacConkey agar (Prodimol Biotechnology, Brazil) followed by detection of common virulence genes for diarrheagenic *E. coli* by PCR (FRANCK et al., 1998), detection of rotavirus and coronavirus by RT-PCR (ASANO et al., 2010) and parasite detection by flotation with Sheather’s sugar solution followed by light microscopy. Additionally, the foals’ samples was submitted to A/B toxin detection using a commercial enzyme immunoassay (*C. difficile* Tox A/B II - Techlab Inc., USA), and isolation of *Clostridium difficile* (DINIZ et al., 2017).

The isolated strains of *Salmonella* spp. from both outbreaks was subjected to antimicrobial susceptibility testing by disk diffusion method according to the Clinical and Laboratory Standards Institute (CLSI) manual VET01-A4 (CLSI, 2013). The following antimicrobials commonly used in animals, including for the treatment of salmonellosis (CLSI, 2015; OIE, 2015; PARVATHI et al., 2011), were tested: oxytetracycline (30µg), marbofloxacin (5µg), florfenicol (30µg), amoxicillin/clavulanic acid (30µg), trimethoprim/sulfamethoxazole (25µg), ceftriaxone (30 µg), and enrofloxacin (5µg) (DME, Brazil).

*Salmonella* spp. strains were assessed using previously described PCR methods to evaluate the presence of relevant, known virulence genes: *invA*, *prgH*, *sopB*, *toIC*, *spaN*, *orgA*, *pefA*, *ironN*, *spaH*, *pagC*, *msgA*, *sipB*, and *spvC* (MOHAMED et al., 2014).

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SKYBERG et al., 2006). Additionally, to evaluate the genetic diversity between isolates from these two outbreaks, the Salmonella strains were fingerprinted by ERIC–PCR using the primers ERIC-1 and ERIC-2 as previously described (LIM et al., 2005; SMITH et al., 2011; VERSALOVIĆ et al., 1991).

RESULTS AND DISCUSSION

Salmonella Ndolo were isolated from both the diseased foals and from four of the six calves tested. Salmonella Typhimurium was isolated from one of the healthy, adult horses. E. coli was isolated in the fecal samples from all calves and foals sampled, but no virulence factors were detected. C. perfringens type A was isolated from three calves. No other enteropathogens were detected. After confirming infection with Salmonella spp., the foals were treated with ceftiofur and the diarrhea resolved within five days of treatment. Treatment with enrofloxacin was initiated for all calves 45 to 90 days old and, the calves with diarrhea were treated with florfenicol in addition to oral and intravenous fluid therapy.

To our knowledge, this is the first report of Salmonella Ndolo infection in domestic animals. We also recently reported fecal shedding of this serovar in two healthy reptiles in Brazil (RAMOS et al., 2018). Furthermore, Salmonella Ndolo has been isolated from snails in Nigeria (OBIO et al., 1980), as well as, from infected humans in several European countries and Brazil (BERTRAND et al., 2013; KAUFFMANN et al., 1950; LEAL et al., 1987). These reports of colonization or infection by Salmonella Ndolo in a broad range of unrelated host species, such as reptiles, calves, foals, and humans, suggesting that it may be a multi-host pathogen (BARROW et al., 2010).

Five clinical isolates of Salmonella Ndolo from foals (n=2) and calves (n=3) were subjected to antimicrobial susceptibility testing by disk diffusion method according to the Clinical and Laboratory Standards Institute (CLSI) manual VET01-A4 (CLSI, 2013). One calf strain was excluded because the isolate could not be recovered after frozen storage, similar to other reports (REIMSCHUESSEL et al., 2017). Both foal isolates were susceptible to all antimicrobials tested, while the three calf strains were susceptible to ceftriaxone, enrofloxacin, and marbofloxacin. Two of the calf isolates were resistant to oxytetracycline, an antimicrobial commonly used in calves (PEZZELLA et al., 2004). One isolate was classified as a multidrug resistant strain (SCHWARZ et al., 2010), based on its resistance to amoxicillin/clavulanic acid, trimethoprim/sulfamethoxazole, and florfenicol. Identification of a multidrug resistant strain is concerning as trimethoprim/sulfamethoxazole and florfenicol are commonly used to treat diarrhea in foals and calves (KUANG et al., 2015) and amoxicillin/clavulanic acid is widely used in both veterinary and human medicine (BELMAR-LIBERATO, 2011; OTEO et al., 2008).

The administration of enrofloxacin immediately after the onset of clinical signs and treatment with florfenicol could have contributed to the clinical improvement of calves. However, despite improvement after antibiotic therapy, one Salmonella Ndolo isolate was resistant to florfenicol. It is important to note that certain bacterial resistance mechanisms to florfenicol can also confer resistance to chloramphenicol, an antimicrobial prohibited from use in animals but often used in humans for salmonellosis treatment (BOLTON et al., 1999).

Identification of genes encoding virulence factors in Salmonella serovars by PCR is considered an alternative to presuming the potential virulence of the strains (GAL-MOR & FINLAY, 2006; HARAGA & MILLER, 2003; JENNINGS et al., 2012). Thus, all Salmonella Ndolo strains (n=8), including the two isolates from the captive reptiles in the previous report (RAMOS et al., 2018), were assessed using previously described PCR methods to evaluate the presence of relevant virulence genes (MOHAMED et al., 2014; SKYBERG et al., 2006). These virulence genes are mainly associated with host cell recognition and invasion (invA, orgA, prsH, tolC, sopB, pefA), intracellular survival and growth in reticuloendothelial tissues (pagC, spvC), survival within macrophages (spidA, msgA), entry into non-phagocytic cells and killing of macrophages (spaN, sipB), and iron acquisition (ironN) (MOHAMED et al., 2014; SKYBERG et al., 2006).

All 13 virulence genes evaluated were detected in all Salmonella Ndolo strains from the foals, calves, and reptiles. This suggested a potential pathogenic role for these isolates, as they were associated with infection and diarrhea in most of these animals (GAL-MOR & FINLAY, 2006; JENNINGS et al., 2012). The virulence genes identified were similar to those detected in other Salmonella serovars isolated from humans and animals (DIONE et al., 2011; JENNINGS et al., 2012; PARVATHI et al., 2011). These results also suggested the presence of at least three Salmonella pathogenicity islands (SPIs) (IOANNIDIS et al., 2013) in these isolates since invA,
orgA, prgH, sipB, and spaN are commonly encoded in SPI-1, spiA in SPI-2, and sopB in SPI-5 (DIONE et al., 2011; SKYBERG et al., 2006). Salmonella pathogenicity island acquisition allows “quantum leaps” to occur in the evolution of Salmonella enterica serovars, therefore, they play a fundamental role in the pathogenesis of salmonellosis (GROISMAN et al., 1996; HENSEL, 2004). Interestingly, the genes spvC and pefA, which are often encoded by the same plasmid (CHU, CHIU, 2006; GULIG et al., 1993), typically are reported only in Salmonella Typhimurium and Salmonella Dublin. Salmonella Typhimurium is known to have a broad range of host species, whereas Salmonella Dublin is a host-adapted serovar commonly infecting bovines (BARROW et al., 2010; CHU & CHIU, 2006; ROTGER & CASADESUS, 1999).

The genetic diversity between the Salmonella Ndolo isolates from these two outbreaks was evaluated by ERIC–PCR using the primers ERIC-1 and ERIC-2 as previously described (LIM et al., 2005; SMITH et al., 2011; VERSALOVIC et al., 1991). The calf sample that was excluded from the antimicrobial evaluation was excluded from the ERIC-PCR testing as well. Also, the two Salmonella Ndolo strains previously isolated from captive reptiles were added for comparison (RAMOS et al., 2018); therefore, seven strains were subjected to the genotyping assay. The genotypic analyses were performed using Bionumerics 7.6 software (Applied Maths, Belgium). Using 90% similarity as a cutoff point (HASHEMI & BAGHBANI, 2015), the molecular characterization of Salmonella Ndolo revealed two major clusters of closely related strains (I and II) (Figure 1).

Although, clustering does not necessarily mean that strains are clonal, the results reflected a high genetic similarity between the strains from the outbreaks in foals (97.1%) and calves (94.3%). This high similarity suggested that the isolates within each outbreak were clonally related; however, strains were different between outbreaks and different from those previously isolated from reptiles. It is also interesting to note that isolates from Minas Gerais (reptile and foal isolates) clustered closer together than those from Goiás (calf isolates), whose municipality is located 600km from Belo Horizonte (Minas Gerais). Together, these results suggested that different strains of Salmonella Ndolo gave rise to each of the outbreaks in the separate locations. Despite this, all strains

![Figure 1](image-url)
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CONCLUSION

In this report, Salmonella Ndolo was isolated for the first time from domestic animals. Two outbreaks involving different species were described; one multidrug resistant strain was isolated. These findings suggested that Salmonella Ndolo is a multi-host pathogen whose pathogenic potential is enhanced by the presence of virulence genes common to Salmonella spp.

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DECLARATION OF CONFLICTING INTERESTS

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

AUTHORS’ CONTRIBUTIONS

ROSS, FCFL and CAGL conceived and designed experiments. CPR, RGX, FMV, IHP and EOL performed the experiments and carried out the lab analyses. EJFF and AUC conducted the flow charts. CPR, RGCX, FMV, IHP and EOL performed the statistical analyses. EJFF, AUC and ROSS prepared the draft of the manuscript.

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