Necrotizing placentitis in a cow caused by *Bacillus cereus*

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ABSTRACT: This report described a case of necrotizing placentitis caused by *Bacillus cereus* in a cow associated with abortion and maternal lethality. The etiological diagnosis of placentitis by *B. cereus* was based on histopathology of placenta, cytology and bacterial isolation from intrauterine amniotic fluid in retained placenta and further characterization of the pathogen by the MALDI-TOF. Although, *B. cereus* abortions are sporadic, the bacterium has the ability to release necrotizing toxins that can lead to placentitis, fetal death and abortion.  

Key words: *Bacillus* sp., placenta, infectious abortion, cattle.

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**ABSTRACT:** Este relato descreve a placentite necrotizante causada por *Bacillus cereus* em uma vaca associada a aborto e mortalidade materna. O diagnóstico etiológico de placentite por *B. cereus* foi baseado na histopatologia da placenta, citologia e isolamento bacteriano do líquido aminiótico em placenta retida e identificação do patógeno pela técnica de MALDI-TOF. Embora abortos por *B. cereus* sejam esporádicos, a bactéria tem a capacidade de liberar toxinas necrotizantes que podem levar a placentite e aborto.  

**Palavras-chave:** *Bacillus* sp., placenta, aborto infeccioso, bovino.

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*Bacillus cereus* is a Gram-positive aerobic or anaerobic facultative spor-forming rod bacterium. It is mobile and found in water, soil, plants and various sediments in the environment (BOTTO, 2010). In cattle, *B. cereus* and *B. licheniformis* are two species considered opportunistic that are associated with mastitis (SCHIEFER et al., 1976; PARKINSON et al., 1999; MAVANGIRA et al., 2013) and abortion (SCHU & WEINSTOCK, 1985; ANDERSON et al., 1990; KIRKBRIDGE, 1993; AGERHOLM et al., 1997). Reports of bovine abortion caused by *Bacillus* spp. have been described in Europe (AGERHOLM et al., 1997; ODSDOTTIR et al., 2004; DI BLASIO et al., 2019; WOLF-JÄCKEL et al., 2020), North America (ANDERSON et al., 1990; KIRKBRIDGE, 1993; CLOTHIER & ANDERSON, 2016), and Oceania (REICHEL et al., 2018) with no previously reported cases in South America.

Necrotic, hemorrhagic and suppurated placentitis are often associated with *Bacillus* sp. infection. *Bacillus* sp.-induced fetal lesions include pneumonia, pericarditis, encephalitis, hepatitis, peritonitis, and inflammation of the thymus (LOGAN, 1988; KIRKBRIDGE, 1993). Silage is considered an important source of contamination for cattle; although, *Bacillus* sp. is widely distributed in nature (LOGAN, 1988).

This report described for the first time in Brazil a bovine abortion in the final third of pregnancy caused by *Bacillus cereus*. A five-year-old Holstein cow, at 32 weeks of gestation, developed apathy, anorexia and pyrexia (41.5 °C). During physical examination, the cow was not in labor and on rectal
palpation the fetus was in an eutopic position with no movement. Antibiotic therapy (penicillin 10,000 IU/Kg and gentamicin 160 mg/kg) was administered every 48 hours in association with anti-inflammatory therapy (phenylbutazone OF 20%, 12 mL) daily. Twenty-four hours later, the cow presented external decubitus and fetal expulsion.

Male fetus with a crown-rump length of 57 cm was expelled with the fragments of the fetal membranes and a large amount of purulent, fetid and bloody fluid. A few hours after fetal expulsion, clinical condition of the cow worsened, progressing to death. Blood samples from the cow were collected for hemogram, and samples of the amniotic fluid from the intrauterine retained placenta were collected for cytology and bacterial isolation. At necropsy, the cow presented an accumulation of fibrinopurulent necrotizing exudate covering the caruncles within the uterus and fetal membranes thickened with pyosanguinolent exudate. Fragments of the placentome were collected for histopathological examination. Necropsy examination of the stillborn was not performed.

The cow had leukocytosis with left deviation. The cytological examination of the amniotic fluid in the panotic stained smears revealed numerous red blood cells and degenerate neutrophils, some of which with high numbers of elongated bacillary microbial structures often sporulated. Gram stained smears demonstrated numerous Gram-positive elongated rods with central endospore, compatible with microorganisms of the genus *Bacillus* both extracellular or within the cytoplasm of macrophages (Figure 1A).

Samples of the placentome were referred to the pathology service at the Veterinary School of

![Figure 1 - (A) Cytology of amniotic fluid. Numerous Gram-positive rods mostly associated with macrophages. Gram staining. (B) Cross section of a bovine placentome fixed in 10% formalin, B. cereus-induced abortion. Chorionic surface covered with a yellowish fibrinopurulent exudate (arrow) and multifocal areas of hemorrhage (arrowheads). (C-D) Bovine placenta, B. cereus-induced abortion. Fibrinous and necrotizing placentitis with mild multifocal neutrophilic inflammatory infiltrate, and bacterial colony. Hematoxylin and eosin, 100X and 400X. (E) Gram-positive rods extra and intracytoplasmic. Good-Pasture, 600x.](image-url)
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In this study, a case of bovine necrotizing placentitis caused by *Bacillus cereus* was reported. The etiological diagnosis of placentitis by *B. cereus* was confirmed by bacterial isolation, which included aerobic and anaerobic protocols. Bacteria were isolated when 5 mL of the aminiotic fluid was included in aerobic and anaerobic protocols. Bacteria *B. cereus* were positive bacteria was established (Figure 1C-E).

The etiological diagnosis of placentitis by *B. cereus* was confirmed by bacterial isolation, which included aerobic and anaerobic protocols. Bacteria were isolated when 5 mL of the aminiotic fluid was heated at 62 °C for 15 minutes, subsequently, plated on sheep blood 5% agar, and incubated at 37 °C for 24-48 hours. Beta hemolytic, irregular, large and matte colonies were submitted to biochemical tests in order to identify the microbial species. The isolates were motility+, catalase+, oxidase-, urease-, citrate+, gelatin+, nitrates reduction+, voges proskauer+, lecithinase+, esculin hydrolysis+. Results were validated according to the taxonomic key of the ABIS ONLINE software (https://www.tgw1916.net/bacteria_logare.html) with an accuracy of 99%. Additionally, when the isolates were cultured in serum agar and incubated with 10% CO2 there was formation of capsule. Cultures in brain heart infusion broth were sent to the Aquacen laboratory (UFMG Veterinary School), where the bacterium was identified as *B. cereus* by the MALDI-TOF (Matrix Associated Laser Desorption-Ionization - Time of Flight) mass spectrometry as previously described (ASSIS et al., 2017).

In addition, other important pathogens associated with bovine abortion or placentitis were investigated by PCR with DNA templates extracted from formalin-fixed placental samples. DNA extraction was performed according to SILVA et al., 2009 and PCR reactions performed as previously described for *Brucella* sp. *Leptospira* sp., *Neospora* caninum (SILVA et al., 2009), *Listeria monocytogenes* (BUBERT et al., 1999), and Herpesvirus Bovine Type 1 (VAN ENGELENBURG, et al., 1993). The quality of the extracted DNA was confirmed by the positive PCR reaction for amplification of the bovine α- actin gene (SILVA et al., 2009). Negative and positive controls were included in each reaction. The etiological diagnosis of placentitis by *B. cereus* in this case was confirmed by bacterial isolation from aminiotic fluid collected directly from the uterine cavity and specific identification by a comprehensive biochemical panel and MALDI-TOF mass spectrometry. Fibrinopurulent placentitis was the probable cause of the clinical changes presented by the cow, which were associated with abortion followed by maternal death.

The cow in this report was approximately at eight months of pregnancy with clinical signs of pyrexia, apathy and anorexia, and absence of fetal movement. Therefore, in this case, fetal death preceded fetal expulsion as reported in previously described cases (SCHUH & WEINSTOCK, 1985). *B. cereus*-induced abortion followed by death of the cow has not been previously described; although, cases of gangrenous mastitis due to *B. cereus* infection are often lethal due to toxemia (SCHIEFER et al., 1976).

In cases of abortion in domestic animals caused by *Bacillus* spp. that the pathogen may be isolated from vaginal swabs, placenta, fetal abomasal contents, heart, liver, lung, bronchial and mediastinum lymph nodes of aborted fetuses (LOGAN et al., 1988). In this case, *B. cereus* was isolated from aminiotic fluid while fetal membranes were still within the uterus, which minimized environmental contamination.

JOHNSON et al. (1994) considered placentome a good tissue to obtain the definitive etiological diagnosis because it allows associating isolation with necrotic placentitis lesion, which may allow a conclusive diagnosis even without access to fetal tissues. There is evidence of abortion with necrotic placentitis by *B. cereus* without fetal injury, even if there is bacterial colonization in the fetus (SCHUH & WEINSTOCK, 1985). In this report, the fetus was not available and the only tissue sent for hispathological examination was a placentome, which along with microbiological examination of the aminiotic fluid supported a conclusive diagnosis of necrotic placentitis with intralesional Gram-positive bacteria (*Bacillus cereus*).

Considering that *B. cereus* is an ubiquitous microorganism, isolation of the bacteria by itself is not sufficient for a conclusive diagnosis since the isolated bacterium may be an environmental contaminant. The diagnosis of bovine abortion by *B. cereus* is usually based on the pure culture of the microorganism isolated from placenta and/or fetus, with presence of microorganisms within microscopic lesions, and the absence of other agents known to cause abortion in cattle (SCHUH & WEINSTOCK, 1985).
this case, no other infectious agent was isolated on microbiological examination or detected by PCR. Furthermore, the visualization of phagocytosed Gram-positive bacteria by macrophages present in the amniotic fluid indicates that this agent was not an environmental contaminant, but it was present within the uterus prior to fetal expulsion and it was associated with the intrauterine inflammatory reaction.

The pathogenesis of *B. cereus*-induced abortion is unknown; although, *B. cereus* secretes several enzymatic or lytic exotoxins, including phospholipases, hemolysins, pore-forming enterotoxins, and cytotoxin K (LOGAN, 1988, BOTTONE, 2010). According to SCHUH & WEINSTOCK (1985), one of the consequences of toxins produced by *B. cereus* is a necrotizing placentitis followed by death and fetal expulsion before fetal injury or even colonization of fetal tissues by *B. cereus* (SCHUH & WEINSTOCK, 1985). Studies suggested that bovine abortions caused by *B. cereus* may be associated to virus-induced immunosuppression as occurs in cases of bovine viral diarrhea virus infection (SCHUH & WEINSTOCK, 1985; KIRKBRIDE, 1993). In conclusion; although, it is an opportunistic agent, *B. cereus* should be considered in the differential diagnosis of sporadic infectious bovine abortion in the country.

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BIOETHICS AND BIOSAFETY COMMITTEE APPROVAL

We authors of the article entitled “Necrotizing placentitis in a cow caused by *Bacillus cereus*” declared, for all due purposes, the project that gave rise to the present data of the same has not been submitted for evaluation to Ethics Committee of the Universidade Federal de Minas Gerais”, but we are aware of the content of the Brazilian resolutions of the National Council for Control of Animal Experimentation - CONCEA “http://www.mct.gov.br/index.php/content/view/310553.html” if it involves animals.

Thus, the authors assume full responsibility for the presented data and are available for possible questions, should they be required by the competent authorities.

DECLARATION OF CONFLICTS OF 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

CEVR, RLS and TAP worked in conception and writing of the manuscript. CEVR, JPLM, JPSM, ACBG, EAC, HCPF carried out the lab analyses. All authors critically revised the manuscript and approved of the final version.

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