The Aggregation of \textit{Brucella abortus} Occurs Under Microaerobic Conditions and Promotes Desiccation Tolerance and Biofilm Formation

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**Abstract:** \textit{Brucella abortus} causes brucellosis mainly in cattle. The infection is transmitted to humans by ingestion of animal products or direct contact with infected material. While the intracellular lifestyle of \textit{Brucella} is well characterized, its extracellular survival is poorly understood. In nature, bacterial persistence is associated with biofilms, where aggregated cells are protected from adversity. The inability of \textit{Brucella abortus} to aggregate under aerobic conditions and that fact that the replicative niche of \textit{Brucella} is characterized by microaerobic conditions prompted us to investigate the capacity of this pathogen to aggregate and grow in biofilms under microaerobic conditions. The results show that \textit{B. abortus} aggregates and produces biofilms. The aggregates tolerate desiccation better than planktonic cells do, adhere and displace even in the absence of the lipopolysaccharide-O antigen, flagella, the transcriptional regulator VjbR, or the enzymes that synthesize, transport, and modify cyclic \(\beta\) (1,2) glucan.

**Keywords:** Aggregation, biofilms, \textit{Brucella}, desiccation, flotation, VjbR.

**INTRODUCTION**

Infection with \textit{Brucella} can result in brucellosis, a chronic lifelong zoonotic disease in humans and cattle. This is a damaging disease, particularly in developing countries with economies that depend on livestock. In humans, transmission occurs mainly through direct contact with infected animals or ingestion of contaminated dairy products, while in wildlife, it is spread due to bacterial persistence in the environment. Although \textit{Brucella spp.} can survive for long periods in soil, dust, slurry, water, and meat, the survival strategies of this pathogen have been mostly studied from within eukaryotic cells.

Bacteria that colonize different surfaces and invade susceptible hosts to cause chronic infections grow predominantly in biofilms [1, 2]. First, bacteria produce an extracellular matrix mainly composed of exopolysaccharides, which surrounds them and plays a key role in keeping them hydrated and helping them adhere to inert or biological surfaces. Even if adhesion is not a population-dependent phenotype [3], it is necessary for the complex process of biofilm growth. Experimental conditions of wild-type \textit{Brucella} growth, as a stable community of cells that form surface-adherent biofilms, have not been demonstrated. Adhesion has been reported only in mutants of \textit{B. melitensis} that present an altered expression of outer-membrane proteins [4-6].

One of these mutants is \textit{B. melitensis} \textit{vjbR}, which cannot express the transcriptional regulator VjbR. As a \textit{B. melitensis} double mutant \textit{vjbR} \textit{omp31} did not aggregate under aerobic conditions, it was postulated that Omp31 is involved in the production and/or secretion of the matrix-forming exopolysaccharides. To support this hypothesis Uzureau \textit{et al.} demonstrated that similar mutations in \textit{B. abortus} \textit{vjbR} do not cause aggregation under the same conditions due to the natural absence of \textit{omp31} in this species [4].

Given the fact that \textit{Brucella} usually survives in niches characterized by low oxygen tension and nutritional scarcity [7, 8] in infected hosts or in the environment and secretes N-dodecanoyl homoserine lactone, which (along with VjbR) belongs to the quorum-sensing system [9-11], we hypothesized that wild-type \textit{B. abortus} produces an extracellular matrix that confers adhesion and stress-resistance properties and promotes the development of biofilms under nutritionally deficient, microaerobic conditions.

**Autoaggregation of \textit{B. Abortus}**

We initiated our work by studying the physiology of \textit{B. abortus} under microaerobic conditions. First, oxygen preference in wild-type \textit{B. abortus} 2308 was assayed in semi-solid media containing thioglycollate. This bacterium behaved as a microaerophile; it produced a zone of growth a few millimetres below the surface of the medium after 48 hours of incubation in a glass tube. However, with prolonged incubation beginning at day 3 to 5, \textit{B. abortus} produced a discontinuous turbidity growing downward towards the bottom of the tube. Though growth was observed in the centre of the tube, it was
prominent along the walls as a film growing downward but unattached to the glass wall (Fig. 1A).

Therefore, we looked for biofilm development in Brucella broth (BB) cultures in glass tubes under microaerobiosis. The microaerobic environment was generated by a 5% CO2 incubator or in 8 mL culture media in 10-mL screw-capped tubes, incubated statically at 37°C. Under these conditions, bacterial growth first occurred at the bottom of the tubes, consistent with its non-motile phenotype. Interestingly, the cellular mass became viscid during incubation. From day 4 to day 7 in static cultures at 37°C, threads spontaneously detached from the viscous bacterial mass were hanging from one end attached to the glass at the air-liquid interface. After 2 weeks, a discontinuous ring appeared and the threads had grown longer (Fig. 1B). This behaviour was not apparent when similar cultures were incubated with shaking for the same period of time to maintain aerobic conditions. The attachment to the glass at the air-liquid interface was weak; the threads easily fell down and the rings were washed away when adhesion was examined by Crystal violet staining [12]. As a result, no confident measurement was possible and no surface pellicle was produced. Although B. abortus barely adheres to glass at the air-liquid interface, it was unable to grow in biofilms on this abiotic surface, under our experimental conditions. To ensure cell viability, samples of threads were stained with the vital stain Acridine orange and observed by fluorescence microscopy [13]. Almost all the cells stained orange, indicating the threads were composed of adherent living bacteria. These results demonstrate that wild-type B. abortus autoaggregates under low oxygen tension and nutrient limitation, acquiring in this form the capacity to move.

One possible explanation for the motility of Brucella could be the involvement of flagella. It is interesting to note that even though B. abortus possesses and expresses flagellar genes, the bacteria have been reported as non-motile in vitro [14]. The inability to synthesize the flagellar hook protein, FlgE, renders a non-flagellated strain. Thus, wild-type B. abortus 2308 and the isogenic flgE [15] were examined in liquid and semi-solid media incubated under microaerobiosis. The results showed a wild-type phenotype for the flgE mutant in both types of cultures, eliminating the possibility of flagellar motility. Changes in buoyancy and coordinated population-migration may explain the bacterial behaviour in liquid and in semi-solid media, respectively.

To further analyse the conditions given by the aggregation phenotype, daily samples starting from overnight cultures were taken from the semi-solid and liquid cultures, incubated under aerobic or microaerobic conditions. Samples were examined microscopically and plated to assess colony morphology. Aggregates were only observed from day 3 of incubation under microaerobiosis. Colony morphology derived from aggregates in all cases was the same as that of non-aggregative cells obtained during log or stationary phase under aerobic conditions. Scanning electron microscopy (SEM) of the aggregates revealed an amorphous matrix surrounding the bacteria (Fig. 2A). The SEM observations were conducted at 10 kV in a Philips XL 30 TMP. Each specimen was mounted on an aluminium stub and coated with gold-palladium.

To verify the polysaccharidic nature of the matrix, samples were stained with calcofluor and Congo red. All samples were fixed by addition of 3.7% formaldehyde solution prior to immobilization. Slides containing the samples were air-dried, then covered with a 2:1 mixture of saturated aqueous Congo red solution and 10% (vol/vol) Tween 80 or freshly prepared 0.1% calcofluor (Cellufluor Polyscience Inc., PA, USA) in distilled water. The appearance of positive fluorescence in the aggregates with both dyes but not in isolated cells suggests the presence of β(1,3) or β(1,4) glucans in the B. abortus extracellular polymeric matrix. Exopolymers such as polysaccharides and fimbria are important components of the extracellular matrix. B. abortus does not express fimbria but produces β(1,2) cyclic glucan. To identify whether this glucan, its transporter (Cgt), or the enzyme that adds succinyl residues to the glucan (Cgm) were involved producing the extracellular matrix, the experiments in liquid and semi-solid media were performed with cgs [16], cgt [17], and cgm [18] mutants. Cgs synthesizes this polymer. These strains exhibit wild-type behaviour, indicating that β(1,2) cyclic glucan, its transporter, and its modifier do

![Fig. (1). B. abortus growth under microaerobic conditions. 10-day old cultures grown in semi-solid (A) or liquid (B) media at 37°C. Bacterial aggregates are visualized at the top of the liquid medium as threads hanging (short arrow) from a discontinuous ring. Note bacterial mass at the bottom of the tube. Large arrows indicate bacterial growth direction.](image-url)
not mediate the matrix production needed to adhere or displace. Since LPS also participates in bacterial adherence [19], \textit{B. abortus} \textit{pgm} [20] was included in our experiments. This strain, deleted for the gene encoding phosphoglucomutase, does not expose lipopolysaccharide-O antigen on the surface, leading to a rough phenotype. Again, this mutant was able to aggregate and displace, showing that the lipopolysaccharide does not play an important role in the adhesion and movement of \textit{B. abortus} under our experimental conditions. These results also demonstrate that matrix production is possible in rough \textit{Brucella} strains.

Absence of the VjbR regulator in \textit{B. abortus} does not lead to the aggregation phenotype observed in the \textit{B. melitensis} vjbR when grown aerobically [4] and wild-type \textit{B. abortus} aggregates under our experimental conditions; therefore, we generated a knock-out mutation in \textit{vjbR} to determine its effect on the \textit{B. abortus} phenotype. Thus, cultures of \textit{B. abortus} \textit{vjbR} [21] in liquid and semi-solid media under microaerobic conditions were examined microscopically and compared to wild type. The mutant was able to aggregate (Fig. 2B) and move as the wild type. These results clearly show that the transcriptional regulator VjbR is not involved in the aggregation and displacement of \textit{B. abortus} under microaerobic conditions. Since it regulates flagellar genes [22], these results also demonstrate that flagella are not involved in the bacterial phenotype under microaerobiosis.

The Desiccation Tolerance of \textit{B. Abortus}

The extracellular matrix usually protects aggregated cells from different stress conditions or even from the proximity of toxic compounds, thus increasing bacterial survival. It has been reported that \textit{B. abortus} can survive in materials such as soil, pasture, or manure and that ingestion of these contaminated materials may be one route by which animals or humans in barns acquire brucellosis [23]. Hence, we investigated if the matrix-embedded cells obtained under starvation and oxygen limitation could encourage \textit{B. abortus} desiccation tolerance. To avoid dealing with the increasing number of dead cells during the stationary phase of growth and taking into account that the matrix was observed at day 3 post-incubation under microaerobic conditions, the experiments were performed with 3-day old cultures grown in BB under aerobic or microaerobic conditions. First, the amount of Congo red bound to cells from each culture was quantified [24] by dissolving the cell-bound stain in DMSO and reading the OD of the supernatant at 490 nm. Standard preparations of the stain in DMSO were made to generate a standard curve of mg Congo red/mL and viable cells were determined by plating serial dilutions of each culture. As shown in Fig. (3), the amount of stained matrix per cell in cells grown under microaerobic conditions was 4 times higher than that observed under aerobic conditions. After observing this significant difference in the extracellular material, small triplicate samples were tested for desiccation tolerance [25]. Briefly, 20-μL samples were taken in triplicate. A total of 480 μL media was immediately added to one of the samples and serial dilutions were plated on BB agar for bacterial counts. The second sample was allowed to dry at 30°C, while the third sample remained unexposed for the same period of time as a control. After total desiccation, 500 or 480 μL media was added and the cells recovered to estimate percent survival. These results (Fig. 3) show that cells provided with an extracellular matrix are better able to survive desiccation than planktonic cells under our experimental conditions. Although extracellular matrix usually protects
Biofilm Development of \textit{B. Abortus}

To investigate \textit{B. abortus} biofilm formation on other abiotic surfaces such as plastic, inocula from cultures containing about $10^8$ CFU/mL were diluted 1:100 into 2 mL BB per well of polyvinyl-chloride 24-well plates (Falcon, Becton Dickinson, France) and into 30 mL of BB in polystyrene-treated 60-mL flasks (Corning Inc., New York, USA). The cells showed very little adherence to the polyvinyl-chloride plates in a microaerobic environment generated by a 5% CO$_2$ incubator for a period of 20 days. However, after the same period of time, biofilms appeared on the sides of the polystyrene flasks. This type of adherence was more stable than that observed on glass or plates since it could withstand the staining procedure. No surface pellicle was observed. To differentiate between attachment of aggregative cells and biofilm growth, 2-day-old \textit{B. abortus} cultures grown in polystyrene flasks were maintained at 37°C for longer under conditions that allow (microaerobiosis) or do not allow (anaerobiosis) bacterial replication. Anaerobiosis was achieved using an AnaeroGen$^\text{TM}$ sachet (Oxoid, Hampshire, UK) to generate less than 1% O$_2$ and 9–13% CO$_2$ in a sealed jar. From days 9 to 11, adherence on the sides of the flasks was detected at the air-liquid interface in both conditions. At the end of the experiment, when bacterial attachment was detected by crystal-violet staining, we observed a blue line at the air-liquid interface in cultures maintained under anaerobiosis. In contrast, the sides of the flasks maintained under low oxygen tension were completely stained (Fig. 4). These results indicate that aggregative cells were able to float and adhere under both conditions but the biofilm was the consequence of downward growth from the attached cells. No surface pellicle was observed. The inability to develop biofilms in plates could be attributed to the components of the plastic. Interestingly, the matrix-embedded cells inside the flask found a better attachment with the pre-treated sides than the others, and this condition promoted biofilm production.

Bacteria in biofilms are not only resistant to desiccation but are also protected from interaction with substances such as antibiotics, antibodies, and disinfectants. Therefore, considering the epidemiology and chronicity of brucellosis, the characteristics of \textit{B. abortus} under these conditions must be analysed at the genetic level to support development of eradication programs. Studies are underway to identify the components of the extracellular matrix and the genes involved in their synthesis and transport.

**CONCLUSION**

The results presented here demonstrate that starved \textit{B. abortus} grown under microaerobiosis produce extracellular matrix, autoaggregate, move, and adhere to abiotic surfaces without the involvement of VjbR, the lipopolysaccharide-O antigen, cyclic β (1,2) glucan, or flagella. Matrix-producing cells are more resistant to desiccation and promote the development of biofilms.

**CONFLICT OF INTEREST**

The authors confirm that this article content has no conflicts of interest.

**ACKNOWLEDGEMENTS**

\textit{B. abortus} mutant strains were kindly provided by R.A.Ugalde. This work was supported by grants from the National Agency for Science and Technology PICT-0651 and PIP-5463 from the National Research Council of Argentina (CONICET), M.A., M.S.R. and N.S. are members of the Research Career, CONICET.
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