RomA, a Periplasmic Protein Involved in the Synthesis of the Lipopolysaccharide Tunes

Down the Inflammatory Response Triggered by *Brucella* 

Ezequiel Valguarnera<sup>1\*</sup>, Juan M. Spera<sup>1</sup>, Cecilia Czibener<sup>1</sup>, Fabiana R. Fulgenzi<sup>1</sup>,

Adriana C. Casabuono<sup>2</sup>, Silvia G. Altabe<sup>3</sup>, Karina A. Pasquevich<sup>1</sup>, Francisco Guaimas<sup>1</sup>,

Juliana Cassataro<sup>1</sup>, Alicia S. Couto<sup>2</sup> and Juan E. Ugalde<sup>1\*</sup>

1 Instituto de Investigaciones Biotecnológicas "Dr. Rodolfo A. Ugalde", IIB-INTECH, CONICET, Universidad Nacional de San Martín, San Martín, Buenos Aires, Argentina.

2 Universidad de Buenos Aires. Facultad de Ciencias Exactas y Naturales. Departamento de Química Orgánica - Consejo Nacional de Investigaciones Científicas y Técnicas. Centro de Investigación en Hidratos de Carbono (CIHIDECAR). Buenos Aires, Argentina.

3 Instituto de Biología Molecular y Celular de Rosario (IBR) and Departamento de Microbiología, Facultad de Ciencias Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Santa Fe, Argentina.

Running title: Immune response during Brucella infection

\*To whom correspondence should be addressed: Instituto de Investigaciones Biotecnológicas "Dr. Rodolfo A. Ugalde", IIB-INTECH. Universidad Nacional de San Martín. Avenida 25 de Mayo y Francia, Campus Miguelete, UNSAM, San Martín (1650), Buenos Aires, Argentina.

Phone: (54)11-4006-1500 (2129). Fax: (54)11-4006-1559. E-mail: jugalde@iibintech.com.ar and evalguarnera@iibintech.com.ar

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**ABSTRACT** 

Brucellaceae are stealthy pathogens with the ability to survive and replicate in the host in the

context of a strong immune response. This capacity relies on several virulence factors that are

able to modulate the immune system, and in their structural components that have low pro-

inflammatory activities. Lipopolysaccharide (LPS), the main component of the outer

membrane, is a central virulence factor of Brucella and it has been well established that

induces a low inflammatory response. We describe here the identification and

characterization of a novel periplasmic protein (RomA) conserved in alpha-proteobacteria,

involved in the homeostasis of the outer membrane. A mutant in this gene showed several

phenotypes, such as membrane defects, altered LPS composition, reduced adhesion and

increased virulence and inflammation. We show that RomA is involved in the synthesis of

LPS, probably coordinating part of the biosynthetic complex in the periplasm. Its absence

alters the normal synthesis of this macromolecule and affects the homeostasis of the outer

membrane, resulting in a strain with a hyperinflammatory phenotype. Our results suggest that

the proper synthesis of LPS is central in order to maximize virulence and minimize

inflammation.

**Keywords:** *Brucella*; lipopolysaccharide; inflammation.

INTRODUCTION

Brucellaceae are wide spread zoonotic intracellular pathogens with the capacity to evade and

modulate the immune response of the infected hosts, a hallmark of their infectious processes

[1]. Many of these immunomodulatory activities are achieved by a plethora of virulence

factors that are able to manipulate the immune system to its benefit, promoting bacterial

proliferation and the establishment of the chronic infectious phase [2-7]. Brucella is

considered a pathogen with a stealthy strategy, meaning that is able to avoid a strong immune

response "hiding" its pathogen-associated molecular patterns (PAMPs). This strategy is

achieved by either downregulating activation of PAMPs, or though the synthesis of structural

components with low pro-inflammatory activities [8]. This last concept raises an interesting

question; are the structural cellular components of Brucella, like lipopolysaccharide (LPS),

silent PAMPs by default or is the bacterium actively modifying them to be non-detected?

Lipopolysaccharide (LPS) is the main component of the outer membrane in all Gram-negative

bacteria. It is composed of lipid A (the lipidic portion inserted in the membrane), a core

oligosaccharide and the O-antigen, which is the most exposed structure [9]. LPS is necessary

for a wide range of functions such as protection against harsh environmental conditions,

selective permeability, immune protection and evasion among others. The synthesis of this

macromolecule is achieved through a complex biosynthetic pathway that starts in the

cytoplasm where the precursors are synthesized, continues in the inner membrane and

periplasmic space where the assembly, polymerization and transport take place, and ends

when the complete molecules are inserted in the outer membrane [9]. Many of these processes

require multiprotein complexes that coordinate their activities in a spatial and temporal way in

order to effectively synthesize, transport and insert this complex macromolecule in its final

organelle [10]. In Brucella, LPS has been shown to be a central virulence factor necessary for

intracellular replication and virulence in mice [8]. Additionally, it has been shown that a

structurally complete LPS is needed for the efficient immune evasion and that it also acts as a

shield against the innate immune response [1, 11].

We describe the identification in  $Brucella\ abortus$  of a gene encoding a protein with no known function but conserved among almost all  $\alpha$ -proteobacteria that we propose is involved in the homeostasis of the outer membrane. The 84 amino-acids protein has a periplasmic localization and its absence results in a strain with several phenotypes: membrane alterations, changes in the LPS composition, defects in the intracellular replication capacity of the bacteria and deregulated inflammatory response in mice. We hypothesize that the absence of this protein alters the proper synthesis of the LPS, probably modifying the assembly of the biosynthetic complex in the periplasm, which in turn affects outer membrane homeostasis and the modulation of the immune response during the infectious cycle.

**METHODS** 

A complete description of the Methods has been included in supplementary information.

Periplasmic and cytoplasmic localization assay

For the periplasmic localization assays the B. abortus strains were grown in TSB for 16-24

hrs at 37°C and 2.5x10<sup>10</sup> bacterial cells were centrifuged 10 min at 3300xg and the

periplasmic and cytoplasmic fractions obtained as we have previously described [12]. For

Western-Blot an anti-FLAG M2, anti GroEL (1:2000) and anti OMP-1 (1:2000) kindly

provided by Dr. Axel Cloeckaert were used as primary antibodies.

**Detergent sensitivity assays** 

For all B. abortus strains, overnight cultures were diluted and seeded onto TSB plates

supplemented with detergents as described [13]. Final detergent concentrations were 125

μg/ml Sarkosyl, 25 μg/ml Zwittergent 3-16, 1 g/ml sodium deoxycholate. Sensitivity was

calculated determining the viable CFU.

**Total lipid extraction** 

For the extraction of total lipids, the Bligh and Dyer method was used [14] on exponentially

grown bacteria.

Cristal violet staining of B. abortus

Serial dilutions of B. abortus cultures were plated in TSB plates and incubated at 37°C until

growth was observed. Plates were stained with a crystal violet solution as described [15].

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LPS purification

LPS was extracted from B. abortus strains using a modification of the phenol-hot water

method [16] from 250 ml of stationary cultures. LPS concentration was determined by the 3-

Deoxy-D-manno-2-octulosonic acid method and analyzed on 12% SDS-PAGE gels and

stained by silver nitrate.

Gentamicin protection assays

Gentamicin protection assays were performed as previously described [17].

**Mice infections** 

Mice infections were performed as previously described [18].

Analysis of fluorescent fusion proteins

Stationary phase cultures of B. abortus strains were diluted in fresh media and grown until

exponential phase was reached. A volume of 3 µl of culture was seeded onto the center of a

PBS 1% agarose pad as previously described [19]. Images were acquired and processed as

described above.

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**RESULTS** 

Bab1\_1280 encodes a periplasmic protein necessary for the homeostasis of the outer

membrane in Brucella.

In a genetic screen to identify genes from Brucella abortus coding for secreted or surface

exposed proteins, we isolated Bab1\_1280. This gene encodes an 84 amino-acid hypothetical

protein conserved in most α-proteobacteria with no known function to date [20] (Fig. S1). We

serendipitously found that an insertion mutant in this gene resulted in a strain with altered

membrane properties. Because the gene was identified in a screen for either

secreted/periplasmic or surface exposed proteins, we performed subcellular fractionation

assays to further determine its localization. We generated a strain expressing FLAG tagged

RomA from a genomic allele and performed a periplasmic extraction protocol [12]. Fig. 1A

shows that the FLAG-tagged protein product of Bab1\_1280 fractioned with the

periplasm/outer membrane as the outer membrane protein 1 (OMP1). Presence of the protein

due to bacterial lysis was discarded since no cytoplasmic contamination was observed

(GroEL). We additionally lysed the strain and determined membrane association by

ultracentrifugation, which indicated that the protein interacts with total membranes (Fig. 1B).

In order to determine the degree of this association, total membranes were resuspended in

different buffers, re-centrifuged and evaluated if the protein remained associated. Fig. 1C

shows that only a mild wash with 10 or 50 mM sodium phosphate was enough to partially

loosen the membrane association, condition that was completely lost with sodium chloride, or

detergent treatments. These results indicate that the product of Bab1\_1280 is a periplasmic

protein with a weak association to either the inner or the outer membranes.

To determine if the product of Bab1\_1280 is involved in maintaining the normal composition

of the outer membrane, we measured the resistance of the mutant to the detergents Sarkosyl,

Zwittergent 3-16 and Sodium Deoxycholate. Fig. 2A shows that the Bab1\_1280 mutant was

less resistant to the three detergents. To further determine if this increased sensitivity to

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detergents is related to the hydrophobicity of the membrane, we performed an N-phenyl-1-

naftilamine (NPN) incorporation assay. As it can be observed in Fig. 2B, the hydrophobic

probe was incorporated more efficiently in the mutant indicating that the outer membrane is

more permeable to hydrophobic molecules. These alterations in the outer membrane

properties of the mutant could be the consequence of differences in the phospholipids or fatty

acid profile. One and two-dimensional thin layer chromatography (TLC) using C<sup>14</sup>-labeled

total bacteria or extracted periplasms (phospholipids of the outer membrane) showed no

differences between strains (Fig. 2C and Fig. S2). These results indicated that the alterations

observed with the mutant in terms of detergent sensitivity as well as membrane

hydrophobicity, are not the result of a differential phospholipid composition but are probably

due to differences in other components of the outer membrane. The localization of the

protein, together with the defects in the membrane properties of the mutant strain, suggested

that RomA might be playing a role in maintaining a degree of homeostasis of the periplasm

and/or outer membrane. For these reasons we renamed Bab1\_1280 as romA, for regulator of

outer membrane.

RomA is implicated in the biosynthesis of LPS

The fact that the  $\Delta romA$  mutant showed altered membrane properties but no differences in the

phospholipid composition raised the possibility that it might have a defective LPS. To test if

this we performed a crystal violet (CV) staining to determine if the strain has a complete LPS

(smooth strain, excludes the staining) or if it lacks the assembled O-antigen (rough strain,

includes the staining). Surprisingly, the  $\Delta romA$  ( $\Delta Bab1\_1280$ ) strain showed a higher degree

of exclusion of CV (Fig. 2D), strongly suggesting a modified LPS, which was further

confirmed by western blot on whole bacteria. Fig. 3A shows that the mutant strain exhibited a

LPS pattern that seemed to have, not only a higher antigenic load, but also higher molecular

weight forms. To further advance in its characterization we performed a western blot on

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antibodies. As it can be observed in Fig. 3B and Fig. 3C, the  $\Delta romA$  mutant showed higher

levels of smooth LPS and lower levels of the rough LPS indicative that the strain might have

an altered equilibrium of the S-LPS:R-LPS ratio in the membrane. To confirm that this

alteration is present in the outer membrane and it is not a consequence of an accumulation of

the S-LPS in the inner membrane, we performed a periplasmic/outer membrane extraction

and analyzed the LPS. As can be observed in Fig. 3C (right panel), the same pattern of S-

LPS:R-LPS was observed in the outer membrane with the mutant, confirming that this strain

exhibits a higher percentage of S-LPS. An additional characteristic that we observed while

analyzing the gels of the  $\Delta romA$  LPS, was that it seemed to have longer O-antigen chains,

although an alternative explanation could be that it was the result of a higher concentration of

smooth LPS and not a chain length issue. To distinguish between these two possibilities we

characterized purified LPS from both strains by Western Blot with the  $\alpha$ -S-LPS and  $\square$ -R-LPS

monoclonal antibodies and silver staining. In Fig. 3D it can be seen that purified LPS showed

the same pattern observed either with whole cells or periplasmic extractions. These

preparations were used to chemically characterize the O-antigen and the core (see Methods).

Monosaccharide analysis by High- Performance Anion-Exchange Chromatography coupled

with Pulsed Amperometric Detection (HPAEC-PAD) of the oligosaccharides released from

the LPSs showed that the  $\Delta romA$  strain LPS presents a significant higher ratio of perosamines

(Rha4N, present in the O-antigen) to N-acetyl-glucosamines (GlcN, present in the core),

indicating that the mutant has a longer O-antigen. (Fig. 4A and Fig. S3).

Furthermore, when the oligosaccharides released after mild acid hydrolysis of the

corresponding LPSs were characterized by MALDI-TOF m.s in the positive mode, significant

differences in the high mass range were detected. Thus, the spectrum corresponding to the

wild type oligosaccharide presented the highest mass signals at m/z 3077.9 and m/z 3100.0

( $\square$ Na) (calc. m/z 3099.2977;  $C_{122}H_{210}N_{15}Na_2O_{73}$ ) corresponding to a structure bearing

KdoGlcQuinMan<sub>2</sub>Rha4N<sub>14</sub>Fo<sub>4</sub>Na<sub>2</sub>. In accordance, signal at m/z 2858.2 (calc. m/z 2857.2574;

 $C_{114}H_{199}N_{15}NaO_{66}$ ) corresponds to the structure GlcQuinMan<sub>2</sub>Rha4N<sub>14</sub>Fo<sub>4</sub>Na. On the other

side, the spectrum of the mutant strain showed not only the signals described above, but also

signals at m/z 4349.3 (calc. m/z 4349.8865;  $C_{173}H_{299}N_{23}NaO_{101}$ ) attributed to a structure

bearing KdoGlcQuinMan<sub>2</sub>Rha $4N_{22}$ Fo<sub>8</sub>Na and m/z 4325.3 (calc. m/z 4325.8630;

C<sub>173</sub>H<sub>296</sub>N<sub>23</sub>Na<sub>2</sub>O<sub>99</sub>) attributed to anhKdoGlcQuinMan<sub>2</sub>Rha4N<sub>22</sub>Fo<sub>7</sub>Na<sub>2</sub>. Furthermore, a signal

at m/z 4835.5 (calc. m/z 4835.0851;  $C_{193}H_{331}N_{26}Na_2O_{111}$ ) corresponds to the latter with three

additional Rham4N and two Fo groups (Fig. 4B). The fact that the ratio of Rha4N to GlcN in

the mutant is increased five times and the O-antigen is almost two folds longer confirms that

the mutant has an altered smooth to rough ratio. Despite that the O-antigen was longer, the

core showed no differences (data not shown).

Altogether these results indicate that RomA affects membrane properties and composition,

and is required for a proper LPS assembly.

RomA is involved in the virulence process and its inactivation profoundly alters the

inflammatory response

The changes in the LPS profile in the  $\Delta romA$  mutant strain led us to evaluate the potential role

of this gene in the virulence of B. abortus. Fig. 5A shows that the  $\Delta romA$  strain exhibited a

significant defect in the intracellular survival capacity in murine bone marrow derived

macrophages during the initial stages of infection (4 and 24 hrs post-infection) but was able to

replicate and, at 48 hrs post-infection, we did not observe any differences with the wild type

parental strain. This intracellular replication pattern was similar when J774 A.1 cells were

used (Fig. S4). Due to the altered membrane of the  $\Delta romA$  mutant we further analyzed if

these early effects were due to a defect in the adhesion of the bacteria to the cells or in their

reduced capacity to exclude the lysosomal marker Lamp-1 during the intracellular trafficking.

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ownloaded from https://academic.oup.com/jid/advance-article-abstract/doi/10.1093/infdis/jiy002/4791945 y quest As can be seen in Fig. S5 and Fig. S6, the mutant showed a statistically significant defect in

the adhesion to J774 A.1 cells as well a reduced capacity to exclude Lamp-1 at 24 hrs post-

infection. These results indicate that the altered membrane structure of the  $\Delta romA$  mutant

probably impacts on several steps of the interaction of the bacterium with the host cells

(adhesion/invasion as well as in the intracellular trafficking).

The results obtained in vitro encouraged us to evaluate the role this gene might play during

pathogenesis in the mouse model. Surprisingly, the  $\Delta romA$  mutant showed a dramatic

increase in the number of bacteria in the spleens of intraperitoneally infected mice at 14 days

post-infection, which correlated with an increased splenomegaly (Fig. 5B, 5C and 5D). An

interesting observation was that the complemented strain exhibited less bacterial load and

splenomegaly than the wild type strain, which led us hypothesize that it might also have an

altered LPS. Fig. S7 shows that the  $\Delta romA$  complemented strain, expressing RomA from a

plasmid, had a LPS with less assembled O-antigen (partly rough).

The increase in the spleen size was the result of an enhanced inflammatory response. Fig. 6,

panel A to D, shows that the mutant induced a significantly higher inflammatory response in

comparison to the wild type, measured by the production and circulation of two pro-

inflammatory cytokines (TNF-α and IFN-γ). This was also confirmed by histological

observation of the spleens that showed a higher cellularity of the red pulp with more

macrophages and neutrophils, as well as more pronounced granulomatous lesions in the

mutant (Fig. S8). Additionally, Fig. S9 shows that the  $\Delta romA$  mutant triggered an increased

inflammatory response as early as 48 hrs post-infection, indicating that the strain per se is

significantly more pro-inflammatory and that the robust inflammation it induced was not the

consequence of an increased bacterial proliferation. To further analyze the kinetics of the

infection in mice; we determined spleen colonization of the mutant in comparison with the

wild type parental strain at 7 and 42 days post-infection. We did not observe differences at

these two time points (Fig. S10) indicating that the increased bacterial load is a phenomenon

restricted to the acute phase of the infection and it does not persist during the chronic phase.

In both time points we observed an enhanced splenomegaly (not shown).

Absence of RomA alters the positioning of LptD, an LPS biosynthetic protein in the

periplasmic space

The localization of RomA as well as the membrane-related phenotypes found in the  $\Delta romA$ 

mutant, suggested that this protein might be involved in the organization/localization of LPS

biosynthetic complexes in the periplasmic space and/or outer membrane. To evaluate if this

was the case we constructed a fusion of the gene lptD that codifies for a protein involved in

the transport of the LPS to the outer membrane [10, 21], with super-folder GFP (sfGFP) [22],

and evaluated its localization in the wild type and  $\Delta romA$  strains. As can be observed in Fig.

7A, LptD showed an altered distribution in the mutant in comparison with the parental strain.

More specifically, while most of the wild type cells showed a single localization spot, the

mutant cells had a more homogenous distribution. This was not observed with the inner

membrane protein responsible for flipping the O-antigen-lipid intermediate to the periplasm,

RfbD [9], (Fig. 7B) indicating that only some of the LPS biosynthesis proteins have an

aberrant distribution in the mutant.

## DISCUSSION

In the present study we have identified a novel gene (romA) in Brucella, and conserved in almost all α-proteobacteria, that codes for a small periplasmic protein with no known function. A mutant in romA is pleiotropic and displays several phenotypes, all related with an altered periplasm and/or outer membrane. The sensitivity of the mutant to several detergents and the fact that its membrane is more permeable to hydrophobic compounds indicated an altered outer membrane. LPS analysis showed that the mutant strain has several modifications. On one side, an altered smooth to rough ratio which results in a strain with significantly more assembled LPS compared to the wild type strain. Additionally, the LPS has an O-antigen with a higher degree of polymerization, with at least twice the amount of perosamines but substituted with formyl residues. These modifications have several implications and raises interesting questions for future studies. To our knowledge this is the first report of a mutant that has an altered smooth:rough LPS ratio, strongly suggesting that Brucella (and probably other members of this group) controls this equilibrium. It is tempting to speculate that a certain level of incomplete LPS is necessary to assemble or expose other outer membrane components that could be affected if the O-antigen is present in all LPS molecules. If this hypothesis is correct the amount of smooth to rough LPS could be determined by a compromise between two needs: to protect against harmful conditions encountered in the environment and to allow the assembly and positioning of a set of proteins or supramolecular structures necessary for motility, virulence, attachment and protein secretion among others. A similar hypothesis has been postulated for the length of the Oantigen and the Type III secretion system in Shigella [23]. The authors proposed that the length of the O-antigen is determined by two opposing necessities, its protective properties and the efficiency of the Type III injectisome. Currently we do not have a molecular explanation of why the  $\Delta romA$  mutant has a longer O-antigen but speculate that the stoichiometry of the LPS biosynthetic machinery is probably altered and that this affects the synthesis. In this regard we have shown that LptD, involved in the transport of the LPS to

the outer membrane [9], but not the flippase located in the inner membrane, showed a

mislocalization in the mutant strongly suggesting that RomA participates in the

organization/assembly of this machinery in the periplasm and/or outer membrane.

It is prompting to hypothesize that the modified LPS in the *romA* mutant probably disturbs the

homeostasis of the outer membrane and this impacts the virulence of the bacterium in several

ways. For example, even though it was less infective in the cellular model of infection it

triggered an exacerbated inflammatory response that actually increased the bacterial load in

the spleens of infected mice during the acute phase of the infectious process. This inability to

tune down the inflammatory response, a hallmark of the *Brucella* infection [1], is probably

the result of a combination of factors, both structural and functional, although we cannot

completely rule out at this stage that the modified LPS is the only component that could

account for all the phenotypes. It has been described that a mutant in the fliC gene in

Brucella, that codes for a component of a flagellar-like structure, induced an increased

splenomegaly and showed higher bacterial loads in the spleens as well as more tissue damage,

similar to the one we observed with the  $\Delta romA$  mutant [24]. Since the flagellum is assembled

in the outer membrane and because we have observed that the expression of fliC is not

affected (not shown) it could be speculated that the altered LPS affects the assembly of this

structure and this results in a similar phenotype as the *fliC* null mutant.

The fact that the mutant induces severe inflammation during the acute phase of the infectious

process indicates that Brucella has active mechanisms to modify its cellular structure in order

to tune down the immune response. This is not trivial, as it has been suggested that the default

cellular structure of Brucella is mainly responsible for its stealthy strategy and it implies that

the bacterium could actually modulate up and down the inflammatory response depending on

the phase of the infectious cycle or its needs, modifying its membrane composition or

structure. It could be speculated that the inflammatory balance and the necessity of the

bacteria to assemble membrane structures needed for virulence must be tightly equilibrated to

establish a successful chronic infection. Under this view, Brucella should be able to finely

counterbalance these two needs to be a successful pathogen.

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**CONFLICT OF INTEREST** 

The authors declare no conflict of interest.

**AUTHOR CONTRIBUTIONS** 

E.V. and J.E.U. conceived and designed research. E.V., J.M.S, C.C., F.R.F., A.C.C., S.G.A.,

K.A.P. and F.G. performed the experiments. J.C. provided input in the cytokine assays

determinations. A.S.C. designed the experiments and analyzed the O-antigen structure

determination. E.V. and J.E.U. wrote the manuscript.

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FIGURE LEGENDS

Figure 1. The protein product of Bab1\_1280 is a membrane-associated periplasmic

protein. A. Cell fractions from the merodiploid strain Bab1\_1280::3xFLAG were analyzed

by Western Blot with a monoclonal α-FLAG antibody. PE is periplasm and PR stands for

protoplasts. Monoclonal  $\alpha$ -Omp1 (porin) and  $\alpha$ -GroEL (Hsp60 homologue) antibodies were

used as outer membrane and cytoplasmic controls, respectively. **B.** Western blot with  $\alpha$ -

OMP1 and α-FLAG antibodies on SDS-PAGE of total membranes prepared from the

Bab1\_1280::3xFLAG strain. C. Total membranes from the Bab1\_1280::3xFLAG strain were

resuspended and incubated in different buffer conditions and ultracentrifuged. Pellet and

supernantant fractions were analyzed by Western Blot with a monoclonal α-FLAG antibody.

Sn stands for supernatant fraction.

Figure 2. The absence of Bab1\_1280 causes pleiotropic membrane defects. A. Detergent

sensitivity assays. Stationary phase cultures of either the wild type 2308 or the  $\Delta Bab1\_1280$ 

strains were grown at 37°C, diluted to OD<sub>600</sub>=1 and serial dilutions were plated in solid media

containing different detergents and incubated at 37°C for CFU determination. Sarkosyl: N-

lauroylsarcosine, Zwittergent: Zwittergent 3-16 and DOC: Sodium deoxycholate. The figure

shows significant differences between the means of both strains for Sarkosyl (\*p<0.01),

Zwittergent (\*p=0.006) and DOC (\*p=0.001). **B**. N-phenyl-1-naphtylamine incorporation

assay. Stationary phase cultures of either the wild type 2308 or the  $\Delta Bab1\_1280$  strains were

grown at  $37^{\circ}$ C, diluted to  $OD_{600}$ =0.1 and grown at  $37^{\circ}$ C until exponential phase was reached.

7.5x10<sup>8</sup> cells (resuspended in 250 µl of PBS) were used per well in 96-well black plates to

measure baseline fluorescence previous to adding NPN (10 µM final concentration) and

measurements were made every 18 seconds during 5 minutes. Relative Fluorescence Units

(RFU) were calculated by dividing each value by the mean obtained for the baseline for each

strain (\*p<0.001). In all cases (A-C), error bars are SD and p-values were calculated by the

unpaired t-tests. C. Two-dimension thin layer chromatography (2D-TLC) of total

phospholipids. Total [14C]acetate-labeled lipids were extracted from cultures in the presence

of choline and analyzed by 2D-TLC and autoradiography. Lipids spots corresponding to CL,

PG, OL, PE, and PC are indicated. CL, cardiolipin; PG, phosphatidylglycerol; OL, ornithine

lipid; PE, phosphatidylethanolamine; PC, phosphatidylcholine. **D.** Crystal violet staining.

Stationary phase cultures of either the wild type 2308 or the  $\Delta Bab1\_1280$  strains were grown

at 37°C, diluted to OD<sub>600</sub>=1 and serial dilutions were plated in solid media and incubated at

37°C until colonies were observed. The crystal violet staining was performed as described in

Experimental Procedures.

Figure 3. RomA is required for LPS homeostasis. A. Whole cell extracts of the wild type

2308 and  $\Delta romA$  strains were analyzed by Western Blot using a rabbit polyclonal  $\alpha$ -Brucella

antibody. Monoclonal  $\alpha$ -GroEL,  $\alpha$ -Omp1,  $\alpha$ -Omp16 and  $\alpha$ -Omp19 antibodies were used for

loading controls. B. Whole cells and periplasmic fractions were analyzed by Western Blot

using a α-Smooth LPS (S-LPS) or a α-Rough LPS (R-LPS). C. Densitometry of gels in panel

b. **D.** LPS was extracted from either the wild type 2308 or the  $\Delta romA$  strains and analyzed by

SDS-PAGE and Western Blot using the  $\alpha$ -S-LPS and  $\alpha$ -R-LPS antibodies (left panel), or

SDS-PAGE and LPS silver staining (right panel).

Figure 4. RomA is required for controlling the O-antigen length. A. Aminosugar analysis

by HPAEC-PAD of the oligosaccharides released after acid hydrolysis of the LPS of the wild

type 2308 or the  $\Delta romA$  strains. Comparison of the Rha4N/GlcN stoichiometry ratio in both

strains. **B.** LPS was purified from the wild type 2308  $\Delta romA$  mutant strains and the O-antigen

was released by acid hydrolysis. The released oligosaccharides were analyzed by MS in the

positive ion mode, and showed that the mutant strain has up to 25 perosamine subunits

(Rha4N) per O-antigen chain (m/z=4835.5) in comparison to the 14 subunits of the wild type

strain (m/z=3100.0). An inset of the wild type strain in the range of m/z=4200-5000 shows

the absence of the peak corresponding to an O-antigen with 25 subunits of perosamine. Fo,

formyl groups.

Figure 5. RomA is important for the intracellular cycle and its absence results in a

**hiperinflammatory strain**. A. Intracellular multiplication of wild type 2308,  $\Delta romA$  and

 $\Delta romA$  complemented strains in bone marrow derived macrophages (BMDM). \*p=0.005 and

\*\*p=0.002. Error bars are SD and p-values were calculated by the unpaired t-test. **B.** Spleens

extracted from the infected mice were homogenized for CFU determination by direct plating

after 15 days post-infection. \*p < 0.0001. Error bars are SEM and p-values were calculated by

the unpaired t-test. C. Spleens weight before homogenization. \*\*\*p=0.0001 and

\*\*\*\*p<0.0001. Error bars are SEM. **D.** Comparative sizes of spleens of mice intraperitoneally

infected with  $1x10^5$  CFU per animal of either the wild type 2308,  $\Delta romA$  and  $\Delta romA$ 

complemented strains and 15 days post-infection.

Figure 6. The mutant  $\Delta romA$  has an increased inflammatory response. A to D.

Inflammatory cytokines determined in the infected animals at 15 days post-infection.

Interferon gamma (IFN-γ) and Tumor necrosis factor alpha (TNF-α) levels were measured in

spleens (A and C) and sera (B and D) by ELISA as described in Experimental Procedures. In

all cases, significant differences were found between the means of the wild type and mutant

strains and between the means of the mutant and the complemented strain. No differences

were found between the wild type and complemented strain. A: \*p=0.03 and \*\*\*p=0.009; B:

\*p=0.02 and \*\*p=0.001; C: \*\*p=0.004 in both groups; **D**: \*\*p=0.001 in both cases. Error

bars are SEM and *p*-values were calculated using ordinary one-way ANOVA multiple comparisons between groups.

Figure 7. RomA participates in the positioning of LptD, a LPS transport protein. A. The outer membrane LPS transporter LptD was expressed with a C-terminal sfGFP from a pBBR4 plasmid in the B. abortus 2308 and \( \Delta romA \) strains. Bacteria were grown at 37°C until exponential phase, placed on agarose pads and analyzed by fluorescence confocal microscopy. B. The inner membrane O-antigen flippase RfbD was expressed with a Cterminal EYFP from pTRC-EYFP in the B. abortus 2308 and  $\Delta romA$  strains. Bacteria were grown at 37°C until exponential phase, placed on agarose pads and analyzed by fluorescence 

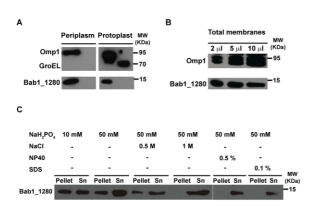


Figure 1



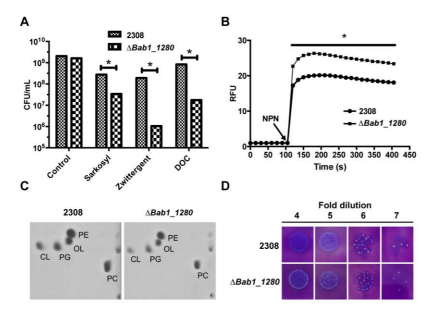


Figure 2

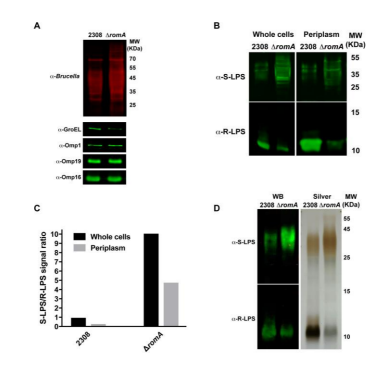
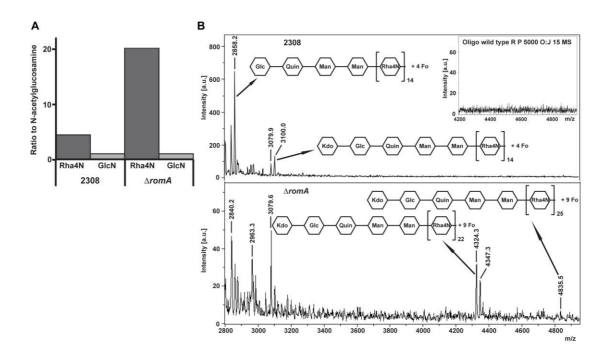


Figure 3



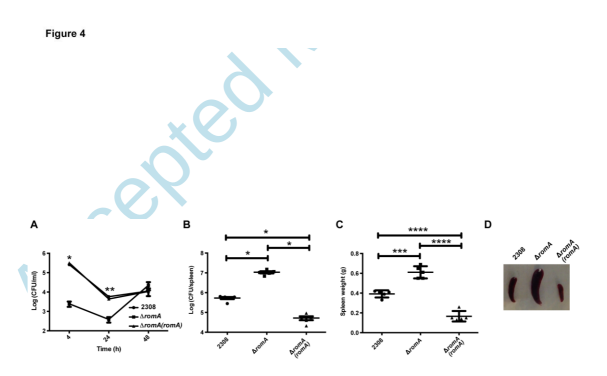


Figure 5

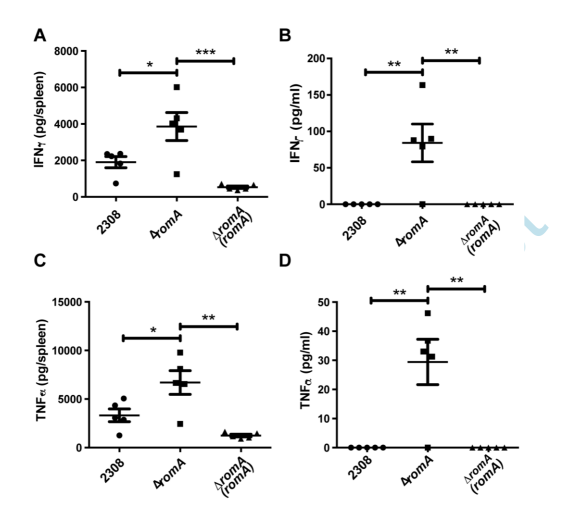


Figure 6



