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Brucella abortus depends on pyruvate phosphate dikinase and malic enzyme but not on Fbp and GlpX fructose-1,6-bisphosphatases for full virulence in laboratory models

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1 ***Brucella abortus* depends on pyruvate phosphate dikinase and malic enzyme but not on**
2 **Fbp and GlpX fructose-1,6-bisphosphatases for full virulence in laboratory models**

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28 Running title: *Brucella* metabolism in virulence models

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40 **ABSTRACT**

41 The brucellae are the etiological agents of brucellosis, a worldwide-distributed zoonosis. These
42 bacteria are facultative intracellular parasites, and thus are able to adjust their metabolism to the
43 extra and intracellular environments encountered during an infectious cycle. However, this
44 aspect of *Brucella* biology is imperfectly known and the nutrients available in the intracellular
45 niche are unknown. Here, we investigated the central pathways of C metabolism used by
46 *Brucella abortus* by deleting the putative fructose-1,6-bisphosphatases (*fbp* and *glpX*),
47 phosphoenolpyruvate carboxykinase (*pckA*), pyruvate phosphate dikinase (*ppdK*) and malic
48 enzyme (*mae*) genes. In gluconeogenic [but not in](#) rich media, growth of mutants $\Delta ppdK$ and
49 Δmae was severely impaired and growth of the double $\Delta fbp\text{-}\Delta glpX$ mutant was reduced. In
50 macrophages, only $\Delta ppdK$ and Δmae showed reduced multiplication, and studies with $\Delta ppdK$
51 confirmed that it reached the replicative niche. Similarly, only $\Delta ppdK$ and Δmae were attenuated
52 in mice, the former being cleared by week 10 and the latter persisting longer than 12 weeks. We
53 also investigated the glyoxylate cycle. Although *aceA* (isocitrate lyase) promoter activity was
54 enhanced in rich medium, *aceA* disruption had no effect *in vitro* or on multiplication in
55 macrophages or mouse spleens. The results suggest that *B. abortus* grows intracellularly using
56 a limited supply of 6 C (and 5 C) sugars that is compensated by glutamate and possibly other
57 amino acids entering the Krebs cycle without a critical role of the glyoxylate shunt.

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67 INTRODUCTION

68 *Brucella* is a genus of gram-negative bacteria that groups the causative agents of
69 brucellosis, a worldwide-extended zoonosis severely affecting animal production and human
70 welfare. Three species, *B. abortus*, *B. melitensis* and *B. suis*, are the most common causes of
71 brucellosis in domestic livestock and humans. These brucellae can grow both *in vitro* and within
72 host cells, and their pathogenicity results largely from their capacity to escape a prompt
73 detection by innate immunity and the use of a type IV secretion system to reach the replicative
74 niche, an endoplasmic reticulum derived vacuole (1-6). In this compartment, these bacteria
75 multiply extensively, which shows their ability to use efficiently substrates provided by the host.
76 However, there is only sparse information on either the nature of these substrates or the
77 metabolic pathways used in the replicative niche (7).

78 *In vitro*, most strains of *B. abortus*, *B. melitensis* and *B. suis* grow in several simple
79 chemically defined media (8). Among these, Gerhardt's medium contains glycerol, lactate,
80 glutamate and mineral salts plus nicotinic acid, thiamine, pantothenic acid and biotin as growth
81 factors. This medium supports growth better than other simple defined media, including those
82 that provide glucose as the C source (8), and this has been attributed to the ability of these
83 bacteria to use glutamate very efficiently through the tricarboxylic acid cycle (TCA) (9). Indeed,
84 growth in this medium is clear proof of the ability of these bacteria to carry out gluconeogenesis
85 *in vitro*. Concerning the catabolism of glucose, it has been accepted that it proceeds through the
86 pentose phosphate pathway (in conjunction with TCA), rather than through the glycolysis or
87 Entner–Doudoroff pathways (Fig. 1). This view, although supported by early radiorespirometric
88 and enzymatic studies (10,11) performed with the attenuated *B. abortus* S19 vaccine, is not
89 consistent with the presence of the genes putatively encoding all the enzymes of the Entner-
90 Doudoroff pathway (12). It is possible that the multiple defects in S19 (13) and/or the

91 experimental conditions in those early experiments precluded detection of key enzymes of the
92 Entner-Doudoroff pathway (12).

93 Concerning intracellular metabolism, some data come from randomly obtained mutants that
94 show attenuation in human or mouse macrophages or in HeLa cells. Genes (putative functions)
95 identified in this manner include, in *B. suis* 1330, a *gguA* homologue (*gluP*, presumably involved
96 in sugar uptake), some erythritol catabolism genes, *gnd* (6-phosphogluconate dehydrogenase,
97 required for using glucose through the pentose phosphate pathway), *rbsk* (ribose kinase), *pyc*
98 (anaplerotic pyruvate carboxylase), *pgi* (phosphoglucose isomerase) and genes related to the
99 biosynthesis of amino acids (14-16); in *B. melitensis* 16M, *dbbA* and *ugpA* (ribose and glycerol-
100 3-phosphate transporters, respectively), *glpD* (glycerol-3-phosphate dehydrogenase) and some
101 erythritol catabolism genes (17,18); and in *B. abortus* 2308, *gluP* (previously shown to encode
102 an active glucose and galactose transporter in this species (19)), *gnd* (6-phosphogluconate
103 dehydrogenase), *gltD* (glutamate synthase) and *gcvB* (glycine dehydrogenase) (20). Also in *B.*
104 *abortus* 2308, *dxs* (an isoprenoid biosynthesis transketolase) and *mocC* (rhizopine or inositol
105 catabolism) were identified as expressed in macrophages using a fluorescent reporter (21).

106 Proteomic analyses have also provided clues on the metabolism of brucellae in the host.
107 Forty-eight h after infection of mouse macrophages with *B. suis* 1330, Al Dahouk et al. (22)
108 found an important reduction of proteins putatively involved in energy, protein and nucleic acid
109 metabolism. Some exceptions were ribitol kinase, glyceraldehyde-3-P-dehydrogenase and the
110 isocitrate lyase (*AceA*) of the glyoxylate cycle. However, other studies in *B. suis* 1330 do not
111 support the use of the glyoxylate cycle within host cells (15). Lamontagne et al. (23) analyzed *B.*
112 *abortus* 2308 protein expression 3, 20 and 44 h after infection of RAW 264.7 macrophages.
113 They found that multiple proteins associated with sugar uptake, TCA, the pentose phosphate
114 shunt and the subsequent generation of pyruvate were down-regulated 3 h after infection. At 24
115 h, several proteins involved in sugar metabolism and transport were also reduced. Enzymes

116 associated with protein and amino acid catabolism were mainly increased early (3 h) but also 24
117 h after infection, when bacteria were already in vacuoles derived from the endoplasmic
118 reticulum. This was also the case of enzymes involved in glutamate synthesis, suggesting
119 conversion of amino acids into glutamate and α -ketoglutarate. Accordingly, amino acid-based
120 alternatives may be the preferred solution for *B. abortus* to derive precursors for the TCA cycle
121 and ancillary routes during the midpoint time course of infection. At later times, the same authors
122 observed an increase in proteins involved in transport, suggesting that the endoplasmic
123 reticulum is able to supply at least some of the substrates required for bacterial growth.
124 Likewise, the pentose phosphate shunt seemed to partially resume its functions.

125 Although the information given by these studies is valuable, the central metabolic pathways
126 used by *Brucella* during infection remain unclear. The results are contradictory in some cases,
127 as for the glyoxylate cycle or the metabolic activity in cells. In addition, some studies suggest the
128 availability of sugars in the replicative niche whereas others indicate that amino acids could be
129 the preferred C source *in vivo*, which may require a gluconeogenic metabolism. Indeed,
130 apparently conflicting data may result from the use of different host cell lines, different times of
131 analysis, polarity of mutations and other experimental conditions. Moreover, there might be
132 some variation among *B. suis* 1330, *B. melitensis* 16M and *B. abortus* 2308, as suggested by
133 the known differences in oxidative rates of sugars and amino acids (24). In this work, we attempt
134 to answer some aspects of the central metabolic pathways used by *B. abortus* in the host. For
135 this purpose, we focused our research on genes putatively involved in classical
136 gluconeogenesis, the anabolic pathways bridging TCA and the triose-phosphate pathway, and
137 the glyoxylate cycle. We constructed in-frame mutants in genes coding for key enzymes and
138 tested them in complex and chemically defined media and for multiplication within cultured cells
139 and for persistence in the mouse model (25). Together with some of the previous analyses, our
140 observations suggest a model of *B. abortus* metabolism in which, although TCA supplies

141 molecules necessary for biosynthesis and subsequent growth, classical fructose-1,6-
142 bisphosphatases Fbp and GlpX are not necessary and 6 and/or 5 C molecules for polymer
143 biosynthesis are obtained mostly from the intracellular milieu.

144 **MATERIAL AND METHODS**

145 **Bacterial strains and growth conditions.** The bacterial strains and plasmids used in this study
146 are listed in Table 1S (supplemental material), and their origin and characteristics are described
147 in previous works (26-28). The strains resulting from the genetic manipulations described below
148 were characterized according to standard *Bruceella* typing procedures: i.e. colonial morphology
149 after 3 days of incubation at 37°C, crystal violet-oxalate exclusion, urease, acriflavine
150 agglutination, sensitivity to Tb, Wb, Iz and R/C phages, agglutination with anti-A and anti-M
151 monospecific sera, CO₂ and serum dependence, and susceptibility to thionin blue, fuchsin and
152 safranin (24). Bacteria were routinely grown in standard Peptone-Yeast Extract-Glucose broth
153 (Biomerieux) or in this media supplemented with agar (TSA). The following antibiotics were used
154 at the indicated concentrations: kanamycin (Km; 50 µg/mL), nalidixic acid (Nal; 25 µg/mL),
155 chloramphenicol (Cm; 20 µg/mL) and/or gentamicin (Genta; 100 µg/mL or 25 µg/mL) (all from
156 Sigma). When needed, media was supplemented with 5% sucrose (Sigma). All strains were
157 stored at – 80°C in skim milk (Scharlau). To study the phenotype of the metabolic mutants,
158 Peptone-Yeast Extract-Glucose or the medium of Gerhardt (henceforth Glutamate-Lactate-
159 Glycerol) were used (29). The components (for 1 L) of the latter were: glycerol (30 g), lactic acid
160 (5 g), glutamic acid (1.5 g), thiamine (0.2 mg), nicotinic acid (0.2 mg), pantothenic acid (0.04
161 mg), biotin (0.0001 mg), K₂HPO₄ (10 g), Na₂S₂O₃·5H₂O (0.1 g), MgSO₄ (10 mg), MnSO₄ (0.1
162 mg), FeSO₄ (0.1 mg), NaCl (7.5 g). The pH was adjusted to 6.8-7.

163 **Growth measurements.** Inocula pre-conditioned to the conditions in the test medium (Peptone-
164 Yeast Extract-Glucose, Glutamate-Lactate-Glycerol, Glycerol-Glutamate, Glycerol-Lactate or
165 Glutamate-Lactate) were prepared as follows. First, the strains to be tested were inoculated into

166 10 mL of Peptone-Yeast Extract-Glucose in a 50 mL flask and incubated at 37°C with orbital
167 shaking for 18 h. These exponentially growing bacteria were harvested by centrifugation,
168 resuspended in 5 mL of the test medium at an optical density (O.D._{600nm}) of 0.1 and incubated at
169 37°C with orbital shaking for 18 h. Then, these pre-conditioned bacteria were harvested by
170 centrifugation, resuspended at an O.D._{600nm} of 0.1 (0.05 starting in the Bioscreen apparatus) in
171 the same test medium in Bioscreen multi-well plates (200 µL/well) and cultivated in a Bioscreen
172 C (Lab Systems) apparatus with continuous shaking at 37°C. Absorbance values at 420-580_{nm}
173 were automatically recorded at 0.5 h intervals over a 120 to 300 h period. All experiments were
174 performed in triplicate. Controls with culture medium and no bacteria were included in all
175 experiments.

176 **DNA manipulations.** Genomic sequences were obtained from the Kyoto Encyclopedia of
177 Genes and Genomes (KEGG) database (<http://www.genome.jp/kegg/>). Searches for DNA and
178 protein homologies were carried out using the National Center for Biotechnology Information
179 (NCBI; <http://www.ncbi.nlm.nih.gov/>) and the European Molecular Biology Laboratory (EMBL) -
180 European Bioinformatics Institute server (<http://www.ebi.ac.uk/>). Primers were synthesized by
181 Sigma-Genosys (Haverhill, UK). DNA sequencing was performed by the “Servicio de
182 Secuenciación del Centro de Investigación Médica Aplicada” (Pamplona, Spain). Restriction-
183 modification enzymes were used under the conditions recommended by the manufacturer.
184 Plasmid and chromosomal DNA were extracted with Qiaprep spin Miniprep (Qiagen) and
185 Ultraclean Microbial DNA Isolation kits (Mo Bio Laboratories), respectively. When needed, DNA
186 was purified from agarose gels using the Qiack Gel extraction kit (Qiagen).

187 In-frame deletion mutants in *fbp* and *glpX* were constructed by polymerase-chain
188 reaction (PCR) overlap using genomic DNA of *B. abortus* 2308 as the DNA template. Primers
189 were designed using the *B. abortus* 2308 sequences available in KEGG
190 (<http://www.genome.jp/kegg/>). For the construction of the *fbp* mutant, two PCR fragments were

191 generated: oligonucleotides *fbp*-F1(5'-GTAGCCAAAAAGCCCAGGT-3') and *fbp*-R2 (5'-
192 GCCAACCCAGAACCAGAGGA-3') were used to amplify a 203 bp fragment including codons 1–
193 14 of the *fbp* ORF as well as a 161 bp fragment upstream of the *fbp* start codon, and
194 oligonucleotides *fbp*-F3 (5'-TCCTCTGGTTCTGGTTGGCGTGGCCGAAGAGGTGGATA-3') and
195 *fbp*-R4 (5'-CATTTGCCGCTTCCATGA-3') were used to amplify a 193 bp fragment including
196 codons 327-341 of the *fbp* ORF and a 148 bp fragment downstream of the *fbp* stop codon. Both
197 fragments were ligated by PCR using oligonucleotides *fbp*-F1 and *fbp*-R4 for amplification, and
198 the complementary regions between *fbp*-R2 and *fbp*-F3 for overlapping. The resulting fragment,
199 containing the *fbp* deletion allele, was cloned into pCR2.1 (Invitrogen) to generate plasmid pAZI-
200 1, sequenced to ensure that the reading frame was maintained, and subcloned into the *Bam*HI
201 and the *Xba*I sites of the suicide plasmid pJQKm (30). The resulting mutator plasmid (pAZI-2)
202 was introduced into *B. abortus* 2308 by conjugation (26). Integration of the suicide vector was
203 selected by Nal and Km resistance, and the excisions (generating both the *fbp* mutant
204 [*BABΔfbp*] and a sibling revertant strain carrying an intact gene [*BABfbp*-sibling revertant]) were
205 then selected by Nal and sucrose resistance and Km sensitivity. The resulting colonies were
206 screened by PCR with primers *fbp*-F1 and *fbp*-R4, which amplified a fragment of 396 bp in the
207 mutant and a fragment of 1332 bp in the sibling revertant strain. The mutation resulted in the
208 loss of about 98% of the *fbp* ORF and the mutant strain was called *BABΔfbp*.

209 The *glpX* mutant was constructed in a similar way. Primers *glpX*-F1 (5'-
210 ACGGTGATTCTGGTGACACA-3') and *glpX*-R2 (5'-CGAGCTCCAGTGTGAGAATG-3') were
211 used to amplify a 576 bp fragment including 61 bp of the *glpX* ORF as well as 515 bp upstream
212 of the *glpX* start codon, and primers *glpX*-F3 (5'-
213 CATTCTCACACTGGAGCTCGATACGACAGATCCGGACGAG-3') and *glpX*-R4 (5'-
214 CATCATACAGTTGCCGATGG-3') were used to amplify a 574 bp fragment including 371 bp of
215 the *glpX* ORF and 203 bp downstream of the *glpX* stop codon. Both fragments were ligated by

216 overlapping PCR using primers *glpX*-F1 and *glpX*-R4, and the fragment containing the deletion
217 allele was cloned into pCR2.1 to generate plasmid pAZI-3, sequenced to confirm that the *glpX*
218 ORF had been maintained, and subcloned in pJQKm to produce the mutator plasmid pAZI-4.
219 This plasmid was then introduced into *B. abortus* 2308 and the deletion mutant generated by
220 allelic exchange was selected by Nal and sucrose resistance and Km sensitivity and by PCR
221 using oligonucleotides *glpX*-F1 and *glpX*-R4, which amplified a fragment of 1150 bp in the
222 deletion strain and a fragment of 1705 bp in the *BABglpX*-sibling revertant strain. The mutation
223 resulted in the loss of approximately 56% of the *glpX* ORF, and the mutant was called
224 *BABΔglpX*.

225 To construct the *BABΔfbpΔglpX* double mutant, the mutator plasmid pAZI-4 was
226 introduced into strain *BABΔfbp*. After allelic exchange, the double mutant was selected as
227 described above using primers *glpX*-F1 and *glpX*-R4.

228 *BABΔaceA* was constructed using the same strategy. Oligonucleotides *aceA*-F1 (5'-
229 TGACAAGATATCGCCAAAACAC-3') and *aceA*-R2 (5'-CGAAGGGATGAGGCTGTAAA-3')
230 amplified a 238 bp fragment, including codons 1-10 of the *aceA* ORF and 208 bp upstream of
231 the *aceA* start codon. Oligonucleotides *aceA*-F3 (5'-TTTACAGCC
232 TCATCCCTTCGGAAACCGCACAGTTCAAGC-3') and *aceA*-R4 (5'-GGATCAAGAGATCA
233 CCCCAGT-3') amplified a 278 bp fragment including codons 420-430 of the ORF *aceA* and 245
234 bp downstream of the *aceA* stop codon. Both fragments were ligated by overlapping PCR using
235 oligonucleotides *aceA*-F1 and *aceA*-R4. The PCR product was cloned into pCR2.1 to generate
236 pAZI-7, sequenced and subcloned into pJQKm to produce the suicide plasmid pAZI-8. *B.*
237 *abortus* 2308 mutants were selected by PCR using oligonucleotides *aceA*-F1 and *aceA*-R4.
238 PCR products were 1743 bp in *BABaceA*-sibling revertant strain and 738 bp in *BABΔaceA*. This
239 mutation eliminated 78% of the *aceA* ORF.

240 For the construction of the *pckA* mutant, oligonucleotides *pckA*-F1 (5'-
241 TGTTTGCAGTTTTCCACACC-3'), *pckA*-R2 (5'-AATCGAAGCGGCCTTATTGT-3'), *pckA*-F3 (5'-
242 ACAATAAGGCCGCTTCGATTGACGGCTCGCTGAACAAT-3') and *pckA*-R4 (5'-TCTTGCGA
243 TAACAGCCAAAA-3') were used. Primers *pckA*-F1 and *pckA*-R2 amplified a 219 bp fragment,
244 which included codons 1-13 of the *pckA* ORF and 180 bp upstream of the *pckA* start codon.
245 Primers *pckA*-F3 and *pckA*-R4 amplified a 319 bp fragment including the last 37 codons of the
246 *pckA* ORF and 208 bp downstream of the *pckA* stop codon. Both PCR products were ligated by
247 overlapping PCR using *pckA*-F1 and *pckA*-R4, cloned into pCR2.1 to generate plasmid pAZI-5
248 and subsequently subcloned into the *Bam*HI and the *Xba*I sites of the suicide plasmid pJQKm.
249 The resulting mutator plasmid pAZI-6 was introduced into *B. abortus* 2308, where it was
250 integrated in the chromosome. A second recombination generated the excision of the plasmid.
251 The resulting colonies were screened by PCR (with *pckA*-F1 and *pckA*-R4) amplifying a
252 fragment of 538 bp in the mutant and a fragment of 1864 bp in the sibling revertant strain. The
253 mutant strain was called *BAB* Δ *pckA* and lacked the 71.14% of the *pckA* ORF.

254 *BAB* Δ *ppdK* was constructed using primers *ppdK*-F1(5'-CTCCCGATTTCATTTTTACAG-3')
255 and *ppdK*-R2 (5'-TGCTCATTTTCAGCCAGGTT-3') to amplify a 288-bp fragment including the
256 first 103 bp of the *ppdK* ORF, as well as 185 bp upstream of the *ppdK* start codon and primers
257 *ppdK*-F3 (5'-AACCTGGCTGAAATGAGCACGGTCTCGACTATGTGTCC-3') and *ppdK*-R4 (5'-
258 TCAACGCATCAAAGCAGAAG-3') to amplify a 220 bp including the last 86 bp of the *ppdK* ORF
259 and 134 bp downstream of the *ppdK* stop codon. Both fragments were ligated by overlapping
260 PCR using primers *ppdK*-F1 and *ppdK*-R4 and the fragment obtained, containing the deletion
261 allele, was cloned into pCR2.1 to generate pMZI-1, sequenced to confirm that the reading frame
262 had been maintained, and subcloned into pJQKm to produce the mutator plasmid pMZI-2. This
263 plasmid was introduced into *B. abortus* 2308 and both the deletion mutant and the sibling
264 revertant strain generated by allelic exchange was selected by Nal and sucrose resistance and

265 Km sensitivity, and by PCR using *ppdK*-F1 and *ppdK*-R4 which amplified a fragment of 508 bp in
266 *BABΔppdK* and a fragment of 2983 bp in *BABppdK*-sibling revertant strain. The mutation
267 generated resulted in the loss of the 93% of *ppdK*.

268 Primers *ppdKII*-F1 (5'-CTCCCGATTCATTTTTTCACG-3'), *ppdKII*-R2 (5'-
269 CTGCTCATTTTCAGCCAGGTT-3'), *ppdKII*-F3 (5'-
270 AACCTGGCTGAAATGAGCAGCGGGTCTCGACTATGTGTCC-3') and *ppdKII*-R4 (5'-
271 TCAACGCATCAAAGCAGAAG-3') were used to obtain the mutator plasmid pAZI-10. This
272 plasmid was introduced into *B. abortus* 2308 to obtain a *ppdK* mutant that maintained only the
273 34 first amino acids of PpdK (*BABΔppdK-II*). This mutant had the same phenotype as the one
274 previously described. Thus, the mutator plasmid pAZI-10 was also introduced into strain
275 *BABΔpckA* carrying the *pckA* mutation to obtain the double mutant *BABΔpckAΔppdK*.

276 *BABΔmae* was constructed using primers *mae*-F1 (5'-TATGACGGCGCACTTGTCTA-3')
277 and *mae*-R2 (5'-TCGGATAGCGATGGAAGAAC -3') to amplify a 341 bp fragment including 76
278 bp of the *mae* ORF, as well as 265 bp upstream of the *mae* start codon, and primers *mae*-F3 (5'-
279 GTTCTTCCATCGCTATCCGAGCGAAGCCAATCTTCTGGTA -3') and *mae*-R4 (5'-
280 CGCCATAAAACGAACCTCAA-3') to amplify a 376 bp including 227 bp of the *mae* ORF and
281 149 bp downstream of the *mae* stop codon. Both fragments were ligated by overlapping PCR
282 using primers *mae*-F1 and *mae*-R4 and the fragment obtained, containing the deletion allele,
283 was cloned into pCR2.1 to generate pMZI-3, sequenced to confirm that the *mae* ORF had been
284 maintained, and subcloned into pJQKm to produce the mutator plasmid pMZI-4. This plasmid was
285 then introduced into *B. abortus* 2308 and the mutant and sibling revertant strains generated by
286 allelic exchange were selected by Nal and sucrose resistance and Km sensitivity and by PCR
287 using oligonucleotides *mae*-F1 and *mae*-R4, which amplified a fragment of 717 bp in the deleted
288 strain and a fragment of 2739 bp in the sibling revertant strain. The mutation generated resulted
289 in the loss of the 87% of the *mae* ORF.

290 To check the different mutations, we used internal primers (*gene*-R5) hybridizing in the deleted
291 regions.

292 For complementation, a plasmid carrying *ppdK* was constructed using the Gateway cloning
293 Technology (Invitrogen). Since the sequence of *ppdK* from *B. abortus* and *B. melitensis* is 99%
294 identical, the clone carrying *ppdK* was extracted from the *B. melitensis* ORFEOMA and the ORF
295 subcloned into plasmid pRH001 (31) to produce plasmid pAZI-19. This plasmid was introduced
296 into *BABΔppdK* by mating with *E. coli* S17 λ*pir* and the conjugants harboring this plasmid
297 (designated as *BABΔppdK* pAZI-19) were selected by plating the mating mixture onto tryptic soy
298 agar (TSA)-Nal-Km plates. [For the construction of *BABΔfbpΔglpX* pAZI-21, gene *fbp* was](#)
299 [amplified from *BAB*-parental using primers *fbp*-Fp \(5'-](#)
300 [GGGATCCATGCTTCTGAAAGGGTGGTACCG-3'\)](#) and [fbp-R4 \(5'-CATTGCGCGCTTCCATGA-](#)
301 [3'\) and cloned into pCR2.1. The resulting plasmid was sequenced, and the *fbp* gene was](#)
302 [subcloned into the *Bam*HI and the *Xho*I sites of the vector pBBR1MCS1. The resulting plasmid](#)
303 [\(pAZI-21\) was introduced into *BABΔfbpΔglpX* by conjugation \(see above\). *BABΔfbpΔglpX* pAZI-](#)
304 [23 was constructed following the same strategy using primers *glpX*-F1 \(5'-](#)
305 [ACGGTGATTCTGGTGACACA-3'\) and *glpX*-R4 \(5'-CATCATACAGTTGCCGATGG-3'\) to amplify](#)
306 [glpX.](#)

307 **Gene expression studies.** To determine whether *aceA* was expressed *in vitro*, its promoter was
308 fused with the luciferase reporter gene. To this end, *aceAp*-F (5'-
309 [GGGATCCTAGTTGCGCTCGATCAGATT-3'\)](#) and *aceAp*-R (5'-
310 [TTCTAGACATTTGGTGTCTCCTCGT-3'\)](#) (respectively containing *Bam*HI and *Xba*I sites;
311 underlined) were used to amplify a 382bp region containing the ATG and the *aceA* promoter
312 from *B. abortus* 2308 genomic DNA. This PCR product was verified by electrophoresis and
313 ligated into the vector pGEM-T Easy (Promega), thereby originating plasmid pAZI-17. Then, the
314 insert was digested with *Bam*HI and *Xba*I and ligated to the pSKOriTKmluxAB to generate

315 plasmid pAZI-18. This plasmid was introduced into *E. coli* S17 λ *pir* Nal^S and then transferred by
316 conjugation to *B. abortus* 2308 Nal^R Km^S. Cells were plated on TSA NalKm. Positive clones
317 gave a 510 bp band when verified by PCR using *aceAp*-F and *luxAB*-R. The resulting strain was
318 called *B. abortus* pBABaceA-*luxAB*. To measure luciferase activity, fresh *B. abortus* pBABaceA-
319 *luxAB* were adjusted (OD_{600nm}) to 0.4 in saline and finally, 50 or 200 μ L were added to flasks with
320 10 mL of Peptone-Yeast Extract-Glucose or Glutamate-Lactate-Glycerol, respectively. Growth
321 was followed by measuring absorbance at OD_{600nm} and 1 mL aliquots were taken at selected
322 intervals to measure the luminescence in Relative Luminescence Units (RLU) after addition of
323 100 μ L ethanol:decanal (1:1).

324 **Cell infections and intracellular trafficking.** *In vitro* infection assays were performed in RAW
325 264.7 macrophages (ATCC TIB-71) and HeLa cells (ATCC CCL-2) cultured in Dulbecco's
326 Modified Eagle Medium (DMEM; Gibco) with 10% (v/v) heat-inactivated fetal bovine serum
327 (Gibco), 1% (v/v) L-glutamine 200 nM (Sigma Aldrich) and 1% (v/v) non-essential amino acids
328 (Gibco). Then, 24-well plates were seeded with 1×10^5 cells/well, and macrophages and HeLa
329 cells were infected at a multiplicity of infection of 50:1 and 200:1 (bacteria:cell), respectively, by
330 centrifuging the plates at $400 \times g$ for 10 min at 4°C. After incubation for 15 min at 37°C under a
331 5% CO₂ atmosphere, extracellular bacteria were removed with four DMEM washes followed by
332 Genta treatment (100 μ g/mL) for 90 min. Then, fresh medium supplemented with 25 μ g/mL of
333 Genta was added and incubation carried on. Two, 24 and 48 h later, cells were washed three
334 times with 100 mM phosphate buffered saline (pH 7.2), lysed with 0.1% (v/v) Triton X-100 in
335 phosphate buffered saline, and plated on TSA to determine the number of intracellular bacteria.
336 All experiments were performed in triplicate (32). Results were expressed as mean and standard
337 error (n=3) of individual log₁₀ CFU/well. Statistical comparison of means was performed by a
338 one-way ANOVA followed by the Fisher's Protected Least Significant Differences (PLSD) tests
339 (33).

340 For immunofluorescence microscopy, RAW 264.7 macrophages and HeLa cells were
341 grown on coverslips and inoculated with bacteria as described above. Cells were fixed in 3%
342 paraformaldehyde in 100 mM phosphate buffered saline (pH 7.2) at 37°C for 10 min. Cells were
343 washed twice with phosphate buffered saline and permeabilized with 0.1% (v/v) Triton X-100
344 and 3% bovine serum albumin (Sigma), for 30 min. Coverslips were incubated with primary
345 antibodies for 45 min at room temperature, washed three times in the same phosphate buffered
346 saline supplemented with 3% bovine serum albumin, and then incubated with the appropriate
347 secondary antibodies. Coverslips were washed three times with phosphate buffered saline and
348 once with H₂O and mounted onto glass slides using Mowiol 4–88. Samples were examined and
349 images acquired using a Leica TCS SP5 laser scanning confocal microscope at the “UNAmur”
350 (Namur, Belgium). The primary antibodies used for immunofluorescence microscopy were rabbit
351 anti-calnexin (SPA-860, Stressgen) and a mouse anti-S-LPS monoclonal antibody
352 (A76/12G12/F12). The secondary antibodies were donkey anti-rabbit IgG conjugated to Alexa
353 Fluor 488, (Invitrogen), and goat anti-mouse IgG conjugated to Alexa Fluor 546 (Invitrogen). For
354 lysosomal labeling, the primary antibody used was rat anti-mouse (DBHS) and the secondary
355 antibody was goat anti-rat anti IgG conjugated to Alexa Fluor 633 (Invitrogen).

356 **Assays in mice.** Female BALB/c mice (Charles River, France) were kept in cages with water
357 and food *ad libitum*, and accommodated under P3 biosafety containment conditions 2 weeks
358 before and during the experiments, in the facilities of the "Instituto de Agrobiotecnología"
359 (registration code ES/31-2016-000002-CR-SU-US). The animal handling and other procedures
360 were in accordance with the current European (directive 86/609/EEC) and Spanish (RD
361 53/2013) legislations, supervised by the Animal Welfare Committee of the "Universidad Pública
362 de Navarra", and authorized by the competent authority of "Gobierno de Navarra". To prepare
363 inocula, TSA grown bacteria were harvested, adjusted spectrophotometrically ($O.D._{600nm} = 0.170$)
364 in 10 mM phosphate buffered saline (pH 6.85) and diluted in the same diluent up to

365 approximately 5×10^5 colony forming units (CFU)/mL (exact doses were assessed
366 retrospectively). For each bacterial strain, five mice were intraperitoneally inoculated with 0.1
367 mL/mouse and the CFU number in spleen was determined at different weeks post-inoculation as
368 described previously (33). The identity of the spleen isolates was confirmed by PCR. The
369 individual number of CFU/spleen was normalized by logarithmic transformation, and the mean
370 log CFU/spleen values and the standard deviation were calculated for each group of mice (n=5).
371 Statistical comparisons were performed by a one-way ANOVA followed by the Fisher's
372 Protected Least Significant Differences (PLSD) tests (33).

373 RESULTS

374 **Dysfunction of *B. abortus fbp*, *glpX*, *ppdK* and *mae* but not of *pckA* or *aceA* homologues**

375 **affects growth on gluconeogenic substrates *in vitro*.** The conversion of fructose-1,6-bisP
376 into fructose-6-P mediated by the cognate bisphosphatase(s) (FBPases) is the only irreversible
377 step among those taking part in gluconeogenesis in *Brucella* (Fig. 1). Therefore, FBPase activity
378 is strictly necessary to grow under gluconeogenic conditions.

379 As in *E. coli* and *Salmonella*, the *B. abortus* 2308 genome presents two ORFs of putative
380 FBPases: BAB2_0364 and BAB1_1292. The former is predicted to encode a protein of 340
381 amino acids that belongs to the class I FBPases (Fbp), and the latter a 328 amino acid protein
382 ortholog to *E. coli* GlpX, a FBPase of class II (Fig. 1). In *E. coli*, Fbp is connected to the
383 production of fructose-6-phosphate for nucleotide, polysaccharide and aromatic amino acid
384 biosynthesis. GlpX, on the other hand, belongs to an operon (*glpFKX*) related to phospholipid
385 biosynthesis (34). The putative *B. abortus glpX*, however, seems isolated and not part of any
386 obvious operon.

387 We constructed mutants carrying in frame deletions in the putative *fbp* (*BABΔfbp* mutant)
388 and *glpX* (*BABΔglpX*) as well as in both genes (*BABΔfbpΔglpX*), and tested their growth in a
389 complex (Peptone-Yeast Extract-Glucose) medium and in the chemically defined medium of

390 Gerhardt (Glutamate-Lactate-Glycerol). In the complex medium, the three mutants grew with
391 generation times and final yields similar to those of *B. abortus* 2308 (*BAB*-parental) (Fig. 2A.1).
392 *BAB*-parental showed reduced growth rates and final yields in Glutamate-Lactate-Glycerol, as
393 expected, and we obtained identical results with either *BABΔfbp* or *BABΔglpX* (Fig. 2A.2). On
394 the other hand, *BABΔfbpΔglpX* produced a markedly lower increase in turbidity in the minimal
395 medium (Fig. 2A.2). [Complementation with plasmid pAZI-21 carrying *fbp* or with pAZI-23](#)
396 [carrying *glpX* restored the ability to grow in Glutamate-Lactate-Glycerol to levels close to that of](#)
397 [the parental strain \(Supplemental Material; Fig. S3\).](#) These experiments strongly suggest that *B.*
398 *abortus* Fbp and GlpX are functional, an interpretation reinforced by their reciprocal
399 complementation (i.e., Fbp complemented *BABΔglpX* and, conversely, GlpX complemented
400 *BABΔfbp*). We then tested combinations of two C sources (Glycerol-Glutamate, Glycerol-Lactate
401 or Glutamate-Lactate). Whereas *BABΔglpX* was not affected, *BABΔfbp* showed retarded growth
402 only in the absence of glycerol (i.e. in Glutamate-Lactate; Fig. 3, upper panels). Since GlpX is
403 the FBPase remaining in *BABΔfbp* and this mutant grew normally when glycerol was present,
404 this result suggests that, as in *E. coli* (34), GlpX is related to glycerol metabolism. Finally,
405 although retarded and diminished, the double *BABΔfbpΔglpX* mutant still showed significant
406 growth (Fig. 3, upper panels).

407 During these experiments we noticed that *BABΔfbpΔglpX* inoculated broths did not produce
408 a homogeneous growth. This was clearly observed when the double mutant was grown in
409 Glutamate-Lactate-Glycerol in side-arm flasks instead of the automated Bioscreen system.
410 Under these conditions, the bacteria formed macroscopic aggregates (Fig. 4) settling on the
411 bottom of the flasks that indicated a profound surface modification consistent with an altered
412 biosynthesis of envelope molecules. In summary, although Fbp and GlpX deficiency did not
413 abrogate bacterial multiplication, they were required not only for full growth but also for
414 production of normal cells.

415 Next, we investigated the pathways that supply pyruvate or PEP for gluconeogenesis. For
416 this purpose, we carried out a genomic search for homologues of the genes encoding the
417 enzymes connecting the TCA cycle and the triose-phosphate pathway in bacteria (Fig. 1). We
418 identified homologues of *pdh* (pyruvate dehydrogenase), *pyk* (pyruvate kinase), *pckA*
419 (phosphoenolpyruvate carboxykinase), *ppdK* (pyruvate-phosphate dikinase), *pyc* (pyruvate
420 carboxylase), and *mae* (malic enzyme) but not of *pyrck* (pyruvate carboxykinase), *pps*
421 (phosphoenolpyruvate synthase) or *ppc* (phosphoenolpyruvate carboxylase) (Fig. 1). Since Pyc,
422 Pdh and Pyk catalyze irreversible catabolic steps, we studied the *pckA*, *ppdK* and *mae*
423 homologues.

424 *B. abortus* 2308 BAB1_2091 is annotated as a pseudogene in some data bases
425 (<http://www.genome.jp/kegg/>) but not in others (<http://biocyc.org>). This ORF encodes a protein
426 of 491 amino acids that bears a 77% similarity with the PckA of the phylogenetically related
427 *Agrobacterium tumefaciens*, a protein of 536 amino acids known to be functional (35). However,
428 *B. abortus* BAB1_2091 is separated by a stop codon from an intergenic region that together with
429 BAB1_2090 encodes the last 45 amino acids present in *A. tumefaciens* PckA. The frame-shift
430 that generates this stop codon is present also in all *B. abortus* strains sequenced so far
431 (<http://www.genome.jp/kegg/>). Nevertheless, *B. abortus* PckA conserves the IGGTSYAGE-KKS
432 domain (amino acids 190 to 202) specifically required for the carboxykinase activity (36) as well
433 as the phosphate binding site (G--G-GKT; amino acids 236 to 243) and ATP and metal binding
434 sites, so that its functionality is not obviously compromised. In contrast, the putative *B. abortus*
435 2308 PpdK (encoded by BAB1_0525) represents a complete protein of 887 amino acids with the
436 PEP-binding (amino acids 19 to 376) and the TIM-barrel (amino acids 530 to 883) domains
437 characteristic of PEP-utilizing enzymes.

438 To test whether these ORFs encoded enzymes are required for growth under gluconeogenic
439 conditions, we constructed the non-polar mutants *BABΔpckA* and *BABΔppdK*. Moreover, since

440 both PckA and PpdK catalyze reactions eventually leading to PEP (Fig. 1), we excluded their
441 reciprocal complementation by constructing the double mutant *BABΔpckAΔppdK*. We then
442 compared the growth of *BAB*-parental and the mutants in Peptone-Yeast Extract-Glucose and in
443 Glutamate-Lactate-Glycerol. Whereas we did not observe differences in the growth of
444 *BABΔpckA* and *BAB*-parental in these two media, both *BABΔppdK* and *BABΔpckAΔppdK* had a
445 markedly reduced growth in Glutamate-Lactate-Glycerol (Fig. 2B). Complementation with
446 plasmid pAZI-19 carrying *ppdK* restored the phenotype (Supplemental Material; Fig. S1), and
447 the sibling revertant control (see Material and Methods) conserved the wild-type phenotype (not
448 shown). These results strongly suggest that PpdK is functional and that, consistent with the
449 frame shift in *pckA*, PckA does not synthesize PEP from oxaloacetate in *B. abortus*. The
450 possibility that the role of *B. abortus* PckA is not detectable under these *in vitro* conditions
451 seems less likely.

452 We followed a similar strategy to study the putative *mae*. BAB1_1036 encodes a protein of
453 774 amino acids annotated as a NADP (or NAD)-dependent enzyme involved in malate
454 metabolism (<http://www.genome.jp/kegg/>; <http://biocyc.org/>). The N-terminal domain (amino
455 acids 28-160) and the NADP (or NAD)-binding domain (amino acids 172-409) characteristic of
456 the malic enzyme are conserved in the BAB1_1036 predicted protein. Mutant *BABΔmae*
457 (deleted in the region encoding amino acids 26 to 673) displayed a small reduction in growth in
458 Peptone-Yeast Extract-Glucose and a more marked one in Glutamate-Lactate-Glycerol (Fig.
459 2C). The impairment was not as accentuated as that of *BABΔppdK* (compare Fig. 2 panels B
460 and C). In these experiments, *mae* mutants with the above-described phenotype were
461 consistently obtained and the control sibling revertants of wild-type phenotype recovered after
462 the last recombination event (see Material and Methods and Supplemental Material; Fig. S2).
463 Taken together, these results support the hypothesis that *B. abortus* Mae supplies pyruvate for
464 PpdK to produce PEP for gluconeogenesis, lactate being a complementary source of pyruvate in

465 Glutamate-Lactate-Glycerol (Fig. 1). We tested this hypothesis further by using media containing
466 only two of the three C substrates of this chemically defined medium. *BABΔmae* did not grow in
467 Glycerol-Glutamate, showed optimal growth in Glycerol-Lactate, and reduced growth in
468 Glutamate-Lactate (Fig. 3, lower panels). These results are fully consistent with the predicted
469 role of Mae (Fig. 1) and suggest that, although *B. abortus* 2308 can use glycerol, glutamate and
470 lactate, provision of the latter cannot completely replace the Mae pathway. This interpretation is
471 also supported by the fact that whereas PpdK dysfunction in *BABΔppdK* severely impaired
472 growth in Glycerol-Lactate and Glutamate-Lactate, this mutant grew in Glycerol-Glutamate (Fig.
473 3, lower panels), two substrates able to act as sources of PEP and pyruvate through the triose-P
474 and Mae pathways, respectively (Fig. 1). Indeed, all these results are in agreement with the
475 early studies that led to the formulation of the simple medium of Gerhardt as well as with the
476 nutritional studies that showed the preferential use of glutamate and the complementary role of
477 glycerol and lactate in *B. abortus* (9). Also, the demonstration by Marr et al. (37) of the ability of
478 *B. abortus* to generate pyruvate (and alanine) from glutamate supports an activity of Mae in this
479 bacterium (Fig. 1).

480 Since the putative Mae was active in *B. abortus* 2308, we investigated whether malate
481 replenishment could occur through the classical glyoxylate pathway (Fig. 1) or the PEP-
482 glyoxylate cycle. The latter cycle combines the operation of *PckA* with the glyoxylate cycle
483 enzymes and operates in *E. coli* under conditions of glucose limitation (38). In these pathways,
484 isocitrate lyase (*AceA*) cleaves isocitrate to yield glyoxylate and succinate, and a malate
485 synthase (*AceB*) condenses glyoxylate and acetyl-CoA to produce malate (39-41). The genome
486 of *B. abortus* 2308 carries only one putative *aceA* (ORF BAB1_1631) and one putative *aceB*
487 (ORF BAB1_1663). The predicted *AceA* is a protein of 429 amino acids with 61% identity and
488 76% similarity to *E. coli* *AceA*, and it conserves the amino acids required for the enzymatic
489 activity and the assembly of the tetrameric enzyme (42-47). The predicted *AceB* has 728 amino

490 acids with 59% identity and 74% similarity to *E. coli* malate synthase G, and conserves the
491 catalytic site and the amino acids interacting with acetyl-CoA (48,49). Accordingly, we
492 constructed a non-polar *BABΔaceA* mutant truncated in the 409 central amino acids. This
493 mutant did not show growth differences with *BAB*-parental in Peptone-Yeast Extract-Glucose
494 and Glutamate-Lactate-Glycerol (data not shown), even though these media contain acetogenic
495 substrates (glucose, glycerol, lactate and serine, threonine and alanine) (50). Since the genomic
496 analysis strongly suggests the presence of the glyoxylate cycle, we examined this point further
497 by constructing a luciferase reporter under the control of the *AceA* promoter. Although growth
498 curves were similar, luciferase activity was considerably higher in Peptone-Yeast Extract-
499 Glucose than in Glutamate-Lactate-Glycerol (Fig. 5), as expected if the glyoxylate cycle
500 becomes active on dependence of the abundance of acetogenic substrates. In such a case, the
501 lack of phenotype in complex media could be explained if the glyoxylate cycle plays only a
502 subsidiary role in this rich medium, and the experiments do not rule out that the possibility that it
503 becomes important under other nutritional conditions.

504 ***B. abortus* mutants in *ppdK* and *mae* but not in *fbp*, *glpX*, *pckA* or *aceA* show lower**
505 **multiplication rates in macrophages.** *B. abortus* is characteristically able to multiply
506 intracellularly in professional phagocytes (51). We thus investigated the ability of the above-
507 described mutants to multiply in macrophages using the parental strain and the attenuated *virB*
508 mutant (unable to reach the replicating vacuole) as controls. Figure 6A shows that
509 *BABΔfbpΔglpX* replicated like *BAB*-parental, even though these bacteria differed in growth
510 under gluconeogenic conditions *in vitro* (see above). *BABΔppdK* (Fig. 6B) and *BABΔmae* (Fig.
511 6C) multiplied in macrophages, although at lower overall rates than *BAB*-parental both 24
512 (p<0,0001) and 48h (p<0,0001) after infection. On the other hand, mutation of *pckA* had no
513 effect either by itself or combined with the *ppdK* deletion (Fig. 6B).

514 These results suggest that, albeit impaired in growth, *BAB* Δ *ppdK* and *BAB* Δ *mae* are still
515 able to reach the replicative intracellular niche. Since Mae and PpdK belong to the same
516 pathway, we selected *BAB* Δ *ppdK* (the mutant blocked in the upper step of the pathway; Fig. 1)
517 to confirm that the metabolic dysfunction did not prevent these bacteria from reaching the
518 endoplasmic reticulum-derived replicating niche. Figure 7A shows that, in contrast to the *virB*
519 mutant, *BAB* Δ *ppdK* and the parental bacteria were similar in intracellular distribution.

520 *B. abortus* can also penetrate and multiply in epithelial cells (51). We found that the behavior
521 of *BAB* Δ *ppdK* in macrophages was reproduced in HeLa cells (Fig 7A and B). In addition, we
522 found that the CFU of the mutant and *BAB*-parental in HeLa cells did not differ 2 h after infection
523 (not shown), indicating that they were similar with regards to penetration.

524 ***B. abortus* mutants in *ppdK* and *mae* but not in *pckA*, *fbp*, *glpX* or *aceA* are attenuated in**
525 **mice.** Virulent *B. abortus* is able to establish spleen infections in mice that characteristically
526 develop in four phases: (i), onset phase (spleen colonization; first 48 h); (ii), acute phase (from
527 the third day to weeks 2-4 when bacteria reach maximal numbers); (iii), chronic steady phase
528 (weeks 2-4 to 12), where the bacterial numbers plateau; and (iv), chronic declining phase, during
529 which brucellae are eliminated. The ability to induce a marked splenomegaly is also a
530 characteristic of virulent brucellae. (25). Using this model, we first studied the *BAB* Δ *fbp* Δ *glpX*
531 mutant and found that it did not differ from *BAB*-parental in either the CFU/spleen profile or the
532 splenomegaly induced (Fig. 8A). Similarly, we did not observe attenuation for *BAB* Δ *pckA* (Fig.
533 8B), which is remarkable because *pckA* expression increases in *B. abortus* mutants in
534 BvrR/BvrS, a master regulator of *B. abortus* virulence (52). Nevertheless, this result is consistent
535 with the genomic features of *pckA* and with the above-described experiments *in vitro* and in
536 macrophages.

537 *BAB* Δ *ppdK* failed to reach the chronic steady phase typical of virulent brucellae yielding
538 significantly lower CFU counts after week 2 (Fig. 8B). Clearly indicative of the strong attenuation

539 of *BABΔppdK*, we did not recover any bacteria from the spleens of 3 out of the 5 mice examined
540 at post-infection week 12, and this mutant induced less splenomegaly than the virulent bacteria
541 (Fig. 8B). Consistent with the observations that showed no additive effect of the mutations of
542 *ppdK* and *pckA* in macrophages or *in vitro*, the results of *BABΔpckAΔppdK* in mice paralleled
543 those of *BABΔppdK* (Fig. 8B). *BABΔmae* produced a CFU/spleen profile that differed from that of
544 either *BAB*-parental or *BABΔppdK*. Although not affected in the first 48 h (onset phase; not
545 shown) this mutant showed a lower multiplication rate during the acute phase (Fig. 8C) that was
546 reminiscent of the lower multiplication rates observed in macrophages. Strikingly, even though
547 CFU/spleen numbers were lower than those of the wild type strains, *BABΔmae* produced a
548 chronic steady phase with reduced splenomegaly.

549 Finally, we tested *BABΔaceA* in mice. In a first experiment, the mutant did not show
550 attenuation at weeks 2, 8 and 12 (not shown). It has been reported that isocitrate lyase is
551 essential for *Salmonella* persistence in mice during chronic infection but dispensable for acute
552 lethal infection (53). Similarly, isocitrate lyase is dispensable in the acute phase of
553 *Mycobacterium tuberculosis* infection in lung macrophages of mice but facilitates persistence
554 during the chronic phase (54). Accordingly, we repeated the mouse infections and determined
555 the number of *BABΔaceA* CFU in spleens 16 and 24 weeks later. However, we did not find any
556 difference between this mutant and the parental strain (not shown). Therefore, even though we
557 did not rule out the possibility that the glyoxylate cycle plays a role under conditions different
558 from those tested *in vitro*, we concluded that this shunt is not essential for *B. abortus*
559 multiplication and persistence in the laboratory models used. Since the PEP-glyoxylate pathway
560 relies on both *AceA* and *PckA*, this conclusion can be extended to this cycle.

561 DISCUSSION

562 *B. abortus* lacks the genes necessary for the metabolism of glycogen or poly-beta-
563 hydroxyalkanoates, the two C reserve materials used by prokaryotes (55). Thus, these bacteria

564 depend on nutrients provided by the host to multiply intracellularly and, accordingly, they need a
565 supply of (at least) 6 C skeletons or to carry out gluconeogenesis. To investigate these
566 possibilities, we deleted ORFs that could be encoding enzymes of critical steps of
567 gluconeogenesis, or of steps providing the necessary precursors. The genomic characteristics of
568 the ORFs analyzed and the phenotypes observed *in vitro* support the hypothesis that they
569 encode FBPsases, a pyruvate phosphate dikinase and a malic enzyme of *B. abortus*. Moreover,
570 the analyses in cells show that the proteins coded for by *ppdK* and *mae* become necessary once
571 the replicative niche is reached, as expected from metabolic mutants.

572 Since FBPsases are essential for gluconeogenesis, the observation that growth in
573 gluconeogenic media was not abolished in the Fbp-GlpX double mutant is intriguing and several
574 hypotheses can be considered to explain these results. The existence of a third FBPsase is the
575 first and more obvious possibility. Up to now, five different types of FBPsases (I to V) have been
576 described in prokaryotes (56). Whereas FBPsases of classes IV and V are restricted to *Archaea*
577 and their close hyperthermophilic *Aquifex* bacterial group, many bacteria have dual
578 combinations of class I (Fbp homologues), class II (GlpX homologues), and class III FBPsases
579 (34,56). However, a genomic search for *Bacillus subtilis* YydE homologues (the prototype of
580 class III FBPsases (57)) in the *Brucellaceae* only revealed an imperfect match (a hypothetical
581 protein of 218 amino acids with a 32% identity with the 671 amino acids in YydE) in
582 *Ochrobactrum anthropi* and none in *Brucella*. This is in agreement with the fact that no bacterial
583 genome has been described to carry a combination of classes I and III (34). Also, it has been
584 described that some *E. coli* carry two class II FBPsases (GlpX and YggF (56)) but genomic
585 analysis of all phosphatases in *B. abortus* fail to identify clear candidates for any phosphatase
586 close to GlpX and Fbp (Supplemental Material; [Fig. S4](#)). This genomic evidence and the Fbp-
587 GlpX reciprocal complementation (i.e. the fact that only the double mutant shows phenotype *in*
588 *vitro* in the absence of glycerol) suggest that these are the main and possibly the only FBPsases

589 in *B. abortus*. An alternative to the third FBPase hypothesis is the existence of an atypical
590 gluconeogenesis less efficient than the classical one. Hypothetically, a fructose-6-P aldolase
591 could take part in gluconeogenesis. This has been described in *E. coli* K-12 where ORF b0825
592 encodes an enzyme that catalyzes the reversible conversion of fructose-6-P to
593 dihydroxyacetone and glyceraldehyde-3-P (58). This aldolase is different from that operating in
594 the pentose-phosphate cycle and its physiological role is uncertain. However, the only
595 homologous in *B. abortus* (ORF BAB1_1813) is annotated as transaldolase and, in addition to
596 the fact that it should represent the enzyme of the pentose-phosphate cycle, the identity (30%) is
597 below the threshold considered to be significant (59). Obviously, a rigorous analysis of these two
598 hypotheses requires enzymatic analysis of double deficient cells. Finally, the significance of the
599 mucoid aggregates produced by the double FBPase mutant cannot be disregarded since this
600 phenotype suggests that the growth observed does not correspond to a natural condition.

601 Bearing in mind that enzymatic analyses are necessary to reach definite conclusions, our
602 results and those of previous works with *B. abortus* 2308 offer insight into some global models of
603 the metabolism of these bacteria during intracellular life in the host.

604 A first model (gluconeogenic model) can be proposed on the ability of *B. abortus* to grow in
605 the defined medium of Gerhardt and on the proteomic studies in macrophages that suggest that
606 *B. abortus* shifts to an amino acid-based metabolism in which the glutamate pool is increased
607 (23). According to this model, molecules like glycerol, lactate or pyruvate and amino acids
608 channeled to oxaloacetate, keto-glutarate or pyruvate are the main substrates, and molecules of
609 6 and 5 C are derived from the latter. In this regard, it is remarkable that dysfunction of two
610 major FBPases did not bring about any perceptible attenuation either in cells or mice. Although
611 the reduced growth of the *BABΔfbpΔglpx* mutant *in vitro* precludes a clear-cut conclusion, the
612 contrasting *in vivo* and *in vitro* multiplication and the mucoid phenotype of the double mutant in
613 the minimal medium are more consistent with models alternative to the gluconeogenic one.

614 Moreover, two lines of evidence indicate that glucose (or closely related hexoses) is available in
615 the host. First, a *B. abortus* 2308 (and *B. suis* 1330) GluP (glucose/galactose transporter)
616 mutant has been identified as attenuated in signature tagged experiments in (15,20,20). Second,
617 it has been reported recently that the multiplication of *B. abortus* in alternatively activated
618 macrophages increases when the intracellular glucose levels are artificially increased (60).
619 Based on these observations, an almost opposite model proposes a main role for 6 C sugars in
620 the replicative niche (and 5 C sugars if we assume that the evidence obtained in *B. melitensis*
621 also applies to *B. abortus*; see Introduction). These sugars would provide trioses-phosphate
622 through the pentose phosphate cycle and serve also as precursors for biosynthesis of envelope
623 polymers. This second model, however, does not account for the attenuation observed for the
624 *ppdK* and *mae* mutants, which strongly suggests that molecules necessary for growth are
625 derived from the TCA *in vivo*.

626 A third model proposes that there is a limited supply of 6 C (and 5 C) sugars that is
627 compensated by glutamate, alanine and other amino acids. Those sugars would be used mostly
628 or exclusively for biosynthesis of envelope polymers and for the pentose-phosphate cycle-
629 dependent biosynthetic reactions. This model is consistent with the results of this and previous
630 works. First, the different phenotype of the *B. abortus* *BABΔfbpΔglpx* double mutant *in vivo* and
631 *in vitro* is more coherent with the hypothesis that classical gluconeogenesis is not extensively
632 used *in vivo*. In addition, the *B. abortus* 2308 *gluP* mutant identified in signature-tagged
633 mutagenesis studies is not clearly attenuated at two weeks post-infection and manifests its
634 attenuation at times (8 weeks) that correspond to the chronic phase (20). This suggests that the
635 bacteria do not depend totally on hexoses for intracellular biosynthetic processes, and that this
636 dependence is manifested at late times perhaps as the result of changes in the replicative
637 vacuole (see below). Indeed, the infection experiments performed in alternatively activated
638 macrophages suggest that, although available, glucose is a limiting factor for *B. abortus* growth

639 at least during the chronic phase (60). Proteomic studies carried out with *B. abortus* 2308 show
640 that expression of two proteins of the dihydroxyacetone kinase complex (Dha) of the PEP-
641 carbohydrate phosphotransferase system (PTS) is reduced throughout infection in macrophages
642 (23). Although this has been interpreted to mean that reduced PTS expression may be the result
643 of a short supply of sugars within the replicative niche, the *Brucella* PTS lacks the sugar
644 permease unit and is likely to act as a regulatory system coordinating C and N metabolism (61).
645 On the other hand, the signature-tagged mutagenesis and proteomic studies show attenuation
646 of a *gltD* (putatively encoding the small subunit of glutamate synthase) mutant (20) and an
647 increment of enzymes involved in increasing the pool of glutamate (23) that are consistent with
648 the model. Finally, this third model accounts for the attenuation of the *ppdK* and *mae* mutants
649 and could also explain in part the differences between the mutants in these two genes that were
650 observed in mice (Fig. 8). Since according to the model sugars are used mostly to construct
651 envelope polymers and for pentose-phosphate cycle derived precursors, additional molecules
652 for biosynthesis must be derived from TCA. PpdK works to produce PEP, which is used to
653 synthesize phenylalanine, tyrosine and tryptophan, glycerolipids and other PEP-derived
654 molecules. Mae supplies pyruvate for PpdK *in vivo* but TCA would not be the only source of
655 pyruvate. This is suggested by the fact that the *mae* mutant was both delayed in reaching the
656 chronic phase of infection and in lower numbers in the spleen during this phase, which contrasts
657 with the inability of the *ppdK* mutant to generate chronic infections. Interestingly, it has been
658 shown recently that, when provided with multiple carbon sources, *Mycobacterium tuberculosis*
659 differentially catabolizes each carbon source through the glycolytic, pentose phosphate, and/or
660 TCA pathways to distinct metabolic fates, and it has been suggested that this ability reflects an
661 adaptation to pathogenicity (62). Indeed, such ability could also be necessary for *B. abortus* to
662 coordinately use the different substrates proposed for this model.

663 Although the last model fits the experimental evidence, it is obvious that it represents only a
664 first approach to the situation in the natural hosts because we cannot assume that the niche is
665 static during a chronic infection or uniform among different types of cells. In the above-
666 mentioned signature-tagged mutagenesis studies, Hong et al. (20) have presented evidence for
667 the hypothesis that different set of genes are required during the onset-acute phases and the
668 chronic steady phase. This study identified three putative metabolic genes (*gluP*, *glD* [see
669 above] and *gcvP*) required during the chronic phase but not markedly during the acute phase of
670 infection. Indeed, both the need of a functional *ppdK* for the infection to progress during the
671 acute phase and the different phenotype of *mae* add further support to the hypothesis that
672 different genes are required to a different extent during the course of infection. Moreover, at
673 least macrophages and trophoblastic cells have been clearly associated with *B. abortus* infection
674 in cattle (63) and the physiological characteristics of these cells are different. Also, different
675 spleen cells become colonized at different times after intraperitoneal inoculation of mice (64).
676 Clearly, research is necessary to investigate these aspects of the relationship between
677 metabolism and intracellular multiplication in *B. abortus* and in other species of the genus.

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875 **FIGURE LEGENDS**

876 **FIG. 1.** Conventional central metabolic pathways (glycolysis, gluconeogenesis, Entner-
877 Doudoroff, pentose-phosphate, TCA and glyoxylate) of bacteria. Dashed arrows indicate steps
878 for which no putative genes can be identified in *B. abortus*. Red arrows indicate the steps
879 studied in this work. Abbreviations used in the figure are: AcCoA (acetyl-CoA), AceA (isocitrate
880 lyase), AceB (malate synthase), Acn (aconitate hydratase), AKG (α -ketoglutarate), Akgdh (α -
881 ketoglutarate dehydrogenase), Cgs (cyclic-glucan synthesis), CIT (citrate), DHAP
882 (dihydroxyacetone phosphate), Eda (keto-deoxy-phosphogluconate aldolase), Edd (6-phospho-
883 D-gluconate dehydratase), E4P (erythrose-4-phosphate), Fba (fructose bisphosphate aldolase),
884 Fbp (fructose-1,6-bisphosphatase), Fum (fumarase), FUM (fumarate), F1,6dP (fructose-1,6-
885 bisphosphate), F6P (fructose-6-phosphate), GalE (UPD-glucose-4 epimerase), GAP
886 (glyceraldehyde-3-phosphate), Gnd (6-phosphogluconate dehydrogenase), Glk (glucokinase),
887 GlpX (fructose-1,6-bisphosphatase), GltA (citrate synthase), GLX (glyoxylate), G1P (glucose-1-
888 phosphate), G6P (glucose-6-phosphate), ICIT (isocitrate), Idh (isocitrate dehydrogenase), KDPG
889 (2-keto-3-deoxyphosphogluconate), LPS (lipopolysaccharide), Mae (malic enzyme), MAL
890 (malate), Mdh (malate dehydrogenase), OAA (oxaloacetate), Pfk (phosphofructokinase), PckA
891 (phosphoenol pyruvate carboxikinase), Pdh (pyruvate dehydrogenase), PEP (phosphoenol
892 pyruvate), Pgi (phosphoglucose isomerase), Pgl (lactonase), Pgm (phosphoglucomutase), Ppc
893 (phosphoenol pyruvate carboxylase), PpdK (pyruvate phosphate dikinase), Pps (phosphoenol
894 pyruvate synthase), Pyc (pyruvate carboxylase), Pyk (pyruvate kinase), PYR (pyruvate), Pyrck
895 (pyruvate carboxikinase), RIB5P (ribulose-5-phosphate), Sdh (succinate dehydrogenase), Stk
896 (succinyl-CoA synthetase), SUC (succinate), SucCoA (succinyl-CoA), Tkt (transketolase), TpiA
897 (triose phosphate isomerase), X5P (xylulose-5-phosphate), Zwf (glucose-6-phosphate
898 dehydrogenase), 6PG (6-phosphogluconate), 6PGL (6-phosphogluconolactone).

899

900 **FIG. 2.** Growth curves in Peptone-Yeast Extract-Glucose (A.1, B.1 and C.1) and Glutamate-
901 Lactate-Glycerol (A.2, B.2 and C.2) of *BAB*-parental and mutants *BABΔfbp*, *BABΔglpX*,
902 *BABΔfbpΔglpX*, *BABΔpckA*, *BABΔppdK*, *BABΔpckAΔppdK* and *BABΔmae*. Each point
903 represents the mean ± standard error (error bars are within the size of the symbols) of optical
904 density (O.D.) values of triplicate samples. The experiment was repeated three times with similar
905 results.

906

907 **FIG. 3.** Growth curves in Glycerol-Glutamate, Glycerol-Lactate and Glutamate-Lactate of *BAB*-
908 parental and mutants *BABΔfbp*, *BABΔglpX*, *BABΔfbpΔglpX*, *BABΔppdK* and *BABΔmae*. Each
909 point represents the mean of triplicate samples (error bars are within the size of the symbols).
910 The experiment was repeated three times with similar results.

911

912 **FIG. 4.** Immunofluorescence analysis of *BAB*-parental and *BABΔfbpΔglpX* grown in Glycerol-
913 Lactate-Glutamate. Cells were labeled with anti-smooth-lipopolysaccharide mouse monoclonal
914 antibody A76/12G12/F12 and Alexa Fluor 546 conjugated goat anti-mouse immunoglobulin.

915

916 **FIG. 5.** Luciferase expression under the control of *B. abortus aceA* promoter in Peptone-Yeast
917 Extract-Glucose and Glutamate-Lactate-Glycerol. The results are representative of three
918 experiments (RLU, relative luminescence units).

919

920 **FIG. 6.** Intracellular multiplication in RAW 264.7 macrophages of *BAB*-parental and mutants
921 *BABΔfbpΔglpX*, *BABΔpckA*, *BABΔppdK*, *BABΔpckAΔppdK* and *BABΔmae* (*virB* is an attenuated
922 mutant used as a control). Values are the mean ± standard error of triplicate infections and the
923 results shown are representative of three independent experiments.

924

925 **FIG. 7.** Trafficking of *BAB*-parental, *BAB* Δ *ppdK* and *virB* in macrophages and HeLa cells. (A),
926 confocal images of infected RAW 264.7 macrophages and HeLa cells labeled with Moabs to
927 either calnexin (in green) or LAMP-1 (in blue) 24 hours after infection (bacteria are
928 immunostained in red). (B), Percentage of calnexin-positive vacuoles of HeLa cells that contain
929 bacteria at 24 and 48 hours post-infection.

930

931 **FIG. 8.** Bacterial multiplication (upper panels) and spleen weights (lower panels) generated in
932 BALB/c mice by *BAB*-parental and mutants *BAB* Δ *fbp* Δ *glpX* (A); *BAB* Δ *pckA*, *BAB* Δ *ppdK* and
933 *BAB* Δ *pckA* Δ *ppdK* (B); and *BAB* Δ *mae* (C). Each point is the mean \pm standard deviation (n=5) of
934 the Log of CFU or grams per spleen. Statistical differences with *BAB*-parental were significant (p
935 < 0.001) from week 4 onwards for *BAB* Δ *ppdK* and from week 2 onwards for *BAB* Δ *mae*.
936 *BAB* Δ *pckA* Δ *ppdK* was also attenuated at weeks 8 and 12 (p < 0.001) (weeks 4 and 6 were not
937 tested).















