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1 Serovar-dependent differences in Hfq-regulated phenotypes in *Actinobacillus*
2 *pleuropneumoniae*

3

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15

16 **Keywords:** *Pasteurellaceae*; RNA chaperone; *Galleria mellonella*; Virulence; Stress.

17 **One-sentence summary:** Serovar-dependent differences identified in regulation of
18 complex phenotypes by the RNA chaperone Hfq in the pig pathogen *Actinobacillus*
19 *pleuroneumoniae* indicate the importance of strain selection and interpretation of results
20 when analysing global gene regulator function.

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24

25 **Abstract**

26 The RNA chaperone Hfq regulates diverse processes in numerous bacteria. In this
27 study, we compared phenotypes (growth rate, adherence, response to different stress
28 conditions, and virulence in *Galleria mellonella*) of wild-type (WT) and isogenic *hfq*
29 mutants of three serovars (1, 8 and 15) of the porcine pathogen *A. pleuropneumoniae*.
30 Similar growth in rich broth was seen for all strains except Ap1 Δ *hfq*, which showed
31 slightly reduced growth throughout the 24 hour time course, and the complemented
32 Ap8 Δ *hfqC* mutant had a prolonged lag phase. Differences were seen between the three
33 serovar WT strains regarding adherence, stress response and virulence in *G. mellonella*,
34 and deletion of *hfq* affected some, but not all of these phenotypes, depending on
35 serovar. Complementation by expression of cloned *hfq* from an endogenous promoter
36 only restored some WT phenotypes, indicating that complex regulatory networks may
37 be involved, and that levels of Hfq may be as important as presence/absence of the
38 protein regarding its contribution to gene regulation. Our results support that Hfq is a
39 pleiotropic global regulator in *A. pleuropneumoniae*, but serovar-related differences
40 exist. These results highlight the importance of testing multiple strains/serovars within a
41 given species when determining contributions of global regulators, such as Hfq, to
42 expression of complex phenotypes.

43

44 **Introduction**

45

46 The Gram-negative bacterium *Actinobacillus pleuropneumoniae* causes porcine
47 pleuropneumonia, a disease that has a negative economic impact on the worldwide
48 swine industry (Sassu *et al.* 2018). Currently, eighteen serovars are recognized based on

49 capsular polysaccharides (Bossé *et al.* 2018). All serovars are pathogenic, but some are
50 more virulent than others, e.g. serovar 3 is rarely pathogenic, but serovar 1 is considered
51 of high virulence (Rogers *et al.* 1990; Frey 2011). In part, this is related to the
52 combinations of RTXs toxins (ApxI-III) present in different serovars (Frey 2011). Other
53 virulence factors, some of which also differ depending on serovar, have been reported
54 for *A. pleuropneumoniae* including: capsule, lipopolysaccharide (LPS), fimbriae, outer
55 membrane proteins, iron-binding proteins, and the ability to form biofilms [reviewed in
56 (Bossé *et al.* 2002; Chiers *et al.* 2010)]. In addition, roles in virulence have been
57 indicated for global regulators of gene expression such as RpoE (Bossé *et al.* 2010),
58 HlyX (Buettner *et al.* 2009), ArcA (Buettner *et al.* 2008), and Hfq (Zhou *et al.* 2008;
59 Subashchandrabose *et al.* 2013), the latter being the subject of this study.

60 Hfq was first identified in 1972 as regulator of phage Q β RNA replication in
61 *Escherichia coli* (Franze de Fernandez *et al.* 1972). It is now known that, through its
62 interactions with small RNAs (sRNAs), Hfq is a major global regulator of gene
63 expression in a wide variety of bacteria (Vogel and Luisi 2011; Sobrero and Valverde
64 2012; Feliciano *et al.* 2016; Dos Santos, Arraiano & Andrade 2019). In *E. coli*, deletion
65 of the *hfq* gene results in pleiotropic changes when compared to WT, including
66 increased cell size, reduced growth rate, increased sensitivity to ultraviolet light and
67 other processes (Tsui *et al.* 1994; Kendall *et al.* 2011). A role for Hfq in virulence, as
68 adjudged *in vivo* or by surrogate markers such as tolerance to stress and ability to form
69 biofilms, has been shown for many Gram-negative bacteria including: *Neisseria*
70 *meningitidis* (Fantappiè *et al.* 2009), *Haemophilus influenzae* (Hempel *et al.* 2013),
71 *Yersinia enterocolitica* (Kakoschke *et al.* 2014), *Brucella melitensis* (Cui *et al.* 2013),
72 *Salmonella enterica* serovar Typhimurium (Behere *et al.* 2016), *Pasteurella multocida*

73 (Mégroz *et al.* 2016), *Xanthomonas campestris* (Lai *et al.* 2018), and *Bordetella*
74 *pertussis* (Hayes *et al.* 2020).

75 With *A. pleuropneumoniae*, it has also been established that Hfq has a role in
76 virulence. Both Zhou *et al.* (2008) and Subashchandrabose *et al.* (2013) demonstrated
77 that *A. pleuropneumoniae* *hfq* mutants of serovar 1 strains Shope 4074 and AP 93-9,
78 respectively, were less virulent in pigs. In addition, an *hfq* mutant of a clinical serovar 8
79 isolate, MIDG2331, was attenuated in the *Galleria mellonella* (wax moth) model of
80 infection (Pereira *et al.* 2015). *In vitro*, the AP 93-9 serovar 1 *hfq* mutant was defective
81 in biofilm formation and was more sensitive to superoxide stress (Subashchandrabose *et*
82 *al.* 2013). In this study, we undertook a comparative analysis of the effect of *hfq*
83 mutagenesis on three different serovars of *A. pleuropneumoniae* to determine if
84 regulation of different Hfq phenotypes is serovar dependent.

85

86 **Materials and Methods**

87

88 **Bacterial strains, growth conditions and maintenance**

89 The *A. pleuropneumoniae* strains used in this study (listed in Table 1) were
90 routinely grown at 37°C with 5% CO₂ in brain heart infusion (BHI; Difco) broth and
91 agar supplement with 10 µg/mL nicotinamide adenine dinucleotide (NAD; Sigma-
92 Aldrich), and the *E. coli* strains in LB broth and agar. Chloramphenicol (1 or 20 µg/mL,
93 for *A. pleuropneumoniae* and *E. coli*, respectively) or kanamycin (75 µg/mL) was added
94 to the medium when required. Salt-free LB agar (10 g tryptone, 5 g yeast extract and 15
95 g agar per L) supplemented with 10% filter-sterilized sucrose, 10% horse serum (TCS
96 Biosciences), and 10 µg/mL NAD (Sigma-Aldrich) was used for counter selection of *A.*

97 *pleuropneumoniae* mutants, as previously described (Bossé *et al.* 2014). *E. coli* MFDpir
98 (Ferrières *et al.* 2010) and Stellar (Clontech) strains were used in conjugation and
99 transformation assays, respectively.

100

101 **Strain construction**

102 Unless otherwise stated, all PCRs were performed using CloneAmp™ HiFi
103 PCR Premix (Takara), and genomic DNA from the serovar 8 strain MIDG2331 (Bossé
104 *et al.* 2016) was used as the template for amplification of *A. pleuropneumoniae* products
105 for cloning. For direct cloning into the T-vector, pGEM-T (Promega), products
106 amplified with the CloneAmp™ HiFi polymerase were first A-tailed by incubation at
107 70°C for 30 minutes with 0.2 mM dATP and 5 U of Taq polymerase (Promega),
108 according to manufacturer's instructions. All initial constructs were transformed into *E.*
109 *coli* Stellar cells (Takara), according to manufacturer's protocol, with selection of
110 clones on media containing chloramphenicol or kanamycin, as appropriate. A
111 description of all primers used in this study is given in Table 2.

112 The Δhfq and *hfq::3XFLAG* strains of *A. pleuropneumoniae* serovars 8 and 15,
113 and the $\text{Ap1}\Delta hfq\text{catsacB}$ and $\text{Ap1}hfq::3XFLAG\text{cat}$ strains of serovar 1, were obtained
114 using the previously described natural transformation technique (Bossé *et al.* 2014).
115 Briefly, the sequence comprising the *hfq* gene, and ~600 bp to either side, was amplified
116 using primers 1 and 2 (Table 2), A-tailed and cloned in pGEM-T (Promega), resulting
117 in p*Thfq*Flank. A selection/countersselection cassette, *catsacB*, was amplified from
118 pUSS*catsac* (Bossé *et al.* 2014) using primers 3 and 4. p*Thfq*Flank was opened by
119 inverse PCR using primers 5 and 6 designed with 15 bp overhangs to allow In-Fusion
120 (Takara) cloning, according to manufacturer's instructions, of the *catsacB* cassette in
121 place of the deleted *hfq* gene to generate plasmid p $\Delta hfq\text{catsacB}$. This plasmid was

122 transformed into *A. pleuropneumoniae* serovars 1, 8, and 15 to obtain $\Delta hfqcatsacB$
123 mutants, as previously described (Bossé *et al.* 2014).

124 An unmarked deletion construct was made by amplifying the flanking regions to
125 either side of *hfq*, using primers 1 and 7 for the left flank, and 2 and 8 for the right flank.
126 Primers 7 and 8 contain 15 bp overhangs to allow direct fusion of the two amplicons by
127 over-lap extension (OE) PCR (Bossé *et al.* 2014). The OE PCR product was cloned into
128 pGEM-T (Promega), resulting in pT Δhfq . A construct containing *hfq* with a 3' fusion to
129 a 3XFLAG tag (3x GAT TAC AAG GAT GAC GAT GAC AGG) was also generated.
130 The 3XFLAG tag was amplified from pDOC-F (accession number GQ889496), a
131 generous gift from S. Wigneshweraraj, using primers 9 and 10. The pT*hfq*F flank
132 construct was opened by inverse PCR using primers 11 and 12, and the 3XFLAG
133 amplicon was inserted by In-Fusion cloning, creating pT*hfq*::3XFLAG.

134 To obtain the unmarked Δhfq and *hfq*::3XFLAG mutant strains, the $\Delta hfqcatsacB$
135 mutants were subjected to a second natural transformation with linearized plasmids,
136 either pT Δhfq or pT*hfq*::3XFLAG, with counterselection on LB-SSN plates (Bossé *et*
137 *al.* 2014). As counterselection with the unmarked deletion constructs was not successful
138 with the Ap1 $\Delta hfqcatsacB$ mutant, an alternate construct, pT*hfq*::3XFLAG*cat*, was used
139 to obtain the FLAG-tagged mutant. Primers 13 and 14 were used for amplification of
140 the *cat* cassette of plasmid pUSS*catsac*. The 3xFLAG tag was amplified from pDOC-F
141 using primers 9 and 15. Primers 14 and 15 contain 15 bp overhangs to allow direct
142 fusion of the two amplicons by OE PCR, as above. The pT*hfq*F flank construct was
143 opened by inverse PCR using primers 11 and 12, and the 3xFLAG*cat* amplicon was
144 inserted by In-Fusion cloning, creating pT*hfq*::3XFLAG*cat*. This plasmid was
145 transformed into *A. pleuropneumoniae* serovar 1 to obtain the Ap1 Δhfq ::3XFLAG*cat*
146 mutant, as previously described (Bossé *et al.* 2014). Deletion of *hfq*, or the presence of

147 FLAG-tagged *hfq*, in the chromosome of respective mutants was confirmed by PCR and
148 sequencing using primers 1 and 2. RT-PCR analysis using cDNA from both WT and
149 Δhfq mutant strains was performed with primer pairs 16 and 17, 18 and 19, as well as
150 20 and 21 (for detection of expression of *hflX*, *miaA*, and *hfq*, respectively) in order to
151 confirm that deletion of *hfq* did not affect expression of the flanking genes. As a
152 positive control for each primer pair, gDNA from the WT strain was used. The presence
153 of expressed FLAG-tagged Hfq was confirmed by Western blot using anti-FLAG
154 antibodies (see below).

155

156 **Hfq promoter analysis and mutation complementation**

157 For a better understanding of the promoter(s) involved in the transcription of the
158 *hfq* gene, a prediction of the *hfq* operon was performed using DOOR (Database for
159 prokaryotic Operons) (Mao *et al.* 2009), followed by prediction of promoters using
160 BPROM (Solovyev *et al.* 2010) and visual analysis of the sequences.

161 Complementation of the Δhfq mutants was achieved by cloning the *hfq* gene,
162 with three of the predicted endogenous promoters, into the low copy plasmid
163 pMIDG100 (O'Dwyer *et al.* 2004; Bossé *et al.* 2009). Primers 22 and 23 were used to
164 amplify the sequence from 850 bp upstream, to 72 bp downstream, of *hfq*. The vector
165 pMIDG100 was digested with *EcoRI* and *BstBI* (New England Biolabs) and the 1.2 kb
166 PCR product was inserted using In-Fusion cloning, as above. The plasmid pMIDG_*hfq*
167 was transformed into *E. coli* MFDpir (Ferrières *et al.* 2010) with selection on LB agar
168 containing kanamycin (75 $\mu\text{g}/\text{mL}$), prior to conjugation into the *A. pleuropneumoniae*
169 Δhfq strains to obtain the complemented ($\Delta hfqC$) strains. Confirmation of the presence
170 of the gene was performed by PCR and sequencing of the intact gene using the same
171 primers described above.

172

173 **Growth rate and Hfq expression**

174 For growth curves, the *A. pleuropneumoniae* WT and mutant strains were
175 cultivated in 20 mL of broth in Erlenmeyer flasks incubated at 37°C for 24 hours with
176 agitation (180 rpm). Optical density at 600 nm (OD₆₀₀) was measured every hour for the
177 first 12 hours, and then at 24 hours, using an Ultrospec 10 (GE Healthcare Life
178 Sciences).

179 In order to verify expression of Hfq during growth in broth culture, each of the
180 three serovar *hfq::3XFLAG* strains were inoculated into 200 mL of broth (initial OD₆₀₀
181 0.01) and then aliquoted into seven flasks of 20 mL each. At time points (1, 2, 3, 4, 6, 8
182 and 12 hours), one of each serovar culture was centrifuged at 9000x g, and the resulting
183 pellets were re-suspended in 1 mL of lysis buffer (20 mM Tris-HCl, 1 mM EDTA, pH
184 7.4) and disrupted by mechanical lysis using Matrix B tubes (MP Biomedicals). For
185 each sample, 10 µg of soluble protein were applied to wells of 4-12% NuPAGE Bolt
186 BisTris Plus (Life Technologies – BG04120BOX) gels. Following electrophoretic
187 separation, the proteins were transferred to nitrocellulose membrane (iBlot 2 NC
188 Regular Stacks; Life Technologies - IB23001) using the iBlot 2 system (Life
189 Technologies - IB21001). The membrane was processed, as previously described
190 (Beddek *et al.* 2004), using an anti-FLAG monoclonal (Sigma) as the primary antibody,
191 and detection using ECL Western blotting detection reagents (GE Healthcare) and
192 Hyperfilm ECL (GE Healthcare).

193

194 **Bacterial adhesion**

195 Bacterial adhesion to three different surfaces was investigated. Adhesion to
196 epithelial A549 cells was determined as previously described by Cuccui *et al.* (2017),

197 and adherence to polystyrene microtiter plates (Kasvi – K12-096) following growth for
198 24 h at 37°C was visualized using crystal violet, as described by Kaplan and Mulks
199 (2005). For the third adhesion assay, strains were inoculated in vials containing 1 cm²
200 steel coupons, as described previously by Moen *et al.* (2015). Briefly, the vials were
201 incubated at 37°C for a period of 24 h. Cultures were then fixed to the steel coupons
202 with 2.5% glutaraldehyde in 0.05 M phosphate buffered saline (PBS) and dehydrated in
203 a graded ethanol series up to 100%. The cells were dried using a CPD 030 critical point
204 dryer (Bal-Tec) and shadowed with gold using a Sputter Coater (Electron Microscopy
205 Sciences) prior to visualization with a scanning electron microscope (VP1430; LEO).

206

207 **Stress tolerance**

208 The following agents and their concentrations were used in BHI-NAD agar to
209 investigate the sensitivity of the Δhfq strains to different stress conditions: 1.5% NaCl;
210 pH 6.0 and 6.5 (adjusted using HCl); 1.25 mM H₂O₂; 4% ethanol; and cultivation at
211 42°C. Bacterial cultures with initial OD₆₀₀ of 1.0 were serially diluted in PBS to 10⁻⁷,
212 and 10 µL of each 10-fold dilution were applied on each selective stress agar in square
213 plates (688 102; Greiner Bio-One). As control, cultures were similarly plated on BHI-
214 NAD agar containing no stress agent. All plates were cultured at 37°C, except the
215 temperature stress plate, which was incubated at 42°C. The growth of strains was
216 compared between the control and test plates.

217

218 **Virulence in *G. mellonella***

219 The *Galleria mellonella* larvae used in this study were reared in our laboratory,
220 kept at 28°C in darkness and fed an artificial diet. On the day of the experiment, last-

221 instar larvae, each weighing 250-300 mg, were selected and kept in the same
222 environmental conditions until inoculation, following our previously described methods
223 (Pereira *et al.* 2015; Pereira *et al.* 2018, Blanco *et al.* 2017). Briefly, *A.*
224 *pleuropneumoniae* cultures were grown to mid-exponential phase and inocula
225 consisting of 10 µl of serially diluted cell suspensions, varying from 10³ to 10⁷ CFU per
226 larva (n=10 larvae per dilution), were injected into the haemocoel of the first right pro-
227 leg. The larvae were incubated at 37 °C, in the dark, and analyzed according to survival
228 at 24, 48, 72 and 96 hours post infection. Larvae were considered as dead if they did not
229 respond to touch stimuli. Survival curves were plotted using the Kaplan-Meier method
230 (Goel *et al.* 2010). For the evaluation of bacterial load, the larval haemolymph was
231 collected at 0, 1, 2, 4 and 24 hours after infection. Thereafter, the CFU/mL were
232 determined. Larvae inoculated with PBS were used as negative controls for the assay.

233

234 **Statistical Analysis**

235 Data from growth curves and adhesion to A549 cells and polystyrene microtiter
236 plates were analyzed by Tukey's test used to compare means using R v.2.13.0. The
237 differences in *G. mellonella* survival were calculated by using the log-rank test using R
238 v.2.13.0. A $p < 0.05$ was considered to be statistically significant. All the assays were
239 done in experimental and biologic triplicates.

240

241 **Results**

242

243 **Construction *A. pleuropneumoniae* hfq mutants**

244 As previously reported (Subashchandrabose *et al.* 2013), the *hfq* gene in *A.*
245 *pleuropneumoniae* is located in the *miaA-hfq-hflX* locus, as it is in *E. coli* (Tsui *et al.*
246 1994). This locus is shown in Fig 1A, with all detected promoters indicated, as well as
247 locations of the priming sites used for PCR amplification of products used to construct
248 the various plasmids. In order to determine the role of Hfq in *A. pleuropneumoniae*
249 serovars 1, 8 and 15, we generated isogenic mutants lacking 220 nucleotides, leaving a
250 truncated *hfq* gene having only 29 nucleotides in the 5' region and 30 nucleotides in the
251 3' region. Clean deletion mutants were generated for serovars 8 and 15, whereas counter
252 selection was not successful with the serovar 1 mutant, leaving the *catsacB* insertion in
253 place of the deleted 220 bases. In order to aid in evaluation of Hfq expression under
254 stress conditions, we also generated isogenic strains where the native *hfq* was replaced
255 with *hfq* additionally encoding a C-terminal 3XFLAG tag (followed by the *cat* gene in
256 the serovar 1 strain). Absence of any polar effects on *miaA* and *hflX* expression in the
257 Δhfq mutants was confirmed by RT-PCR, with representative results for serovar 8
258 shown in Fig1B. Expression of Hfq by the serovar 1, 8 and 15 Hfq::3XFLAG strains
259 during growth in broth culture was confirmed by Western blotting (Fig1C).

260

261 **Growth rate and Hfq expression**

262 Only the Ap1 Δhfq strain had a reduced growth rate in BHI-NAD, and the
263 complemented Ap8 Δhfq C strain had a prolonged lag phase, compared to its isogenic
264 WT strain, which was significant using the Tukey's test ($p < 0.05$) (Fig2A). Western blot
265 results for the FLAG-tagged mutants of each serovar showed that Hfq expression was
266 detectable at all time points assayed (Fig 2B), with apparent slight increase in Hfq
267 expression for the Ap8 $hfq::3XFLAG$ and Ap15 $hfq::3XFLAG$, but not
268 Ap1 $hfq::3XFLAG$ strains over the time course.

269

270 **Bacterial adhesion**

271 The WT strains of the different serovars tested showed marked differences in
272 adhesion to A549 epithelial cells, with serovar 8 being most, and serovar 15 least,
273 adherent (Fig 3A). Reduction of adherence was significant for the Ap1 Δ *hfq* and
274 Ap15 Δ *hfq* strains ($p < 0.05$), but not Ap8 Δ *hfq*, relative to their isogenic WT strains (Fig
275 3A). Instead of restoring WT levels, expression of *hfq* from the complementation vector
276 further reduced adherence for all serovars (Fig 3A), though the difference was only
277 significant for Ap15 Δ *hfq*C ($p < 0.05$). All Δ *hfq* mutants had reduced adhesion to
278 polystyrene compared to their WT strains ($p < 0.05$) (Fig 3B), with the Ap8 Δ *hfq* strain
279 showing the greatest reduction. In contrast to the assay using A549 epithelial cells, all
280 of the complemented strains showed increased adhesion to polystyrene compared to
281 their respective Δ *hfq* mutants, however these increases were not significant and did not
282 restore WT levels. The images of steel coupons obtained by electron microscopy
283 indicated that all Δ *hfq* strains had lower adherence capacity to this surface than their
284 respective WT strains (Fig 3C). This was particularly marked with Ap1 Δ *hfq* where
285 there were, in contrast to WT, few adherent cells, with clear complementation in the
286 Ap1 Δ *hfq*C strain. Although adherence of the Ap8 Δ *hfq* and Ap15 Δ *hfq* mutants was not
287 completely abolished, complementation did not restore WT levels (Fig 3C).

288

289 **Stress tolerance**

290 We investigated the responses of *A. pleuropneumoniae* serovars 1, 8 and 15, and
291 their respective Δ *hfq* mutants, to a variety of stress inducing agents or physical stress
292 (higher temperature) whilst growing on BHI-NAD-agar plates, as shown in Fig 4. The

293 WT strains of serovars 1, 8, and 15 showed different levels of resistance to the different
294 stresses, with serovar 8 being more sensitive to NaCl, and serovar 15 being more
295 resistant to ethanol, but more sensitive to elevated temperature (42°C) and pH 6.0, than
296 the other serovars. Furthermore, the respective Δhfq mutants also showed differences.
297 Unlike Ap8 Δhfq , no growth of Ap1 Δhfq and Ap15 Δhfq was found in the presence of
298 1.25 mM H₂O₂. Compared to the WT strains, all Δhfq mutants were more sensitive to
299 1.5% NaCl, although with differing degrees of growth reduction. Ap8 Δhfq and
300 Ap15 Δhfq were sensitive to the presence of ethanol, whereas only Ap15 Δhfq showed a
301 slight reduction in growth at 42°C. Ap1 Δhfq and Ap15 Δhfq , in contrast to Ap8 Δhfq ,
302 were more sensitive to growth at pH 6.5 than their WT strains. Except in the case of the
303 serovar 8 WT and mutant strains grown in the presence of NaCl, restoration of WT
304 levels of growth was achieved by complementation for all other conditions where
305 deletion of *hfq* resulted in increased sensitivity.

306

307 **Virulence in *G. mellonella***

308 The results of the virulence assay using the *G. mellonella* infection model are
309 shown in Fig 5. The concentration of 1.0×10^5 CFU per larva was chosen for graphic
310 representation, as it was found to be the best dose to allow visualization of the
311 differences in the virulence profiles between WT, Δhfq and $\Delta hfqC$ for each serovar. At
312 24 hours, larvae inoculated with serovar 1 strains showed 4% survival for the WT, 22%
313 for Δhfq and 25% for $\Delta hfqC$ indicating that the serovar 1 WT was highly virulent in this
314 infection model, and the Δhfq mutant only slightly attenuated compared to the WT
315 ($p < 0.05$), but complementation did not restore the WT level of virulence (Fig 1A).
316 Similar survival rates were found at 96 hours. The serovar 8 WT strain was not as
317 virulent as the serovar 1 WT, with survival of *G. mellonella* of 60% at 24, and 26% at

318 96 hours (Fig 1A). However, the Ap8 Δ hfq mutant was fully attenuated, with 100%
319 survival of *G. mellonella* through the 96 hours test period. Partial complementation was
320 seen for Ap8 Δ hfqC, with 51% survival of *G. mellonella* at 96 hours. The serovar 15 WT
321 did not appear to be virulent in the *G. mellonella* infection model (over 90% survival
322 through 96 hours), and no difference was seen for the AP15 Δ hfq and AP15 Δ hfqC
323 strains (Fig 1A). Analysis of bacterial load also showed a similar decrease over time in
324 the larvae infected with the serovar 15 WT, Δ hfq and Δ hfqC strains (Fig 5B). The
325 Ap1 Δ hfq and Ap8 Δ hfq strains both showed a five-log decrease in the number of
326 colonies per larva in the course of 24 hours, and few bacterial (approximately 10¹) cells
327 were observed in the haemolymph after of 24 hours of the experiment (Fig 5B). In
328 contrast, larvae infected with the serovar 1 and 8 WT strains showed an increase of
329 bacterial load between 1 and 4 hours, followed by less than a two-log reduction by 24
330 hours post-infection. In contrast to the results for the *G. mellonella* survival assay, the
331 Ap1 Δ hfqC showed partial, whereas the Ap8 Δ hfqC showed no, complementation in
332 regard to bacterial load.

333

334 **Discussion**

335 The role of the RNA chaperone Hfq in different bacterial species can be
336 variable. For example, *Francisella novicida* (Chambers and Bender 2011) and
337 *Cronobacter sakazakii* (Kim *et al.* 2015) hfq mutants are less resistant to oxidative
338 stress, by contrast *Staphylococcus aureus* mutants are more resistant to oxidative stress
339 (Liu *et al.* 2010). As Hfq mediates the interaction of many sRNAs with their target
340 mRNAs, in some cases leading to repression and in others activation of target gene
341 expression (Vogel and Luisi, 2011; Feliciano *et al.* 2016). The distribution of specific

342 genes and sRNAs involved in encoding and regulating expression of complex
343 phenotypes such as growth, biofilm formation, stress resistance, and virulence can vary
344 between different serovars/strains of the same species, so it is not surprising to find the
345 effects of global regulators can be significantly strain as well as species dependent.

346 In this study, we compared the effects of *hfq* mutation in serovars 1 (Shope
347 4074; reference strain), 8 (MIDG2331; clinical isolate) and 15 (HS143; reference strain)
348 of *A. pleuropneumoniae*, an important swine pathogen for which there are 18 known
349 serovars (Bossé *et al.* 2018) that can vary in their degree of virulence in pigs (Rogers *et*
350 *al.* 1990; Sassu *et al.* 2018). Serovar 1 isolates, expressing Apx toxins I and II, are
351 typically characterized by high virulence, whereas serovars 8 and 15, expressing ApxII
352 and III, are characterized by moderate virulence (Frey 2011). In addition, factors other
353 than RTX toxins, some of which are serovar specific, also contribute to virulence
354 (Bossé *et al.* 2002; Chiers *et al.* 2010). Initially, our goal had been to characterize the
355 influence of Hfq on several aspects of the physiology of *A. pleuropneumoniae* serovar
356 8, using MIDG2331 as a model, with serovar 1 and 15 strains used as controls for *hfq*
357 mutation (Subashchandrabose *et al.*, 2013) and natural transformation (Bossé *et al.*,
358 2009), respectively. However, as different phenotypes for the mutants became apparent,
359 along with differences in the virulence profiles and other features of the WT strains, we
360 shifted our efforts towards comparing the differential influence of the lack of Hfq in
361 strains from these distinct serovars.

362 A previous study by Subashchandrabose *et al.* (2013), characterizing an *hfq*
363 mutant of a clinical serovar 1 strain (AP 93-9), showed a slight reduction in growth rate
364 compared to the WT during cultivation in rich broth, which could be complemented by
365 expression of *hfq* from a plasmid. In our current study, we found similar results for a
366 Δhfq mutant of the serovar 1 reference strain (Shope 4074), however deletion of *hfq* in

367 the serovar 8 and 15 strains tested had no effect on growth in rich broth, indicating a
368 possible serovar-related effect.

369 The majority of clinical isolates of *A. pleuropneumoniae* readily form biofilms,
370 but this phenotype tends to be lost after passage in broth culture, suggesting repression
371 *in vitro* (Kaplan and Mulks 2005). Of the twelve serovar reference strains tested, only
372 the serovar 5b and 11 strains (L20 and 56513, respectively) retained the ability to adhere
373 to glass tubes or polystyrene plates, indicating possible serovar-related differences in
374 regulation of this phenotype (Kaplan and Mulks 2005). Production of a poly-1,6-*N*-
375 acetylglucosamine (PNAG) exopolysaccharide matrix has been shown to be the main
376 contributor to *A. pleuropneumoniae* biofilm formation on abiotic surfaces (Kaplan *et al.*
377 2004; Izano *et al.* 2007), with the O-antigen component of LPS also shown to contribute
378 (Hathroubi *et al.* 2015). Components of LPS, PNAG, pili, outer membrane proteins, and
379 glycoproteins have also been implicated in binding of *A. pleuropneumoniae* to various
380 cell lines (Cuccui *et al.* 2017; Rioux *et al.* 1999; Paradis *et al.* 1994; Van Overbeke *et*
381 *al.* 2002; Auger *et al.* 2009; Li *et al.* 2012; Liu *et al.* 2015; Liu *et al.* 2018), indicating a
382 more complex phenotype than binding to abiotic surfaces.

383 Deletion of *hfq* in the AP 93-9 clinical serovar 1 strain, a strong biofilm former,
384 was shown to reduce expression of *pgaC*, encoding the glycosyltransferase involved in
385 PNAG biosynthesis, and completely abrogated the ability to adhere to polystyrene
386 (Subashchandrabose *et al.* 2013). In our study, we further investigated the contribution
387 of Hfq to regulation of adherence of *A. pleuropneumoniae* to biotic and abiotic surfaces.
388 We found that the serovar 1, 8, and 15 WT strains tested showed different levels of
389 adhesion to various surfaces. The WT serovar 8 clinical isolate (MIDG2331) showed
390 the highest level of adherence to the A549 human alveolar basal epithelial cell line,
391 which we have previously used for *A. pleuropneumoniae* adhesion assays (Cuccui *et al.*

392 2017), as well as to polystyrene plates. Furthermore, the biofilm formed by the serovar
393 8 WT on steel coupons showed a more mature 3-D architecture, compared to those of
394 the serovar 1 and 15 WT strains. Although the serovar 15 reference strain showed
395 similar adherence to polystyrene and steel coupons when compared to the serovar 1
396 reference strain, it also showed the lowest level of adherence to A549 cells.

397 Deletion of *hfq* resulted in varying degrees of adherence reduction to the
398 different surfaces depending on the serovar. All three serovar Δhfq mutants showed
399 slight, but significant ($p < 0.05$), reduction of adherence to polystyrene, with some
400 restoration (not significant) of binding in each following expression of the *hfq* gene
401 from the complementation vector. It is not clear why we did not see complete
402 abrogation of binding to polystyrene with our *hfq* mutants, or full complementation, as
403 was seen in the study by Subashchandrabose *et al.* (2013), but this may have been due
404 to differences in the isolates and/or how the assays were performed. Results of the
405 adhesion assay using steel coupons showed all Δhfq strains had lower adherence
406 capacity to this surface than their respective WT strains, but restoration of the WT
407 adherence phenotype was only seen with the Ap1 Δhfq C strain. In the assay using A549
408 cells, all three Δhfq mutants showed reduced adherence compared to their respective
409 WT strains, but the level of reduction was only significant ($p < 0.05$) for the serovar 1
410 and 15 mutants. As opposed to no, or partial, complementation of the binding
411 phenotypes, each of the three serovar Δhfq mutants expressing the plasmid encoded *hfq*
412 gene showed even further reductions in binding to A549 cells, though this was only
413 significant ($p < 0.05$) for the Ap15 Δhfq C strain.

414 It is difficult to determine from the current studies whether the different *in vitro*
415 adhesion phenotypes are due to serovar related differences in encoded adhesion genes,
416 or to changes in gene regulation following passage of clinical isolates in the laboratory,

417 or both. Clearly the clinical serovar 8 WT strain showed the greatest adherence to all of
418 the surfaces tested compared to the WT serovar 1 and 15 reference strains, and deletion
419 of *hfq* reduced adherence in all cases, but to different extents. These data support a role
420 for Hfq in regulating at least some of the gene products contributing to adherence to the
421 different surfaces in each of the serovar strains tested, but other regulators such as RpoE
422 and H-NS (Bossé *et al.* 2010) and the two component systems CpxA/CpxR (Li *et al.*
423 2018) and QseA/QseB (Liu *et al.* 2015) have also been shown to be involved, and their
424 relative contributions to regulating this complex phenotype are unresolved.

425 The ability to respond to and repair damage caused by a variety of stresses is
426 important for the survival of *A. pleuropneumoniae* within its host, especially during
427 acute disease (Sheehan *et al.* 2003; Klitgaard *et al.* 2012). Numerous genes involved in
428 stress response have been identified, and their expression has been shown to be
429 regulated by factors including RpoE and (p)ppGpp (Bossé *et al.* 2010; Li *et al.* 2015). In
430 the study by Subashchandrabose *et al.* (2013), Hfq was shown to contribute to
431 resistance of their clinical serovar 1 isolate to oxidative stress, but other sources of
432 stress were not investigated. Here we have shown that there was variation in the
433 response of the three different serovar WT strains, as well as their Δhfq mutants, to
434 different stressors. For example, the serovar 8 WT was more sensitive to NaCl stress,
435 whereas the serovar 15 WT was more sensitive to heat stress at 42°C, but more resistant
436 to ethanol stress, than the other two WT strains. The Δhfq mutants of serovars 1 and 15,
437 but not serovar 8, were more sensitive to H₂O₂ and pH 6.5 compared to their WT
438 parental strains. In contrast to the adhesion experiments, the stress susceptible
439 phenotypes were complemented by expression of the plasmid-encoded *hfq* gene.

440 Although Subashchandrabose *et al.* (2013) previously reported that deletion of
441 *hfq* in *A. pleuropneumoniae* did not result in increased sensitivity to H₂O₂ or cumene

442 hydroperoxide, it did increase sensitivity to methyl viologen and potassium tellurite –
443 both known to generate superoxide radicals within bacterial cells. In their study,
444 sensitivity to these agents was tested using a disk diffusion assay, and they used a strong
445 biofilm forming clinical isolate of serovar 1 (AP 93-9) of *A. pleuropneumoniae*. Both of
446 these factors could explain the differences in results found in our current study. Overall,
447 the data indicate that, as for adhesion and biofilm formation, Hfq plays a role in stress
448 resistance, but there are serovar- or even strain-dependent differences in regulation of
449 this complex phenotype.

450 Finally, we compared virulence of serovar 1, 8 and 15 *A. pleuropneumoniae* WT
451 and Δhfq mutants in the *G. mellonella* infection model that we previously described
452 (Pereira *et al.* 2015). As for the other phenotypes tested, there were variations between
453 the different WT serovars and Δhfq mutant strains with regards to virulence in this
454 model. Since the serovar 15 WT was avirulent under the conditions tested, no difference
455 was seen following deletion of the *hfq* gene in this strain. The serovar 1 WT was the
456 most virulent, but only showed a slight decrease, whereas the moderately virulent
457 serovar 8 WT was completely attenuated following deletion of its *hfq* gene.
458 Furthermore, complementation was not successful for the serovar 1 mutant expressing
459 the plasmid-encoded *hfq* gene, but restored almost full WT level of virulence for the
460 serovar 8 mutant, indicating possible differences in genes (and possible differences in
461 gene regulation) contributing to virulence of these serovars in this infection model.

462 Complementation of mutated phenotypes using cloned genes expressed from
463 shuttle plasmids is always challenging, as factors including level of expression (due to
464 copy number of plasmid and strength of promoter used for expression) and indirect
465 effects of interaction of the expressed gene with regulatory network(s) can influence the
466 overall success of restoring the WT phenotype. This is especially true for

467 complementation of genes encoding global regulators, such as Hfq. We tried account
468 for possible confounding issues by cloning the *hfq* gene into a low copy number
469 plasmid (pMIDG100), with expression possible from the endogenous sigma 70 and/or
470 sigma E promoters included in the upstream sequence. However, we still found that
471 pMIDG_*hfq* was able to complement some, but not all of the Δ *hfq* mutant phenotypes,
472 and this was sometimes serovar-dependent. In the case of binding to A549 epithelial
473 cells, expression of the plasmid copy of *hfq* resulted in further reductions rather than
474 restoration of WT levels in adherence for all 3 serovar strains. Each of the phenotypes
475 analyzed in this study are complex, and result from coordinated expression of different
476 genes, some of which may be regulated by Hfq-dependent sRNAs and others not.
477 Adding to this complex network, other regulators such as sigma factors and DNA
478 binding proteins may also be involved, and the balance of these factors likely
479 determines the resulting phenotype. Similar observations of partial complementation
480 and/or exacerbation of phenotype have been made by others when expressing *hfq* on
481 plasmids, either from its own promoter or an inducible one (Fantappiè *et al.* 2009;
482 Schilling *et al.* 2009; Bai *et al.* 2010; Chambers *et al.* 2011). It is possible that, even
483 though we deliberately cloned the *hfq* gene along with three possible endogenous
484 promoters, on a low copy number plasmid, the intracellular levels of Hfq were either
485 lower or higher than those present in the WT strains, resulting in differential regulation
486 of genes affecting the different phenotypes.

487 In summary, we found that Hfq contributes to regulation of adhesion to biotic
488 and abiotic surfaces, resistance to various stress conditions, and virulence in a surrogate
489 model of infection, to differing extents in the three serovars of *A. pleuropneumoniae*
490 studied. The full set of genes and sRNAs contributing to each of these phenotypes, and
491 how these differ between serovar/strains of *A. pleuropneumoniae*, remain to be

492 determined. We conclude the need for caution in extrapolating the effects of deletion of
493 global regulators, and *hfq* in particular, to other strains of the same species, especially
494 regarding complex phenotypes.

495

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Table 1. Strains and plasmids used in this study.

Strains and plasmids	Description	Source or reference
<i>Actinobacillus pleuropneumoniae</i>		
Serotype 1		
Shope 4074 WT	Shope 4074 Wild-type	ATCC 27088
Ap1Δ <i>hfgcatsacB</i>	Δ <i>hfg</i> mutant of Shope 4074	This study
Ap1Δ <i>hfgcatsacBC</i>	Complemented strain	This study
Ap1 <i>hfg</i> ::3XFLAG <i>cat</i>	WT containing a 3XFLAG tag replacing the last codon of the <i>hfg</i> gene.	This study
Serotype 8		
MIDG2331 WT	Serotype 8 clinical isolate from UK	(Bossé <i>et al.</i> 2016)
Ap8Δ <i>hfg</i> ^a	Δ <i>hfg</i> mutant of MIDG2331	This study
Ap8Δ <i>hfgC</i>	Complemented strain	This study
Ap8 <i>hfg</i> ::3XFLAG	WT containing a 3XFLAG tag replacing the last codon of the <i>hfg</i> gene.	This study
Serotype 15		
HS143 WT	HS143 Wild-type	(Blackall <i>et al.</i> 2002)
Ap15Δ <i>hfgcat</i>	Δ <i>hfg</i> mutant	This study
Ap15Δ <i>hqc</i>	Complemented strain	This study
Ap15 <i>hfg</i> ::3XFLAG	WT containing a 3XFLAG tag replacing the last codon of the <i>hfg</i> gene.	This study
<i>Escherichia coli</i>		
Stellar	Competent cell: F ⁻ , <i>endA1</i> , <i>supE44</i> , <i>thi-1</i> , <i>recA1</i> , <i>relA1</i> , <i>gyrA96</i> , <i>phoA</i> , Φ80 <i>Δ lacZΔ M15</i> , Δ (<i>lacZYA - argF</i>) U169, Δ (<i>mrr-hsdRMS-mcrBC</i>), Δ <i>mcrA</i> , λ ⁻	Takara
MFDpir	Conjugative cell: MG1655 RP4-2-Tc::[Δ <i>Mu1::aac(3)IV- ΔaphA- Δnic35- ΔMu2::zeo</i>] Δ <i>dapA::(erm-pir) ΔrecA</i> . Strain used to introduce pMIDG_ <i>hfg</i> in <i>A. pleuropneumoniae</i> Δ <i>hqc</i> strains.	(Ferrières <i>et al.</i> 2010)
Plasmids		
pUSS <i>catsac</i>	Template DNA for amplification of <i>catsacB</i> cassette which contains DNA uptake sequences for natural transformation into <i>A. pleuropneumoniae</i> .	(Bossé <i>et al.</i> 2014)
p <i>Thfg</i> Flank	Plasmid pGEM-T containing 600 nucleotides upstream <i>hfg</i> gene, <i>hfg</i> gene and 600 nucleotides downstream <i>hfg</i> gene.	This study
pTΔ <i>hfgcatsacB</i>	Plasmid pTΔ <i>hfg</i> containing the <i>hfg</i> gene disrupted by <i>catsacB</i> cassette.	This study
p <i>Thfg</i> 3XFLAG	Plasmid pGEM-T containing the <i>hfg</i> gene with a 3XFLAG tag in the region 3' of the <i>hfg</i> gene.	This study
p <i>Thfg</i> 3XFLAG <i>cat</i>	Plasmid pGEM-T containing the <i>hfg</i> gene with a 3XFLAG tag in the region 3' of the <i>hfg</i> gene followed by the <i>cat</i> gene.	This study
p <i>Thfg</i> ::3XFLAG <i>catsacB</i>	Plasmid pGEM-T containing the <i>hfg</i> gene with a 3XFLAG tag in the region 3' of the <i>hfg</i> gene disrupted by <i>catsacB</i> cassette.	This study
pMIDG_ <i>hfg</i>	pMIDG plasmid (Bossé <i>et al.</i> , 2009) containing the <i>hfg</i> gene under promoter inside of the <i>miaA</i> gene. Strain used to complement the Δ <i>hfg</i> strains.	This study

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507

^a Although this mutant strain has been used in a previous study by our group (Pereira *et al.*, 2015), this is the first description of the generation of the mutation.

Table 2. Primers used in this study.

N°	Primer	Oligonucleotide sequence (5' to 3')	Description
1	<i>hfq</i> flank_for	TTCCGGTGGAAGTAATTAGCGTAGA	For amplification of the <i>hfq</i> cassette.
2	<i>hfq</i> flank_rev	ATATCCGCTTTCTGACGAGTTTTGC	
3	<i>cat_delta</i> <i>hfq</i>	<u>ATCTTTACAAGATCCT</u> TACAAGCGGT CGGCAATAAGTTACC	For amplification of <i>catsacB</i> cassette containing 15 bp overhangs (underlined) complimentary to p <i>Thfq</i> Flank opened by inverse PCR with primers 5 and 6.
4	<i>sac_delta</i> <i>hfq</i>	<u>CGCAACCGCTTCAACGAATTGCGTG</u> AAGCTCGAGGTATG	
5	<i>delta</i> <i>hfq_inv</i> <i>cat</i>	<u>TGCCGACCGCTTGTAGGATCTTGTA</u> AAGATTGACCTTTTGC	For inverse PCR amplification of p <i>Thfq</i> Flank to remove all but 59 bp of the <i>hfq</i> gene and adding 15 bp overhangs (underlined) complementary to the <i>catsacB</i> cassette generated with primers 3 and 4.
6	<i>delta</i> <i>hfq_inv</i> <i>sac</i>	<u>AGCTTCACGCAATTCGTTGAAGCGG</u> TTGCGGATAAAGC	
7	<i>delta</i> <i>hfq_1</i>	<u>CGCAACCGCTTCAACGGATCTTGTA</u> AAGATTGACCTTTTGC	For generation of Δ <i>hfq</i> construct. Addition of 15 bp overhangs (underlined) allow direct fusion of left flank amplified using primers 1 and 7 to right flank generated using primers 2 and 8.
8	<i>delta</i> <i>hfq_2</i>	<u>ATCTTTACAAGATCCGTTGAAGCGG</u> TTGCGGATAAAGC	
9	FLAG_ <i>hfq</i>	<u>GTTGCGGATAAAGCGGGTACCGAC</u> TACAAAGACCATGAC	For amplification of the 3XFLAG cassette containing 15 bp overhangs (underlined) complementary to p <i>Thfq</i> Flank opened by inverse PCR with primers 11 and 12.
10	FLAG_ <i>hflX</i>	<u>TTGGTATCTGATCGGCTCCAGCCTA</u> CATTACTATTTATCG	
11	<i>hfq_inv</i> 1	CGCTTTATCCGCAACCGCTTCAAC	For generation of p <i>Thfq</i> ::3XFLAG
12	<i>hfq_inv</i> 2	CCGATCAGATACCAAATACAGATG	
13	<i>cat</i> _FLAG	<u>TAATGTAGGCTGGAGGTACAAGCG</u> GTCGGCAATAGTTACC	For amplification of the <i>cat</i> cassette containing 15 bp overhangs (underlined) complimentary to 3XFLAG and p <i>Thfq</i> Flank opened by inverse PCR with primers 11 and 12.
14	<i>cat_hflX</i>	<u>TTGGTATCTGATCGGGAAGTGCGGT</u> ATGCCGTCTGAAC	
15	FLAG_ <i>cat</i>	<u>GCCGACCGCTTGTACCTCCAGCCTA</u> CATTACTATTTATCG	For amplification, in combination with primer 9, of the 3xFLAG cassette containing 15 bp overhangs (underlined) complementary to <i>cat</i> cassette and p <i>Thfq</i> Flank opened by inverse PCR with primers 11 and 12.
16	<i>hflX_for</i>	CACGAGCTTAGTCCGTCACA	
17	<i>hflX_rev</i>	AATGCTACCCGCTGTATGCT	For RT-PCR analysis of <i>hflX</i> expression.
18	<i>miaA_for</i>	TAATGGGTCCAACGGCTTCG	For RT-PCR analysis of <i>miaA</i> expression.
19	<i>miaA_rev</i>	CACTGTTCCAACCTCGCAGCCAAG	
20	EcoRI_ <i>hfq</i>	GCGCGAATTCAGGAAAAGAAAATG GCAAAGGTCAATCT	For RT-PCR analysis of <i>hfq</i> expression.
21	<i>hfq_SacI</i>	GCGCGAGTCTATTATCCGCTTTAT CCGCAACCGC	
22	<i>hfq</i> MIDG_for	GCTCAAGCTTCGAATTCGAGCTTGC CCCTCACCGCTTGATTG	For amplification of <i>hfq</i> gene with its own promoter region and containing 15

23 *hfq*MIDG_rev TTGGGATCTTTCGAAGCGTTTTTCAT
CTGTATTTGGTATCTG

bp overhangs complementary to
pMIDG100 cut with *Eco*RI and *Bst*BI.

509

510 **Figure legends:**

511

512 **Fig 1. Generation and confirmation of *hfq* mutant strains.** (A) Genomic
513 organization of the *miaA*, *hfq* and *hflX* genes in *A. pleuropneumoniae*. Predicted
514 promoter sequences are indicated by the bent arrows labelled P1 to P4, and a predicted
515 transcriptional terminator is indicated by a stem-loop structure downstream of *hfq*. The
516 primers used in mutant construction, cloning and RT-PCR, are represented by arrows
517 below their targets, numbered according to their identification in Table 2. (B) RT-PCR
518 analysis of possible polar effects due to deletion of *hfq*, showing representative results
519 for MIDG2331. PCR was performed with the products of cDNA synthesis from RNA
520 template of either the WT or Δhfq strain (as indicated), both with (RT+) and without
521 (RT-) the addition of reverse transcriptase. Note that the genomic DNA (gDNA) control
522 used to confirm primer function and product size for each primer pair was from the WT
523 strain only, thus a product for *hfq* amplification is seen as a comparison for the lack of
524 amplification by RT-PCR from the Δhfq strain. Amplification of the target sequences in
525 *hfq*, *miaA* and *hflX* was achieved with the primer pairs 20/21, 18/19 and 16/17,
526 respectively. M = molecular weight marker (DNA Marker Quick-load 100bp DNA
527 ladder, Neb Biolabs). (C) Western blot showing the detection of the 14 kDa
528 Hfq::3XFLAG protein. For each *hfq*::3XFLAG strain of *A. pleuropneumoniae* serovars
529 1 (Ap1), 8 (Ap8) and 15 (Ap15), ten micrograms of soluble protein from early
530 stationary phase culture were separated by SDS-PAGE and transferred to nitrocellulose
531 membrane for detection using an anti-Flag antibody. The molecular weight marker lane

532 (M = SeeBlue Plus2; Invitrogen) from the corresponding stained gel is shown next to
533 the blot.

534

535 **Fig 2. Growth of *A. pleuropneumoniae* WT, *hfq* mutants and complemented**
536 **strains.** (A) Growth curve of *A. pleuropneumoniae* serovars 1, 8 and 15 strains. (B)
537 Hfq::3XFLAG expression analysis during the growth curve of the *A. pleuropneumoniae*
538 strains. WT (wild-type), *hfq*::3XFLAG (strain that express Hfq::3XFLAG), Δhfq (*hfq*
539 mutant), $\Delta hfqC$ (complemented strain). Error bars are shown for all points in the graphs,
540 but may not be visible in some cases.

541

542 **Fig 3. Effect of Hfq on adherence of *A. pleuropneumoniae* serovars 1, 8 and 15 to**
543 **biotic and abiotic surfaces.** (A) The adherence to eukaryotic cells. (B) The adherence
544 to polystyrene microplate was examined by crystal violet reading in OD₆₀₀ and the
545 adherence capacity was determined according to WT strains. (C) The adherence to steel
546 coupons was examined by scanning electron microscopy (SEM). Bars: 10 μ m. Different
547 letters inside of the Fig represent statistical significance difference among the strains in
548 relation the cell length. The statistical analysis was performed using Tukey's test with
549 $p < 0.05$. All the assays were conducted in experimental and biological triplicates. WT
550 (wild-type), Δhfq (*hfq* mutant), $\Delta hfqC$ (complemented strain).

551

552 **Fig 4. Effect of Hfq on stress tolerance in *A. pleuropneumoniae* serovars 1, 8 and**
553 **15.** Exponentially growing *A. pleuropneumoniae* strains (OD₆₀₀ = 1.0; $\sim 10^8$ cell/mL)
554 were exposed to different stress conditions: oxidative (1.25 mM H₂O₂), osmotic (1.5%
555 NaCl), alcoholic (4% ethanol), temperature (42°C), pH (6.5 and 6.0). As control, the
556 strains were grown in BHI-NAD-agar at 37°C, 5% CO₂ and no stressor agent. The

557 numbers 1, 2 and 3 indicate the WT (wild-type), Δhfq (*hfq* mutant) and $\Delta hfqC$
558 (complemented strain), respectively.

559

560 **Fig 5. Effect of Hfq on the virulence of *A. pleuropneumoniae* serovars 1, 8 and 15 in**

561 ***G. mellonella*.** (A) Killing was monitored after larval infection with 1×10^5 CFU of WT,

562 Δhfq and $\Delta hfqC$ *A. pleuropneumoniae* strains from serovars 1, 8 and 15. The virulence

563 attenuation was verified in the Δhfq strains of serovars 1 and 8 ($p < 0,05$). (B)

564 Determining of the bacterial load in *G. mellonella* hemolymph at 0, 1, 2, 4 and 24 hours

565 of the assay of (A) in three biological replicates. Larvae inoculated with PBS 1X were

566 used as negative control. WT (wild-type), Δhfq (*hfq* mutant) and $\Delta hfqC$ (complemented

567 strain).

568

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