

1 **Supplementary feeding stations for conservation of vultures could be an important**
2 **source of monophasic *Salmonella* Typhimurium 1,4,[5],12:i:-**

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22 **ABSTRACT**

23 Vultures are nature's most successful scavengers, feeding on the carcasses of dead animals
24 present in the field. Availability of domestic carrion has been unstable due to rapidly
25 changing agro-grazing economies and increasing sanitary regulations that may require burial
26 or burning of livestock carcasses. Thus, several griffon vulture (*Gyps fulvus*) recoveries are
27 based on European legislation that guarantees the animals' welfare, avoids intense
28 persecution of the vultures and allows the feeding of threatened wildlife in supplementary
29 feeding stations (SFS). However, in recent years, many studies have speculated on the
30 likelihood that avian scavengers may be infected by feeding on pig carcasses at SFS from
31 intensive livestock. In this context, the present study evaluated whether free-living griffon
32 vultures and pig farms share zoonotic *Salmonella* strains to test the hypothesis that vulture
33 are infected during consumption of carcasses provided at SFS. Here, the occurrence,
34 serotypes and genomic DNA fingerprinting (phage typing and pulsed-field gel
35 electrophoresis) of isolated strains were carried out in griffon vultures and pig farms
36 authorised to provided carcasses at SFS in Castellón province (eastern Spain). The
37 bacteriological analyses revealed that 21.1% of vultures and 14.5% for pig farms samples
38 tested were *Salmonella*-positive. Monophasic *S. Typhimurium* 1,4,[5],12:i:- was the most
39 frequently isolated serovar. Comparison of *Salmonella* strains isolated from vultures and pig
40 farms revealed that monophasic *S. Typhimurium* 1,4,[5],12:i:-, *S. Derby* and *S. Rissen* strains
41 were highly genetically homogeneous (similar DNA fingerprint). In conclusion, the current
42 study indicates that free-living griffon vultures and pig farms that provide the carcasses at
43 SFS share several zoonotic *Salmonella* strains. On this basis, and although transmission could
44 be bidirectional, our result seems to corroborate the pig carcasses-to-vulture transmission and
45 cross-infection at SFS. As an immediate *Salmonella* control strategy in wild avian

46 scavengers, we suggest the implementation of a programme to guarantee that solely pig
47 carcasses from *Salmonella*-free farms arrive at SFS.

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50 **Keywords:** Avian scavengers; intensive farming; pig; environment; *Salmonella* Derby;
51 *Salmonella* Rissen.

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54 **1. Introduction**

55

56 Wild birds have repeatedly been highlighted as a source in the dissemination of *Salmonella*
57 spp. (Tizard, 2004; Hilbert et al., 2012; Krawiec et al., 2015) and links have been documented
58 between *Salmonella* contamination of modern pig production and wild birds (Andrés et al.,
59 2013; Andrés-Barranco et al., 2014). Furthermore, the number of wildlife species acting as
60 reservoirs, amplifiers and disseminators is unknown (Molina-López et al., 2011). Hence, the
61 role of wildlife as a *Salmonella* reservoir is of increasing interest (Hilbert et al., 2012).
62 *Salmonella* stands out as one of the most most common causes of human bacterial food
63 poisoning (EFSA, 2017).

64

65 During the past century, the availability of domestic carrion has been unstable due to rapidly
66 changing agro-grazing economies and increasing sanitary regulations that may require burial
67 or burning of livestock carcasses. The conservation and reintroduction of avian scavengers
68 would therefore not have been possible without European Regional legislation to ensure their

69 welfare and avoid their intense persecution, as well as the European ruling that permitted the
70 feeding of threatened wildlife in SFS (Camiña-Cardenal et al., 2004; Margalida et al., 2011;
71 Cortés-Avizanda et al., 2016). In the late 1960s, conservationists created “vulture
72 restaurants” or SFS as a way to increase the availability of food resources (Bijleveld, 1974;
73 Gilbert et al., 2007; Donázar et al., 2009; Fielding et al., 2014). At community level, SFS has
74 been widely accepted as an effective management tool among conservationists and managers
75 (Cortés-Avizanda et al., 2016). Encouraging fallen stock to be left in situ is ecologically
76 harmonious, inexpensive and an efficient management method for the conservation of
77 scavengers (Donázar et al., 2009).

78

79 *Salmonella* has been isolated in vultures in several studies, but an especially remarkable
80 finding is the unexpected abundance of *Salmonella* ser. Typhimurium, one of the most
81 common *Salmonella* serovars in foodborne illness outbreaks related with pork consumption
82 (Millán et al., 2004; Molina-López et al., 2011:2015; Marin et al., 2014; Jurado-Tarifa et al.,
83 2016; Blanco, 2018). Besides, to date it remains unknown whether *Salmonella* can cause
84 clinical illness in avian scavengers (Blanco, 2018), which could have potential implications
85 for conservation. Notably, *S.* Typhimurium, including monophasic variants (1,4,[5],12:i- and
86 1,4,12:i-), represented 21.8% of all reported serovars of confirmed human cases in 2016 in
87 the EU (EFSA, 2017). In particular, *S.* Typhimurium accounted for 63.6% of the isolates
88 reported in pig samples (EFSA, 2017). After the prion crisis, pig carcasses have been the
89 scavengers’ main foodstuff provided at SFS (Blanco et al., 2016; Green et al., 2016; Blanco,
90 2018). Thus, a recent study carried out by Blanco (2018) in Segovia province (central Spain)
91 supports the role of pig carcasses as a primary source of *Salmonella*, and the risk of scavenger
92 infection in SFS, based on the concordance of serotypes and resistance patterns in an obligate

93 scavenger partially relying on pig carcasses. Our driving hypothesis was that pig farms could
94 be one source of vulture transmission and a cross-infection route of *Salmonella* at SFS. In
95 this context, the present study evaluated whether free-living griffon vultures and pig farms
96 share zoonotic *Salmonella* strains to test the hypothesis that vultures could be infected during
97 consumption of pig carcasses provided at SFS.

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100 **2. Material and methods**

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102 **2.1. Animals**

103 This study was conducted within the conservation project for endangered species in the
104 Valencia Region. The study population is located at the Cincorres observatory (Castellón
105 province, eastern Spain). In 2008, 236 breeding pairs were found in this area (93% of the
106 breeding pairs in the Community of Valencia) (GVA, 2008). All the experimental procedures
107 used in this study were performed in accordance with Directive 2010/63/EU EEC on animal
108 experiments. The Department of Infrastructure, Planning and Environment of the Valencian
109 Regional Government (Generalitat Valenciana) granted the ethical and animal welfare
110 permission to take samples.

111

112 **2.2. Sample collection**

113 The vultures were live-captured in two sessions in September and in October 2016, during
114 the observatory's normal ringing schedule as part of the reserve's monitoring programme. A
115 total of 104 free-living griffon vultures were captured using a remotely activated purpose-
116 built cage (for more details, see Marin et al., 2014). The age of the animals was determined

117 according to the plumage characteristics and the colour of the bill and eye, classified as
118 juvenile (less than 2 years), sub-adult (from 2 to 5 years) and adult (more than 5 years). Base
119 on our previous results where there were no age-related differences in relation to the presence
120 of *Salmonella* (Marin et al., 2014), data from all individuals was pooled. From each vulture,
121 one cloacal sample was obtained using sterile cotton swabs (Cary Blair sterile transport
122 swabs, Deltalab, Barcelona, Spain). The cotton swab was inserted 1 to 2 cm into the cloaca
123 to collect a suitable sample. At each sampling day, samples of the facilities in close contact
124 with pig carcasses during supplementary feeding (warehouse of cadavers where farmers
125 legally dispose of dead animals, trucks that transport the carcasses from the warehouse to the
126 SFS and pig carcasses at SFS) were collected. A total of 20 sterile cotton swabs were taken
127 from 20 pig carcasses deposited in the SFS (10 samples per day). In addition, before the
128 animals were loaded and delivered to the SFS point, 30 sterile cotton swabs samples were
129 taken directly from the surface of the 2 trucks (floor and wall) that transport the carcasses
130 from the warehouse of cadavers to the SFS (15 samples per truck and day). Moreover, 20
131 sterile cotton swabs were taken from the warehouse of cadavers (container walls) where
132 farmers disposed of dead livestock (2 samples per container and day). Finally, during the
133 week after the intensive sampling described above, a total of 11 pig farms that provide the
134 carcasses were sampled to determine the potential transfer of *Salmonella* isolates from pig
135 carcasses to vultures. Five pens (four in the corners and one in the middle of the barn) were
136 chosen in each farm. Briefly, 500 gr of faeces were collected in pools from different points
137 of the pens in sterile containers and transported under refrigeration to the laboratory. All
138 samples were analysed within 24 h of collection. The experimental design of this study is
139 shown in Fig. 1.

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141 **2.3. *Salmonella* spp. isolation and identification**

142 The procedure was based on the official method ISO 6579: 2002 recommendations (Annex
143 D). Cotton swab samples were pre-enriched in 1:10 vol/vol Buffered Peptone Water 2.5%
144 (BPW, Scharlau, Barcelona, Spain). Faeces samples were homogenised and 25 gr were
145 transferred into 225 mL of BPW. All BPW enrichments were incubated at 37 ± 1 °C for 18 ± 2
146 h. Next, x ul of these enrichments were inoculated onto Modified Semi-Solid Rappaport
147 Vassiliadis agar plates (MSRV, Difco, Valencia, Spain), which were incubated at 41.5 ± 1 °C
148 for 24–48 h. Suspicious growths on MSRV plates were selected for inoculation onto Xylose–
149 Lysine–Deoxycholate (XLD, Liofilchem, Valencia, Spain) and ASAP (ASAP, bioMerieux,
150 Madrid, Spain) agar plates and incubated at 37 ± 1 °C for 24–48 h. After the incubation period,
151 5 presumptive *Salmonella* colonies were selected and streaked onto nutrient agar plates
152 (Scharlab, Barcelona, Spain) 37 ± 1 °C for 24 ± 3 h. *Salmonella* isolates were serotyped
153 according to the Kauffman-White-Le Minor scheme (Grimont and Weill, 2007) and was
154 carried out at the Laboratori Agroalimentari (Cabrils, Spain) of the Departament
155 d’Agricultura, Ramaderia, Pesca i Alimentació.

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157 **2.4. Molecular typing of *Salmonella* strain isolates**

158 Enterobacterial repetitive intergenic consensus (ERIC)-PCR of all *Salmonella* isolates was
159 performed and representative isolates from the different ERIC-PCR patterns and different
160 origin were further analysed by pulsed-field gel electrophoresis (PFGE). ERIC-PCR was
161 performed as previously described, except that a 50°C annealing temperature was used
162 (Antilles et al., 2015). Primer pairs used were ERIC-F (5’-AAG TAA GTG ACT GGG GTG
163 AGC G-3’) and ERIC-R (5’-ATG TAA GCT CCT GGG GAT TCA C-3’) (Versalovic *et al.*,
164 1991). PFGE typing was performed according to the standard operating procedure of

165 PulseNet (www.pulsenetinternational.org). Genomic DNA was digested with the restriction
166 enzyme XbaI (Roche Applied Science, Indianapolis, IN) and the restriction fragments were
167 separated by electrophoresis in a CHEF-DR III System (Bio-Rad, Hercules, CA, USA).
168 Fingerprinting II v3.0 software (Bio-Rad, Hercules, CA, USA) was used to compare the
169 PFGE patterns by cluster analysis using Dice coefficient and unweighted pair group method
170 with arithmetic averages (UPGMA dendrogram type).

171

172 **2.5. Statistical analysis**

173

174 We tested whether occurrence of *Salmonella spp.* was related to sampling point. To do so,
175 we fitted a generalised linear model (GLM) where occurrence of *Salmonella spp.* was the
176 response variable and the sampling point (pig faeces, warehouse of cadavers, trucks that
177 transport the carcasses from the warehouse to the SFS, carcasses in SFS and vultures), session
178 (1 and 2) and their interaction were fixed effects. For this analysis, the error was designated
179 as having a binomial distribution and the probit link function was used. Binomial data for
180 each sample were assigned a 1 if *Salmonella spp.* was isolated or a 0 if not. The sampling
181 point x session interaction effect was included in the analysis, but this was discarded because
182 it was not significant. In addition, we tested whether occurrence of *Salmonella spp.* was
183 related to ages of vultures, using a GLM as previously. To do so, we fitted GLM where
184 occurrence of *Salmonella spp.* was the response variable, and age of vultures (juveniles, sub-
185 adult and adult) was the fixed effect. As estimators of the relative quality of the model,
186 Akaike information criterion (AIC) and Bayesian information criterion (BIC) were
187 considered. A P value <0.05 was considered to indicate a statistically significant difference.

188 Analyses were carried out using a commercially available software program (SPSS 21.0
189 software package; SPSS Inc., Chicago, IL, 2002).

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192 **3. Results**

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194 **3.1. *Salmonella* occurrence**

195 *Salmonella spp.* was detected in all of the sampling points. The proportion of *Salmonella*-
196 positive samples were in decreasing order: 82.8% for trucks that transport the carcasses from
197 the warehouse to the SFS (100% of the trucks), 40.0% for pig carcasses disposed of in SFS,
198 32.3% for warehouse of cadavers (100% of the containers), 21.1% for vultures and 14.5%
199 for pig faeces at farm. The sampling point clearly has a significant effect on occurrence of
200 *Salmonella spp.* (Table 1), as revealed by the model analyses (deviance of 88.05% with AIC
201 and BIC values of 43.546 and 64.043 respectively). No significant differences in occurrence
202 of *Salmonella spp.* were found between ages of vultures (Table 2). P values for this difference
203 did not achieve significance (Bonferroni test, P=0.617).

204 Serovar identification was obtained for 69 pooled samples (95.8%), with 3 isolates remaining
205 undetermined. All belonged to one of two subspecies: enterica (93.9%) and salamae (6.1%).

206 A total of 8 serotypes were identified (pooling all sampling point positive samples, Table 3).

207 The most predominant serotype was monophasic *S. Typhimurium* 4,12:i:- detected in 49.3%
208 of positive samples (pooling all sampling point positive) followed by *S. Panama* (23.2%), *S.*

209 *London* (13.0%), *S. 4,12:b[-]* (5.8%), *S. Derby* (2.9%), *S. Rissen* 6,7: f,g: [-](2.9%), *S.*

210 *Typhimurium* 4,12:i: 1,2 (1.4%) and *S. Kedougou* (1.4%).

211

212 **3.2. Genetic characterisation of *Salmonella* isolates**

213

214 All isolates were first screened by ERIC-PCR and subsets of 46 were selected according to
215 their different profiles and origin for further analysis by PFGE. Thus, a total of 16 isolates
216 from vultures, 5 from pooled faecal samples from the farms, 7 from the warehouse of
217 cadavers, 12 from the trucks that transport the carcasses from the warehouse to the SFS and
218 6 from carcasses deposited in SFS were examined by PFGE.

219 Monophasic *S. Typhimurium* 1,4,[5],12:i:- isolates were identical (>90% genetic homology)
220 and belonged to samples from vultures and pig faeces from farms and carcasses disposed of
221 in SFS (Fig. 2). In addition, *S. Derby* isolates were identical (>90% genetic homology) and
222 belonged to samples from vultures and pig faeces from farms (Fig. 2). *Salmonella* Rissen
223 isolates were identical (>85% genetic homology) and belonged to samples from vultures and
224 pig faeces from farms (Fig. 2). Finally, *Salmonella* Kedougou isolates were identical (>90%
225 genetic homology) and belonged to samples from vultures and carcasses disposed of in SFS
226 (Fig. 2).

227

228 **4. Discussion**

229

230 This study demonstrated that free-living vulture strains (some monophasic *S. Typhimurium*
231 1,4,[5],12:i:-, *S. Derby* and *S. Rissen*) displayed genomic DNA fingerprinting patterns similar
232 to those observed in *Salmonella* strains from pig farms, suggesting that pig farms would
233 introduce *Salmonella* infection into vultures at SFS. This is further supported by the

234 hypothesis proposed by several authors (Millán et al., 2004; Marin et al., 2014; Vela et al.,
235 2015; Blanco et al., 2016; Blanco, 2018). Besides, *Salmonella* serovars isolated in vultures
236 in the current study have frequently been recorded in vultures in Spain (Marin et al., 2014;
237 Blanco, 2018), and are also often seen in modern pig production (EFSA, 2017). Interestingly,
238 Blanco (2018) not only found similar *Salmonella* serotypes between faeces of vultures
239 feeding on pig carcasses and the pig carcasses, but also identified similar antimicrobial
240 multiresistant patterns between these serotypes. However, no studies have based the results
241 on a molecular identification of DNA polymorphisms to differentiate strains and accurately
242 trace their diffusion. Today, the PFGE system is considered the gold standard for use in
243 epidemiological studies of *Salmonella* (Zou et al., 2013). Thus, monophasic *S. Typhimurium*
244 1,4,[5],12:i:-, *S. Derby* and *S. Rissen* strains isolated in free-living vultures and pig farms
245 authorised to provide carcasses at SFS were highly genetically homogeneous (similar DNA
246 fingerprint). This supports the idea that cross-infection and contamination occurs between
247 pig farms and free-living vultures. Additionally, monophasic *S. Typhimurium* and *S. Derby*
248 are included in the top five most commonly reported serovars in human salmonellosis cases
249 acquired in EU during 2016 (EFSA, 2017). This highlights the role of SFS in the potentiation
250 of griffon vultures as reservoirs, amplifiers and disseminators of *Salmonella*, but also for
251 conservation and reintroduction of avian scavengers, as it remains unknown today whether
252 *Salmonella* can cause clinical illness in this species (Blanco, 2018). Indeed, several studies
253 have reported on the role of pig farms in *Salmonella* transmission among wild birds (Andrés
254 et al., 2013; Andrés-Barranco et al., 2014). Furthermore, different pathways whereby wildlife
255 can be involved in human salmonellosis have been documented (Hilbert et al., 2012).

256

257 *Salmonella* occurrence in the current study doubled that of previous studies, where the
258 *Salmonella*-positive rate was lower than 10% in captive scavengers (Millán et al., 2004;
259 Molina-López et al., 2011:2015; Jurado-Tarifa et al., 2016), but was reduced compared to a
260 recent study carried out in central Spain on free-living scavengers, where 61.0% of griffon
261 vultures were *Salmonella*-positive (Blanco, 2018). Strikingly, our previous study carried out
262 in the same observatory and with a similarly large number of samples showed a high level of
263 the bacterium in comparison with the current study (Marin et al., 2014). In this context,
264 *Salmonella* determination is challenging due to intermittent day-to-day shedding and within-
265 day shedding by particular individuals, which could explain the slight differences in
266 occurrence among experiments (Tizard, 2004; Daoust and Prescott, 2007). Nevertheless,
267 cloacal swab is the preferable method to determine the identity of each individual host and
268 prevent cross-contamination by vectors, as well as environmental factors. In spite of this
269 particular point, some *Salmonella* serovars, such as *S. Typhimurium*, monophasic *S.*
270 *Typhimurium* 1,4,[5],12:i:- and *S. Derby*, have frequently been recorded in vultures
271 throughout different regions of Spain (Millán et al., 2004; Molina-López et al., 2011:2015;
272 Marin et al., 2014; Jurado-Tarifa et al., 2016; Blanco, 2018). In this scenario, one might
273 suggest that our results do not seem to be specific to our area of study. Nevertheless, further
274 research is required to assess the contribution of pig production as a primary source of
275 *Salmonella* in scavenger infection in SFS compared with zoonotic agents in other
276 geographical areas. In fact, this situation should not be considered exclusive to swine
277 production, as poultry and beef production have recently been implicated in large outbreaks
278 of multi-drug-resistant *Salmonella* both in Europe and North America (Mindlin et al., 2013;
279 Laufer et al., 2015; CDC, 2016).

280 In Spain, carcasses generally disposed of in SFS often come from intensive livestock farming
281 with pigs (Camiña and Montelío, 2006; Blanco, 2018), mainly because Spain is the second
282 largest swine producer in the EU and fourth worldwide (Marquer et al., 2014). In particular,
283 7 out of 11 farms analysed in the current study were *Salmonella*-positive, where monophasic
284 *S. Typhimurium* 1,4,[5],12:i:- was isolated in 5 of them. Currently, monophasic variants of
285 *S. Typhimurium* (1,4,[5],12:i- and 1,4,12:i-) have emerged as a public health threat, as it is
286 the third most frequently isolated serovar from human cases of salmonellosis in Europe,
287 representing 8.3% of confirmed human cases in 2015 (Andres and Davies, 2015).
288 Monophasic *S. Typhimurium* constitutes a high proportion of the multi-drug-resistant
289 *Salmonella* isolates and its occurrence in pigs has been increasing since 2010 (Andres and
290 Davies, 2015). The worldwide spread of monophasic *S. Typhimurium* 1,4,[5],12:i:- in swine
291 populations is likely related to the selective advantage offered by multi-drug-resistant
292 profiles associated with stable genetic elements, also carrying virulence features. These
293 bacterial lineages are well adapted to the porcine host and are prevalent in human infections
294 as a result of contaminated pig meat (EMA, 2017). In Spain, monophasic *S. Typhimurium*
295 serovar accounted for 31.3% of the isolates from pigs in 2015 (Andres and Davies, 2015).
296 Matching with this, a more recent study found that several serotypes isolated from egyptian
297 and griffon vultures faeces at an SFS presented a resistance pattern simultaneously resistant
298 to aminopenicillins, aminoglycosides and tetracyclines, including *S. Typhimurium* 4,12:i:1,2
299 and the monophasic *S. Typhimurium* 4,12:i:- serotypes (Blanco, 2018). This observation is
300 in line with other studies, which have associated pig carcasses with avian scavenger
301 contamination with veterinary pharmaceuticals and the creation of new resistances and the
302 amplification of these acquired pathogens (Blanco et al., 2016; 2017a; Blanco, 2018). In
303 addition, different studies have highlighted the potential impact of pig carcasses disposed in

304 the SFSs on development of fungal and parasitic infections in wild avian scavengers (Blanco
305 et al., 2017b; 2017c; Pitarch et al., 2017), although conceptually food security and food safety
306 can potentially be better assured in the SFSs (Margalida et al., 2014). To circumvent this
307 problem, in France conservationists, vets and stakeholders promoted the development of
308 individual SFS, with the principle that each farmer directly recycles their carcasses at their
309 own SFS, avoiding carcass displacement and limiting potential dissemination of pathogens,
310 and furthermore providing carcasses spread more spatially for vultures, in a more natural way
311 (Dupont et al., 2012). In this context, it is worth noting that the *Salmonella* status of the
312 facilities in close contact with griffon vultures during supplementary feeding in this study
313 clearly demonstrated that both the trucks that transport the carcasses from the warehouse to
314 the SFS and the warehouse of cadavers could be an important source of cross-contamination
315 (Dorr et al., 2009). As a measure for practical implementation, if each farm directly recycles
316 its carcasses at its own SFS, authorities should be taking into account sanitary assurances that
317 these farms are pathogen-free. In Spain, some Regional Governments have restricted the
318 supply of feeding stations with swine carcasses (Blanco et al., 2018). The repercussions of
319 this change on avian scavenger populations should be evaluated.

320 In conclusion, the current study indicates that free-living griffon vultures and pig farms that
321 provide the carcasses at SFS share several zoonotic *Salmonella* strains based upon their DNA
322 fingerprint, including monophasic *S. Typhimurium* and *S. Derby*. Taken together with
323 previous studies and although transmission and cross-infection could be bidirectional, our
324 result seems to corroborate the pig carcasses-to-vulture transmission and cross-infection at
325 SFS. However, the current study contains some important biases and limitations. Our results
326 were located at only one SFS. In addition, bidirectional transmission of *Salmonella* has not

327 been evaluated. Under this scenario, there is an urgent need to avoid infection risk and
328 prevent the spread of *Salmonella*, but also to find new strategies to keep the feeding stations
329 as a useful tool for scavenger conservation and assess the potential role of these wild fauna
330 in *Salmonella* epidemiology. Nowadays, initiatives promoting low-intensity farming
331 practices and the use of carcasses from free-ranging ruminants left in the countryside for
332 scavenger consumption are being proposed (Blanco, 2018). As an immediate *Salmonella*
333 control strategy in wild avian scavengers, we suggest the implementation of a programme to
334 ensure that only pig carcasses from *Salmonella*-free farms arrive at SFS. Moreover, we
335 emphasise the need for continuous local surveillance programmes to identify the potential
336 risk to wildlife and the environment.

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484 **Figure legends**

485

486 **Fig 1.** Experimental scheme design to evaluate whether free-living griffon vultures captured
487 during the observatory's normal ringing programme and pig samples from farms authorised
488 to provided carcasses share zoonotic *Salmonella* strains. Thus, the occurrence, serotypes and
489 genomic DNA fingerprinting (phage typing and PFGE) of *Salmonella* spp. strains isolated at
490 different sampling points: (A) pig faeces on farms; (B) warehouse of cadavers where farmers
491 legally dispose of dead animals; (C) trucks that transport the carcasses from the warehouse
492 to the supplementary feeding station; and (D) pig carcasses disposed in the supplementary
493 feeding station and griffon vultures in Cincorres observatory located in Castellón province
494 (eastern Spain).

495

496 **Fig 2.** Dendrogram showing the XbaI profiles of *Salmonella* spp. strains identified from free-
497 living vultures, pig faeces samples from farms authorised to provided carcasses at SFS and
498 several discrete samples obtained from facilities in close contact with pig carcasses during
499 supplementary feeding (warehouse of cadavers where farmers legally dispose of dead
500 animals, trucks that transport the carcasses from the warehouse to the SFS and pig carcasses
501 at SFS) in Cincorres observatory, located in Castellón province (eastern Spain).