

1 Genome-Wide Fitness Analyses of the Foodborne Pathogen

2 *Campylobacter jejuni* in *In Vitro* and *In Vivo* Models

3 Stefan P.W. de Vries^{1§}, Srishti Gupta^{1§}, Abiyad Baig^{1a}, Elli Wright², Amy Wedley²,
4 Annette Nygaard Jensen³, Lizeth LaCharme Lora², Suzanne Humphrey^{2c}, Henrik
5 Skovgård⁴, Kareen Macleod⁵, Elsa Pont¹, Dominika P Wolanska¹, Joanna
6 L'Heureux¹, Fredrick M. Mobegi^{6b}, David G.E. Smith⁷, Paul Everest⁵, Aldert Zomer⁸,
7 Nicola Williams⁹, Paul Wigley^{2,3}, Thomas Humphrey¹⁰, Duncan J Maskell¹, Andrew J
8 Grant^{1*}

9 ¹ Department of Veterinary Medicine, University of Cambridge, Cambridge, United
10 Kingdom.

11 ² Department of Infection Biology, Institute of Infection and Global Health, University
12 of Liverpool, Leahurst Campus, Neston, United Kingdom.

13 ³ Technical University of Denmark, National Food Institute, Copenhagen, Denmark

14 ⁴ Department of Agroecology, University of Aarhus, Slagelse, Denmark.

15 ⁵ University of Glasgow, Veterinary School, Glasgow, United Kingdom.

16 ⁶ Department of Paediatric Infectious Diseases, Radboud Institute for Molecular Life
17 Sciences, Radboud University Medical Centre, Nijmegen, the Netherlands

18 ⁷ Heriot-Watt University, School of Life Sciences, Edinburgh, Scotland, United
19 Kingdom.

20 ⁸ Department of Infectious Diseases and Immunology, Faculty of Veterinary
21 Medicine, Utrecht University, Utrecht, The Netherlands.

22 ⁹ Department of Epidemiology and Population Health, Institute of Infection and Global
23 Health, University of Liverpool, Leahurst Campus, Neston, United Kingdom.

24 ¹⁰ School of Medicine, Institute of Life Sciences, Swansea University, Swansea,
25 United Kingdom.

26 ^a Current address: School of Veterinary Medicine and Science, University of
27 Nottingham, Sutton Bonnington, Leicestershire, United Kingdom.

28 ^b Current address: Division of Molecular Carcinogenesis, The Netherlands Cancer
29 Institute, Amsterdam, The Netherlands.

30 ^c Current address: Institute of Infection, Immunity and Inflammation, University of
31 Glasgow, Glasgow, United Kingdom

32 * To whom correspondence should be addressed: E-mail ajg60@cam.ac.uk

33 § These authors contributed equally to this work.

34 **Short title:** *Campylobacter jejuni* Fitness Analyses

35

36 Abstract

37 Infection by *Campylobacter* is recognised as the most common cause of foodborne
38 bacterial illness worldwide. Faecal contamination of meat, especially chicken, during
39 processing represents a key route of transmission to humans. There is currently no
40 licenced vaccine and no *Campylobacter*-resistant chickens. In addition, preventative
41 measures aimed at reducing environmental contamination and exposure of chickens
42 to *Campylobacter jejuni* (biosecurity) have been ineffective. There is much interest in
43 the factors/mechanisms that drive *C. jejuni* colonisation and infection of animals, and
44 survival in the environment. It is anticipated that understanding these mechanisms
45 will guide the development of effective intervention strategies to reduce the burden of
46 *C. jejuni* infection. Here we present a comprehensive analysis of *C. jejuni* fitness
47 during growth and survival within and outside hosts. A comparative analysis of
48 transposon (Tn) gene inactivation libraries in three *C. jejuni* strains by Tn-seq

49 demonstrated that a large proportion, 331 genes, of the *C. jejuni* genome is
50 dedicated to (*in vitro*) growth. An extensive Tn library in *C. jejuni* M1cam (~10,000
51 mutants) was screened for the colonisation of commercial broiler chickens, survival in
52 houseflies and under nutrient-rich and –poor conditions at low temperature, and
53 infection of human gut epithelial cells. We report *C. jejuni* factors essential throughout
54 its life cycle and we have identified genes that fulfil important roles across multiple
55 conditions, including *maf3*, *fliW*, *fliD*, *pflB* and *capM*, as well as novel genes uniquely
56 implicated in survival outside hosts. Taking a comprehensive screening approach
57 has confirmed previous studies, that the flagella are central to the ability of *C. jejuni*
58 to interact with its hosts. Future efforts should focus on how to exploit this knowledge
59 to effectively control infections caused by *C. jejuni*.

60

61 Author Summary

62 *Campylobacter jejuni* is the leading bacterial cause of human diarrhoeal disease. *C.*
63 *jejuni* encounters and has to overcome a wide range of “stress” conditions whilst
64 passing through the gastrointestinal tract of humans and other animals, during
65 processing of food products, on/in food and in the environment. We have taken a
66 comprehensive approach to understand the basis of *C. jejuni* growth and
67 within/outside host survival, with the aim to inform future development of intervention
68 strategies. Using a genome-wide transposon gene inactivation approach we
69 identified genes core to the growth of *C. jejuni*. We also determined genes that were
70 required during the colonisation of chickens, survival in the housefly and under
71 nutrient-rich and –poor conditions at low temperature, and during interaction with
72 human gut epithelial tissue culture cells. This study provides a comprehensive
73 dataset linking *C. jejuni* genes to growth and survival in models relevant to its life

74 cycle. Genes important across multiple models were identified as well as genes only
75 required under specific conditions. We identified that a large proportion of the *C.*
76 *jejuni* genome is dedicated to growth and that the flagella fulfil a prominent role in the
77 interaction with hosts. Our data will aid development of effective control strategies.
78

79 Introduction

80 Infection by *Campylobacter* is the most common cause of foodborne bacterial
81 diarrhoeal disease worldwide, responsible for ~96 million foodborne illnesses and
82 ~21,000 foodborne deaths in 2010 [1]. While most cases are self-limiting, for some,
83 campylobacteriosis is a particularly serious infection, and it is also associated with
84 severe post-infection complications, including irritable bowel and Guillian-Barré
85 syndromes. Consumption of undercooked poultry, unpasteurised dairy products and
86 contaminated water represent the most common sources of human infection [2, 3].
87 *Campylobacter jejuni* has a broad range of environmental reservoirs that include
88 water, birds and other domestic animals [3]. In addition, flies have been implicated as
89 a transmission vector for *C. jejuni* for both chicken flocks and possibly also for
90 humans as well [4-7].

91 *C. jejuni* encounters and has to overcome various stress conditions whilst
92 passing through the gastrointestinal tract of humans and other animals, during
93 processing of food products (e.g. slaughter process of poultry), on/in food (e.g. in
94 milk or poultry meat, generally stored at low temperature) and in the environment
95 (e.g. in surface water, soil, or in flies, the latter representing a transmission vector [4-
96 7]) [8]. However, compared to other enteric pathogens such as pathogenic
97 *Escherichia coli* and *Salmonella* spp, the survival mechanisms used by *C. jejuni* to
98 cope with these stresses are less well-understood [9].

99 Based on genome analysis, the capacity of *C. jejuni* to survive outside the
100 host and adapt to environmental stress conditions appears to be limited due to the
101 lack of key stress regulators found in other enteric pathogens [10], however survival
102 at low temperature in water has been reported for up to four months [9].
103 Establishment of colonisation and infection of host organisms by *C. jejuni* is a

104 multifactorial process with key roles for “swimming” motility, chemotaxis, interaction
105 with gut epithelial cells, toxin production, and adaptation to oxidative and metabolic
106 stress [11, 12]. Factors involved in these key processes include the flagella, capsule,
107 glycosylation systems, and two-component regulatory systems. These have all been
108 identified as being important for chicken colonisation, human infection and
109 environmental survival [9, 13-15].

110 Despite considerable research in the field, this has not led to the development
111 and/or implementation of effective control strategies. Here, we describe the
112 generation of extensive *C. jejuni* transposon (Tn) gene inactivation mutant libraries,
113 and their use to comprehensively assess which genes contribute to bacterial fitness
114 during *in vitro* growth, colonisation of commercial broiler chickens, survival in the
115 housefly, and survival during exposure to low temperature in nutrient-rich and –poor
116 conditions, and infection of human gut epithelial cells. This study reinforces the
117 importance of flagella for host interactions, and identifies genes required for survival,
118 colonisation and infection in multiple phases of the bacterium’s life cycle. Some of
119 these may represent an Achilles’ heel of the pathogen and be a possible target for
120 novel intervention strategies.

121

122 Results

123 Identification of Genes Required for Fitness in *C. jejuni*

124 To assess the genetic basis of *C. jejuni* growth and survival, genes were
125 randomly inactivated using Tn mutagenesis in three well-characterized *C. jejuni*
126 strains [M1cam [16, 17], NCTC 11168 (hereafter referred to as 11168) [10] and 81-
127 176 [18]]. Tn mutant libraries were characterised by Tn insertion site sequencing (Tn-
128 seq [19]) (Table S1), providing a measure for the relative abundance of each Tn
129 mutant in the library. Genes that are required for growth and survival, hereafter
130 referred to as “fitness” genes, cannot tolerate Tn insertions, or Tn mutants in these
131 genes are severely underrepresented in the libraries. To identify fitness genes,
132 23,334 unique chromosomal Tn insertions were analysed in M1cam, 15,008 in
133 11168, and 17,827 in 81-176 (Table S1), reaching near-saturation in terms of the
134 number of genes that could be inactivated (Fig 1a). In addition to chromosomal Tn
135 insertions, 2,009 and 1,919 unique insertions were in the 81-176 plasmids pVir [20]
136 and pTet [21], respectively (Table S1). No apparent Tn insertion bias was observed
137 (Fig S1) and each Tn library predominately yielded unique Tn insertions (Fig S2).

138

139 **Fig 1. Gene fitness analysis during *in vitro* growth of *C. jejuni* M1cam, 11168**
140 **and 81-176.** (A) Rarefaction analysis of intragenic Tn insertions. (B) Density plot
141 fitness score (Log_2 fold-change measured reads/expected reads) per gene. Dots
142 indicate fitness score cut-off values. (C) Overlap of genes required for fitness in *C.*
143 *jejuni* M1cam, 11168, and 81-176. (D) Functional class (COG; Cluster of Orthologous
144 Genes) enrichment analysis of fitness genes. Fisher exact test with Q-value multiple
145 testing correction; * $Q < 0.05$, ** $Q < 0.01$ and *** $Q < 0.001$.

146 Gene fitness score (Log_2 fold-change between the observed vs expected of
147 sequence reads [22]) density plots followed a bimodal distribution with the “left”
148 population representing genes required for *in vitro* growth and survival (Fig 1b). In
149 total, 445 genes were required for fitness in M1cam, 413 genes in 81-176 and 499 in
150 11168 (Table S2). Interestingly, the pTet plasmid in 81-176 harboured a single gene
151 of unknown function (*cpp13*) that appeared to contribute to fitness. A variant of *C.*
152 *jejuni* 81-176 that has lost pTet exists (D. Hendrixson, personal communication),
153 which implies that *cpp13* is not obligate essential but may contribute to fitness, or it
154 could be antitoxin of an uncharacterised toxin-antitoxin system. Unexpectedly, Tn
155 insertions were observed in *dnaA* (1,323 bp) at bp position 352 in M1cam and at
156 position 1,115 in 81-176. Consequently, *dnaA* did not pass the stringent fitness gene
157 criteria. Disruption of *dnaA* may be tolerated at the 3' end of the gene, in the rare
158 event of a secondary site mutation [23], or due to the existence of merodiploids [24].

159 With the aim of providing a comprehensive analysis of fitness genes in the
160 species *C. jejuni*, homologous genes were compared for the three *C. jejuni* strains
161 (Table S2). Out of the 1,424 identified homologous gene clusters, 331 genes were
162 required for fitness in all three strains tested and 486 genes were required in two or
163 more strains (Fig 1c). We found that 845 gene clusters were not required for fitness
164 in any of the analysed strains. Genes implicated in fitness were relatively dispersed
165 across the *C. jejuni* genomes, but there were regions that were (almost) devoid of
166 fitness genes, for example, the flagellar glycosylation gene cluster (Fig S3). The high
167 percentage of genes required for fitness in *C. jejuni* is likely to be the consequence of
168 its relative “minimal” genome and its proportionally large core genome [25] as well as
169 the large number of transcriptionally coupled genes [26, 27].

170

171 The number of fitness genes shared by *C. jejuni* M1cam, 81-176 and 11168
172 was substantially larger, at 331, than the 175-233 genes previously reported to be
173 obligately essential [28-31]. This may be partially due to the inclusion of genes, in our
174 study, whose inactivation is lethal (obligate essential) as well as genes which when
175 inactivated by a Tn result in severely compromised growth and/or survival. Further,
176 this could be related to *C. jejuni* strain differences, growth conditions, the Tn element
177 used, the number of Tn mutants analysed and the read-out technology. To assess
178 the “validity” of our analysis, a systematic review of the literature on *C. jejuni* 11168
179 defined gene deletion and Tn mutants was performed. This analysis revealed that for
180 38 out of 486 (7.8%) fitness genes (required in two or more strains) identified in this
181 study, mutants have been reported in *C. jejuni* 11168, indicating a low false-positive
182 rate in our study. Of importance, although mutants in some of these genes could be
183 generated, these may still result in a growth defect, *e.g.* as reported for a *C. jejuni*
184 11168 *pycB* mutant [32], especially if the mutant is compared against other ‘more-fit’
185 mutants or mutants with wild-type fitness, as is the case in the Tn screen. Our
186 comprehensive fitness analysis indicated that a large part of the *C. jejuni* genome is
187 dedicated to growth and survival under the condition tested, and implies that there
188 could be opportunities for targeting some of these genes for novel intervention
189 strategies, *e.g.* as previously reported by Mobegi *et al* [33].

190 Cluster of Orthologous Genes (COG) enrichment analysis showed that genes
191 implicated in translation, lipid metabolism and cell cycle control were overrepresented
192 amongst the genes required for fitness during *in vitro* growth/survival in all three
193 tested *C. jejuni* strains. Extending the analysis to fitness genes required in two or
194 more strains also showed overrepresentation of coenzyme metabolism, nucleotide
195 metabolism and intracellular trafficking and secretion genes (Fig 1d). Required for

196 fitness were, amongst others, genes implicated in replication, transcription,
197 translation (40 out of 49 ribosomal protein genes), purine and pyrimidine metabolism,
198 energy metabolism (ATP and NAD synthase, NADH-quinone oxidoreductase),
199 isoprene biosynthesis, protein secretion (Sec and Tat pathway), as well as genes
200 involved in cofactor biosynthesis (thiamine, folic acid and heme) and oxidative stress
201 (see [Table S2](#) for a complete overview). Further, the complete gluconeogenic
202 pathway was found to be required for fitness whereas the majority of the enzymes of
203 the tricarboxylic acid (TCA) cycle were not, except for aconitase (*acnB*), probably
204 reflecting flexibility in this part of the bacterium's metabolism.

205 *C. jejuni* expresses surface structures such as flagella, lipooligosaccharide
206 (LOS) and capsular polysaccharides (CPS). Inactivation of genes in these pathways
207 severely attenuated fitness. This included flagellar basal body rod proteins (encoded
208 by *flgAC* and *fliEL*), the MS-ring (*fliF*) and C-ring (*fliG*) as well as components of the
209 flagellar type III secretion system (*fliQH*). Genes required for formation of the LOS
210 lipid A molecule (*lpxABCDL*), KDO (*kdsAB*) and the first L-glycero-D-manno-heptose
211 residue (*waaC*) were required for fitness, whereas the remainder of the genes
212 responsible for the core oligosaccharide were not. Of the capsular biosynthesis gene
213 cluster only the inactivation of the last two genes (*kpsDF*) resulted in impaired fitness.
214 Genes required for cell envelope generation were also important for fitness including
215 fatty acid biosynthesis genes (*accABCD* and *fabDFGHLZ*), peptidoglycan (*dapADEF*,
216 *ddl*, *murABCDEFGF*, *pbpABC*), and the rod-shape determining protein genes
217 (*mreBCD*). Protein glycosylation is tightly linked to virulence, of which the N-linked
218 protein glycosylation pathway genes *pglACD* were required for fitness. This is in
219 contrast to flagella glycosylation genes, which had no conserved role in fitness during
220 *in vitro* growth and survival ([Table S1](#)).

221 Quantitative Analysis of Genes Implicated in the Life Cycle of *C.*

222 *jejuni*

223 The same extensive Tn library in strain M1cam ([Table S1](#); 9,951 unique Tn
224 insertions with 1,124 genes harbouring Tn insertions) was screened in various *in vivo*
225 and *in vitro* models as a proxy for some of the conditions that *C. jejuni* might
226 encounter during its life cycle from chicken-to-human. Screening the same Tn mutant
227 library through all of the different models facilitated a comparative analysis across the
228 models. The M1cam library was screened during the colonisation of commercial
229 broiler chickens (natural host), survival in the housefly (transmission vector), survival
230 under nutrient-rich- and nutrient-poor conditions at low temperature, and in models
231 that mimic (stages of) infection of humans, *i.e.* adhesion and invasion of human gut
232 epithelial cells. For comparative purposes we included data that we obtained in a
233 study analysing the infection of gnotobiotic piglets (de Vries *et al.*, submitted). Tn-seq
234 analysis after exposure to a challenge, compared with the control condition, provided
235 a quantitative measure for the contribution of a gene to fitness in each of the models
236 ([Fig 2](#) and [Table S3](#) and [S4](#)).

237

238 **Fig 2. Identification of conditional essential genes in *C. jejuni*.** (A) Effect of Tn
239 insertions on the ability of *C. jejuni* M1cam to colonize commercial broiler chickens,
240 infect gnotobiotic piglets, adhere and invade gut epithelial tissue culture cells, survive
241 in flies and at 4°C in various media (chicken juice, BHI, tap water, rain water, and
242 sterile water). Genes of which Tn mutants showed significantly attenuated or
243 enriched fitness in the experimental models are shown and are grouped according to
244 their COG functional classification. Data represented as Log₂ fold-change is also

245 presented in [Table S4](#). (B) Gene-model interaction network showing attenuated
246 (orange lines) and enriched (blue lines) Tn-seq scores linked to their respective
247 models; the thickness of the connecting lines corresponds to the Log₂ fold-change
248 (input/output). The gene numbers correspond to the *C. jejuni* M1cam locus-tags [16]
249 and are color-coded according to their COG functional class. The size of the genes
250 and the models displayed increases with the number of interactions.

251 The number of Tn mutants recovered from each model confirmed that the
252 complexity of the Tn library was maintained in survival in the housefly, under nutrient-
253 rich and –poor conditions at 4°C and during adhesion and invasion of gut epithelial
254 cells ([Fig S4](#)). The Tn mutant library complexity was reduced after colonisation of
255 chickens, *i.e.* 23% of the input Tns were recovered from chickens ([Fig S4](#)), which is
256 likely due to the existence of a population bottleneck ([Fig S4](#)). Therefore, we applied
257 more stringent criteria for selection of candidate genes required in this model ([Table](#)
258 [S3](#) and Materials and Methods). For comparison, 72% of the Tn's were recovered
259 after infection of gnotobiotic piglets ([Fig S4](#)) (de Vries *et al.*, submitted). A detailed
260 analysis for each of the models in this study is provided below.

261

262 Genes Required for Colonisation of Commercial Broiler 263 Chickens

264 Broilers in general become colonised with *Campylobacter* spp. at about 3-4
265 weeks of age [34]. We have screened the M1cam Tn library 'C' in a relevant chicken
266 colonisation model, *i.e.* in 3-week-old Ross 308 commercial broiler birds. Four cages,
267 each with 5-7 birds, were inoculated with the M1cam Tn library and 6 days post-
268 inoculation (p.i.) colonising Tn mutants were recovered. Due to the coprophagic

269 behaviour of chickens, each individual cage was considered a colonisation unit [35].
270 We hypothesised that a group-level approach would improve the robustness of the
271 analysis and reduce any bias introduced by random dropout of Tn mutants as
272 observed in our previous work using a signature-tagged mutagenesis approach [36]
273 and using wild-type isogenic-tagged strains (WITS) that have indistinguishable
274 phenotypes in pure culture [37]. Both these approaches indicated the highly complex
275 and dynamic nature of chicken colonisation by *C. jejuni*.

276 On average, cages harboured $1,641 \pm 226$ Tn mutants (> 10 reads) compared
277 to $7,325 \pm 538$ Tns in the input (Fig S4), revealing the existence of a population
278 bottleneck, even at the cage level. However, the average read count of the four
279 cages demonstrated recovery of 3,111 Tn mutants (> 10 reads) covering 701 genes.
280 This indicated that, despite a population bottleneck, a large part of the genome could,
281 with additional stringent filtering steps (see Materials and Methods), be analysed for
282 its role during colonisation of chickens.

283 Tn mutants of 172 genes were significantly under-represented in colonised
284 chickens, whereas 24 appeared to enhance fitness during colonisation (Fig 2 and
285 Table S3 and S4). Functional class enrichment analysis revealed that genes linked to
286 the COG class “cell motility” were significantly enriched amongst those required
287 during colonisation (Fig 2 and Fig S5), underscoring the prominent role of motility for
288 colonisation of chickens.

289 Previously, work by Johnson *et al.*, reported a Tn library screen in a 1-day-old
290 chicken colonisation model [29]. Screening 1,155 *C. jejuni* 81-176 Tn mutants
291 identified 130 genes that were required for colonisation and 30 genes that appeared
292 to enhance colonisation [29]. Interpretation of the importance of the colonisation
293 genes identified by Johnson *et al.*, is complicated due to limited validation of the Tn

294 screen; only the importance of a single candidate gene, *mapA*, was tested and
295 confirmed [29]. We found a limited overlap of 23 chicken colonisation genes between
296 the two studies, 11 of which were linked to motility and the flagellar system. The
297 datasets had one gene (CJM1_0420; hypothetical protein) in common that was
298 beneficial for colonisation. Differences between identified colonisation genes are
299 likely the result of the experimental model employed, e.g. older birds harbouring a
300 more mature intestinal microbial community and younger birds being more
301 permissive for colonisation by *C. jejuni*, e.g. as reflected in a lower inoculation dose
302 to establish colonisation [38].

303 For validation of the chicken colonisation Tn library screen, defined deletion
304 mutants in 19 genes were tested individually for their ability to colonise chickens. The
305 importance in chicken colonisation was confirmed for genes involved in chemotaxis
306 (*mcp4_1*), the flagellar system (*pflB*, *fliD*, *fliW*, and *maf3*), N-linked protein
307 glycosylation system (*capM*, also referred to as *pglH*), and phosphate transport
308 (*pstA*) (Fig 3a). Although motility is considered a driving factor in chicken
309 colonisation, we have found that mutants in *maf3*, *capM* and *pstA*, that have a defect
310 in colonisation, displayed wild-type motility (Fig S6). In the Tn library screen, mutants
311 in *eptA* (also referred to as *eptC* [39]), *glnP*, *jlpA*, *fdhA* and CJM1cam_1125
312 (uncharacterised glycosyltransferase) showed reduced colonisation. However,
313 validation with defined deletion mutants revealed slightly higher bacterial loads in
314 chickens when compared to the wild-type (Fig 3a); motility of these defined gene
315 deletion mutants did not differ significantly from the wild-type (Fig S6). In contrast to
316 our findings, previous work by Cullen *et al.*, reported a slight reduction in motility after
317 deletion of *eptC* in *C. jejuni* 81-176. EptC catalyses the phosphoethanolamine
318 modification of FlgG and lipid A, [39], suggesting that the importance of *eptA* could

319 be strain dependent. No difference to the wild-type was found in the colonisation
320 proficiency of defined mutants in *moaA*, CJM1cam_0303 (hypothetical protein),
321 CJM1cam_0438 (hypothetical protein), *flaG*, *ilvB*, *engD* and *gltA*. Identification of
322 these genes as attenuated in the Tn library screen might be attributed to the
323 competition effect of Tn mutants in other genes being present during the Tn library
324 screen, whilst validation experiments were conducted with single mutant inocula.
325 Here we opted for validation experiments with single mutant inocula due to the
326 unpredictable outcome of competition experiments, even in the simplest case of two
327 phenotypically indistinguishable (in pure culture) WITS [37]. It is possible that some
328 candidate colonisation genes represent false positives due to the random loss of
329 mutants, *i.e.* an infection bottleneck that was observed in the chicken model.

330

331 **Fig 3. Validation of chicken colonisation Tn mutant library screen.** (A)
332 Colonisation levels of *C. jejuni* M1cam gene deletion mutants 6 d pi. ($n \geq 7$) (B)
333 Colonisation of gene deletion mutants and complemented gene deletion mutants ($n \geq$
334 7). (C) Colonisation kinetics of *C. jejuni* M1cam wild-type and defined mutants in *fliW*
335 and *capM* ($n \geq 9$). Chickens being colonised vs not colonised were 2/8, 2/8 and 3/6
336 for the *fliW* mutant and 4/5, 4/6 and 5/5 for the *capM* mutant at day 3, 6 and 10 p.i.,
337 respectively. Statistical significance was analysed using a Mann-Whitney test with * P
338 < 0.05 , ** $P < 0.01$ and *** $P < 0.001$. The orange arrows represent an attenuated
339 phenotype while the blue arrows represent an enriched chicken colonisation
340 phenotype.

341

342 Genetic complementation of mutants in *pstA*, *capM* and *maf3* did significantly
343 increase their colonisation capacity compared to their respective gene deletion

344 mutants (Fig 3b), confirming their role in chicken colonisation. Deletion of *fliD*,
345 encoding the flagellar cap protein, resulted in severely attenuated chicken
346 colonisation (Fig 3a). However, attempts to complement the *fliD* deletion failed (see
347 Fig S8 for details). When analysing colonisation of the genetically complemented
348 *mcp4_1* mutant in chickens, we found that, in contrast to initial validation experiments
349 (Fig 3a), the *mcp4_1* mutant was able to colonise chickens; which could be the result
350 of intrinsically lower colonisation resistance in the batch of chickens used in this
351 experiment.

352 Despite the existence of a colonisation bottleneck and the incomplete
353 reproducibility when comparing defined deletion mutants with the results obtained in
354 the Tn library screen, this work has identified *pstA*, *capM*, *maf3* and *mcp4_1* as novel
355 colonisation factors in 3-week-old broiler chickens. Previously, phosphate transport
356 system (*pstACS*) genes in *C. jejuni* 81-176 were shown to be expressed at higher
357 levels during colonisation of 1-day-old chickens compared to *in vitro* grown bacteria
358 [40], which can likely be attributed to low levels of phosphate in the chicken gut.
359 Deletion of the *N*-linked protein glycosylation gene *capM* in *C. jejuni* 81116 was
360 previously reported to reduce colonisation of chickens [41, 42]. The genomic location
361 of *maf3* (motility accessory factor) suggests a role in flagellar glycosylation [43]. In
362 addition to the role of *mcp4_1* in the colonisation of chickens identified in our study,
363 deletion of other methyl-accepting chemotaxis proteins (Mcp) also resulted in
364 reduced colonisation by *C. jejuni* of mice [44].

365 In validation experiments (Fig 3a) we observed an 'all or nothing' colonisation
366 effect in some chickens infected with particular defined mutants, *e.g.* *pflA*, *fliD* and
367 *fliW*, whereas for other deletion mutants, such as *capM* and *mcp4_1*, colonisation
368 levels varied considerably between the birds. This indicates that there might be

369 differences in the colonisation permissiveness/resistance between birds, within and
370 between experiments. To investigate this in more detail, the colonisation of the wild-
371 type, *fliW* and *capM* mutants were measured at different time intervals (3, 6 and 10
372 days p.i.) [Fig 3c](#)). Chickens were colonised at high levels from 3 days onwards and
373 neither an increase in the levels of colonisation nor the number of colonised birds
374 was observed. Surprisingly, the level of colonisation of the *fliW* and *capM* mutants did
375 not increase between 3 and 10 days p.i. and there was also no obvious increase in
376 the number of colonised chickens ([Fig 3c](#)). We hypothesise that the colonisation
377 responses observed in our validation experiments were potentially confounded by
378 variation in gut microbiota composition or differential inflammatory responses elicited
379 during colonisation [40, 45, 46].

380

381 Genes Required for Survival in the Housefly

382 As a transmission vector model for *C. jejuni* [4-7], survival of the M1cam Tn
383 library 'C' was examined 4 hours after individual inoculation of houseflies *via* their
384 proboscis. Tn mutants in 48 genes showed reduced survival and no genes were
385 identified for enhanced survival ([Fig 2](#) and [Table S3](#) and [S4](#)). Genes of the COG
386 class "nucleotide transport and metabolism" were overrepresented amongst the
387 genes linked to survival in the housefly ([Fig 2](#) and [Fig S5](#)). These were the non-
388 essential genes in the purine (*purLMN*) and pyrimide (*pyrC_1/2* and *pyrDF*)
389 biosynthesis pathways. Although the Tn library screen identified a number of
390 candidate survival genes, validation experiments with 7 defined gene deletion
391 mutants (inoculated as single mutants) were non-confirmative ([Fig 4a](#)). Our inability
392 to confirm the role of identified candidate genes might be the result of low levels of
393 attenuation, or be due to the lack of competition with other mutants. However, the

394 attenuated survival of a *capM* deletion mutant approached significance ($P = 0.0513$,
395 two-tailed Mann-Whitney) when compared to the wild-type. As the importance of
396 *capM* in chicken colonisation was confirmed *via* genetic complementation of the
397 deletion mutant, we also tested the *capM* mutant alongside its genetically
398 complemented mutant for survival in the housefly. This confirmed that *capM* is also
399 involved in survival in the housefly (Fig 4b).

400

401 **Fig 4. Validation of Tn mutant library screen during survival in the housefly.** (A)
402 Survival (Log_{10} decrease inoculum vs recovered) of defined *C. jejuni* M1 gene
403 deletion mutant in the housefly after 4 h. (B) Survival of wild-type, *capM* defined gene
404 deletion mutant and genetically complemented mutant. Data shown are Log_{10}
405 decrease of CFU per fly relative to the inoculum and plotted as means with SEM ($n \geq$
406 4). Significance was analysed using a Mann-Whitney test with * $P < 0.05$.

407

408 Survival in Nutrient-Rich and –Poor Conditions at Low 409 Temperature

410 To identify *C. jejuni* genes involved in survival in nutrient-rich and –poor
411 conditions at low temperature (4°C), the M1cam Tn library ‘C’ was incubated in
412 “chicken juice” (liquid obtained after thawing chicken carcasses), BHI broth and in
413 water (sterile, tap and rain), with the “water” models being included to assess the
414 survival under more general environmental conditions. The Tn mutants that survived
415 were recovered after 7 days incubation and compared to the Tn library composition
416 at $T = 0$ days. Ten genes were implicated in survival in chicken juice and 6 genes in
417 BHI medium. Considerable variation was found for the number of genes implicated in

418 survival at 4°C in nutrient-poor conditions, with 13 genes identified in tap water and
419 57 in sterile water. However, only one candidate gene, encoding the heat shock
420 response protein ClpB, was identified as being required for survival in rain water (Fig
421 2 and Table S3 and S4). We suspect that these variations could arise due to
422 differences in chemical composition, pH and potentially other bacteria present in
423 water samples [47, 48]. Previous work in *C. jejuni* 11168 showed that *clpB* is involved
424 in regulating a stress response to acidic pH while transiting the human stomach [49].
425 In addition, ClpB could facilitate the solubilisation and renaturing of aggregated
426 proteins at low temperature under which translation is repressed [50]. This could be
427 translated to our findings that ClpB may act as a stress response regulator and aid in
428 the survival of *C. jejuni* under conditions of stress. Given the relatively low number of
429 genes associated with survival at low temperature and the relatively mild attenuation,
430 as expressed by Tn-seq fitness score, we hypothesise that survival may be a passive
431 rather than active mechanism [51].

432 At low temperature, an oxidative stress response is induced in *C. jejuni* [9, 52].
433 In our Tn library screen, the gene encoding the oxidoreductase TrxC was found to be
434 required for survival in sterile water and BHI at low temperature. In addition, the
435 regulator of oxidative stress, PerR, was also linked to survival in sterile water. PerR
436 plays a role in controlling oxidative stress resistance and survival under aerobic
437 conditions [52, 53]. The RacRS two-component system is important for chicken
438 colonisation and is part of a temperature-dependent signalling pathway [54]. In our
439 study, Tn mutants in *racS* had reduced survival in chicken juice and tap water at low
440 temperature. Chemotaxis has also been suggested to play a role in survival at low
441 temperature [55]. In line with this, Tn mutants of *mcp4_2* were attenuated for survival
442 in chicken juice and tap water. In addition, Tn mutants in *kefB* and *czcD* (antiporters),

443 *fabI* (fatty acid metabolism) and CJM1cam_0181 to CJM1cam_0183 (*purN*, *nnr*, and
444 a gene encoding a hypothetical protein) were linked to survival in both chicken juice
445 and tap water.

446 For validation of the Tn-seq screen, 10 defined gene deletion mutants were
447 assayed under all conditions described above. Interestingly, survival of the wild-type
448 strain was significantly lower in chicken juice compared to any of the other conditions
449 analysed (Mann-Whitney test, $P < 0.001$). We found that several deletion mutants
450 were attenuated for survival after 3 days in BHI and water (sterile, tap and rain).
451 However, at 7 days their survival did not differ significantly to that of the wild-type.
452 This is most likely due to further decreasing levels of the wild-type between day 3
453 and 7 (Fig 5a).

454

455 **Fig 5. Validation of Tn mutant library screen during survival at low temperature**
456 **under nutrient-rich and –poor conditions.** (A-E) Survival of defined *C. jejuni*
457 M1cam gene deletion mutant in (A) BHI, (B) chicken juice, (C) sterile water, (D) tap
458 water, (E) rain water. Survival of *trxC* defined gene deletion mutant and
459 complemented mutant (F), at 4°C in different media. Data shown are Log₁₀ decrease
460 of CFU/ml relative to the 0 day time-point and plotted as means with SEM ($n \geq 4$).
461 Statistical significance was analysed using a Mann-Whitney test with * $P < 0.05$, ** P
462 < 0.01 and *** $P < 0.001$.

463

464 Confirmatory experiments with deletion mutants highlighted the contribution of
465 *hisC* (aromatic amino acid aminotransferase) and *trxC* (thiol-disulphide
466 oxidoreductase) for survival at low temperature (Fig 5b-e). The *hisC* mutant showed
467 attenuated survival in BHI (7 days), sterile water (3 days), tap water (3 and 7 days)

468 and rain water (3 days) (Fig 5b-e). Deletion of *trxC*, however, resulted in reduced
469 survival of *C. jejuni* in BHI, sterile water and tap water after 7 days (Fig 5b-d) and in
470 chicken juice and rain water after 3 days (Fig 5a,e). Although the deletion of *trxC*
471 resulted in a prolonged lag-phase during growth *in vitro* in BHI broth (Fig S7), genetic
472 complementation of the *trxC* deletion restored its phenotype under all conditions
473 tested, confirming the importance of *trxC* in the survival of *C. jejuni* at 4°C, both in
474 nutrient-rich and –poor conditions (Fig 5b). Thus far, the function of TrxC in *C. jejuni*
475 is unknown. The closest ortholog is TrxC in *Helicobacter pylori*, which is required for
476 protection against oxidative stress [56]. We found that the sensitivity of the M1cam
477 *trxC* mutant to hydrogen peroxide was increased, and that sensitivity was restored to
478 near wild type levels in the genetically complemented mutant (Fig 6). Therefore, our
479 data support a role for *C. jejuni* TrxC in an oxidative stress response, and could be a
480 good target for intervention strategies.

481

482 **Fig 6. Sensitivity to hydrogen peroxide of *C. jejuni* M1cam wild-type, *trxC***
483 **defined gene deletion mutant and *trxC* complemented mutant.** Data shown are
484 zones of inhibition and plotted as means with SEM ($n \geq 4$). Statistical significance
485 was analysed using a two-way ANOVA test with *** $P < 0.001$.

486

487 Adhesion and Invasion of Human Gut Epithelial Cells

488 The M1cam mutant library 'C' was used to infect Caco-2 human gut epithelial
489 cells. To identify genes required for adhesion, Tn mutants that adhered to Caco-2
490 epithelial cells were compared to the non-adherent fraction rather than to the
491 inoculum. This was to compensate for the survival of the Tn mutant library in the
492 infection medium, *i.e.* to restrict the number of false positives due to attenuated

493 survival in the infection medium. Comparing the Tn mutants that invaded the
494 epithelial cells with the non-adherent fraction identified 57 candidate genes involved
495 in cellular invasion, whereas only two genes passed our filtering criteria for adhesion
496 (Fig 2 and Table S3 and S4).

497 As anticipated, genes linked to the COG class “cell motility” were significantly
498 overrepresented amongst the genes required for invasion of Caco-2 cells (Fig 2 and
499 Fig S5). Previous work has shown that flagellar motility plays an important role in the
500 pathogenesis of *C. jejuni* and is considered a driving factor for host interaction [3, 28,
501 57, 58]. Employing a Cos-1 monkey kidney fibroblast cells invasion model, Gao *et al.*,
502 screened a Tn mutant library in *C. jejuni* 81-176, which led to the identification of 36
503 invasion genes [28], of which 19 genes were also identified in this study using Caco-
504 2 human gut epithelial tissue culture cells. A total of 38 genes were only required for
505 invasion by M1cam and 17 only in 81-176 [28]. Of the M1cam unique invasion genes,
506 14 were linked to the flagellar system, including *fliK*, *flaG*, *fliD*, *fliW*, and *maf3*. The
507 differences in invasion requirements are likely to be caused by the different cell-types
508 used and might vary between different strains of *C. jejuni*.

509 Validation of invasion genes identified by Tn-seq was performed using a panel
510 of 15 defined gene deletion mutants, this included genes not previously reported to
511 be linked to motility and the flagellar system (Fig 7a,b). In these experiments, the role
512 of 13 out of 15 selected genes in invasion was confirmed (Fig 7b). Although Tn-seq
513 only detected two genes of which Tn mutants showed reduced adhesion, 14 out of
514 15 deletion mutants selected for confirmatory experiments displayed attenuated
515 adhesion (Fig 7a). This apparent discrepancy might be due to differences in the
516 experimental set-up between the Tn library screen and the validation experiments. In
517 the Tn library screen, interactions between adhesion-deficient and -proficient Tn

518 mutants may have compensated for adhesion of otherwise adhesion-deficient Tn
519 mutants. However, it is also well recognised that competition between Tn mutants is
520 a confounding factor in Tn library screens [29, 58]. This is in contrast to the validation
521 experiments presented here, in which defined gene deletion mutants were allowed to
522 interact with Caco-2 cells as single mutant inocula.

523

524 **Fig 7. Validation of Tn mutant library screen during adhesion and invasion of**
525 **human gut epithelial tissue culture cells.** Adhesion to (A), and invasion (B) of,
526 Caco-2 cells by *C. jejuni* M1cam defined gene deletion mutants. Mutants in *luxS* and
527 *pstA*, which were not identified in our Tn-seq screen with Caco-2 cells, were included
528 as negative controls and *rpoN*, which was previously shown to be required for
529 adhesion and invasion [59, 60], served as a positive control. Caco-2 adhesion (C),
530 and invasion (D), of gene deletion and genetically complemented mutants. Data is
531 represented as percentage of wild-type ($n \geq 3$) and plotted as means and SEM. The
532 competitive index (CI) was calculated by dividing the ratio of mutant to wild-type
533 bacteria recovered upon (E) adhesion to, and (F) invasion of, Caco-2 cells by the
534 ratio of mutant to wild-type bacteria that were used in the inoculum. Statistical
535 significance was calculated using a Mann-Whitney test where * $P < 0.05$, ** $P < 0.01$
536 and *** $P < 0.001$.

537

538 Amongst the genes that were confirmed to play a role in interacting with Caco-
539 2 cells were *livM* (amino acid metabolism), *fabL* (fatty acid metabolism) and *engD*
540 (GTP-dependent nucleic acid-binding protein) that have no known link to the flagellar
541 system. We assessed the motility of these mutants in semi-solid agar (Fig S6) and
542 found that deletion of *livM* resulted in ~50% reduced motility compared to the wild-

543 type. WGS analysis did not reveal any genomic variations in the *livM* mutant linked to
544 motility (Table S5). The role of *engD* in colonisation of chickens was not confirmed in
545 validation experiments with a defined deletion mutant (Fig 3). However, we confirmed
546 the contribution of *engD* in adhesion and invasion of Caco-2 tissue culture cells (Fig
547 7). The *maf3* gene deletion mutant was motile (Fig S6) but had a reduced capacity to
548 colonise chickens (Fig 3a,b) and also lacked the ability to adhere to, and invade,
549 Caco-2 cells (Fig 7a-d).

550 Genetic complementation of the defined gene deletions in *fabL*, *livM*, *fliW*,
551 *maf3* and *flgK* mutants restored their adhesion and invasion capacity to wild type-
552 levels, confirming their role in the interaction with human gut epithelial cells (Fig
553 7c,d). Genetic complementation of the *flaG* deletion mutant significantly enhanced
554 adhesion and invasion levels, however these levels were still lower than the wild-
555 type. Although not confirmed, this may be due to deregulated *fliD* or *fliS* expression
556 as observed for the *fliD* mutant (see Fig S8 for details). Genetic complementation of
557 the *engD* deletion mutant was also unsuccessful due to the lack of expression of
558 *engD* (Fig S8). Further, the *capM* and *fdhA* Tn-seq phenotypes were confirmed when
559 mutants were assayed in competition with the wild-type (Fig. 7ef), however in mono-
560 infection no phenotypes were observed (Fig. 7a-d). The *luxS* negative control mutant
561 was not attenuated for adhesion and invasion in competition with the wild-type (Fig.
562 7ef).

563

564

565 Discussion

566 This study presents a comprehensive analyses of gene fitness in *C. jejuni* and
567 underlines various molecular mechanisms that are critical in its life cycle. Profiling the
568 genes required for *in vitro* growth of three *C. jejuni* strains underscored that a large
569 part of the genome (~27%) is vital to bacterial fitness (Table S2). Most likely as a
570 consequence of a low redundancy in the relatively minimal genome of *C. jejuni* and
571 transcriptional coupling of genes, as demonstrated by Dugar *et al.*, and Porcelli *et al.*,
572 [26, 27].

573 Elements of the flagellar system, *i.e.* the flagellar base and periplasmic rod
574 structure and T3SS (Fig 8), encoded by indispensable genes or genes which when
575 inactivated by a Tn have a severe impact on fitness, provide a potential focus for
576 intervention along with genes required for the LOS lipid A and KDO moieties. We
577 also found that inclusion of the first L-glycero-D-manno-heptose residue (catalyzed
578 by WaaC) was required for fitness. These structural elements have a critical role
579 during host interaction and therefore represent promising targets for developing
580 intervention strategies [57, 61]. Our gene fitness analysis revealed that components
581 of the gluconeogenesis pathway were essential for the growth of *C. jejuni*. The
582 gluconeogenic pathway is required for biosynthesis of glucose-(derivatives) that
583 serve as building blocks for both LOS and CPS as well as the N- and O-linked
584 protein glycosylation pathways [14]. In addition, genes (*fabZ*, *fabF*, *fabH*, *fabD* and
585 *fabG*) that are part of the type II fatty acid synthesis pathway (FASII) were required
586 for fitness in all three *C. jejuni* strains. There are several classes of FASII pathway
587 inhibitors that have potent antimicrobial properties and consequently fatty acid
588 synthesis is also considered a lucrative target for antibiotics [33, 62].

589

590 **Fig 8. Overview of flagellar system genes required for fitness during *in vitro***
591 **growth and in model representing host interaction.** Genes involved in the
592 flagellar system that were required for fitness during *in vitro* growth of *C. jejuni* strains
593 M1cam, 81-176 and 11168 are indicated in blue. Genes that passed the fitness score
594 cut-off but did not pass the 0.95 probability for Tn inactivation are indicated with blue
595 dashed boxes. Genes which in the Tn library screens were shown to be required for
596 chicken colonisation, gnotobiotic piglet infection, invasion of human gut epithelial
597 tissue culture cells or survival in houseflies are indicated in orange.

598

599 The same *C. jejuni* M1cam Tn library was screened in all of the experimental
600 models presented in this study and at the same time compared with data derived
601 from screening the same Tn library during infection of gnotobiotic piglets (De Vries *et*
602 *al.*, submitted). This enabled comparative analyses of genes required in *in vitro* and
603 *in vivo* experimental models relevant to the life cycle of *C. jejuni*. Comparing genes
604 required for colonisation of chickens and infection of gnotobiotic piglets as well as the
605 genes required for interaction with human gut epithelial cells, not only enabled us to
606 identify genes associated with propagation and survival across different host species
607 but also of genes specific to different hosts.

608 Twenty-eight genes were found to be required in all three host interaction
609 models (chicken, pig, and cellular invasion), of which 21 genes belong to the flagellar
610 system. This was in-line with our COG analysis, which revealed that motility-related
611 genes were significantly overrepresented during host interaction (chicken, pig, and
612 cellular invasion) (Fig 2 and Fig S5). Amongst the genes that were required across
613 three host models and that did not belong to the flagellar system were *engD*, *livM*
614 and *capM* (see Fig 3 and Fig 7). The EngD ortholog in *E. coli* (YchF) belongs to the

615 GTPase family and is a negative regulator of the oxidative stress response [63].
616 YchF proteins have been implicated in pathogenesis of other bacterial species [63,
617 64]. With branched-chain amino acids (BCAA) being linked to chemotaxis [65], *livM*,
618 part of the ABC-type BCAA transport system (*livM*, *livH*, *livK*, and *livJ*), may be
619 implicated in chemotaxis within host organisms. Previous studies have shown a role
620 for the *N*-linked protein glycosylation gene *capM* in chicken colonisation [41, 42],
621 along with other members of the surface protein glycosylation locus, including *pglB*,
622 *pglE* and *pglF* [66]. Our Tn-seq screen in the chicken colonisation model also
623 identified Tn mutants in *pglB*, *pglF* and *pglI* exhibiting reduced colonisation (Table
624 S3), whereas inactivation of *pglACD* resulted in reduced fitness during *in vitro* growth
625 (Table S2). The PglH protein plays a crucial role in the glycan assembly process as it
626 has a polymerase activity and adds the final *N*-acetylgalactosamine residues during
627 surface decoration [67].

628 A total of 96 genes were only identified in the chicken colonisation screen and
629 not during infection of gnotobiotic piglets, among these genes were the phosphate
630 metabolism genes (*pstABS* and *phoR*). *PstA*, a gene that appears to be unrelated to
631 motility (Fig S6), did not affect adhesion and invasion of human gut epithelial cells
632 (Fig 7), and is vital for the capacity of *C. jejuni* to colonise chickens (Fig 3a,b). The
633 expression of the phosphate regulon in *C. jejuni* 81-176 was increased in 1-day-old
634 chickens, which suggests that phosphate levels might be low in chickens, thereby
635 triggering the activation of the phosphate regulon [40]. The phosphate regulon might
636 operate in a second messenger system that (in)directly regulates expression of
637 chicken colonisation genes [68]. However, it remains to be investigated which *C.*
638 *jejuni* genes are under control of this system. Tn mutants in genes involved in
639 chemotaxis, *mcp4_1* and *cheV*, were uniquely attenuated during chicken

640 colonisation, whereas Tn mutants in *mcp4_3* and *mcp4_4* were uniquely attenuated
641 during gnotobiotic piglet infection, suggesting the existence of host- or substrate-
642 specific chemotaxis. A large number of genes involved in the utilisation of amino
643 acids and organic acids such as lactate, pyruvate, acetate and tricarboxylic acid
644 (TCA) cycle intermediates only appear to be required during gnotobiotic piglet
645 infection. These host specific (central) metabolic requirements are most likely related
646 to dietary differences between chickens and piglets. Metabolic pathways required
647 across different hosts may be considered potential targets to develop antimicrobials.
648 Our Tn library screens indicated that pyruvate kinase (*pyk*) is important in both
649 chickens and piglets. Pyk is part of the Embden-Meyerhof-Parnas pathway, and
650 shifting the metabolic flux at the level of Pyk has previously been suggested as a
651 potential antimicrobial strategy [69].

652 A detailed analysis of the flagellar system across various *in vitro* and *in vivo*
653 experimental models indicated that the extracellular part of the flagellum, the filament
654 and hook structure, play a vital role in *C. jejuni* (Fig 8). This included genes encoding
655 the filament chaperone gene FliW [16, 70] and the flagellar cap FliD. A FliD subunit
656 vaccine has been shown to induce a transient but significant reduction (~2 log) of *C.*
657 *jejuni* loads in the chicken caecum [71], demonstrating its potential as a target for
658 intervention. Linked to the flagellar system is also the motility accessory factor 3
659 (*maf3*). We found that a *maf3* deletion mutant was motile, but displayed attenuated
660 chicken colonisation and adhesion and invasion of Caco-2 cells. The *maf3* gene is
661 located in the flagellar glycosylation locus [43]. However, the function of this gene is
662 still unknown, and interestingly, isoelectric focusing analysis of a flagellar preparation
663 from a *C. jejuni* 81-176 *maf3* deletion mutant did not indicate altered glycosylation of
664 the flagellar filament [72].

665 Experimental models of *C. jejuni* survival and transmission, *i.e.* in the housefly
666 and under low temperature conditions, were shown to be less stringent than the *in*
667 *vivo* and *in vitro* host-pathogen models. This was reflected by subtler fold-changes
668 compared to the host interaction models. This led to a lower number of candidate
669 genes being identified and a lower conformation in validation experiments with
670 defined gene deletion mutants (Fig 4 and Fig 5). Interestingly, this led to a novel
671 potential target for intervention that is not linked to host interactions. The *trxC* gene
672 was required for survival at low temperature, both in nutrient-rich and –poor
673 conditions (Fig 5) and for resistance against peroxide (Fig 6), indicating a role in the
674 oxidative stress response.

675 Although *C. jejuni* is considered the leading cause of bacterial gastroenteritis,
676 our understanding of its biology is rather limited. Thus far, intervention strategies are
677 unsuccessful in controlling *C. jejuni* in poultry and its transmission to humans. The
678 development of future (novel) intervention strategies might best be aided by a
679 thorough understanding of the biology of *C. jejuni* in its life cycle. The generated Tn
680 mutant libraries in three well-characterised *C. jejuni* isolates represent a valuable tool
681 for the *Campylobacter* research community. Our work in this study pointed out that
682 many genes/pathways make indispensable contributions to the ability of *C. jejuni* to
683 thrive in the host and environment. We anticipate that the use of the Tn mutant
684 libraries in future studies will provide a continued insight into the mechanisms
685 required for growth as well as survival within and outside host organisms. Perhaps
686 most importantly, our comprehensive screening approach has again clearly shown
687 that the flagella drive *C. jejuni* interaction with its hosts. Therefore, future efforts
688 should focus on how to exploit this to effectively control infections caused by *C.*
689 *jejuni*.

690

691 Materials and Methods

692 Ethics Statement

693 All chicken infection work was conducted in accordance with UK legislation
694 governing experimental animals under project licence 40/3652 and was approved by
695 the University of Liverpool ethical review process prior to the award of the licence.

696

697 Bacterial Strains and Growth Conditions

698 Wild-type strains, defined gene deletion mutants, genetically complemented
699 mutants and plasmids are summarised in [Table S6](#). *C. jejuni* strains M1cam
700 (derivative of M1 used in our laboratory) [16, 17], 11168 [10], and 81-176 [18] were
701 cultured in Brain Heart Infusion (BHI) broth or on BHI agar supplemented with 5%
702 (v/v) defibrinated horse blood in the presence of 5 µg/ml trimethoprim (TrM). Tn
703 mutant libraries and defined gene deletion mutants were grown in the presence of 10
704 µg/ml chloramphenicol (Cm), whereas 50 µg/ml kanamycin (Km) was added to the
705 media for culturing genetically complemented mutant strains. *C. jejuni* were grown
706 under microaerophilic conditions (5% O₂, 10% CO₂, 85% N₂) in a MACS VA500
707 Variable Atmosphere Work Station (Don Whitley, Shipley, United Kingdom). Liquid
708 cultures were grown with agitation (200 rpm). For use in the experimental models, *C.*
709 *jejuni* M1cam wild-type, defined gene deletion mutants and genetically
710 complemented mutants were cultured for ~48 h, re-plated on fresh plates and grown
711 for another 16 h. For Tn mutant libraries, freezer stocks were plated on 9 x 90 mm
712 BHI blood agar plates (200 µl per plate) and grown for 16 h. *E. coli* NEB 5α or 10-β
713 (New England Biolabs) were used for cloning and were cultured in Luria Bertani (LB)
714 medium supplemented with appropriate antibiotics, at 37 °C.

715

716 Construction of Defined Gene Deletion Mutants and Genetically 717 Complemented Mutants

718 *C. jejuni* M1cam gene deletion mutants were constructed by allelic
719 replacement of the gene with a chloramphenicol (*cat*) resistance cassette as
720 described in de Vries *et al.* [16]. Gene deletion mutants were also subjected to
721 phenotypic and genotypic characterisation, including motility (Fig S6) and *in vitro*
722 growth in liquid culture (Fig S7). Whole genome sequencing (WGS) based variant
723 analysis (single nucleotide polymorphisms [SNP] and insertion/deletions [INDELS]) of
724 defined deletion mutants was used to screen for second-site mutations that might
725 have affected the phenotypes under investigation. Variants were detected at 96
726 positions relative to the *C. jejuni* M1cam reference genome [16] (Table S5). The
727 variant database was cross-referenced with data obtained in Tn mutant library
728 screens to assess whether the gene affected by the variant had a potential impact on
729 the phenotype under investigation. In addition, phenotypes of deletion mutants
730 sharing a variant were compared to predict possible confounding effects of the
731 second-site mutation. This in-depth analysis did not identify confounding effects of
732 second-site mutations. For selected candidates, the respective defined gene deletion
733 mutant was genetically complemented (Fig S8) and tested.

734 Genetic complementation of *mcp4_1*, *fliK*, *flaG*, Δ *fliD*, *pstA*, *fabL*, *engD*, *livM*,
735 *trxC*, *capM*, *maf3* and *flgK* mutants was performed using the pSV009 plasmid as
736 described in de Vries *et al.* [16]. Predicted promoter region(s), including a ribosome-
737 binding site (RBS) were derived from the Dugar *et al.* dataset [26]. For genes
738 arranged in an operon, the promoter region was mostly identified upstream of the 5'
739 end of the transcript. In such cases, the expected promoter region was synthesised

740 as a DNA string (GeneArt, Life Technologies, UK) flanked with appropriate restriction
741 sites. Open reading frames (ORFs) of *fliK*, *fliD*, *pstA*, *engD* and *maf3* were cloned
742 into *XbaI* and *BamHI* sites in pSV009, following which, respective promoter regions
743 were inserted within the *XhoI* and *XbaI* sites, yielding the pSV009 derived plasmids
744 containing the gene of interest fused to a suitable promoter. For *mcp4_1*, *flaG*, *fabL*,
745 *trxC*, *capM* and *flgK*, a promoter consensus region was identified upstream of the
746 start codon. A fragment containing the ORF plus ~200 bp upstream was cloned in
747 the *XhoI* and *BamHI* sites of the plasmid pSV009. For *livM* no obvious promoter
748 region could be located and therefore its transcription was placed under the control of
749 a *cat* promoter by cloning the ORF into *XbaI* and *BamHI* sites of pSV009. Genetic
750 complementation fragments, *i.e.* sequences for homologous recombination, a
751 suitable promoter and ORF of the gene to be complemented, were amplified by PCR
752 from the respective pSV009-derived plasmids using primers pSV009_GCamp1
753 _FW1/RV1. The genetic complementation fragments were introduced into respective
754 M1cam defined gene deletion mutants by electroporation as described in de Vries *et*
755 *al.* [16]. Oligonucleotide sequences are listed in [Table S7](#).

756

757 Gene Expression Analysis in Genetically Complemented Gene 758 Deletion Mutants

759 For expression analysis of genes in complemented gene deletion mutants,
760 total RNA was isolated as previously described in de Vries *et al.* [73]. DNA-free total
761 RNA (500 ng) was reverse transcribed into cDNA using the QuantiTect reverse
762 transcription kit (Qiagen). Real-time quantitative PCR (RT-qPCR) was performed
763 using the SensiFast SYBR No-ROX Kit (Bioline) on a Rotor-Gene Q machine
764 (Qiagen). Expression fold-changes relative to wild-type were calculated using the

765 $\Delta\Delta\text{Ct}$ method [74] with the *gyrA* gene as a reference. For the isolation of total RNA,
766 each bacterial strain was cultured in duplicate and RT-qPCR reactions were
767 performed in duplicate.

768

769 Analysis of Growth Kinetics

770 *C. jejuni* wild-type and gene deletion mutants were harvested from BHI blood
771 agar plates in 2 ml of BHI broth. Culture suspensions were diluted to $\text{OD}_{600\text{nm}} \sim 0.2$
772 and were used to analyse growth kinetics and motility. This suspension was used to
773 inoculate 5 ml pre-warmed BHI broth at an $\text{OD}_{600\text{nm}}$ of ~ 0.005 . Bacterial growth was
774 monitored by optical density after 4, 8 and 24 h incubation at 200 rpm at 42°C under
775 microaerophilic conditions ($n \geq 3$). Ten-fold serial dilutions were made and plated
776 onto BHI blood agar plates.

777

778 Analysis of Motility

779 Motility assays ($n = 3$) were performed in semi-solid agar as described in de
780 Vries *et al.*[16]. Bacterial suspensions that were diluted to $\text{OD}_{600\text{nm}} \sim 0.2$ (see previous
781 section) were used to stab 90 mm petri dishes containing 25 ml of BHI broth
782 supplemented with 0.4% (w/v) agar. These plates were incubated under
783 microaerophilic conditions at 42°C for 24 h before measuring the diameter of
784 growth/motility.

785

786 Genome Sequencing of Defined Gene Deletion Mutants

787 Libraries for Illumina sequencing were prepared using the NEBNext Ultra or
788 Ultra II DNA library prep kit (New England Biolabs) and sequenced on the MiSeq

789 platform as described in de Vries *et al.* [16]. For variant analysis, reads were mapped
790 to the M1cam reference genome (accession no. CP012149 [16]) using Stampy [75],
791 variants were identified with Samtools [76] and the effect at the protein level was
792 predicted using SnpEff [77]. Gene deletion mutants in *pflA* and *flaD* were analysed
793 previously in de Vries *et al.*[16].

794

795 Construction of Tn Mutant Libraries in *C. jejuni*

796 A Tn donor plasmid suitable for Tn-seq [19] was constructed by amplifying the
797 *mariner* Tn encoding the Cm-resistance cassette from pAJG39 [36] using a single 5'-
798 phosphorylated primer PBGSF20. This introduced *MmeI* restriction sites within the
799 inverted repeats of the Tn element. The Tn element was sub-cloned into pJET1.2
800 (Thermo Scientific) and the resulting plasmid pSV006, was used for *in vitro* Tn
801 mutagenesis as described in Holt *et al.*, [78] with minor adjustments. Briefly, 2 µg of
802 *C. jejuni* DNA was incubated for 5 h at 30°C with 1 µg of pSV006 and ~250 ng
803 Himar1-C9 transposase, which was purified as described in Akerley *et al.* [79]. After
804 end-repair with T4 DNA polymerase and *E. coli* DNA ligase the mutagenesised DNA
805 was transferred to *C. jejuni* by natural transformation [16]. Tn transformants were
806 harvested from plates and pooled. The pooled library was used to inoculate 100-200
807 ml BHI-TrM-Cm broth to an OD_{600nm} of ~0.1 and grown overnight). This yielded the
808 “working stocks” that were used in further experiments. Chromosomal DNA was
809 isolated with Genomic-tip columns (Qiagen).

810

811 Gene Fitness Analysis

812 Tn insertion sites were identified using Tn-seq [19], essentially as described in
813 Burghout *et al.*, [80]. Briefly, Tn mutant library DNA was fragmented by *MmeI*

814 restriction digestion. Adapters containing inline barcodes were ligated to the *Mmel*
815 fragments, and amplified using the primers PBGSF29 and PBGSF30 with NEBNext
816 high fidelity polymerase (New England Biolabs). Tn insertion sites were sequenced
817 using single-end 40-50 bp sequencing on the Illumina HiSeq 2500 platform.
818 Sequence reads were demultiplexed using the FastX toolkit barcode splitter and
819 analysed further with the ESSENTIALS pipeline [22]. Sequence reads were aligned
820 to the *C. jejuni* genomes [16, 18, 81] with a match of ≥ 16 nt. Kernel density plots
821 were generated in R to distinguish “true” Tn insertions from “noise” sequencing
822 reads, yielding a read count cut-off per Tn library. Insertion “hot spotting” was
823 analysed by plotting the Log_2 read count per chromosomal position using an in house
824 Perl script. As a measure of gene fitness, the Log_2 fold-change of observed vs
825 expected reads was calculated per gene, with Kernel density plots allowing accurate
826 delineation of fitness (required for *in vitro* growth) and non-fitness genes [22].
827 Additional criteria were: a Benjamini & Hochberg adjusted $P < 0.05$ and a probability
828 that the gene was inactivated by a Tn insertion of > 0.95 , as calculated using a
829 derivative of Poisson’s law; $1 - e^{-N \times \text{Ln}(1-f)}$, with N = number of unique Tn insertion
830 mutants and f being the gene size divided by the size of the genome. In addition,
831 genes for which no sequence reads were detected and the probability of inactivation
832 was > 0.95 were considered to be required for fitness. For functional class
833 enrichment analysis COGs were assigned to M1cam, 11168 and 81-176 proteins and
834 consensus COGs were assigned to homologous groups (HGs; see next section). The
835 overrepresentation of COG classes was assessed using a Fisher exact test with Q-
836 value multiple testing correction [82].

837

838 **Identification of Homologs in *C. jejuni* Strains**

839 Protein sequences of *C. jejuni* M1cam, 11168 and 81-176 were clustered into
840 putative homologous groups (HGs) with OrthAgogue [83]. For this, a collective
841 database was generated and proteins of the three strains were queried against this
842 database. The reciprocal best-hit protein pairs were identified by applying an e-value
843 filter cut-off of 1e-5 to the “all against all” BLAST output (only proteins > 50 amino
844 acids in length were included). The putative HGs were identified by clustering with
845 MCL by applying different values of an inflation parameter that defines cluster
846 tightness [84]. After testing different inflation parameters, an inflation value of 2.6 was
847 selected. This ensured that the majority of core HGs had only one representative
848 gene, keeping the number of HGs with duplicated genes from an isolate to a
849 minimum.

850

851 Systematic Literature Review on *C. jejuni* 11168 Defined Gene 852 Deletion and Tn Mutants

853 To assess which *C. jejuni* genes have been deleted in strain 11168, a PubMed
854 search was conducted in July 2015 with the following search terms: “*Campylobacter*
855 *jejuni*” and “mutant”. Publications were screened for the description of defined gene
856 deletion mutants.

857

858 Chicken Colonisation Experiments

859 One-day-old Ross 308 broiler chicks were obtained from a commercial
860 hatchery. Chicks were housed in the University of Liverpool, High Biosecurity Poultry
861 unit. Chicks were individually tagged with leg rings or wing bands and maintained in
862 floor pens at UK legislation recommended stocking levels allowing a floor space of

863 2,000 cm² per bird at 25°C on wood-shavings litter that was changed weekly prior to
864 inoculation, and were given *ad libitum* access to water and a pelleted laboratory
865 grade vegetable protein-based diet (SDS). Prior to experimental infection, all birds
866 were confirmed as *Campylobacter*-free by taking cloacal swabs, which were streaked
867 onto selective blood-free agar (mCCDA) (Lab M) supplemented with *Campylobacter*
868 Enrichment Supplement (SV59) and grown for 48 h at 42°C under microaerophilic
869 conditions.

870 At 21 days of age, birds were inoculated by oral gavage with $\sim 1.8 \times 10^8$ CFU
871 *C. jejuni* M1cam Tn library 'C'. The birds were split into five cages ($n = 6, 7, 7, 6, 6$;
872 group 1-5, respectively). Six days post-inoculation (p.i.), birds were killed by cervical
873 dislocation. At necropsy the caeca were removed aseptically and caecal content
874 was collected. Next, the caecal contents were diluted 5- and 50-fold with Maximum
875 Recovery Diluent (MRD) and 0.5 ml was plated per mCCDA-Cm (10 plates in total
876 per dilution) for recovery of Tn mutants that successfully colonised the birds. Ten-fold
877 dilution series were plated on mCCDA-Cm for enumeration. After 2 days growth, *C.*
878 *jejuni* M1cam Tn mutants were recovered from the plates by scraping in 2 ml MDR
879 and pelleted by centrifugation. The resulting pellets were stored at -80°C. DNA was
880 isolated from plate harvest pellets with Genomic-tip columns (Qiagen).

881 The composition of the input Tn mutant library ($n = 3$) and the library
882 recovered per cage of birds ($n = 4$, cage 1-4), a cage was considered as a single unit
883 of colonisation, was analysed by Tn-seq [19]. For this, 1 µg of DNA was pooled per
884 group of birds ($n = 6, 7, 7, 5$ for groups 1 to 4, respectively). One bird from group 4
885 and 2 birds from group 5 could not be analysed due to heavy contamination on the
886 recovery plates. As a result of this group 5 was eliminated from further analysis.

887 For validation with *C. jejuni* M1cam defined gene deletion mutants and
888 genetically complemented mutants, birds were inoculated with $\sim 1.8 \times 10^8$ CFU, at 6
889 days p.i. chickens were killed by cervical dislocation, the caeca were removed
890 aseptically and the caecal contents plates onto mCCDA plates for enumeration.

891

892 Infection of Houseflies

893 Newly pupated female houseflies (*Musca domestica*) were inoculated with 1 μ l
894 *C. jejuni* suspension *via* their proboscis [85]. For screening of Tn mutant survival, 5
895 groups of 10 flies were inoculated with $\sim 10^6$ CFU M1cam Tn library 'C' on four
896 different days. The flies were incubated for 4 h at 20°C (in the dark) after which flies
897 were homogenized using a Drigalsky spatula and 5 ml BHI was added. Large debris
898 was removed through a low speed (700 x g) spin and the supernatant was adjusted
899 to 5 ml with BHI before plating 0.5 ml per mCCDA-Cm (10 plates in total). Ten-fold
900 dilution series were plated on mCCDA-Cm for enumeration. After 24 h growth, Tn
901 mutants were recovered from plates by scraping in 2 ml BHI medium, centrifuged
902 and pellets were stored at -80°C for DNA isolation with Genomic-tip columns
903 (Qiagen). Chromosomal DNA from the 5 groups of 10 flies per day was pooled;
904 resulting in a single pooled sample for the library recovered in each of the 4 replicate
905 experiments. The composition of the input Tn mutant library ($n = 4$) and the library
906 recovered per group of flies ($n = 4$) was analysed by Tn-seq [19].

907 For validation experiments with *C. jejuni* M1cam gene deletion mutants ($n \geq 4$)
908 and genetically complemented mutants ($n = 6$), groups of 5 flies were inoculated with
909 $\sim 10^6$ CFU, killed 4 h p.i., and the bacterial load was quantified from pools of five flies
910 or five flies individually (only for 2 replicates with gene deletion mutants), in both

911 cases the average CFU per fly was used to calculate the Log₁₀ decrease of CFU per
912 fly relative to the inoculum.

913

914 Cold Survival Assay

915 A suspension was prepared from *C. jejuni* M1cam Tn mutant library 'C' plate
916 harvests, to an OD_{600nm} ~ 0.5 (~5 x 10⁸ CFU/ml). The Tn mutant library suspension
917 was harvested by centrifugation and resuspended in either BHI or sterile tissue-
918 culture grade water (experiment 1) or chicken juice, tap water or rain water
919 (experiment 2). The chicken juice was prepared as follows: 10 frozen whole chickens
920 were purchased from a commercial supplier and were allowed to defrost within the
921 packaging for at least 16 h, as described by Brown *et al.*[86]. Concentrated liquid
922 was recovered (> 200 ml), sterilised through a 0.2 µM filter and stored at -20 °C. Tap
923 water was obtained from a mains-fed tap in our laboratory that was allowed to run for
924 at least 2 min prior to collecting the water used in experiments. To collect rain water,
925 a large tub was left outside our laboratory overnight on a rainy evening. The rain
926 water and tap water were passed through a 0.22 µM filter unit and stored at 4°C.
927 Chicken juice, tap water and rain water were also screened for the presence of
928 *Campylobacter* spp. by plating on mCCDA plates. The Tn library samples were
929 incubated at 4°C and aliquots were taken at 0 h, 6 h, 1 and 7 days for experiment 1
930 (*n* = 3) or at 1, 3, and 7 days for experiment 2 (*n* = 4) and plated for recovery of Tn
931 mutants. Ten-fold dilution series were plated on BHI-TrM-Cm for enumeration. After 2
932 days incubation, Tn mutants were recovered from plates by scraping in 2 ml BHI per
933 plate, centrifuged and pellets were stored at -80°C for DNA isolation with Genomic-tip
934 columns (Qiagen). The Tn library composition at time = 0 h was compared to the

935 library recovered after incubation at 4°C under the above mentioned conditions by
936 Tn-seq [19].

937 For validation experiments, the survival of gene deletion mutants and
938 genetically complemented mutants was assayed as described above ($n = 4$).

939

940 Infection of Human Gut Epithelial Cells

941 Caco-2 cells (ATCC CC-L244 HTB-37), were cultured in DMEM (Life
942 Technologies) supplemented with 10% (v/v) heat inactivated FBS (Gibco), and 1%
943 (v/v) non-essential amino acids (Sigma Aldrich), at 37 °C with 5% CO₂.

944 The *C. jejuni* M1cam Tn mutant library 'C' was used to infect seven 143 cm²
945 dishes per replicate ($n = 4$) containing a monolayer of Caco-2 cells at a multiplicity of
946 infection (MOI) of 100 in low phosphate HEPES buffer (10 mM HEPES, 5.4 mM KCl,
947 145 mM NaCl, 5 mM glucose, 1 mM MgCl₂, 1 mM CaCl₂, 2 mM phosphate buffer pH
948 7.4) [87]. Cells were incubated for 2 h after which the non-adherent fraction from two
949 dishes was recovered onto BHI blood agar plates. For adherence, two dishes were
950 washed three times in Dulbecco's PBS (D-PBS), Caco-2 cells were lysed in 10%
951 (v/v) Triton-X100 solution in D-PBS, and bacteria were recovered on BHI blood agar
952 plates. The remaining five dishes were washed three times in D-PBS and incubated
953 for an additional 2 h in DMEM with 250 µg/ml gentamycin. Tn mutants that invaded
954 Caco-2 cells were recovered after washing three times in D-PBS and Caco-2 cell
955 lysis in 10% (v/v) Triton-X100 in D-PBS. After two days of growth on plates, Tn
956 mutants were recovered from the plates in 2 ml BHI, centrifuged and pellets were
957 stored at -80°C. DNA was isolated from harvested pellets using Genomic-tip columns
958 (Qiagen).

959 Validation of the screen was performed using 24-well plates, for which Caco-2
960 cells were infected with various *C. jejuni* M1cam defined gene deletion mutants and
961 genetically complemented strains ($n \geq 4$) as described above. Bacteria recovered
962 from different fractions were serially diluted and plated on BHI agar plates containing
963 appropriate antibiotics. Adhesion and invasion of Caco-2 cells by various *C. jejuni*
964 strains was calculated relative to the matched wild-type ($n \geq 3$).

965 To test the effect of competition, Caco-2 cells were infected at an MOI of 100
966 with a mix of the wild-type strain to a defined gene deletion mutant at a ratio of 100:1.
967 A competitive index (CI) score was calculated by dividing the ratio of mutant to wild-
968 type recovered from adherent and invaded fractions by the ratio of mutant to wild-
969 type bacteria in the inoculum ($n \geq 4$).

970

971 Analysis of Conditionally Essential Genes

972 Tn-seq data from the different experimental models (conditionally essential
973 genes screens) was processed as described in “gene fitness analysis”. To identify
974 genes of which Tn mutants were attenuated or enriched in the tested models, read
975 counts were collected per gene and compared between output (recovered) and the
976 input or control conditions (as defined above). Only genes covered by > 100 reads in
977 the control condition were considered, allowing assessment of 809 ± 58 ($67 \pm 5\%$)
978 non-fitness genes (define ‘non-fitness genes’) in the selected models. The following
979 filter steps were applied: a Log_2 fold-change (FC) below the attenuated cut-off value
980 or higher than the enriched cut-off value, Benjamini & Hochberg false discovery rate
981 < 0.05 , and two or more Tn mutants showing a Log_2 fold-change below the
982 attenuated cut-off value or higher than the enriched cut-off value (analysed using a
983 custom Python script). The Log_2 fold-change cut-offs were selected based on MA-

984 plots. In addition, the 514 genes that were obligate essential or required for fitness
985 (defined above) were eliminated from the analysis, see “gene fitness analysis”. COG
986 functional class enrichment was analysed using a Fisher exact test with Q-value
987 multiple testing correction [82].

988

989 Hydrogen Peroxide Sensitivity Assay

990 *C. jejuni* strains were added to pre-cooled BHI agar (~45°C) to a calculated
991 OD_{600nm} ~0.005 and 25 ml of this media-bacterial suspension was poured into 90 mm
992 petri dishes. Blank filter paper discs, 6 mm, were loaded with 10 µl of 0.05, 0.1, 0.25,
993 0.5, 1, 2.5, 5 or 7.5 M H₂O₂ solution. The discs were allowed to air dry and were then
994 placed in the center of the solidified agar plates. The plates were incubated for 24 h,
995 after which time the inhibition zone diameter was measured using a ruler (*n* = 4).

996

997 Sequencing Data

998 Tn-seq and genome sequencing data has been deposited in the European
999 Nucleotide Archive (<http://www.ebi.ac.uk/ena>) and are available *via* study accession
1000 number [This will be openly available after acceptance and can be made available for
1001 review].

1002

1003 Statistical Analysis

1004 Statistical analysis was performed in GraphPad Prism v6.

1005

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1011

1012 **Author contributions**

1013 Conceptualization: NW, PW, TH, DJM, AJG. Methodology: SPWdV, SG, AB, EW,
1014 AW, ANJ, LLL, SH, HK, KM, PE, NW, PW, AJG. Software: AB, FMM. Validation:
1015 SPWdV, SG, AB, AJG. Formal analysis: SPWdV, SG, AB, ANJ, AJG. Investigation:
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1017 Resources: FMM, AZ. Data curation: SPWdV, SG, AB. Writing original draft: SPWdV,
1018 SG, AG. Writing review and editing: SPWdV, SG, DJM, AG. Visualization: SPWdV,
1019 SG, AG. Supervision: AJG. Project administration: AJG. Funding acquisition: TH,
1020 DJM.

1021

1022

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1316

1317 **Competing interests**

1318 The authors have declared that no competing interests exist.

1319

1320 Supporting Information

1321 **Table S1. Overview of Tn mutant libraries constructed in *C. jejuni* M1cam,**
1322 **11168, and 81-176.**

1323

1324 **Table S2. Overview gene fitness analysis in *C. jejuni* M1, 11168 and 81-176.** As
1325 a measure of gene fitness, the Log₂ fold-change of observed vs expected reads was
1326 calculated per gene, with Kernel density plots allowing accurate delineation of fitness
1327 and non-fitness genes in each *C. jejuni* strain. Additional selection criteria were: a
1328 Benjamini & Hochberg adjusted $P < 0.05$ and a probability that the gene was
1329 inactivated by a Tn insertion of > 0.95 , as calculated using a derivative of Poisson's
1330 law. In addition, genes for which no sequence reads were detected and the
1331 probability of inactivation was > 0.95 were included. For comparative analysis of
1332 fitness genes in the three *C. jejuni* strains, homologs were identified; see Materials
1333 and Methods for a detailed description.

1334

1335 **Table S3. Overview of conditionally essential gene analysis.** To identify genes of
1336 which Tn mutants were attenuated or enriched in the tested experimental models,
1337 read counts were collected per gene and compared between output (recovered) and
1338 the input or control conditions. The following filter steps were applied: genes
1339 represented > 100 reads in the input/control condition, a Log₂ fold-change (FC) below
1340 the attenuated cut-off value or higher than the enriched cut-off value, Benjamini &
1341 Hochberg false discovery rate < 0.05 , and two or more Tn mutants showing a Log₂
1342 fold-change below the attenuated cut-off value or higher than the enriched cut-off
1343 value. The Log₂ fold-change cut-offs were selected based on MA-plots. In addition,

1344 the 514 genes that were obligate essential or required for fitness in *C. jejuni* M1cam
1345 ([Table S2](#)) were eliminated from this analysis.

1346

1347 **Table S4. Conditionally essential genes grouped per functional (COG)**
1348 **category.** Effect of Tn insertions on the ability of *C. jejuni* M1cam to colonize
1349 commercial broiler chickens, infect gnotobiotic piglets, adhere and invade human gut
1350 epithelial tissue culture cells, survive in houseflies and at 4°C in various media
1351 (chicken juice, BHI, tap water, rain water, and sterile water). Genes of which Tn
1352 mutants showed significantly attenuated or enriched fitness in the tested
1353 experimental models are listed and are grouped according to their COG functional
1354 classification. Data represented as Log₂ fold-change is also presented in [Fig 2a](#).
1355 Orange = significantly attenuated, Blue = significantly enriched, and Grey = no
1356 significant Tn-seq Log₂ fold-change.

1357

1358 **Table S5. Whole genome sequencing (WGS) based variant analysis of *C. jejuni***
1359 **M1cam defined gene deletion mutants.** For variant analysis, reads were mapped
1360 to the *C. jejuni* M1cam reference genome [16] with Stampy, variants (single
1361 nucleotide polymorphisms [SNP] and insertion/deletions [INDELS]) detected using
1362 Samtools, and the effect at the protein level was predicted using SnpEff. Cross-
1363 referencing the variants with data obtained in the Tn mutant library screens as well
1364 as phenotypes of defined gene deletion mutants sharing a variant did not reveal any
1365 confounding effects of second-site mutations, suggesting that the observed
1366 phenotypes for gene deletion mutants were true.

1367

1368 **Table S6. Bacterial strains and plasmids used in this work.**

1369 **Table S7. Oligonucleotides used in this study.**

1370

1371 **Fig S1. Distribution of Tn insertion sites in *C. jejuni* M1cam, 11168 and 81-176**

1372 **Tn mutant libraries.** Plotted are Log₂ reads per Tn insertion above the read count
1373 cut-off (see [Table S1](#) and Materials and Methods).

1374

1375 **Fig S2. Overlap of the Tn insertions per mutant library in *C. jejuni* M1cam,**
1376 **11168, and 81-176 Tn libraries.** Only Tn insertion sites above the read count cut-off
1377 are included in this analysis (see [Table S1](#) and Materials and Methods).

1378

1379 **Fig S3. Circular genome visualisation indicating fitness genes in *C. jejuni***
1380 **M1cam, 11168 and 81-176.** Genes required for fitness in one, two or all three of the
1381 strains are coloured-coded accordingly (See [Table S2](#)). A detailed description of the
1382 selection criteria is provided in the Materials and Methods section.

1383

1384 **Fig S4. Complexity of the *C. jejuni* M1cam Tn mutant library ‘C’ (Table S1) in**
1385 **the conditionally essential gene screens as analysed by Tn-seq.** Tn mutants
1386 represented by > 10 reads were considered to be present. Of note, in the “housefly
1387 survival” and “survival under nutrient-rich/poor conditions” screens, a higher number
1388 of Tn mutants were detected in the recovered (output) samples, which was the result
1389 of a lower sequence depth for some replicates of the input/inocula. Data is shown as
1390 individual data points and bars representing the mean with SEM. Statistical
1391 significance was analysed using a Mann-Whitney test with * $P < 0.05$ and ** $P < 0.01$.

1392

1393 **Fig S5. Functional class (COG) enrichment analysis of genes required during**
1394 **colonisation of chickens, infection of gnotobiotic piglets, invasion of human**
1395 **gut epithelial tissue culture cells, or survival in houseflies.** The
1396 overrepresentation of COG classes was assessed using a Fisher exact test with Q-
1397 value multiple testing correction; * $Q < 0.05$ and *** $Q < 0.001$.

1398

1399 **Fig S6. Motility of *C. jejuni* M1cam defined gene deletion mutants.** The data is
1400 represented as percentage relative to the wild-type ($n \geq 4$). Statistical significance
1401 was calculated using Kruskal-Wallis with Dunn's correction for multiple comparisons
1402 with * $P < 0.05$, ** $P < 0.01$ and *** $P < 0.001$.

1403

1404 **Fig S7. Growth kinetics of *C. jejuni* M1cam defined gene deletion mutants.**
1405 Growth of *C. jejuni* M1cam wild-type and gene deletion mutants were recorded as
1406 viable counts over 24 h at 42°C under microaerophilic conditions. Data is
1407 represented as mean and SD ($n \geq 2$).

1408

1409 **Fig S8. Genetic complementation of *C. jejuni* M1cam defined gene deletion**
1410 **mutants.** RT-qPCR analysis revealed the restored expression of *mcp4_1*, *flaG*, *pstA*,
1411 *fabL*, *livM*, *fliW*, *maf3* and *flgK* in genetically complemented stains. Lower expression
1412 of *fliD*, *trxC* and *capM* was observed in respective genetically complemented strains,
1413 compared to the wild-type. Although the *fliD* gene was expressed in the genetically
1414 complemented mutant, albeit at lower levels than the wild-type, it was unable to
1415 restore motility (data not shown). This might be (partially) caused by deregulated
1416 flagellar assembly due to an increased expression (2.9-fold compared to wild-type) of
1417 *fliS* (a flagellar chaperone [88]) located downstream of *fliD* (data not shown). WGS

1418 analysis did not reveal any genomic variations in the *fliD* mutant linked to motility
1419 (Table S5). The data is represented as Log₂ fold-change relative to the expression
1420 levels of the candidate gene in wild-type. Statistical significance ($n \geq 4$) of mutant vs
1421 wild-type and genetically complemented strains vs wild-type was calculated using a
1422 one-way ANOVA test with Bonferroni correction for multiple comparisons. For all
1423 experiments, data shown are means and SD. * $P < 0.05$, *** $P < 0.001$
1424















