- 2 virulence and physiology of *Clostridioides difficile* **R20291**
- 3 Running title: Coregulations of Fliw, CsrA, and FliC in *C. difficile*
- 4 Authors: Duolong Zhu<sup>a</sup>, Shaohui Wang<sup>a</sup>, Xingmin Sun<sup>a#</sup>
- 5 Affiliations: <sup>a</sup>Department of Molecular Medicine, Morsani College of Medicine, University of
- 6 South Florida, Tampa, FL, United States
- <sup>7</sup> <sup>#</sup> Corresponding author.
- 8 Address: 12901 Bruce B. Downs Blvd Tampa, FL 33612, United States
- 9 Fax: +1-8139747357
- 10 E-mail: sun5@usf.edu
- 11 Phone: Phone: +1-8139744553
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# 24 ABSTRACT

Clostridioides difficile is a Gram-positive, spore-forming, and toxin-producing anaerobe that can 25 cause nosocomial antibiotic-associated intestinal disease. In C. difficile, the expression of 26 flagellar genes is coupled to toxin gene regulation and bacterial colonization and virulence. The 27 flagellin FliC is responsible for pleiotropic gene regulation during *in vivo* infection. However, 28 29 how *fliC* expression is regulated is unclear. In *Bacillus subtilis*, flagellin homeostasis and motility are coregulated by flagellar assembly factor FliW, Flagellin Hag (FliC homolog), and 30 CsrA (Carbon storage regulator A), which is referred to as partner-switching mechanism "FliW-31 32 CsrA-Hag". In this study, we characterized FliW and CsrA functions by deleting or overexpressing fliW, csrA, and fliW-csrA in C. difficile R20291. We showed that both fliW 33 deletion or csrA overexpression in R20291, and csrA complementation in R20291AWA (fliW-34 csrA codeletion) dramatically decreased FliC production, however, *fliC* gene transcription was 35 unaffected. While suppression of *fliC* translation by *csrA* overexpression was mostly relieved 36 when *fliW* was coexpressed, and no significant difference in FliC production was detected when 37 only *fliW* was complemented in R20291AWA. Further, loss of *fliW* led to increased biofilm 38 formation, cell adhesion, toxin production, and pathogenicity in a mouse model of C. difficile 39 40 infection (CDI), while *fliW-csrA* codeletion decreased toxin production and mortality in vivo. Taken together, these data suggest that CsrA negatively modulates *fliC* expression and FliW 41 42 indirectly affects *fliC* expression through inhibition of CsrA post-transcriptional regulation, 43 which seems similar to the "FliW-CsrA-Hag" switch in B. subtilis. Our data also suggest that "FliW-CsrA-*fliC*/FliC" can regulate many facets of C. *difficile* R20291 pathogenicity. 44

45 **IMPORTANCE** 

46	C. difficile flagellin FliC is associated with toxin gene expression, bacterial colonization and
47	virulence, and is also involved in pleiotropic gene regulation during in vivo infection. However,
48	how fliC expression is regulated remains unclear. In light of "FliW-CsrA-Hag" switch
49	coregulation mechanism reported in <i>B. subtilis</i> , we showed that <i>fliW</i> and <i>csrA</i> play an important
50	role in flagellin synthesis which affects C. difficile motility directly. Our data also suggest that
51	FliW-CsrA-fliC/FliC" can regulate many facets of C. difficile R20291 pathogenicity. These
52	findings further aid us in understanding the virulence regulation in C. difficile.
53	KEYWORDS Clostridioides difficile, FliW, FliC, CsrA, R20291, virulence
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## 69 **INTRODUCTION**

Clostridioides difficile (formerly Clostridium difficile) (1, 2) is a Gram-positive, spore-forming, 70 toxin-producing, anaerobic bacterium that is a leading cause of nosocomial antibiotic-associated 71 diarrhea in the developed countries (3). C. difficile infection (CDI) can result in a spectrum of 72 symptoms, ranging from mild diarrhea to pseudomembranous colitis and potential death (4). C. 73 74 difficile has many virulence factors, among which toxin A (TcdA) and toxin B (TcdB) are the major ones (5, 6). These toxins can disrupt the actin cytoskeleton of intestinal cells through 75 76 glucosylation of the Rho family of GTPases, and induce mucosal inflammation and symptoms 77 associated with CDI (7).

The carbon storage regulator A (CsrA) has been reported to control various physiological 78 79 processes, such as flagella synthesis, virulence, central carbon metabolism, quorum sensing, motility, biofilm formation in pathogens including *Pseudomonas aeruginosa*, *Pseudomonas* 80 syringae, Borrelia burgdorferi, Salmonella typhimurium, and Proteus mirabilis (8-14). It is a 81 widely distributed RNA binding protein that post-transcriptionally modulates gene expression 82 through regulating mRNA stability and / or translation initiation of target mRNA (13, 15). CsrA 83 typically binds to multiple specific sites that are located nearby or overlapping the cognate 84 85 Shine-Dalgarno (SD) sequence in the target transcripts (16, 17). The roles of CsrA in *Bacillus* subtilis have been also reported (17-20). Yakhnin et al. (17) first reported that CsrA in B. subtilis 86 can regulate translation initiation of the flagellin (hag) by preventing ribosome binding to the hag 87 88 transcript. Meanwhile, two CsrA binding sites (BS1: A51 to A55; BS2: C75 to G82) were identified in the hag leader of mRNA, among which BS2 overlaps with the hag mRNA SD 89 90 sequence. Mukherjee et al. (18) elucidated that the interaction between CsrA and FliW could 91 govern flagellin homeostasis and checkpoint on flagellar morphogenesis in B. subtilis. FliW, the

92 first protein antagonist of CsrA activity was also identified and characterized in B. subtilis. They elegantly demonstrated a novel regulation system "a partner-switching mechanism" (Hag-FliW-93 CsrA) on flagellin synthesis in B. subtilis. Briefly, following the flagellar assembly checkpoint of 94 hook completion, FliW was released from a FliW-Hag complex. Afterward, FliW binds to CsrA 95 which will relieve CsrA-mediated hag translation repression for flagellin synthesis concurrent 96 97 with filament assembly. Thus, flagellin homeostasis restricts its own expression on the translational level. Results also suggested that CsrA has an ancestral role in flagella assembly 98 and has evolved to coregulate multiple cellular processes with motility. Oshiro et al. (19) further 99 quantitated the interactions in the Hag-FliW-CsrA system. They found that Hag-FliW-CsrA<sup>dimer</sup> 100 functions at nearly 1:1:1 stoichiometry. The Hag-FliW-CsrA<sup>dimer</sup> system is hypersensitive to the 101 102 cytoplasmic Hag concentration and is robust to perturbation.

103 Recently, the role of CsrA on carbon metabolism and virulence associated processes in C. difficile 630 $\Delta$ erm was analyzed by overexpressing the csrA gene (20). Authors showed that the 104 csrA overexpression can increase motility ability, toxin production, and cell adherence, and 105 induce carbon metabolism change. C. difficile flagellin gene fliC is associated with toxin gene 106 expression, bacterial colonization, and virulence, and is responsible for pleiotropic gene 107 108 regulation during *in vivo* infection (21-25). The delicate regulations among *fliC* gene expression, toxin production, bacterial motility, colonization, and pathogenicity in C. difficile are indicated. 109 Though the important roles of CsrA in flagellin synthesis and flagellin homeostasis have been 110 111 studied in other bacteria (17-19), the regulation of FliW, CsrA, and FliC and the function of *fliW* in C. difficile remain unclear. 112

In this communication, we aimed to study the involvement of FliW and CsrA in *fliC*expression and *C.difficile* virulence and physiology by constructing and analyzing *fliW* and *fliW*-

115 csrA deletion mutants of C. difficile R20291. We evaluated these mutants in expression of fliC,

motility, adhesion, biofilm formation, toxin production, sporulation, germination, andpathogenicity in a mouse model of CDI.

118 **RESULTS** 

## 119 Construction of *fliW* and *fliW-csrA* deletion mutants and complementation strains

The *C. difficile* R20291 flagellar gene operon was analyzed through the *IMG/M* website (https://img.jgi.doe.gov/), and the late-stage flagellar genes (F1) were drawn as Fig. 1A (21). Among them, *fliW* and *csrA* genes have a 10 bp overlap and were demonstrated as cotranscription by RT-PCR (Fig. S1).

To analyze the role of *fliW* and *csrA* in R20291, CRISPR-AsCpfI based plasmid pDL1 124 (pMTL82151-Ptet-AscpfI) was constructed for gene deletion in C. difficile (26, 27). pDL1-fliW 125 and pDL1-csrA gene deletion plasmids were constructed, and the fliW gene (288 bp deletion) 126 (R20291 $\Delta$ fliW, referred hereafter as R20291 $\Delta$ W) was deleted successfully. However, after 127 several trials, we couldn't get the csrA gene deletion mutant possibly due to its small size (213 128 bp). Therefore, *fliW-csrA* codeletion plasmid pDL1-*fliW-csrA* was constructed and the *fliW-csrA* 129 (445 bp deletion) codeletion mutant (R20291 $\Delta fliW$ -csrA, referred hereafter as R20291 $\Delta$ WA) was 130 131 obtained (Fig. 1 B and C). To study the role of *csrA* in R20291, the single gene complementation strain R20291AWA-W and R20291AWA-A were constructed. R20291, R20291-pMTL84153 132 R20291ΔW-pMTL84153 (R20291ΔW-E), 133 (R20291-E), and R20291\DeltaWA-pMTL84153 134 (R20291 $\Delta$ WA-E) were used as control strains when needed.

The effects of *fliW* and *fliW-csrA* deletion on R20291 growth were evaluated. Fig. 1D showed that there was no significant difference in bacterial growth between wild type strain and mutants in BHIS media.

# 138 Effects of *fliW* and *fliW-csrA* deletions on *C. difficile* motility and biofilm formation

To characterize the effects of *fliW* and *fliW-csrA* deletions on *C. difficile* motility, swimming (Fig. 139 2A; Fig. S2) and swarming (Fig. S2) motilities of R20291, R20291AWA, and R20291AW were 140 first analyzed at 24 h and 48 h post-inoculation, respectively. The diameter of the swimming halo 141 of R20291 $\Delta$ WA increased by 27.2% (p < 0.05), while that of R20291 $\Delta$ W decreased by 58.4% (p142 143 < 0.05) compared to that of R20291. Next, we examined the motility of the complementation strains (Fig. 2B; Fig. S2), and similar results were obtained among R20291-E, R20291∆WA-E 144 (with the swimming halo increased by 74.8%, p < 0.05), and R20291 $\Delta$ W-E (with the swimming 145 146 halo decreased by 59.2%, p < 0.05) (Fig. 2B). No significant difference was detected between complementation strain R20291AWA-WA, R20291AWA-W, R20291AW-W, and the parent 147 strain R20291-E except R20291 $\Delta$ WA-A which decreased by 52.0% (p < 0.05) in swimming 148 halo (Fig. 2B). The swarming (48 h) and swimming (24 h) motilities analyzed on agar plates 149 were shown in Fig. S2. 150

The effects of *fliW* and *fliW-csrA* deletions on *C. difficile* biofilm formation were also analyzed. In comparison with R20291, the biofilm formation of R20291 $\Delta$ W increased by 49.5% (*p* < 0.01), and no significant difference in biofilm formation was detected in R20291 $\Delta$ WA (Fig. 2C). The biofilm formation of R20291 $\Delta$ W-E increased 1.12 fold (*p* < 0.001) and R20291 $\Delta$ WA-A increased by 79.9% (*p* < 0.001) compared to R20291-E (Fig. 2D). Meanwhile, the biofilm formation of R20291 $\Delta$ WA-WA and R20291 $\Delta$ WA-W decreased by 42.8% (*p* < 0.01) and 25.2% (*p* < 0.05), respectively.

Together, these data indicate that loss of FliW impairs *C. difficile* motility, and increases biofilm production. The decrease of motility and increase of biofilm production were also detected in R20291 $\Delta$ WA-A, which was largely restored by coexpressing *fliW* with *csrA* in 161 R20291 $\Delta$ WA (Fig. 2B; Fig. 2D), indicating that *fliW* works together with *csrA* to regulate 162 bacterial motility and biofilm production.

### 163 Effects of *fliW* and *fliW-csrA* deletions on bacterial adherence *in vitro*

The ability of *C. difficile* vegetative cells to adhere to HCT-8 cells *in vitro* was analyzed. Fig. 2E 164 showed that the mean adhesion number of R20291 was  $2.40 \pm 0.70$  bacteria / cell, while that of 165 R20291 $\Delta$ W was 7.17  $\pm$  0.61, which was 3.0 fold (P < 0.0001) of R20291. No significant 166 difference was detected between R20291∆WA and R20291. In the complementation strains, we 167 detected a similar result which showed that the mean adhesion number of R20291 $\Delta$ W-E (6.17 ± 168 0.64) was 3.20 fold (P < 0.0001) of R20291-E (1.93 ± 0.25) (Fig. 2F). The adhesion ability of 169 complementation strains nearly recovered to that of wild type strain except for R20291 $\Delta$ WA-A 170  $(7.13 \pm 0.66, P < 0.0001)$  which was 3.69 fold of R20291-E in the mean adhesion number (Fig. 171 172 2F).

To visualize the adhesion of *C. difficile* to HCT-8 cells, the *C. difficile* vegetative cells were labeled with the chemical 5(6)-CFDA. Fig. 2G and 2H showed that the fluorescence intensity of R20291 $\Delta$ W was 3.50 fold (*P* < 0.0001) of that in R20291, and the fluorescence intensity of R20291 $\Delta$ W-E was 2.36 fold (*P* < 0.001) and R20291 $\Delta$ WA-A was 4.08 fold (*P* < 0.0001) of that in R20291-E, respectively, which is consistent with the results showed in the Fig. 2E and 2F. Meanwhile, the adherence of *C. difficile* to HCT-8 cells was also visualized by fluorescence microscopy (Fig. S3).

180 Our data showed that FliW negatively affects bacterial adherence. CsrA complementation in 181 R20291 $\Delta$ WA increased adherence, while the phenotype change can be recovered partially when 182 *fliW* was coexpressed with *csrA* in R20291 $\Delta$ WA, suggesting that *fliW* works together with *csrA* 183 to regulate bacterial adherence. The results from bacterial adherence analysis were consistent with biofilm production analysis indicating the close relation between biofilm production andadherence in *C. difficile*.

### 186 Effects of deletion and overexpression of *fliW* and *fliW-csrA* on *fliC* expression

In B. subtilis, FliW interacts with CsrA to regulate hag (a homolog of fliC) expression. We 187 reasoned that FliW and CsrA would also regulate *fliC* expression in *C. difficile*. As shown in Fig. 188 3A, the transcription of *fliC* in R20291 $\Delta$ WA increased 1.12 fold (p < 0.05), while the *fliW* 189 deletion impaired the *fliC* transcription slightly while no significant difference. Fig. 3B showed 190 the production of FliC in R20291 $\Delta$ W dramatically decreased (10.4 fold reduction, p < 0.001), 191 192 while that of R20291 $\Delta$ WA increased significantly (increased by 27.5%, p < 0.05). To further determine the role of the single gene csrA on FliC synthesis, the csrA and fliW were 193 complemented into R20291AWA or overexpressed in R20291, respectively. Results showed that 194 the significant difference of *fliC* transcription could only be detected in R20291 $\Delta$ WA-E 195 (increased by 32.3%, p < 0.05) (Fig. 3C) and R20291-W (increased by 69.8%) compared to 196 R20291-E (Fig. 3E). Interestingly, the FliC production of R20291 $\Delta$ WA-A was 4.2 fold (p <197 0.001) of that in R20291-E, while that of R20291 $\Delta$ WA-WA only decreased by 14.3% (p < 0.05) 198 and no significant difference of FliC production in R20291 $\Delta$ WA-W was detected (Fig. 3D). As 199 200 shown in Fig. 3E and 3F, the *fliC* transcription of R20291-A was not affected compared to R20291-E, but the FliC production in R20291-A decreased 5.3 fold (p < 0.0001). The decrease 201 of FliC production in R20291-A can be partially recovered when *fliW* was coexpressed with *csrA* 202 203 (R20291-WA decreased by 16.2%, p < 0.05).

Collectively, our data indicate that CsrA negatively modulates *fliC* expression posttranscriptionally and FliW works against CsrA to regulate *fliC* expression possibly through inhibiting CsrA-mediated negative post-transcriptional regulation.

### 207 Effects of *fliW* and *fliW-csrA* deletions on toxin expression

It has been reported that the expression of csrA could affect toxin expression in C. difficile (20). 208 To evaluate the effects of *fliW* and *fliW-csrA* deletions on toxin production, the supernatants of C. 209 difficile cultures were collected at 24 and 48 h post-inoculation, and the toxin concentration was 210 determined by ELISA. Fig. 4A showed that the TcdA concentration of R20291AWA decreased 211 by 28.6% (P < 0.05), while R20291 $\Delta$ W increased by 65.1% (P < 0.01) compared to R20291 at 212 24 h post-inoculation. However, after 48 h incubation, no significant difference was detected. In 213 Fig. 4B, TcdB concentration of R20291 $\Delta$ WA decreased by 26.4% (P < 0.05) at 24 h post-214 inoculation, while that of R20291 $\Delta$ W increased by 93.6% (P < 0.01) at 24 h and 33.0% (P < 215 0.05) at 48 h. Similar results were also detected in the complementation strains group (Fig. 4C 216 and 4D). As shown in Fig. 4C and 4D, after 24 h post-inoculation, TcdA (Fig. 4C) concentration 217 218 of R20291 $\Delta$ WA-E and R20291 $\Delta$ WA-W decreased by 33.0% (\*P < 0.05) and 47.7% (P < 0.01), and TcdB (Fig. 4D) concentration of R20291AWA-E and R20291AWA-W decreased by 37.9% 219 (P < 0.05) and 31.3% (P < 0.05), respectively. While TcdA concentration of R20291 $\Delta$ W-E, 220 R20291 $\Delta$ WA-A, and R20291 $\Delta$ W-W increased by 83.1% (*P* < 0.01), 64.7% (*P* < 0.05), and 56.5% 221 (P < 0.05), respectively. Meanwhile, TcdB concentration of R20291 $\Delta$ W-E increased by 100.2% 222 223 (P < 0.01). At 48 h post-inoculation, though no significant difference in TcdA production was detected among different C. difficile strains, TcdB concentration of R20291 $\Delta$ WA-A increased by 224 28.5% (*P* < 0.05) compared to R20291-E. 225

To analyze the transcription of *tcdA* and *tcdB* in the complementation strains, RT-qPCR was performed. As shown in Fig. 4E and 4D, the transcription of *tcdA* and *tcdB* of R20291 $\Delta$ WA-E and R20291 $\Delta$ WA-W decreased significantly (*P* < 0.05), while that of R20291 $\Delta$ W-E increased significantly (*P* < 0.05). Interestingly, the *tcdA* transcription of R20291 $\Delta$ WA-A also showed a significant increase (P < 0.05) compared to the wild type strain. Our data indicate that FliW negatively regulates toxin expression, while CsrA plays a positive regulation role in toxin expression.

# 233 Effects of *fliW* and *fliW-csrA* deletions on sporulation and germination

To assay the sporulation ratio of *C. difficile* strains, R20291, R20291 $\Delta$ WA, and R20291 $\Delta$ W were cultured in Clospore media for 48 and 96 h, respectively. Results (Fig. S4A) showed that no significant difference in the sporulation ratio was detected between the wild type strain and the mutants. The germination ratio of *C. difficile* spores was evaluated as well. Purified spores of R20291, R20291 $\Delta$ WA, and R20291 $\Delta$ W were incubated in the germination buffer supplemented with taurocholic acid (TA). As shown in Fig. S4B, there was no significant difference in the germination ratio between the wild type strain and the mutants.

# Evaluation of *fliW* and *fliW-csrA* deletions on bacterial virulence in the mouse model of CDI

To evaluate the effects of *fliW* and *fliW-csrA* deletions on *C. difficile* virulence *in vivo*, the mouse 243 model of CDI was used. Thirty mice (n=10 per group) were orally challenged with R20291, 244 R20291 $\Delta$ WA, or R20291 $\Delta$ W spores (1  $\times$  10<sup>6</sup> spores / mouse) after antibiotic treatment. As 245 246 shown in Fig. 5A, the R20291 $\Delta$ W infection group lost more weight at post challenge days 1 (P < 0.05) and the R20291 $\Delta$ WA infection group lost less weight at post challenge days 3 (P < 0.05) 247 compared to the R20291 infection group. Fig. 5B showed that 60% of mice succumbed to severe 248 249 disease within 4 days in the R20291AW infection group and 20% in the R20291AWA infection group compared to 50% mortality in the R20291 infection group (no significant difference with 250 251 log-rank analysis). Meanwhile, 100% of mice developed diarrhea in both the R20291 $\Delta$ W and 252 R20291 infection groups versus 80% in the R20291 $\Delta$ WA infection group at post challenge days

253 2 (Fig. 5C). As shown in Fig. 5D, the CFU of the R20291 $\Delta$ W infection group increased in the 254 fecal shedding samples at post challenge days 1 and 2 (*P* < 0.05), while the CFU of the 255 R20291 $\Delta$ WA infection group decreased at post challenge days 1, 5, and 6 (*P* < 0.05) compared 256 to the R20291 infection group.

To evaluate the toxin level in the gut, the concentration of TcdA and TcdB in the feces was 257 258 measured. In comparison with the R20291 infection group, the TcdA of the R20291 $\Delta$ W infection group increased significantly at post challenge days 1 (P < 0.05), 2 (P < 0.05), 3 (P < 0.01), and 259 5 (P < 0.05) (Fig. 5E). While the TcdA of the R20291 $\Delta$ WA infection group was decreased 260 significantly at post challenge days 1 (P < 0.05) and 4 (P < 0.05) (Fig. 5E). As shown in Fig. 5F, 261 the TcdB concentration of the R20291 $\Delta$ W infection group decreased significantly at post 262 challenge days 1 (P < 0.05), 2 (P < 0.05), and 3 (P < 0.05), and that of the R20291 $\Delta$ WA 263 264 increased significantly at post challenge days 1 (P < 0.05), 2 (P < 0.01), and 3 (P < 0.01). Taken together, our results indicate that the FliW defect increases R20291 pathogenicity in vivo, while 265 266 the *fliW-csrA* codeletion impairs R20291 pathogenicity.

### 267 **DISCUSSION**

In this study, we sought to characterize the impacts of FliW, CsrA, and FliC on C. difficile 268 269 pathogenicity. Our data suggest that CsrA negatively modulates *fliC* expression post-270 transcriptionally and FliW affects *fliC* expression possibly through inhibiting CsrA-mediated negative post-transcriptional regulation. Our data also indicate that FliW negatively affects C. 271 272 difficile pathogenicity possibly by antagonizing CsrA in vivo. Based on our current pleiotropic phenotype analysis, a similar partner-switching mechanism "FliW-CsrA-fliC/FliC" is predicted 273 274 in C. difficile, though more direct experimental data are needed to uncover the molecular 275 interactions of CsrA, FliW, and *fliC*/FliC in C. difficile (Fig. S5).

276 It has been reported that overexpression of the *csrA* gene could result in flagella defects, poor motility, and increased toxin production and adhesion in C. difficile 630 $\Delta$ erm (20). We 277 278 found that *fliW* and *csrA* genes are broadly found in the C. *difficile* genomes, among them 10 279 different C. difficile strains from ribotype 106 (RT106), RT027, RT001, RT078, RT009, RT012, RT046, and RT017 were selected and compared to R20291 (Table S2). CsrA and FliW widely 280 281 exist in C. difficile, even in the C. difficile strains without flagellar like C. difficile M120 (28), 282 indicating a potentially important role of FliW-CsrA in C. difficile. Interestingly, while there is 283 no flagellar in C. difficile M120, but 6 flagellar structure genes (fliS, fliN, flgK, flgL, fliC, and 284 *fliD*) are still found in the genome, which inspired us to explore the potential roles of *fliW*, *csrA*, and *fliC* in C. *difficile* by deleting or overexpressing *fliW*, csrA, and *fliW*-csrA genes. The 285 important roles of CsrA in flagellin synthesis and flagellin homeostasis have been reported (17-286 20). A previous study had shown that the overexpression of the *csrA* gene can cause a dramatic 287 motility reduction and a significant Hag decrease, suggesting that CsrA represses the Hag 288 expression (17). FliW (the first protein regulator of CsrA activity) deletion abolished the B. 289 subtilis swarming and swimming motility and decreased the number of flagella and flagellar 290 length (18, 29). In this study, we obtained similar results that FliW defect impaired R20291 291 292 motility significantly (Fig. 2A) and increased biofilm formation (Fig. 2C and 2D). Interestingly, the csrA gene complementation in R20291 $\Delta$ WA dramatically suppressed bacterial motility and 293 showed a similar result to R20291 $\Delta$ W. Inversely, the *fliW-csrA* codeletion increased R20291 294 295 motility. Meanwhile, no significant difference was detected between R20291AWA-W and R20291AWA, but there was a significant change between R20291AWA-W and R20291-E, 296 297 indicating that CsrA can suppress C. difficile motility and increase biofilm production, while 298 FliW needs to work together with *csrA* to regulate bacteria motility and biofilm formation.

The partner-switching mechanism "Hag-FliW-CsrA" on flagellin synthesis was elucidated 299 in B. subtilis and the intracellular concentration of the flagellar filament protein Hag is restricted 300 301 tightly by the Hag-FliW-CsrA system (18). To investigate whether FliW and CrsA coregulate the fliC expression in C. difficile, we evaluated both the transcriptional and translational expression 302 level of *fliC* gene. Our data (Fig. 3) showed that the *fliW* deletion resulted in a 10.4 fold decrease 303 304 of FliC accumulation, while the *fliW-csrA* codeletion increased FliC production, indicating that CsrA could suppress the *fliC* translation and FliW works against CsrA to regulate FliC 305 production. In csrA, fliW, and fliW-csrA overexpression experimental groups, we found that the 306 307 csrA overexpression dramatically decreased FliC production (5.3 fold reduction) and the reduction of FliC production in R20291-A can be partially recovered when *fliW-csrA* was 308 coexpressed. The FliW complementation in R20291AWA didn't affect FliC production, but the 309 *fliW* overexpression in R20291 increased FliC production. Taken together, our data suggest that 310 311 CsrA negatively modulates *fliC* expression post-transcriptionally and FliW works against CsrA to regulate *fliC* expression through inhibiting CsrA-mediated negative post-transcriptional 312 regulation, indicating a similar partner-switching mechanism "FliW-CsrA-FliC" in C. difficile 313 (Fig. S5). In B. subtilis, two CsrA binding sites (BS1: A51 to A55; BS2: C75 to G82) were 314 315 identified in the hag leader of the mRNA (17). Based on the hag 5' untranslated region (5'-UTR) sequence and CsrA conserved binding sequence, a 91 bp 5'-UTR structure with two potential 316 CsrA binding sites (BS1: 5'-TGACAAGGATGT-3', BS2: 5'-CTAAGGAGGG-3') of *fliC* gene 317 318 was predicted (Fig. S6) (30). Recently, it was also reported that cytoplasmic Hag levels play a central role in maintaining proper intracellular architecture, and the Hag-FliW-CsrA<sup>dimer</sup> system 319 320 works at nearly 1:1:1 stoichiometry(19). Further studies on the exquisite interactions of CsrA, 321 FliW, and *fliC*/FliC in *C. difficile* are still needed.

322 Flagella play multiple roles in bacterial motility, colonization, growth, toxin production, and survival optimization (21, 31, 32). Recently, several papers have reported that the flagellar genes 323 324 can affect toxin expression in C. difficile, but results from different research groups were 325 controversial (21-23). It was hypothesized that the regulation of the flagellar genes on toxin expression could be caused by the direct change or loss of flagellar genes (such as *fliC* gene 326 327 deletion) rather than loss of the functional flagella (21). Future study about *fliC* deletion in M120 will be very interesting and will further address the *fliC* gene function in *C*. *difficile* as there is no 328 329 flagellar in RT078 strains. In our study, our data indicate that CsrA negatively modulates *fliC* 330 expression and also plays a positive regulation in toxin expression. Inversely, FliW works against CsrA to regulate *fliC* expression which can negatively regulate toxin production. While 331 studies of flagellar effects on motility and toxin production in C. difficile from different groups 332 were controversial, the role of the flagella in C. difficile pathogenicity can not be overlooked. 333 Dingle et. al (33) and Baban et. al (23) both showed higher mortality of the *fliC* mutant in the 334 335 animal model of CDI compared to the wild type strains. Our study showed results similar to the published data suggesting that R20291AW whose FilC production was dramatically suppressed 336 exhibited higher fatality, while R20291 $\Delta$ WA showed a decreased pathogenicity compared to 337 338 R20291 (Fig. 5). In 2014, Barketi et al. (24) examined the pleiotropic roles of the *fliC* gene in R20291 during colonization in mice. Interestingly, the transcription of fliW and csrA in the fliC339 mutant was 2.03 fold and 4.36 fold, respectively, of that in R20291 in vivo experiment (24), 340 341 which further corroborated that there is a coregulation among *fliC*, *fliW*, and *csrA*. Surprisingly, transcription of *treA*, a trehalose-6-phosphate hydrolase, increased 177.63 fold (24). Recently, 342 343 Collins et al. (34) hypothesized that dietary trehalose can contribute to the virulence of epidemic 344 C. difficile. The relationship of FliW, CsrA, FliC, and trehalose metabolization is another 345 interesting question in C. difficile and some other carbon metabolism affected by the fliC mutation could also facilitate C. difficile pathogenesis in vivo. Previous studies have also 346 highlighted that the flagella of C. difficile play an important role in toxin production, biofilm 347 formation, and bacterial adherence to the host (22, 23, 25, 33, 35). In this study, we showed that 348 the FliW defect led to a significant motility decrease, while the biofilm, adhesion, and toxin 349 350 production increased significantly. Inversely, R20291 $\Delta$ WA-W, which can imitate the *csrA* gene deletion, showed an increase in motility and a decrease in biofilm formation, toxin production, 351 352 and adhesion (Fig. 2, Fig. S2, and Fig. S3).

In conclusion, we characterized the function of FliW and CsrA and showed the pleiotropic functions of FliW and CsrA in R20291. Our data suggest that *fliW* and *csrA* play important roles in flagellin (FliC) synthesis which could contribute to *C. difficile* pathogenicity. Currently, *in vitro* study of the interactions of CsrA, FliW, and *fliC*/FliC in *C. difficile* is underway in our group.

### 358 EXPERIMENTAL PROCEDURES

Bacteria, plasmids, and culture conditions Table 1 lists the strains and plasmids used in this 359 study. C. difficile strains were cultured in BHIS media (brain heart infusion broth supplemented 360 with 0.5% yeast extract and 0.1% L-cysteine, and 1.5% agar for agar plates) at 37 °C in an 361 anaerobic chamber (90% N<sub>2</sub>, 5% H<sub>2</sub>, 5% CO<sub>2</sub>). For spores preparation, C. difficile strains were 362 cultured in Clospore media and purified as described earlier (36). Escherichia coli DH5a and E. 363 364 coli HB101/pRK24 were grown aerobically at 37 °C in LB media (1% tryptone, 0.5% yeast extract, 1% NaCl). E. coli DH5a was used as a cloning host and E. coli HB101/pRK24 was used 365 366 as a conjugation donor host. Antibiotics were added when needed: for E. coli, 15  $\mu$ g/ml 367 chloramphenicol; for *C. difficile*, 15 μg/ml thiamphenicol, 250 μg/ml D-cycloserine, 50 μg/ml
368 kanamycin, 8 μg/ml cefoxitin, and 500 ng/ ml anhydrotetracycline.

### 369 **DNA manipulations and chemicals**

DNA manipulations were carried out according to standard techniques (37). Plasmids were 370 conjugated into C. difficile as described earlier (38). The DNA markers, protein markers, PCR 371 372 product purification kit, DNA gel extraction kit, restriction enzymes, cDNA synthesis kit, and 373 SYBR Green RT-qPCR kit were purchased from Thermo Fisher Scientific (Waltham, USA). 374 PCRs were performed with the high-fidelity DNA polymerase NEB Q5 Master Mix, and PCR 375 products were assembled into target plasmids with NEBuilder HIFI DNA Assembly Master Mix (New England, UK). Primers (Supporting Information Table S1) were purchased from IDT 376 (Coralville, USA). All chemicals were purchased from Sigma (St. Louis, USA) unless those 377 stated otherwise. 378

# 379 Construction of R20291 mutant strains of gene deletion, complementation, and 380 overexpression

The Cas12a (AsCpfI) based gene deletion plasmid pDL-1 was constructed and used for C. 381 difficile gene deletion (26). The target sgRNA was designed with an available website tool 382 383 (http://big.hanyang.ac.kr/cindel/) and the off-target prediction was analyzed on the Cas-OFFinder website (http://www.rgenome.net/cas-offinder/). The sgRNA, up and down homologous arms 384 were assembled into pDL-1. Two target sgRNAs for one gene deletion were selected and used 385 386 for gene deletion plasmid construction in C. difficile, respectively. Briefly, the gene deletion plasmid was constructed in the cloning host E. coli DH5a and was transformed into the donor 387 388 host E. coli HB101/pRK24, and subsequently was conjugated into R20291. Potential successful 389 transconjugants were selected with selective antibiotic BHIS-TKC plates (15 µg/ml 390 thiamphenicol, 50  $\mu$ g/ml kanamycin, 8  $\mu$ g/ml cefoxitin). The transconjugants were cultured in BHIS-Tm broth (15 µg/ml thiamphenicol) to log phase, then the subsequent cultures were plated 391 inducing plates (BHIS-Tm-ATc: 15 µg/ml thiamphenicol and 500 ng/ml 392 on the anhydrotetracycline). After 24 - 48 h of incubation, 20 - 40 colonies were used as templates for 393 colony PCR test with check primers for correct gene deletion colony isolation. The correct gene 394 395 deletion colony was sub-cultured into BHIS broth without antibiotics and was passaged several 396 times to cure the deletion plasmid, then the cultures were plated on BHIS plates and subsequent 397 colonies were replica plated on BHIS-Tm plates to isolate pure clean gene deletion mutants 398  $(R20291\Delta W \text{ and } R20291\Delta WA)$ . The genome of  $R20291\Delta W$  and  $R20291\Delta WA$  were isolated and used as templates for the PCR test with check primers, and the PCR products were 399 400 sequenced to confirm the correct gene deletion.

The *fliW* (396 bp) (primers 3-F/R), *csrA* (213 bp) (primers 4-F/R), and *fliW-csrA* (599 bp) 401 (primers 5-F/R) genes were amplified and assembled into SacI-BamHI digested pMTL84153 402 403 plasmid, yielding the complementation plasmid pMTL84153-fliW, pMTL84153-csrA, and pMTL84153-fliW-csrA, and were subsequently conjugated into R20291ΔWA, R20291ΔW, and 404 R20291 yielding complementation strain R20291\DeltaWA/pMTL84153-fliW (referred as 405 406 R20291∆WA-W), R20291*Δ*WA/pMTL84153-*csrA* (R20291∆WA-A), R20291\DeltaWA/pMTL84153-fliW-csrA (R20291∆WA-WA), R20291*Δ*W/pMTL84153*-fliW* 407 R20291/pMTL84153-fliW overexpression 408 (R20291∆W-W) and strain (R20291-W), 409 R20291/pMTL84153-csrA (R20291-A), R20291/pMTL84153-fliW-csrA (R20291-WA).

410 Growth profile, motility, and biofilm assay

411 *C. difficile* strains were incubated to an optical density of  $OD_{600}$  of 0.8 in BHIS media and were 412 diluted to an  $OD_{600}$  of 0.2. Then, 1% of the culture was inoculated into fresh BHIS, followed by 413 measuring  $OD_{600}$  for 32 h.

To examine the effect of *fliW* and *fliW-csrA* deletion on *C. difficile* motility, R20291, R20291 $\Delta$ WA, and R20291 $\Delta$ W were cultured to an OD<sub>600</sub> of 0.8. For swimming analysis, 2 µl of *C. difficile* culture was penetrated into soft BHIS agar (0.175%) plates, meanwhile, 2 µl of culture was dropped onto 0.3% BHIS agar plates for swarming analysis. The swimming assay plates were incubated for 24 h and the swarming plates were incubated for 48 h, respectively.

419 For biofilm formation analysis, wild type and mutant C. difficile R20291 strains were cultured to an OD<sub>600</sub> of 0.8, and 1% of C. difficile cultures were inoculated into Reinforced 420 Clostridial Medium (RCM) with 8 well repeats in a 96-well plate and incubated in the anaerobic 421 chamber at 37 °C for 48 h. Biofilm formation was analyzed by crystal violet dye. Briefly, C. 422 423 difficile cultures were removed by pipette carefully. Then 100 µl of 2.5% glutaraldehyde was added into the well to fix the bottom biofilm, and the plate was kept at room temperature for 30 424 min. Next, the wells were washed with PBS 3 times and dyed with 0.25% (w/v) crystal violet for 425 10 min. The crystal violet solution was removed, and the wells were washed 5 times with PBS, 426 427 followed by the addition of acetone into wells to dissolve the crystal violet of the cells. The dissolved solution was further diluted with ethanol 2 - 4 times and biomass was determined at 428 OD<sub>570</sub>. 429

## 430 Adherence of *C. difficile* vegetative cells to HCT-8 cells

431 *C. difficile* adhesion ability was evaluated with HCT-8 cells (ATCC CCL-244) (39). Briefly, 432 HCT-8 cells were grown to 95% confluence  $(2 \times 10^5/\text{well})$  in a 24-well plate and then moved into 433 the anaerobic chamber, followed by infecting with  $6 \times 10^6$  of log phase of *C. difficile* vegetative 434 cells at a multiplicity of infection (MOI) of 30:1. The plate was cultured at 37 °C for 30 min. 435 After incubation, the infected cells were washed with 300  $\mu$ l of PBS 3 times, and then suspended 436 in RPMI media with trypsin and plated on BHIS agar plates to enumerate the adhered *C. difficile* 437 cells. The adhesion ability of *C. difficile* to HCT-8 cells was calculated as follows: CFU of 438 adhered bacteria / Total cell numbers.

To visualize the adherence of C. difficile to HCT-8 cells, C. difficile vegetative cells were 439 labeled with the chemical 5(6)-CFDA (5-(and -6)-Carboxyfluorescein diacetate) (40). Briefly, C. 440 *difficile* strains were cultured to an  $OD_{600}$  of 0.8, then washed with PBS 3 times and resuspended 441 442 in fresh BHIS supplemented with 50 mM 5(6)-CFDA, followed by incubation at 37 °C for 30 min in the anaerobic chamber. After post-incubation, the labeled C. difficile cells were collected 443 and washed with PBS 3 times, and then resuspended in RPMI medium. Afterward, the labeled C. 444 difficile cells were used for the infection experiment as described above. After 30 min post-445 infection, the fluorescence of each well was scanned by the Multi-Mode Reader (excitation, 485 446 nm; emission, 528 nm), the relative fluorescence unit (RFU) was recorded as F0. Following, the 447 plates were washed with PBS 3 times to remove unbound C. difficile cells, then the plates were 448 scanned and the RFU was recorded as F1. The adhesion ratio was calculated as follows: F1/F0. 449 450 After scanning, the infected cell plates were further detected by the fluorescence microscope.

### 451 *fliC* expression assay

For *fliC* transcription analysis, 2 ml of 24 h post inoculated *C. difficile* cultures were centrifuged at 4 °C, 12000×g for 5 min, respectively. Then, the total RNA of different strains was extracted with TRIzol reagent. The transcription of *fliC* was measured by RT-qPCR with primers Q*-fliC*-F/R. All RT-qPCRs were repeated in triplicate, independently. Data were analyzed by the comparative CT ( $2^{-\Delta\Delta CT}$ ) method with 16s rRNA as a control. 457 To analyze the FliC protein level, C. difficile cell lysates from overnight cultures were used for Western blot analysis. Briefly, overnight C. difficile cultures were collected and washed 3 458 times with PBS and then resuspended in 5 ml of distilled water. The suspensions were lysed with 459 TissueLyser LT (Qiagen), followed centrifuged at  $4^{\circ}$ C,  $25000 \times g$  for 1h. The final pellets were 460 resuspended in 30 µl of PBS and the total protein concentration was measured by using a BCA 461 462 protein assay (Thermo Scientific, Suwanee, GA). Protein extracts were subjected to 10% SDS-PAGE. Sigma A protein (SigA) was used as a loading control protein in SDS-PAGE. FliC and 463 464 SigA proteins on the gel were detected with anti-FliC and anti-SigA primary antibody (1:1000) 465 and horseradish peroxidase-conjugated secondary antibody goat anti-mouse (Cat: ab97023, IgG, 1:3000, Abcam, Cambridge, MA) by Western blot, respectively. Anti-FliC antibody used in the 466 467 Western blot analysis is an anti-FliCD serum, generated in the lab. FliCD is a fusion protein containing *C.difficile* FliC and FliD (41). 468

### 469 Toxin expression assay

To evaluate toxin expression in *C. difficile* strains, 10 ml of *C. difficile* cultures were collected at 24 and 48 h post incubation. The cultures were adjusted to the same density with fresh BHIS. Then the collected *C. difficile* cultures were centrifuged at 4 °C,  $8000 \times g$  for 15 min, filtered with 0.22 µm filters, and used for ELISA. Anti-TcdA (PCG4.1, Novus Biologicals, USA) and anti-TcdB (AI, Gene Tex, USA) were used as coating antibodies for ELISA, and HRP-Chicken anti-TcdA and HRP-Chicken anti-TcdB (Gallus Immunotech, USA) were used as detection antibodies.

For toxin transcription analysis, 2 ml of 24 and 48 h post inoculated *C. difficile* cultures were centrifuged at 4 °C,  $12000 \times g$  for 5 min, respectively. Next, the total RNA of different strains was extracted with TRIzol reagent. The transcription of *tcdA* and *tcdB* was measured by

480 RT-qPCR with primers Q-*tcdA*-F/R and Q-*tcdB*-F/R, respectively. All RT-qPCRs were repeated 481 in triplicate, independently. Data were analyzed by using the comparative CT  $(2^{-\Delta\Delta CT})$  method 482 with 16s rRNA as a control.

# 483 Germination and sporulation assay

C. difficile germination and sporulation analysis were conducted as reported earlier (42). Briefly, 484 for C. difficile sporulation analysis, C. difficile strains were cultured in Clospore media for 4 days. 485 Afterward, the CFU of cultures from 48 and 96 h were counted on BHIS plates with 0.1% TA to 486 detect sporulation ratio, respectively. The sporulation ratio was calculated as CFU (65 °C heated) 487 488 / CFU (no heated). For C. difficile germination analysis, C. difficile spores were collected from 2- week Clospore media cultured bacteria and purified with sucrose gradient layer (50%, 45%, 489 35%, 25%, 10%). The heated purified spores were diluted to an  $OD_{600}$  of 1.0 in the germination 490 buffer [10 mM Tris (pH 7.5), 150 mM NaCl, 100 mM glycine, 10 mM taurocholic acid (TA)] to 491 detect the germination ratio. The value of  $OD_{600}$  was monitored immediately (0 min,  $t_0$ ), and was 492 detected once every 2 min (t<sub>x</sub>) for 20 min at 37 °C. The germination ratio was calculated as 493  $OD_{600}$  (tx) /  $OD_{600}$  (T<sub>0</sub>). Spores in germination buffer without TA were used as the negative 494 control. 495

# 496 Evaluation of R20291, R20291 $\Delta$ WA, and R20291 $\Delta$ W virulence in the mouse model of *C*. 497 *difficile* infection

C57BL/6 female mice (6 weeks old) were ordered from Charles River Laboratories, Cambridge,
MA. All studies were approved by the Institutional Animal Care and Use Committee of
University of South Florida. The experimental design and antibiotic administration were
conducted as described earlier (43). Briefly, 30 mice were divided into 3 groups in 6 cages.
Group 1 mice were challenged with R20291 spores, group 2 mice with R20291ΔWA spores, and

group 3 mice with R20291 $\Delta$ W spores, respectively. Mice were given an orally administered antibiotic cocktail (kanamycin 0.4 mg/ml, gentamicin 0.035 mg/ml, colistin 0.042 mg/ml, metronidazole 0.215 mg/ml, and vancomycin 0.045 mg/ml) in drinking water for 4 days. After 4 days of antibiotic treatment, all mice were given autoclaved water for 2 days, followed by one dose of clindamycin (10 mg/kg, intraperitoneal route) 24 h before spores challenge (Day 0). After that, mice were orally gavaged with 10<sup>6</sup> of spores and monitored daily for a week for changes in weight, diarrhea, and mortality.

# 510 Evaluation of *C. difficile* spores and determination of toxin level in feces

511 Fecal pellets from post infection day 0 to day 7 were collected and stored at -80 °C. To enumerate C. difficile numbers, feces were diluted with PBS at a final concentration of 0.1 g/ml, 512 followed by adding 900 µl of absolute ethanol into 100 µl of the fecal solution, and kept at room 513 temperature for 1 h to inactivate vegetative cells. Afterward, 200 µl of vegetative cells 514 inactivated fecal solution from the same group and the same day was mixed. Then, fecal samples 515 were serially diluted and plated on BHIS-CCT plates (250 µg/ml D-cycloserine, 8 µg/ml 516 cefoxitin, 0.1% TA). After 48 h incubation, colonies were counted and expressed as CFU/g feces. 517 To evaluate toxin tilter in feces, 0.1 g/ml of the fecal solution was diluted two times with PBS, 518 519 followed by examining TcdA and TCdB ELISA.

### 520 Statistical analysis

The reported experiments were conducted in independent biological triplicates except for the animal experiment, and each sample was additionally taken in technical triplicates. Animal survivals were analyzed by Kaplan-Meier survival analysis and compared by the Log-Rank test. One-way analysis of variance (ANOVA) with post-hoc Tukey test was used for more than two

groups comparison. Results were expressed as mean  $\pm$  standard error of the mean. Differences were considered statistically significant if *P* < 0.05 (\*).

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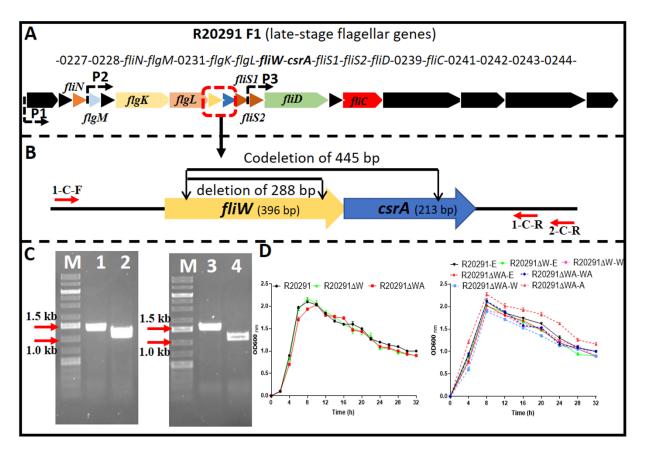
# **TABLES**

# 672 Table 1. Bacteria and plasmids utilized in this study

Strains or plasmids	Genotype or phenotype	Reference
Strains		
E. coli DH5a	Cloning host	NEB
E. coli HB101/pRK24	Conjugation donor	(44)
C. difficile R20291	Clinical isolate; ribotype 027	(45)
R20291∆W	R20291 deleted <i>fliW</i> gene	This work
R20291∆WA	R20291 deleted fliW-csrA genes	This work

R20291-E	R20291 containing blank plasmid pMTL84153	This work
R20291ΔW-E	R20291 $\Delta$ W containing blank plasmid pMTL84153	This work
R20291∆WA-E	R20291ΔWA containing blank plasmid pMTL84153	This work
R20291∆W-W	R20291 $\Delta$ W complemented with pMTL84153- <i>fliW</i>	This work
R20291∆WA-WA	R20291AWA complemented with pMTL84153-fliW-csrA	This work
R20291∆WA-W	R20291ΔWA complemented with pMTL84153-fliW	This work
R20291∆WA-A	R20291ΔWA complemented with pMTL84153-csrA	This work
R20291-W	R20291 containing pMTL84153-fliW	This work
R20291-A	R20291 containing pMTL84153-csrA	This work
R20291-WA	R20291 containing pMTL84153-fliW-csrA	This work
Plasmids		
<b>Plasmids</b> pDL1	AsCpfI based gene deletion plasmid	This work
	AsCpfI based gene deletion plasmid sRNA promoter template	This work This work
pDL1		
pDL1 pUC57-PsRNA	sRNA promoter template	This work
pDL1 pUC57-PsRNA pDL1 <i>-fliW</i>	sRNA promoter template fliW gene deletion plasmid	This work This work
pDL1 pUC57-PsRNA pDL1 <i>-fliW</i> pDL1- <i>csrA</i>	sRNA promoter template <i>fliW</i> gene deletion plasmid <i>csrA</i> gene deletion plasmid	This work This work This work
pDL1 pUC57-PsRNA pDL1 <i>-fliW</i> pDL1- <i>csrA</i> pDL1- <i>fliW-csrA</i>	sRNA promoter template <i>fliW</i> gene deletion plasmid <i>csrA</i> gene deletion plasmid <i>fliW-csrA</i> gene deletion plasmid	This work This work This work This work
pDL1 pUC57-PsRNA pDL1 <i>-fliW</i> pDL1 <i>-csrA</i> pDL1 <i>-fliW-csrA</i> pMTL84153	sRNA promoter template <i>fliW</i> gene deletion plasmid <i>csrA</i> gene deletion plasmid <i>fliW-csrA</i> gene deletion plasmid Complementation plasmid	This work This work This work This work (46)
pDL1 pUC57-PsRNA pDL1 <i>-fliW</i> pDL1 <i>-csrA</i> pDL1 <i>-fliW-csrA</i> pMTL84153 pMTL84153 <i>-fliW-csrA</i>	sRNA promoter template <i>fliW</i> gene deletion plasmid <i>csrA</i> gene deletion plasmid <i>fliW-csrA</i> gene deletion plasmid Complementation plasmid pMTL84153 containing <i>fliW-crsA</i> genes	This work This work This work (46) This work

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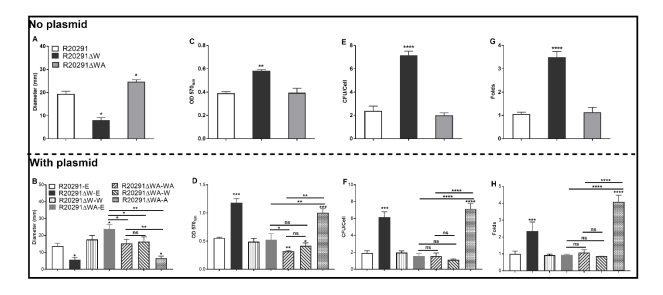


**Fig. 1. R20291** late-stage flagellar genes (F1) and *fliW* and *fliW-csrA* deletions

(A) Schematic representation of late-stage flagellar genes (F1). Dotted arrows (P1, P2, and P3) 693 694 indicate the potential promoters in F1. (B) Deletion of *fliW* and *fliW-csrA* genes. 1-C-F/R were used to verify *fliW* deletion, and 1-C-F and 2-C-R were used to test *fliW-csrA* codeletion. (C) 695 Verification of *fliW* and *fliW-csrA* deletions by PCR. M: DNA ladder; 1: R20291 genome as 696 697 PCR template; 2: R20291∆W genome as PCR template; 3: R20291 genome as PCR template; 4: R20291 $\Delta$ WA genome as PCR template. (D) Growth profile of parent strain and gene deletion 698 mutants. Experiments were independently repeated thrice. Bars stand for mean  $\pm$  SEM. One-way 699 700 ANOVA with post-hoc Tukey test was used for statistical significance.

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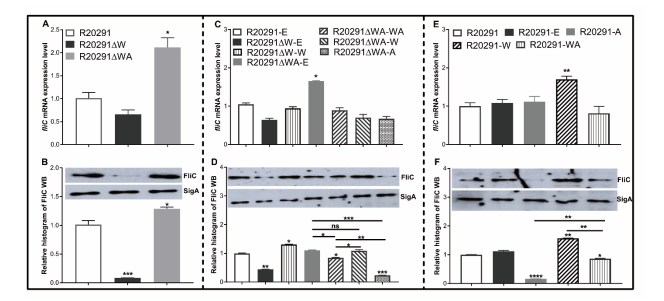
# Fig. 2. Motility, biofilm, and adhesion analysis

(A) and (B): Halo diameter of motility (swimming analysis on 0.175% agar plate). (C) and (D): 706 Biofilm formation analysis. (E) an (F): Adherence of C. difficile vegetative cells to HCT-8 cells 707 in vitro. (G) and (H): Adhesion analysis with 5(6)-CFDA dye. The fluorescence intensity was 708 scanned by the Multi-Mode Reader (excitation, 485 nm; emission, 528 nm). The original relative 709 fluorescence unit (RFU) was recorded as F0, after PBS wash, the RFU was recorded as F1. The 710 adhesion ratio was calculated as follows: F1/F0. Experiments were independently repeated thrice. 711 712 Bars stand for mean  $\pm$  SEM (\*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001, \*\*\*\*P < 0.0001). One-way ANOVA with post-hoc Tukey test was used for statistical significance. \* directly upon the 713 column means the significant difference of the experimental strain compared to R20291 or 714 715 R20291-E. 716

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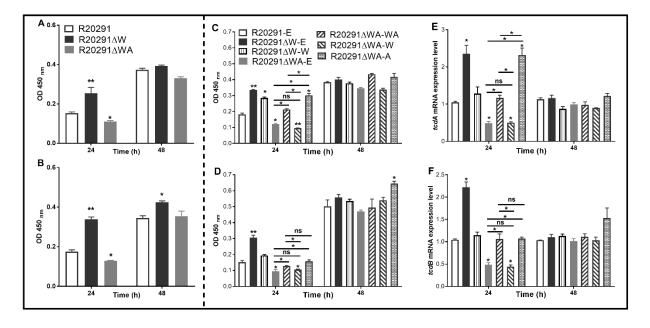
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# 722 Fig. 3. *fliC* expression analysis

(A), (C), and (E) Analysis of *fliC* expression on transcription level. (B), (D), and (F) Analysis of *fliC* expression on translation level by Western blot. SigA protein was used as a loading control. Experiments were independently repeated thrice. Bars stand for mean  $\pm$  SEM (\**P* < 0.05, \*\**P*<0.01, \*\*\**P* < 0.001, \*\*\*\**P* < 0.0001). One-way ANOVA with post-hoc Tukey test was used for statistical significance. \* upon the column directly means the significant difference of experimental strain compared to R20291 or R20291-E.

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### 733 Fig. 4. Toxin expression analysis

(A) TcdA concentration in the supernatants of R20291, R20291 $\Delta$ WA, and R20291 $\Delta$ W. (B) 734 TcdB concentration in the supernatants of R20291, R20291∆WA, and R20291∆W. (C) TcdA 735 concentration in the supernatants of parental and gene complementation strains. (D) TcdB 736 concentration in the supernatants of parental and gene complementation strains. (E) 737 Transcription of *tcdA* in the supernatants of parental and gene complementation strains. (F) 738 739 Transcription of *tcdB* in the supernatants of parental and gene complementation strains. Experiments were independently repeated thrice. Bars stand for mean  $\pm$  SEM (\*P < 0.05, \*\*P <740 0.01). One-way ANOVA with post-hoc Tukey test was used for statistical significance. \* upon 741 742 the column directly means the significant difference of experimental strain compared to R20291 or R20291-E. 743

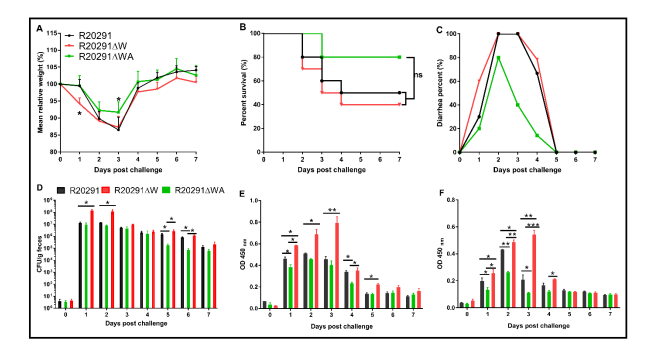
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751 Fig. 5. Effects of *fliW* and *fliW-csrA* deletion on *C. difficile* virulence in mice

(A) Mean relative weight changes. (B) Survival curve. (C) Diarrhea percentage. (D) *C. difficile* in feces. (E) TcdA titer of fecal sample. (F) TcdB titer of fecal sample. Bars stand for mean  $\pm$ SEM (\**P* < 0.05, \*\**P* < 0.01). One-way ANOVA with post-hoc Tukey test was used for statistical significance. Animal survivals were analyzed by Kaplan-Meier survival analysis with a log-rank test of significance.

