A mathematical model of flagellar gene regulation and construction in *Salmonella enterica*

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Abstract

Millions of people worldwide develop foodborne illnesses caused by *Salmonella enterica* (*S. enterica*) every year. The pathogenesis of *S. enterica* depends on flagella, which are appendages that the bacteria use to move through the environment. Interestingly, populations of genetically identical bacteria exhibit heterogeneity in the number of flagella. To understand this heterogeneity and the regulation of flagella quantity, we propose a mathematical model that connects the flagellar gene regulatory network to flagellar construction. A regulatory network involving more than 60 genes controls flagellar assembly. The most important member of the network is the master operon, *flhDC*, which encodes the FlhD⁴C₂ protein. FlhD⁴C₂ controls the construction of flagella by initiating the production of hook basal bodies (HBBs), protein structures that anchor the flagella to the bacterium. By connecting a model of FlhD⁴C₂ regulation to a model of HBB construction, we investigate the roles of various feedback mechanisms. Analysis of our model suggests that a combination of regulatory mechanisms at the protein and transcriptional levels induce bistable FlhD⁴C₂ levels and heterogeneous numbers of flagella. Also, the balance of regulatory mechanisms that become active following HBB construction is sufficient to provide a counting mechanism for controlling the total number of flagella produced.

Author summary

*Salmonella* causes foodborne illnesses in millions of people worldwide each year. Flagella, which are appendages that the bacteria use to move through the environment, are a key factor in the infection process. Populations of genetically identical bacteria have been observed to contain both motile cells, generally with 6–10 flagella, and nonmotile cells, with no flagella. In this paper, we use mathematical models of the gene network that regulates flagellar construction to explore how the bacteria controls the number of flagella produced. We suggest that a bacterium must accumulate a threshold amount of a master regulator protein to initiate flagella production and failure to reach the threshold results
in no flagella. Downstream mechanisms that impact the amount of master regulator protein are sufficient to determine how many flagella are produced.

**Introduction**

Millions of people worldwide develop foodborne illnesses caused by *Salmonella enterica* (*S. enterica*) every year [1]. The pathogenesis of *S. enterica* depends on flagella, which are appendages that the bacteria use to move through the environment. Flagella are located peritrichously on the bacterium’s surface and generally number between 6 and 10 per cell [2]. However, not all bacteria produce flagella. Partridge and Harshey describes the distributions of the number of flagella per bacterium under different growth conditions [3]. It was observed under each growth condition that some bacteria did not produce any flagella, while the number of flagella on bacteria that produced them roughly follows a normal distribution, where the mean depended on the growth condition. This phenotypic heterogeneity enables the evasion of the host immune system during acute infection [4]. It is not yet clear what mechanisms determine flagella heterogeneity and how the bacteria regulate the number of flagella produced.

Structurally, a flagellum is divided into three parts: basal body, hook, and filament. The basal body anchors the flagellum to the cell membrane, the hook is a flexible joint that links the rigid basal body with the filament, and the filament is a rigid helical structure which rotates to drive the cell forward. Assembly of functional flagella involves the temporally coordinated expression of more than 60 genes [5]. In particular, these genes are divided into three classes: Class I, Class II, and Class III depending on the timing of their activation. The *flhDC* operon is the only member of Class I. This operon encodes the master regulator proteins FlhD and FlhC, which form a transcriptional activation complex FlhD$_4$C$_2$ [6]. The FlhD$_4$C$_2$ complex is responsible for activation of Class II genes. Class II promoters control the expression of all the proteins required for the assembly of a functional hook-basal body (HBB) structure [7]. The HBB includes the flagellar type III secretion system, which exports flagellar proteins from the cytoplasm through the growing structure during assembly [8]. Once the HBB assembly is complete, FlhD$_4$C$_2$ activates expression of *flgM*, *fliA*, and other Class III genes. *fliA* encodes the flagella-specific sigma factor, $\sigma^{28}$, which is essential for Class III promoter activation. *flgM* encodes an anti-sigma factor FlgM. Prior to HBB completion FlgM and $\sigma^{28}$ form a complex that inhibits the activity of $\sigma^{28}$. After the HBB is completed, the specificity of the secretion apparatus switches, so filament subunits and FlgM are secreted from the cell. $\sigma^{28}$ facilitates the secretion of FlgM through the HBB, thereby acting as the FlgM type III secretion chaperone [9]. The secretion of FlgM frees $\sigma^{28}$ inside the cell, which leads to the activation of Class III genes. Class III genes control the assembly of the filament, the chemotaxis system, and motor proteins.

While the components of the gene network have been identified, understanding the additional details and dynamics of the system requires further study. In particular, genetically identical bacteria grown in the same environment exhibit bistable expression of multiple flagellar genes [3]. Expression of Class II and Class III genes, *flhB* and *fliC*, respectively, is bimodal, and this bimodality is likely due to bistability upstream in the network [10–13]. Other factors also impact the flagella gene network, including nutrient availability, which enhances the expression of flagellar and motility genes [14]. Nutrients repress RfIP, formerly YdiV, expression through the protein CsrA [15]. Low nutrient levels correspond to high RfIP expression, and high nutrient levels correspond to low RfIP expression. The tuning of RfIP expression
influences the flagellar gene network because RflP promotes the degradation of FlhD$_4$C$_2$ through the protease ClpXP [16].

A series of mathematical models have been proposed to explore the dynamics of the flagellar gene network [17–19]. While these models investigated the roles of some of the feedback mechanisms in the network, they fail to describe the heterogeneity observed in bacterial populations, since all bacteria are predicted to grow flagella. Further, the number of HBBs was modeled as a continuous variable, although HBBs are discrete entities.

Meanwhile, Koirala et al. presented a model that attempted to explain how nutrients tune flagellar gene expression dynamics in *Salmonella enterica* [13]. Using experimental observations and a deterministic model, the authors demonstrated that environmental nutrient levels can regulate RflP expression and that this expression can lead to bistable expression of Class II and Class III genes in the bacteria [13]. The model does not include HBB construction, or, consequently, the additional feedback mechanisms that are activated following HBB completion.

To investigate the heterogeneous flagella quantities within populations of clonal bacteria, we propose two mathematical models of the flagellar gene regulatory network. We first consider a sub-system of the network to focus on FlhD$_4$C$_2$ production and degradation. We use bifurcation analysis of an ordinary differential equation model to explore the role of flhDC and rflP expression rates on overall FlhD$_4$C$_2$ concentration. We find bistability in FlhD$_4$C$_2$ concentration as a function of both expression rates. We then extend the model to consider HBB construction and additional regulatory mechanisms. We retain the bistability observed in the sub-system model and investigate factors that impact the bistable regime. We also propose a mechanism for how *S. enterica* controls the number of flagella produced via FliT:FliD dynamics and Class III regulation of FliZ.

**Models**

**Sub-system model**

We first consider a model of the key components that regulate flhDC expression and FlhD$_4$C$_2$ protein concentrations. The model includes the FliZ-RflP-FlhD$_4$C$_2$ feedback loop and RflM activity. Expression of fliZ is up-regulated by FlhD$_4$C$_2$ and FliZ represses expression of rflP [20]. RflP inhibits the activity of FlhD$_4$C$_2$ by binding to the FlhD subunit and targeting FlhD$_4$C$_2$ for proteolysis via ClpXP [16]. Thus, FliZ is involved in a negative-negative feedback loop through RflP on the FlhD$_4$C$_2$ protein. FlhD$_4$C$_2$ up-regulates expression of rflM, and RflM represses the transcription of FlhD$_4$C$_2$ [21–23]. Fig 1 summarizes the mechanisms included in this model.

We assume that on the time scale of flagellar gene regulation and construction, the bacterial cytoplasm is well-mixed. The diffusion coefficients for proteins in the bacterial cytoplasm range from 1–10 $\mu$m$^2$s$^{-1}$, and since *S. enterica* are 2–5 microns long, proteins can diffuse across the bacterium on the order of seconds [24, 25]. Since HBBs are completed 30 minutes after induction of flhDC expression and nascent flagellar filaments become visible at 45 minutes, spatial non-uniformity of the proteins will not be significant [2].

Using the well-mixed assumption, we set $x = [\text{FlhD}_4C_2]$, $z = [\text{FliZ}]$, $y = [\text{RflP}]$, $c_1 = [\text{RflP}: \text{FlhD}_4C_2]$ and $m = [\text{RflM}]$, and we construct the following model

$$\frac{dx}{dt} = \frac{\rho_1}{1 + \eta_1 m} - \kappa_1 xy - \gamma_1 x$$  (1)
We model transcription and translation processes as Michaelis-Menten functions and protein binding, unbinding, and degradation processes linearly. FlhD$_4$C$_2$ is produced at a basal rate of $\rho_1$, and the production rate is divided by the concentration of RflM, scaled by $\eta_1$. FlhD$_4$C$_2$ binds to RflP at rate $\kappa_1$. FliZ is produced as a function of FlhD$_4$C$_2$ concentration. RflP is produced at a basal rate of $\rho_2$, and the production is repressed by FliZ. External nutrient availability also represses $rflP$ expression, so $\rho_2$ is a proxy for nutrient availability [15]. RflP forms a complex with FlhD$_4$C$_2$ and is recycled at rate $\delta_1$ where FlhD$_4$C$_2$ is degraded by ClpXP. We assume that the complex does not dissociate spontaneously. RflM is produced as a function of FlhD$_4$C$_2$ concentration. All components, including the RflP:FlhD$_4$C$_2$ complex, are assumed to degrade at a small, basal rate, $\gamma_i$, where $i = 1, 2, \ldots, 5$. Table 1 lists the model parameters with descriptions and values.

**Bifurcation analysis of sub-system model.** Since some bacteria produce flagella and others do not, we suspect that FlhD$_4$C$_2$ concentration is bistable. We use resultant analysis of the model to determine what parameter regimes support bistability [26]. To apply resultant analysis, we take the model to steady state, reduce the system to a single polynomial in one variable, and perform a rescaling to reduce the number of parameters. We then solve the equation for a
rescaled parameter of interest. To find double roots of the first derivative, we then take the resultant of the first and second derivatives of the right side of the equation. The curve in parameter space on which the first and second derivatives have a common root corresponds to the separation between monotonic and triphasic behavior. We first take the model to steady state and find

\[ y = \frac{1}{\kappa_1 x} \left( \frac{\rho_1}{1 + \eta_m} - \gamma_1 x \right) \]

\[ z = \frac{\gamma_2 x}{\gamma_2 (k_1 + x)} \]

\[ c_1 = \frac{\kappa_1 x y}{\delta_1 + \gamma_4} \]

\[ m = \frac{\gamma_4 x}{\gamma_5 (k_2 + x)} . \]

Substituting into \( 0 = \frac{\rho_2 y \gamma_2 (k_1 + x)}{\gamma_2 (k_1 + x) + \gamma_2 \eta_2 x} - \left( \frac{\gamma_4}{\delta_1 + \gamma_4} + \frac{\gamma_3}{\kappa_1 x} \right) \left( \frac{\rho_1 \gamma_2 (k_2 + x)}{\gamma_3 (k_2 + x) + \eta_2 x} - \gamma_1 x \right) \), we find

We reduce the number of parameters by taking the rescaling \( x = k_1 w \) and letting \( K = \frac{k_1}{k_2} \),

\[ A_2 = \frac{\gamma_2}{\gamma_5}, \quad P_1 = \frac{\rho_1}{\delta_1}, \quad D = \frac{\gamma_1 \gamma_2 k_1}{(\delta_1 + \gamma_4) \gamma_3}, \quad P_2 = \frac{\gamma_1 \rho_2}{\delta_1 \gamma_3}, \quad \text{and} \quad A_1 = \frac{\gamma_2}{\gamma_5}. \]

We find

\[ 0 = \frac{P_2 w (1 + w)}{1 + w + A_1 w} - (Dw + 1) \left( \frac{P_1}{1 + A_2 w} - w \right). \]
We solve for $P_2$ to find
\[
P_2 = \frac{(1 + w + A_1 w)(Dw + 1)}{w(1 + w)} \left( \frac{P_1}{1 + A_2 \frac{w}{w + D}} - w \right).
\]

We define $f(w)$ as
\[
f(w) = \frac{(1 + w + A_1 w)(Dw + 1)}{w(1 + w)} \left( \frac{P_1}{1 + A_2 \frac{w}{w + D}} - w \right).
\]

We next use resultant analysis of $f^3(w)$ and $f''(w)$ to determine when $f(w)$ is monotone and when it is triphasic. The triphasic regions correspond to bistability, and the monotone regions correspond to monostability.

We first consider the case in which $\eta_1 = 0$ ($A_2 = 0$), which means that RflM does not impact $flhDC$ expression. $f(w)$ simplifies to
\[
f_{fA_2=0}(w) = \frac{(1 + w + A_1 w)(Dw + 1)(P_1 - w)}{w(1 + w)}.
\]

We next calculate the resultant of the numerators of the first and second derivatives of $f_{fA_2=0}(w)$ using Maple (2019; Maplesoft, Waterloo, ON), and we find
\[
R_{f,A_2=0} = C_3 A_1^3 + C_2 A_1^2 + C_1 A_1 + C_0,
\]
where
\[
C_3 = (-1 + (D - 1)P_1)^3,
C_2 = -3(D - 1)^2 P_1^3 + (-3D^3 - 21D^2 + 30D - 6)P_1^2 + (-21D^2 + 21D - 3)P_1 - 3D,
C_1 = (3D - 3)P_1^3 + (-21D^2 + 21D - 3)P_1^2 + (3D^2 - 30D + 21D)P_1 - 3D^2,
C_0 = -P_1^3 - 3P_1^2 D + 3P_1 D^2 - D^3.
\]

The function $R_{f,A_2=0} = 0$ separates parameter space into regions for which $f_{fA_2=0}(w)$ is monotone increasing ($R_{f,A_2=0} < 0$) and triphasic ($R_{f,A_2=0} > 0$). In the triphasic region, there is bistability, which will be verified later. The solid curve in Fig 2 shows the zero level contour for $R_{f,A_2=0}$.

$R_{f,A_2=0}$ is negative to the left of the curve and is positive to the right of the curve. $f(w)$ is monotone for values of $P_1$ and $A_1$ that fall to the left of the curve, whereas for values that fall to the right of the curve, $f(w)$ is triphasic, and there is bistability. Thus, bistability is possible under constitutive expression of $flhDC$ (i.e. when $A_2 = 0$).

We note that in the case that $D = 0$, the resultant reduces to
\[
R_{f,A_2=0,D=0} = -16A_1^2 (P_1 + 1)^5 (A_1 P_1 + A_1 + P_1)^3 P_1^2.
\]

For positive values of $A_1$ and $P_1$, $R_{f,A_2=0,D=0}$ is always negative, and either $A_1 = 0$ or $P_1 = 0$ satisfy $R_f = 0$. This indicates that we must have $D > 0$ to have bistability. In terms of the original parameters, this means that $\gamma_4 > 0$ is required to have bistability.

We next consider the case in which $\eta_1 \neq 0$ ($A_2 \neq 0$), meaning RflM represses the expression of $flhDC$. We calculate the resultant of the numerators of the first and second derivatives of $f(w)$, but it is too long to show here. The dashed curve in Fig 2 is the zero level contour for $R_f$. 

Flagellar regulation
when RfM activity is included. We see that the $R_f = 0$ curve moves up and to the right as $A_2$ increases, which decreases the size of the parameter space in which there is bistability. In particular, the repressive activity of RfM requires an increase in the baseline production rates of FlhD$_4$C$_2$ ($P_1$) and FliZ ($A_1$) to have bistability.

We next use the parameters in Table 1 and XPPAUT (8.0) to visualize the bistable regime identified by the resultant analysis. Fig 3 shows the steady state concentration of FlhD$_4$C$_2$ plotted as a function of the parameters $\rho_1$ and $\rho_2$, demonstrating bistability for certain regions of

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**Fig 2.** Zero level contours of $R_f$, where $D = 4$ and $K = 0.2$. The solid curve indicates the $A_2 = 0$ case, and the dashed curve indicates the $A_2 = 0.2$ case. $R_f < 0$ below and to the left of the curve and $R_f > 0$ above and to the right of the curve.

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**Fig 3.** Steady state solution for concentration of FlhD$_4$C$_2$ (black) and FlhD$_4$C$_2$ production rate (blue) plotted as functions of $\rho_1$ and $\rho_2$. (A) Bistability is exhibited as $\rho_1$ varies, where $\rho_2 = 30$. (B) Bistability is also found as $\rho_2$ varies and $\rho_1 = 45$ is fixed.

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\( \rho_1, \rho_2 \) parameters. Fig 3 also shows that the value of the FlhD-C2 production term, \( \frac{\rho}{1 + \eta_1 \gamma_4} \), is multivalued for the same regions of \( \rho_1 \) and \( \rho_2 \).

Depending on the values of \( \rho_1 \) and \( \rho_2 \), the steady state value of FlhD-C2 concentration has one solution or three solutions. As shown in Fig 3A, the FlhD-C2 concentration is low for small \( \rho_1 \) values. This means that the bacteria do not produce enough FlhD-C2 to initiate flagella production. With \( \rho_1 \) increased sufficiently, a saddle node bifurcation occurs, and bistability arises. The lower solution represents the case in which the bacterium produces no flagella, and the upper solution represents the case in which the bacterium produces flagella. Increasing \( \rho_1 \) further results in another saddle node bifurcation, and a return to a single steady state value for FlhD-C2 concentration. The expectation is that for high flhDC production rates, the bacterium will have a high FlhD-C2 concentration, and, consequently, produce flagella. Saddle node bifurcations are also observed for \( \rho_2 \), as shown in Fig 3B. In this case, small \( \rho_2 \) values result in a high steady state FlhD-C2 concentration, intermediate \( \rho_2 \) values result in bistability, and large \( \rho_2 \) values result in a low FlhD-C2 concentration.

Many factors control flhDC expression, so it is reasonable to expect that clonal bacteria may have different values for \( \rho_1 \). In this case, the variation in \( \rho_1 \) values could contribute to heterogeneity in flagella number, assuming the bacteria on the lower branch have no flagella and bacteria on the upper branch have enough FlhD-C2 to produce flagella. Also, since nutrient availability represses rflP expression, small variations in nutrient availability could result in differing values for \( \rho_2 \). Variability in \( \rho_2 \) would result in different concentrations of FlhD-C2, and, consequently, the presence or absence of flagella. We note that, as with the case \( A_1 = 0, \gamma_4 > 0 \) is required for bistability.

We also consider the \( \eta_1 = 0 \) case, which corresponds to a knockout of RflM or constitutive expression of flhDC via an inducible promoter. Under this condition, the FlhD-C2 production term reduces to \( \rho_1 \). Fig 4 shows the steady state concentration of FlhD-C2 is bistable for certain regions of \( \rho_1 \) and \( \rho_2 \), but the FlhD-C2 production term is not multivalued.

In this case, constitutive expression of flhDC can lead to bistable FlhD-C2 levels. In particular, the \( \eta_1 = 0 \) case shows bistability in FlhD-C2 even when the production of flhDC is not multivalued. Therefore, we suggest that RflM knockout cells should still exhibit bistability in the protein level of FlhD-C2.

\[ \text{Fig 4. Steady state solution for concentration of FlhD-C2 (black) and FlhD-C2 production rate (blue), where } \eta_1 = 0. \text{ Bistability is exhibited for FlhD-C2 concentration, but not FlhD-C2 production, as } \rho_1 \text{ varies, where } \rho_2 = 30 \text{ (A), and as } \rho_2 \text{ varies and } \rho_1 = 49.5 \text{ (B).} \]

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In anticipation of a need to understand the role of FlhD(C)_2 degradation, we next investigate the impact of the basal degradation rate of FlhD(C)_2 on FlhD(C)_2 concentration. Fig 5A shows bistability in FlhD(C)_2 concentration as a function of the parameter γ_1. A saddle node bifurcation is observed for FlhD(C)_2 concentration as a function of γ_1. Small γ_1 values result in bistable steady state FlhD(C)_2 concentration, and increasing γ_1 sufficiently results in a low FlhD(C)_2 concentration. We next investigate the impact of γ_1 on the bistability observed in FlhD(C)_2 as a function of ρ_2. Fig 5B shows a cusp bifurcation in the γ_1-ρ_2 plane. This demonstrates that the bistability previously observed in FlhD(C)_2 as a function of ρ_2 is sensitive to γ_1.

Extended model

To explore how the bacteria regulate the total number of flagella constructed, we extend the model to consider the construction of HBBs and additional regulatory mechanisms that become active following HBB completion. In particular, FlhD(C)_2 up-regulates the flidST operon, which encodes the FliT and FliD proteins. Before HBB completion, FliT and FliD are bound together, which inhibits the activity of FliT. Following HBB completion, FliD is secreted through the HBB and is assembled as the cap of the flagellum. The secretion of FliD frees FliT in the cell to bind to FlhD(C)_2 and target FlhD(C)_2 for proteolysis via ClpXP [27–29]. FliT is recycled during FlhD(C)_2 degradation [27]. FlhD(C)_2 also up-regulates production of FlgM and σ^{28}, which form a complex, and complete HBBs secrete FlgM, resulting in free σ^{28} in the bacterium. σ^{28} up-regulates the expression of Class III genes, including flIC. FlIC is also secreted through the HBB to become the flagellar filament. σ^{28} also up-regulates flIZ, which means flIZ is under both Class II and Class III regulation. The gene network for the full model is shown in Fig 6.

For simplicity, we assume that FliT and FliD are produced as a complex (FliT:FliD), rather than modeling the production of each protein and formation of the FliT:FliD complex. Similarly, σ^{28} and FlgM are assumed to be produced as the complex σ^{28}:FlgM.
We use the variables defined in the sub-system model and set $v = [\text{FliT}], c_2 = [\text{FliT}:\text{FlhD}_4\text{C}_2], c_3 = [\text{FliT}:\text{FliD}], c_4 = [\alpha^{28}:\text{FlgM}], a = [\alpha^{28}],$ and $f = [\text{FliC}].$ Referring to Fig 6, we construct the following model

$$\frac{dx}{dt} = \frac{\rho_1}{1 + \eta_1 m} - \kappa_{1}xy - \kappa_{2}xv - \gamma_{1}x$$  \hspace{1cm} (6)$$

$$\frac{dz}{dt} = \frac{\alpha_{2}x}{k_{1} + x} + \frac{\beta_{1}a}{k_{s} + a} - \gamma_{2}z$$  \hspace{1cm} (7)$$

$$\frac{dy}{dt} = \frac{\rho_2}{1 + \eta_{2}z} - \kappa_{3}xy + \delta_{1}c_{1} - \gamma_{3}y$$  \hspace{1cm} (8)$$

$$\frac{dc_{1}}{dt} = \kappa_{4}xy - \delta_{1}c_{1} - \gamma_{4}c_{1}$$  \hspace{1cm} (9)$$

$$\frac{dm}{dt} = \frac{\alpha_{2}x}{k_{2} + x} - \gamma_{5}m$$  \hspace{1cm} (10)$$

$$\frac{dv}{dt} = f = \frac{\kappa_{5}c_{3}}{k_{3} + c_{3} + c_{4} + f} - \kappa_{2}xv + \delta_{2}c_{2} - \gamma_{6}v$$  \hspace{1cm} (11)$$

$$\frac{dc_{2}}{dt} = \kappa_{4}xy - \delta_{2}c_{2} - \gamma_{7}c_{2}$$  \hspace{1cm} (12)$$
\[
\frac{dc_3}{dt} = \frac{\alpha_3 x}{k_3 + x} - f \frac{\sigma c_3}{k_3 + c_3 + f} - \gamma_4 c_3
\] (13)

\[
\frac{dc_4}{dt} = \frac{\alpha_4 x}{k_4 + x} - f \frac{\sigma c_4}{k_4 + c_4 + f} - \gamma_4 c_4
\] (14)

\[
\frac{da}{dt} = f \frac{\sigma c_4}{k_4 + c_4 + f} - \gamma_4 a
\] (15)

\[
\frac{df}{dt} = \beta_2 a - f \frac{\sigma f}{k_i + c_4 + f} - \gamma_4 f
\] (16)

As before, we assume that the system is well-mixed, and we model transcription and translation processes as Michaelis-Menten functions and protein binding, unbinding, and degradation processes linearly. \(J\) is the number of complete HBBs that are capable of secreting FlgM, FliD, and FliC, and we assume that the secretion of the three components, FlgM, FliD, and FliC, through the HBBs is competitive and saturating. We assume that these three components have the same secretion affinity, \(\sigma\). As in the sub-system model, we assume that every compound, including the complexes, degrade at some basal rate.

**Model reduction.** The model involves both protein level dynamics, including binding and degradation, and transcription and translation. We assume that the protein level dynamics occur on a faster time scale than transcription and translation, so we take a quasi-steady-state approximation for the protein level dynamics. We assume that \(c_1 = [RflP:FlhD] C_2\), \(y = [RflP]\), and \(c_2 = [FliT:FlhD] C_2\) are fast equilibrating, so that they are at equilibrium. We therefore assume that

\[
k_1 x y - \delta_1 c_1 - \gamma_4 c_1 = 0,
\]

meaning that

\[
c_1 = \frac{k_1 x y}{\delta_1 + \gamma_4}.
\]

We also assume that

\[
\frac{\rho_2}{1 + \eta_2 x} - k_1 x y + \delta_1 c_1 + \gamma_4 y = 0,
\]

meaning that

\[
y = \frac{\rho_2}{1 + \eta_2 x} \left( \frac{1}{\gamma_3 + k_1 x \left( 1 - \frac{\delta_1}{\delta_1 + \gamma_4} \right)} \right).
\]

We now let \(u = v + c_2\), the sum of \([FliT]\) and \([FliT:FlhD] C_2\), so we have

\[
\frac{du}{dt} = f \frac{\sigma c_4}{k_4 + c_4 + f} - \gamma_4 u - \gamma_4 c_2.
\]

We then assume that

\[
k_2 x v - \delta_2 c_2 - \gamma_7 c_2 = 0.
\]
meaning that
\[ c_2 = \frac{\kappa_2 x v}{\delta_2 + \gamma_7}, \]

Then we have
\[ v = \frac{u}{1 + \frac{\kappa_2 x}{\delta_2 + \gamma_7}}, \]

so that
\[ \frac{du}{dt} = \frac{\sigma c_i}{k_i + c_3 + c_4 + f} - \frac{u}{1 + \frac{\kappa_2 x}{\delta_2 + \gamma_7}} \left( \gamma_6 + \frac{\gamma_7 \kappa_2 x}{\delta_2 + \gamma_7} \right). \]

Using the quasi-steady-state approximation, the model no longer includes equations for \( c_1 = [\text{RflP}\text{FlhD}_4\text{C}_2], y = [\text{RflP}], \) and \( c_2 = [\text{FliT}\text{FlhD}_4\text{C}_2]. \) The reduced model with competitive, saturating secretion, is
\[
\begin{align*}
\frac{dx}{dt} &= \frac{\rho_1}{1 + \eta_1 m} - \kappa_1 x \left( \frac{\rho_2}{1 + \eta_2 z} \left( \frac{1}{\gamma_3 + \kappa_1 x \left( 1 - \frac{\beta_1}{\delta_3 + \gamma_4} \right)} \right) \right) - \left( \frac{\kappa_2 x u}{1 + \frac{\kappa_2 x}{\delta_2 + \gamma_7}} \right) - \gamma_3 x \\
\frac{dz}{dt} &= \frac{\alpha x}{k_1 + x} + \frac{\beta_1 a}{k_3 + a} - \gamma_3 z \\
\frac{dm}{dt} &= \frac{\alpha x}{k_3 + x} - \gamma_3 m \\
\frac{dc_3}{dt} &= \frac{\alpha x}{k_3 + x} - f \frac{\sigma c_3}{k_3 + c_3 + c_4 + f} - \gamma_3 c_3 \\
\frac{du}{dt} &= f \frac{\sigma c_3}{k_3 + c_3 + c_4 + f} - \frac{u}{1 + \frac{\kappa_2 x}{\delta_2 + \gamma_7}} \left( \gamma_6 + \frac{\gamma_7 \kappa_2 x}{\delta_2 + \gamma_7} \right) \\
\frac{dc_4}{dt} &= \frac{\alpha x}{k_4 + x} - f \frac{\sigma c_3}{k_3 + c_3 + c_4 + f} - \gamma_3 c_4 \\
\frac{da}{dt} &= f \frac{\alpha x}{k_3 + x} - \frac{\sigma c_3}{k_3 + c_3 + c_4 + f} - \gamma_3 a \\
\frac{df}{dt} &= \frac{\beta_2 a}{k_6 + a} - f \frac{\sigma c_3}{k_3 + c_3 + c_4 + f} - \gamma_3 f.
\end{align*}
\]

**HBB model.** Since little is understood about HBB initiation and construction, we propose a simple model that assumes the environment within the bacteria is well-mixed. We assume that a general HBB protein is produced at a rate that depends on the concentration of FlhD4C2 in the bacterium. Mouslim and Hughes showed that flagella construction requires a threshold of flhDC expression, and, consequently, Class II expression [21]. This finding suggests that the concentration of HBB proteins must reach some threshold before initiation of HBB
construction. We assume that once the concentration of the HBB protein reaches a threshold, nucleation of a HBB occurs, and the free HBB protein concentration drops to zero. The newly nucleated HBB grows by recruiting free HBB proteins until it reaches a threshold level, at which point the HBB is considered complete [30, 31]. Complete HBBs do not recruit free HBB proteins but secrete FlgM, FliD, and FliC, which results in free $\sigma^{28}$ and FliT within the bacterium. Depending on the HBB protein production rate and recruitment rate of incomplete HBBs, the bacteria are capable of growing multiple HBBs simultaneously. Let $h$ be the concentration of free HBB proteins in the bacterium, and $b_i$ be the number of HBB proteins in the $i^{th}$ incomplete HBB. We then have

$$\frac{dh}{dt} = \alpha_5 x - K \frac{rh}{k_h + h} - \gamma_{12} h$$

$$\frac{db_i}{dt} = g_i \frac{rh}{k_h + h} ,$$

where $\alpha_5$ is the production rate of free HBB proteins, $k_7$ is the Michaelis constant for HBB protein expression, $r$ is the recruitment rate of free HBB proteins to incomplete HBBs, $k_h$ is the Michaelis constant for HBB protein recruitment, $\gamma_{12}$ is the degradation rate of free HBB proteins, and $g_i$ is zero or one, depending on whether the $i^{th}$ HBB is growing. The nucleation and completion thresholds are $h_{nuc}$ and $h_{max}$, respectively.

**Simulation methods.** We use the parameters in Tables 1 and 2 to simulate the model in MATLAB (2019a; MathWorks, Natick, MA).

We use all zero initial conditions, unless specified otherwise, and set $K = 0$ and $J = 0$. We then numerically integrate Eqs 17–26 until $h$ reaches the nucleation threshold, $h_{nuc}$, which indicates the first nucleation event. Using the state of the system just prior to nucleation and setting $h \to 0$, $b_i \to h_{nuc}$, $g_i \to 1$, $K \to 1$, we numerically integrate the system again until $h = h_{nuc}$ or $b_i = h_{max}$. If $h = h_{nuc}$, another HBB is nucleated, meaning $h \to 0$, $b_j \to h_{nuc}$, $g_j \to 1$, $K \to K + 1$, where the $j^{th}$ HBB is newly nucleated. If $b_i = h_{max}$, we complete construction of the $i^{th}$ HBB, meaning $b_i \to 0$, $g_i \to 0$, $K \to K - 1$, $J \to J + 1$. The process continues until the system reaches steady-state.

**Results**

The bistability observed in the sub-system model is maintained in the extended model. Depending on the initial conditions, HBBs are either produced or not. Since we do not explicitly model the construction of the filament, the number of HBBs is a proxy for the number of flagella. We first assume an initial condition of zero for all components of the gene network. Due to basal production of FlhD$_4$C$_2$, there is some expression of Class II genes (Fig 7A), but an insufficient concentration of FlhD$_4$C$_2$ is accumulated to initiate HBB production (Fig 7B).

In contrast, when we assume a small initial concentration of FliZ and FlhD$_4$C$_2$, FlhD$_4$C$_2$ accumulates sufficiently to initiate HBB production (Fig 8A), resulting in the production of seven HBBs (Fig 8B). This value falls within the experimentally observed range of 6–10 per bacterium [2].

The differing outcomes confirm that the FlhD$_4$C$_2$-FliZ-RfIP feedback loop, as predicted by the sub-system model, determines if HBBs are produced. We then compute the separatrix in [FlhD$_4$C$_2$]-[FliZ] initial condition space to determine what initial conditions are necessary to stimulate HBB production (solid curve in Fig 9).
We next consider the impact of nutrient availability on HBB production by changing the RFP production rate, $\rho_2$. We use $\rho_2$ as a proxy for environmental nutrient conditions in which increased nutrient availability decreases $\rho_2$ and decreased nutrient availability increases $\rho_2$. A 20% decrease in $\rho_2$ results in the production of eight HBBs (Fig 10A), which is one more in comparison to the baseline case, whereas increasing $\rho_2$ by 20% results in no HBB production (Fig 10B).

Since changing $\rho_2$ results in a different number of HBBs produced, we know that $\rho_2$ impacts the separatrix between no HBB production and HBB production. Increasing $\rho_2$ by 20% shifts the curve to require larger initial concentrations of FlhD$_4$C$_2$ and FliZ to produce HBBs (dashed curve in Fig 9). A 20% decrease in $\rho_2$ lowers the curve out of the positive parameter quadrant, so that HBBs are produced under any biologically reasonable initial conditions.

We next investigate the impact of the FliT:FliD production rate, $\alpha_3$, on HBB production. Decreasing $\alpha_3$ results in more HBBs (Fig 11A), while increasing $\alpha_3$ results in fewer HBBs (Fig 11B), in comparison to the baseline $\alpha_3$.

### Table 2. Extended model parameters.

| Parameter | Description | Value |
|-----------|-------------|-------|
| $\rho_1$  | Maximum production rate of FlhD$_4$C$_2$ | 78 min^{-1} |
| $\rho_2$  | Production rate of RFP | 37.5 min^{-1} |
| $\alpha_3$ | Production rate of FliT:FliD complex | 30 min^{-1} |
| $\alpha_4$ | Production rate of $\sigma^28$:FlgM complex | 30 min^{-1} |
| $\alpha_5$ | Production rate of $H$ proteins | 30 min^{-1} |
| $\beta_1$ | Class III production of FliZ | 60 min^{-1} |
| $\beta_2$ | Production rate of FliC | 30 min^{-1} |
| $K_3$ | Michaelis constant for FliT:FliD expression | 0.1 conc. |
| $K_4$ | Michaelis constant for $\sigma^28$:FlgM expression | 0.1 conc. |
| $K_5$ | Michaelis constant for Class III FliZ expression | 5 conc. |
| $K_6$ | Michaelis constant for FliC production | 1 conc. |
| $K_7$ | Michaelis constant for $H$ protein expression | 5 conc. |
| $K_8$ | Rate of binding of FliT and FlhD$_4$C$_2$ | 60 min$^{-1}$ |
| $\gamma_6$ | Degradation rate of FliT | $3 \text{ min}^{-1}$ |
| $\gamma_7$ | Degradation rate of FliT:FlhD$_4$C$_2$ complex | $0.3 \text{ min}^{-1}$ |
| $\gamma_8$ | Degradation rate of FliT:FliD complex | $0.3 \text{ min}^{-1}$ |
| $\gamma_9$ | Degradation rate of $\sigma^28$:FlgM complex | $0.3 \text{ min}^{-1}$ |
| $\gamma_{10}$ | Degradation rate of $\sigma^28$ | $3 \text{ min}^{-1}$ |
| $\gamma_{11}$ | Degradation rate of FliC | $3 \text{ min}^{-1}$ |
| $\gamma_{12}$ | Degradation rate of free $H$ proteins | $0.3 \text{ min}^{-1}$ |
| $\delta_2$ | ClpXP degradation rate of FlhD$_4$C$_2$ in FliT complex | $3 \text{ min}^{-1}$ |
| $r$ | Recruitment rate of free $H$ proteins to incomplete HBBs | $3 \text{ min}^{-1}$ |
| $K_s$ | Michaelis constant for recruitment of free $H$ proteins | 1 conc. |
| $\sigma$ | Total secretion rate through the HBBs | 1.8 min$^{-1}$ |
| $K_s$ | Michaelis constant for total secretion through the HBBs | 1 conc. |
| $h_{\text{sec}}$ | HBB nucleation threshold of free $H$ proteins | 15 |
| $h_{\text{max}}$ | HBB completion threshold of $H$ proteins | 100 |

Parameter values were chosen so that the first HBB is completed after approximately 30 minutes, as observed experimentally [2]. Concentration has arbitrary units.

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The change in HBB production in response to changes in $\alpha_3$ suggests that the FliT:FliD concentration, and, consequently, the free FliT concentration, is a key factor in controlling the termination of HBB production. This is exemplified by setting $\alpha_3$ to zero (Fig 12). In this case, HBB construction continues indefinitely, which means the repressive activity of FliT on FlhD$_4$C$_2$ is required to stop HBB production.

To further consider the impact of FliT on FlhD$_4$C$_2$ concentration, we recall the sub-system model (Eqs 1–5). Even though FliT is not explicitly included in the model, increasing the basal rate of degradation of FlhD$_4$C$_2$, $\gamma_1$, can be interpreted as a proxy for the repressive activity of FliT on FlhD$_4$C$_2$. As described earlier, Fig 5A shows that FlhD$_4$C$_2$ concentration is bistable as a function of $\gamma_1$. This means that as HBBs are completed and free FliT levels increase, $\gamma_1$
increases, which results in a saddle-node bifurcation and collapse of the system to a single, low steady state value. Similarly, Fig 5B shows that the bistability of FlhD$_4$C$_2$ concentration as a function of $\gamma_1$ is sensitive to $\gamma_1$. As $\gamma_1$ increases, the system moves outside the region in parameter space that supports bistability. These results suggest that there is a threshold level of free FliT required to shut-down HBB production.

Fig 9. Separatrices in FlhD$_4$C$_2$-[FliZ] initial condition space. The solid curve indicates the $\rho_2 = 37.5$ case and the dashed curve indicates the $\rho_2 = 45$ case. Initial conditions to the right of the curve lead to HBB production, while initial conditions to the left of the curve do not.

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Fig 10. The tally of incomplete and complete HBBs with perturbed $\rho_2$ values. (A) Eight HBBs are completed when $\rho_2 = 30$. (B) HBBs are not produced when $\rho_2 = 45$.

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Finally, we investigate the role of Class III regulation of FliZ. In particular, we set the Class III production rate of FliZ, $\beta_1$, to zero. Without Class III regulation of FliZ, the FliZ concentration drops dramatically following the first HBB completion (Fig 13A), and, as a result, only five HBBs are produced (Fig 13B). The reduction in the number of HBBs produced, in comparison to the baseline case, is due to the lack of HBB nucleation following the first HBB completion event. Following HBB completion, FliT activity decreases the concentration of FlhD$_4$C$_2$. This slows the production of FliZ, and, without Class III regulation, the FliZ concentration drops dramatically. The drop in FliZ concentration further contributes to the decrease in FlhD$_4$C$_2$ concentration. FlhD$_4$C$_2$ then drops to a level that does not support further HBB nucleation.

**Fig 11. The tally of incomplete and complete HBBs with perturbed $\alpha_3$ values.** (A) Eight HBBs are completed when $\alpha_3 = 24$. (B) Six HBBs are completed when $\alpha_3 = 36$.

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Finally, we investigate the role of Class III regulation of FliZ. In particular, we set the Class III production rate of FliZ, $\beta_1$, to zero. Without Class III regulation of FliZ, the FliZ concentration drops dramatically following the first HBB completion (Fig 13A), and, as a result, only five HBBs are produced (Fig 13B). The reduction in the number of HBBs produced, in comparison to the baseline case, is due to the lack of HBB nucleation following the first HBB completion event. Following HBB completion, FliT activity decreases the concentration of FlhD$_4$C$_2$. This slows the production of FliZ, and, without Class III regulation, the FliZ concentration drops dramatically. The drop in FliZ concentration further contributes to the decrease in FlhD$_4$C$_2$ concentration. FlhD$_4$C$_2$ then drops to a level that does not support further HBB nucleation.

**Fig 12. Model dynamics with $\alpha_3 = 0$.** (A) The protein concentrations increase indefinitely and FliT concentration remains zero. (B) The tally of incomplete and complete HBBs increases indefinitely.

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Together, these results suggest that the FlhD$_4$C$_2$-FliZ-RfIP feedback loop determines whether flagella production will be initiated, and the balance of FliT:FliD dynamics and Class III regulation of FliZ control the total number of flagella produced.

**Discussion**

*S. enterica* is a common foodborne pathogen that impacts millions of people each year, and the flagellum is an essential component of *S. enterica* pathogenesis. Motility is critical in enabling bacteria to reach the site of infection and establish disease. Therefore, understanding how *S. enterica* regulate and construct flagella is of interest for disease prevention and treatment. An interesting feature of flagella regulation in *S. enterica* is the presence of a heterogeneous quantity of flagella per bacterium [3].

Here, we developed mathematical models of the gene network that regulates flagella construction to improve our understanding of flagella heterogeneity. Analysis of a sub-system model (Eqs 1–5) of the network identified the possibility of bistable FlhD$_4$C$_2$ concentrations. The bistability was determined by the production rates of FlhD$_4$C$_2$ and RfIP, and could explain why some bacteria within a clonal population grow flagella while others do not. This finding could be tested by measuring the expression of *flhDC* in a population of *rifM* knockout bacteria.

We then extended the sub-system model to include the construction of HBBs and additional regulatory mechanisms. Simulations of this model (Eqs 17–26) suggest that FliT activity and Class III regulation of FliZ provide a counting mechanism by which the bacteria can control how many flagella are produced. Class III production of FliZ prevents underproduction of flagella, and FliT prevents overproduction. Hence, FlhD$_4$C$_2$ is regulated by multiple negative feedback loops to ensure that protein levels in the bacteria can be tightly controlled, and both under- and overproduction of flagella is avoided.

The *flhDC* operon is primarily transcribed from two promoters, P1$_{flhDC}$ and P5$_{flhDC}$, and many additional factors are shown to positively and negatively regulate *flhDC* expression, including HilD, RcsDBC, RtsB, and LrhA [21, 32–34]. Our model includes the main components involved in the expression of *flhDC* from P1$_{flhDC}$. Expression from the P5$_{flhDC}$, which is...
regulated via a FlhZ-HilD positive feedback loop [21], plays a role in virulence rather than motility. Therefore, we focus on expression from P1_{flhDC}. Future models could include expression from P5_{flhDC} and additional regulatory mechanisms.

The limited availability of data meant that parameter values were selected manually, rather than measured or estimated. This study investigated the possible behaviors of the gene network, and, thus, does not rely on accurate parameter estimates. As additional data becomes available, the parameter values could be updated. Further, due to a limited understanding of HBB construction, we have proposed a simple model of the process. As we learn more about HBB construction, the model could also be updated.

Our results explain the bimodality of flagella count data and how the bacteria regulate the total number of flagella produced. However, since the model is deterministic, it cannot capture the spread of 6–10 flagella in the count data. Future models could incorporate sources of stochasticity, such as cell division, to understand the variability in the number of flagella. Since the system is assumed to be well-mixed but HBBS are discrete entities, cell division could be modeled by equally dividing the proteins in the parent bacterium and randomly splitting the HBBS between the two daughter cells. The randomness of HBB inheritance could be sufficient to generate the observed flagella distributions.

This paper has proposed two mathematical models of the flagellar gene network in *S. enterica*. We predict that the FlhDC_{2}-FliZ-RfP feedback loop determines whether flagella are produced, and the combination of FliT activity and Class III regulation of FliZ determines the total number of flagella produced.

**Author Contributions**

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**References**

1. Eng SK, Pusparajah P, Ab Mutalib NS, Ser HL, Chan KG, Lee LH. *Salmonella*: A review on pathogenesis, epidemiology and antibiotic resistance. Front Life Sci. 2015; 8(3):284–293. [https://doi.org/10.1080/21553769.2015.1051243](https://doi.org/10.1080/21553769.2015.1051243)

2. Karlinsen JE, Tanaka S, Bettenworth V, Yamaguchi S, Boos W, Aizawa SI, et al. Completion of the hook-basal body complex of the *Salmonella typhimurium* flagellum is coupled to FlgM secretion and fliC transcription. Mol Microbiol. 2000; 37(5):1220–1231. [https://doi.org/10.1046/j.1365-2958.2000.02081.x](https://doi.org/10.1046/j.1365-2958.2000.02081.x) PMID: 10972838

3. Partridge JD, Harshey RM. More than motility: *Salmonella* flagella contribute to overriding friction and facilitating colony hydration during swarming. J Bacteriol. 2013; 195(5):919–929. [https://doi.org/10.1128/JB.02064-12](https://doi.org/10.1128/JB.02064-12)

4. Stewart MK, Cummings LA, Johnson ML, Berezow AB, Cookson BT. Regulation of phenotypic heterogeneity permits *Salmonella* evasion of the host caspase-1 inflammatory response. Proc Natl Acad Sci. 2011; 108(21):20742–20747. [https://doi.org/10.1073/pnas.1108963108](https://doi.org/10.1073/pnas.1108963108)

5. Chevance FF, Hughes KT. Coordinating assembly of a bacterial macromolecular machine. Nat Rev Microbiol. 2008; 6(6):455–65. [https://doi.org/10.1038/nrmicro1887](https://doi.org/10.1038/nrmicro1887)
6. Wang S, Fleming RT, Westbrook EM, Matsumura P, McKay DB. Structure of the *Escherichia coli* FlhDC complex, a prokaryotic heteromeric regulator of transcription. J Mol Biol. 2006; 355(4):798–808. https://doi.org/10.1016/j.jmb.2005.11.020

7. Macnab RM. How bacteria assemble flagella. Annu Rev Microbiol. 2003; 57(1):77–100. https://doi.org/10.1146/annurev.micro.57.030502.090832

8. Macnab RM. Type III flagellar protein export and flagellar assembly. Biochim Biophys Acta—Mol Cell Res. 2004; 1694(1-3):207–217. https://doi.org/10.1016/j.bbrc.2004.04.005

9. Aldridge PD, Karlinsky JE, Aldridge C, Birchall C, Thompson D, Yagasaki J, et al. The flagellar-specific transcription factor, ς^II^, is the Type III secretion chaperone for the flagellar-specific anti-ς^II^ factor FlgM. Genes Dev. 2006; 20:2315–2326. https://doi.org/10.1101/gad.380406 PMID: 16912280

10. Cummings LA, Wilkerson WD, Bergsbaken T, Cookson BT. In vivo, *flIC* expression by *Salmonella enterica* serovar Typhimurium is heterogeneous, regulated by ClpX, and anatomically restricted. Mol Microbiol. 2006; 61(3):795–809. https://doi.org/10.1111/j.1365-2958.2006.05271.x

11. Stewart MK, Cookson BT. Mutually repressing repressor functions and multi-layered cellular heterogeneity regulate the bistable *Salmonella* *flIC* census. Mol Microbiol. 2014; 94(6):1272–1284. https://doi.org/10.1111/mmi.12828

12. Saini S, Koirala S, Floess E, Mears PJ, Chemla YR, Golding I, et al. FlI-Z induces a kinetic switch in flagellar gene expression. J Bacteriol. 2010; 192(24):6477–6481. https://doi.org/10.1128/JB.00751-10 PMID: 20935096

13. Koirala S, Mears P, Sim M, Golding I, Chemla YR, Aldridge PD, et al. A nutrient-tunable bistable switch controls motility in *Salmonella enterica* serovar Typhimurium. MBio. 2014; 5(5):2–7. https://doi.org/10.1128/mBio.01611-14

14. Wada T, Morizane T, Abo T, Tominaga A, Inoue-Tanaka K, Kutsukake K. EAL domain protein YdiV acts as an anti-FlhDC2 factor responsible for nutritional control of the flagellar regulon in *Salmonella enterica* serovar Typhimurium. J Bacteriol. 2011; 193(7):1600–1611. https://doi.org/10.1128/JB.01494-10

15. Jonas K, Edwards AN, Ahmad I, Romeo T, Römling U, Melefors Ö. Complex regulatory network encompassing the Csr, c-di-GMP and motility systems of *Salmonella Typhimurium*. Environ Microbiol. 2010; 12(2):524–540. https://doi.org/10.1111/j.1462-2920.2009.02097.x

16. Takaya A, Erhardt M, Karata K, Winterberg K, Yamamoto T, Hughes KT. YdiV: A dual function protein that targets FlhDC for ClpXP-dependent degradation by promoting release of DNA-bound FlhDC complex. Mol Microbiol. 2012; 83(6):1268–1284. https://doi.org/10.1111/j.1365-2958.2012.08007.x

17. Saini S, Floess E, Aldridge C, Brown J, Aldridge PD, Rao CV. Continuous control of flagellar gene expression by the ς^II^ factor in the *Salmonella* flagellar regulon. *Salmonella enterica*. Environ Microbiol. 2010; 12(19):5191–5198. https://doi.org/10.1111/j.1462-2947.2010.01611-08

18. Jain K, Pradhan A, Mokashi C, Saini S. Mathematical model of flagella gene expression dynamics in *Salmonella enterica* serovar Typhimurium. Syst Synth Biol. 2015; 9(1-2):19–31. https://doi.org/10.1007/s11699-015-9160-3

19. Das C, Mokashi C, Mande SS, Saini S. Dynamics and control of flagella assembly in *Salmonella typhimurium*. Front Cell Infect Microbiol. 2018; 8(FEB):36. https://doi.org/10.3389/fcimb.2018.00036 PMID: 29473025

20. Wada T, Tanabe Y, Kutsukake K. FlI-Z acts as a repressor of the ydiV Gene, which encodes an anti-FlhDC2 factor of the flagellar regulon in *Salmonella enterica* serovar Typhimurium. J Bacteriol. 2011; 193(19):5191–5198. https://doi.org/10.1128/JB.05441-11

21. Mouslim C, Hughes KT. The effect of cell growth phase on the regulatory cross-talk between flagellar and *Spi1* virulence gene expression. PLoS Pathog. 2014; 10(3):e1003987. https://doi.org/10.1371/journal.ppat.1003987

22. Wozniak CE, Lee C, Hughes KT. T-POP array identifies EcNR and Pefl-SrgD as novel regulators of flagellar gene expression. J Bacteriol. 2009; 191(5):1498–1508. https://doi.org/10.1128/JB.01177-08

23. Singer HM, Erhardt M, Hughes KT. RIM functions as a transcriptional repressor in the autogenous control of the *Salmonella* flagellar master operon *flhDC*. J Bacteriol. 2013; 195(18):4274–4282. https://doi.org/10.1128/JB.00728-13

24. Barlq B, Beutel O, Janning D, Czamiak F, Richter CP, Kommnick C, et al. Single molecule super-resolution imaging of proteins in living *Salmonella enterica* using self-labelling enzymes. Sci Rep. 2016; 6(2015):1–14. https://doi.org/10.1038/srep31601 PMID: 27534893

25. Andino A, Hanning I. *Salmonella enterica*: Survival, colonization, and virulence differences among serovars. Sci World J. 2015; 2015. https://doi.org/10.1155/2015/520179
26. Gelfand IM, Kapranov MM, Zelevinsky AV. Discriminants, Resultants, and Multidimensional Determinants. Boston, MA: Birkhäuser Boston; 1994. Available from: http://link.springer.com/10.1007/978-0-8176-4771-1.

27. Sato Y, Takaya A, Mouslim C, Hughes KT, Yamamoto T. FlIT selectively enhances proteolysis of FlhC subunit in FlhD2C2 complex by an ATP-dependent protease, ClpXP. J Biol Chem. 2014; 289(47):33001–33011. https://doi.org/10.1074/jbc.M114.593749

28. Aldridge C, Poonchareon K, Saini S, Ewen T, Soloyva A, Rao CV, et al. The interaction dynamics of a negative feedback loop regulates flagellar number in Salmonella enterica serovar Typhimurium. Mol Microbiol. 2010; 78(6):1416–1430. https://doi.org/10.1111/j.1365-2958.2010.07415.x PMID: 21143315

29. Yamamoto S, Kutsukake K. FlIT acts as an anti-FlhD2C2 factor in the transcriptional control of the flagellar regulon in Salmonella enterica serovar Typhimurium. J Bacteriol. 2006; 188(18):6703–6708. https://doi.org/10.1128/JB.00799-06

30. Keener JP. A molecular ruler mechanism for length control of extended protein structures in bacteria. J Theor Biol. 2010; 263(4):481–489. https://doi.org/10.1016/j.jtbi.2009.12.015

31. Erhardt M, Singer HM, Wee DH, Keener JP, Hughes KT. An infrequent molecular ruler controls flagellar hook length in Salmonella enterica. EMBO J. 2011; 30(14):2948–2961. https://doi.org/10.1038/emboj.2011.185

32. Lin D, Rao CV, Slauch JM. The Salmonella SPI1 type three secretion system responds to periplasmic disulfide bond status via the flagellar apparatus and the RcsCDB system. J Bacteriol. 2008; 190(1):87–97. https://doi.org/10.1128/JB.01323-07

33. Ellermeier CD, Slauch JM. RtsA and RtsB coordinately regulate expression of the invasion and flagellar genes in Salmonella enterica serovar Typhimurium. J Bacteriol. 2003; 185(17):5096–5108. https://doi.org/10.1128/JB.185.17.5096-5108.2003

34. Lehnen D, Blumer C, Polen T, Wackwitz B, Wendisch VF, Unden G. LrhA as a new transcriptional key regulator of flagella, motility and chemotaxis genes in Escherichia coli. Mol Microbiol. 2002; 45(2):521–532. https://doi.org/10.1046/j.1365-2958.2002.03032.x