Adaptation at the output of the chemotaxis signalling pathway

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In the bacterial chemotaxis network, receptor clusters process input1–3, and flagellar motors generate output4. Receptor and motor complexes are coupled by the diffusable protein CheY-P. Receptor output (the steady-state concentration of CheY-P) varies from cell to cell5. However, the motor is ultrasensitive, with a narrow operating range of CheY-P concentrations6. How the match between receptor output and motor input might be optimized is unclear. Here we show that the motor can shift its operating range by changing its composition. The number of FliM subunits in the C-ring increases in response to a decrement in the concentration of CheY-P, increasing motor sensitivity. This shift in sensitivity explains the slow partial adaptation observed in mutants that lack the receptor methyltransferase and methylesterase7,8 and why motors show signal-dependent FliM turnover9. Adaptive remodelling is likely to be a common feature in the operation of many molecular machines.

The chemotaxis signalling pathway allows bacterial cells to sense and respond to changes in concentrations of chemical attractants or repellents10–12. Binding of chemicals by receptors modulates the activity of an associated histidine kinase, CheA, thereby changing the level of phosphorylation of the response regulator, CheY. CheY-P binds to FliM, a component of the switch complex at the base of the flagellar motor and modulates the direction of motor rotation. A phosphatase, CheZ, dephosphorylates CheY-P. The chemotaxis pathway is well known for its high gain10,11, wide dynamic range11,12 and robust adaptation13, mediated by receptor methylation and demethylation (by CheR and CheB).

The output of the chemotaxis pathway, the flagellar motor, is ultrasensitive to the intracellular concentration of CheY-P, with a Hill coefficient of about 10, imposing a narrow operational range for [CheY-P]6. Whereas precise adaptation is a robust property of the chemotaxis pathway for certain attractants, for example aspartate, the steady-state concentration of CheY-P is not7. Given cell-to-cell variations in the concentration of CheY-P and the fact that different cells can maintain their chemotactic sensitivity14, it has been suggested that cells might have additional molecular mechanisms to adjust the CheY-P concentration around the operational value of approximately 3 μM6. One possibility is a feedback mechanism that allows a cell to adjust its kinase activity in response to motor output. This mechanism would increase the kinase activity if cells only ran, and would decrease the kinase activity if cells only tumbled. In earlier work, we looked for such a mechanism by monitoring the kinase activity with a fluorescence (TIRF) microscopy of green fluorescent protein (GFP)-labelled protein14, and a similar technique has been used to demonstrate FliM9 and FliN20 exchange in cells containing CheY-P. The present work addresses the functional consequences of FliM exchange. We studied cheR cheB cells, which are defective in methylation and demethylation, and monitored motor and kinase responses to step-addition of the non-metabolizable attractant α-methylaspartate (MeAsp), using bead21 and FRET15 assays. These experiments cannot be done with wild-type cells because their adaptation to aspartate is robust, so that the steady-state concentration of CheY-P does not change. Motor adaptation occurs on a minute rather than on a second timescale and does not play a direct role in sensing temporal gradients. Instead, it helps to match the operating point of the motor to the output of the chemotaxis receptor complex, obviating the requirement for fine-tuning of that output.

Figure 1 | Motor responses to stepwise addition of chemical attractants monitored by the bead assay. The attractants were applied at the times indicated by the arrows. Error bounds for standard errors of the mean are shown as dotted lines. a. Averaged responses of seven cheR cheB cells (JY35 carrying pKAF131) to 1 mM MeAsp (weak attractant). b. Averaged responses of four cheR cheB cheZ cells (JY32 carrying pVS7 and pKAF131) to 2 mM MeAsp + 0.5 mM L-serine (strong attractant).

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Using a bead assay, we found partial adaptation in cheR cheB cells within 1 min following the initial response, Fig. 1a, which shows the averaged responses of seven motors on different cells to stepwise addition of 1 mM MeAsp. These results are similar to those obtained previously with tethered cells. A recent model suggests that partial adaptation might be due to dynamic localization of CheZ. To test this hypothesis, we repeated the bead experiments using cheR cheB cheZ cells. The results were essentially the same; Fig. 1b shows the averaged responses of four motors on different cells of a cheR cheB cheZ strain to stepwise addition of 2 mM MeAsp + 0.5 mM 1-serine (a stronger stimulus needed because of the lower sensitivity of cheZ strains). So CheZ is not required for this partial adaptation.

CheY-P concentrations were monitored by measuring FRET between cyan fluorescent protein (CFP)-conjugated CheZ (CheZ–CFP) and yellow fluorescent protein (YFP)-conjugated CheY (CheY–YFP). We measured responses in cheR cheB cells to stepwise addition of 1 mM MeAsp, Fig. 2a. The response shown in Fig. 2a is similar to that obtained previously. No adaptation is apparent. To rule out possible complications due to CheZ oligomerization, we also measured CFP–FliM/CheY–YFP FRET in cheR cheB cells following stepwise addition of 2 mM MeAsp + 0.5 mM 1-serine (a stronger stimulus needed because of the lower sensitivity of CFP–FliM/CheY–YFP FRET), as shown in Fig. 2b. No adaptation is apparent in either panel of Fig. 2, so the partial adaptation shown in Fig. 1 does not occur upstream of CheY-P. It must occur at the level of the flagellar motor.

Clockwise (CW) biases of motors were measured before addition of attractant, immediately after addition of attractant, and after time was allowed for partial adaptation. We focused on motors with pre-stimulus CW biases around 0.8 (ranging from 0.70 to 0.95). Owing to cell-to-cell variation, the lowest biases following stimulation ranged from 0.7 to 0.75. The concentration of CheY-P in a given cell was estimated from its CW bias at the time of the lowest bias, using the response curve measured previously, shown by the red line in Fig. 3 (Hill coefficient \( N_H = 10.3 \), CheY-P dissociation constant \( K = 3.1 \mu M \)). Then the CW bias found after that cell had adapted was plotted as a function of this concentration, as shown by the blue data points in Fig. 3. The measurements were carried out for 49 motors on different cheR cheB cells with stepwise addition of 0.5 or 1 mM MeAsp. Following adaptation, the relationship between the CW bias and the concentration of CheY-P shifted to lower concentrations of CheY-P, increasing motor sensitivity to CheY-P.

How the motor accomplishes this shift is intriguing. We sought to explain the shift in the motor response curve by using a Monod-Wyman–Changeux (MWC) type model, which has been used previously to explain the motor switching kinetics. In this model, the C-ring is considered to be an allosteric switch, stochastically switching between two conformational states, counterclockwise (CCW) and CW, with \( N \) independent binding sites for CheY-P, corresponding to \( N \) units of FliM in the C-ring. The CW state has a higher affinity to CheY-P than the CCW state. The CW bias of the motor is given by \( B_{CW} = (1 + [\text{CheY-P}]/K)^n(1 + [\text{CheY-P}]/K)^m + L(1 + [\text{CheY-P}]/K)^m) \), where \( L \) is the ratio of the probability that the motor is in the CCW state to the probability that it is in the CW state in the absence of CheY-P, \( K \) is the CheY-P dissociation constant for the CW state, and \( n \) is the ratio of dissociation constants for the CCW and CW states, respectively. With reasonable values for the parameters, for example, \( N = 34 \), \( L = 10^3 \), and \( K = 3.1 \mu M \), the model can be fit to the ultrasonic sensitivity data of ref. 6 with a best-fit value of \( C = 4.1 \), as shown by the red curve in Fig. 3. With these values for the parameters \( L, K, C \), and assuming that the number of FliM units \( N \) varies with [CheY-P], we can fit the data measured for the adapted motor using the MWC model with \( N = N_{sw} + \alpha \cdot [\text{CheY-P} - 2.7] \), where \( N \) is written as a Taylor expansion about the average value of [CheY-P], 2.7 \( \mu M \). We obtain a two-parameter fit with \( N_{sw} = 36 \) and \( \alpha = -1.2 \mu M^{-1} \), shown by the green curve in Fig. 3. The average value of \( N \) has increased from 34 to 36. Sensitivity of an MWC complex is known to increase with \( N \) for fixed values of \( L, K, K \) and \( C \) (ref. 28). Equivalently, the motor bias versus [CheY-P] curve shifts to smaller [CheY-P] with larger \( N \) as shown in Fig. 3. Intuitively, the fact that increasing the number of FliM units causes an increase in CW bias can be understood by considering the energetics of the switch. Each CheY-P binding decreases the energy level of CW state by a specific amount. With the values for parameters \( L, K, C \) fixed, increasing the number of FliM, that is CheY-P binding sites, increases the number of CheY-P bound to the motor.

**Figure 2** | FRET responses (Y/C ratio) of cheR cheB cells to stepwise addition of chemical attractants. The attractants were added at the times indicated by the arrows. a. CheY–YFP/CheZ–CFP FRET responses to 1 mM MeAsp (weak attractant). b. CheY–YFP/CFP–FliM FRET responses to 2 mM MeAsp + 0.5 mM 1-serine (strong attractant).

**Figure 3** | CW bias as a function of CheY-P concentration. The red curve is for pre-stimulus wild-type motors, as measured in ref. 6 and fit with an MWC model with 34 FliM units. The blue dots are data for motors that have partially adapted to stepwise addition of attractant, with a two-parameter fit to the MWC model with \( N_{sw} = 36 \) FliM units shown by the green curve; see text.
This decreases the energy level of the CW state, thereby increasing the CW bias. To test directly for this increase of the number of FliM units, we fused YFP to the carboxy terminus of FliM and monitored the fluorescence intensity of single motors using TIRF microscopy. To minimize shifts in motor position, we tethered a cheR cheB strain that lacks the flagellar filament to glass via single hooks with anti-hook antibody.

Changes in fluorescence were measured upon addition of 2 mM MeAsp + 0.5 mM l-serine, which should saturate the chemoreceptors and eliminate CheY-P, or of 1 mM MeAsp, which should simply reduce the concentration of CheY-P. The results are shown in Fig. 4. Figure 4a is the averaged response of 20 motors to the strong attractant, and Fig. 4b is the averaged response of 22 motors to the weak attractant. In either case, the fluorescence intensity increased following the addition of attractant on a time scale consistent with partial adaptation, by a larger amount for the stronger attractant. We compensated for fluorescence bleaching by subtracting a control curve (Fig. 4c) and fitting the results to a model in which the FliM off rate decreases when the CheY-P concentration decreases; see Methods. Steady state is reached when \( N_{\text{off}} = (M - N)k_{\text{on}} \), where \( N \) is the number of FliM molecules in the motor, and \( M \) is the maximum number of FliM binding sites in the motor. The fits are shown in magenta and the final values for \( N \) are given in each panel (assuming an initial value of 34). These values agree with those presented in Fig. 3. So the motor increases the number of FliM units as it partially adapts to a decrement in the concentration of CheY-P. By doing so, it increases its dynamic range.

We eliminated the concern that binding of CheY-P to FliM—YFP might be different than binding to wild-type FliM by using the bead assay to compare the biases, switching rates and speeds for motors of cheR cheB cells expressing FliM—YFP or wild-type FliM: the biases were 61 ± 16% or 58 ± 19%, the switching rates were 3.8 ± 1.2 or 3.4 ± 1.0 s⁻¹, and the speeds were 50.4 ± 8.4 or 51.5 ± 8.2 Hz, respectively.

The motor adaptation mechanism observed here is related to the turnover of motor C-ring components discovered recently, where exchange of FliM was found to be signal-dependent and exhibited a similar timescale. The detailed mechanism should involve changes in FliM on/off rates dependent upon either CheY-P binding or rates of motor switching. As noted earlier, the timescale for motor adaptation (1 min) is much slower than that for receptor methylation/demethylation (1 s), which enables cells to make rapid temporal comparisons; thus, motor adaptation does not play a critical role in that aspect. Instead, it helps match the operating point of the motor to the output of the chemotaxis receptor complex.

**METHODS SUMMARY**

All strains used in this study were derivatives of *Escherichia coli* K12 strain RP437. Cells were grown at 33 °C in 10 ml T-broth supplemented with the appropriate antibiotics and inducers and an \( A_{600nm} \) of 0.45 to 0.50. Cells were collected by centrifugation (10 min at 1,300g), washed twice in 10 ml of motility medium (10 mM potassium phosphate/0.1 mM EDTA/1 μM methionine/10 mM lactic acid, pH 7.0), and resuspended in 10 ml of this medium. They were used immediately for experiments or stored at 4 °C for up to 2h. All experiments were carried out with a custom-made flow chamber at room temperature.

For the bead assay, cells were sheared to truncate flagella, and 1.0-μm-diameter polystyrene latex beads were attached to the filament stubs. Rotation of the bead was monitored with a laser dark-field setup described previously. Rotational velocity as a function of time was determined for each motor and smoothed with a 25-point running average. CW bias was calculated over a 20-s interval every 2 s, leading to a plot of CW bias versus time.

FRET measurements of bacterial populations were carried out as described previously.

For TIRF measurements, cells were tethered to the bottom window of the flow chamber by single hooks using anti-FliG antibody, following a protocol adapted from ref. 21. The fluorescence intensity of the motors was monitored with a TIRF microscope (Nikon Eclipse Ti-U), and images were recorded with a back-illuminated, cooled (−55 °C), electron-multiplying CCD camera (DV887ECS-BV, Andor Technology). Image analysis of the motor spots was carried out using a Gaussian mask method described previously.

**Full Methods** and any associated references are available in the online version of the paper at www.nature.com/nature.

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METHODS

Strains and plasmids. All strains used in this study are derivatives of E. coli K12 strain RP437 (ref. 31): JY32 (cheb cheb chey chef fil), JY35 (cheb cheb fil), RP2893 (A2206[chez-yfP]), JY37 (cheb cheb chey fil), and JY40 (cheb cheb filM filC). The flm-eyfpGFPknox fusion with a 3× glycine linker was cloned into pTrc99A131 under an isopropyl-β-D-thiogalactoside (IPTG)-inducible promoter, yielding PRW72. pDFB72 carrying wild-type flm on pTrc99A was a gift from D. Blair. pVS7 carrying wild-type pRWB7, pDFB72 carrying wild-type pVS54 carrying pBAD33 (ref. 33), and pVS54 carrying chez-eyfp on pBAD33 (ref. 33), were described previously29. For studies of cheb cheb cells with the bead assay, JY35 carrying pKAF131 was used. For studies of cheb cheb cells with the bead assay, JY32 carrying pVS7 was used. For CFPL–CheY–YFP FRET studies of cheb cheb cells, RP2893 carrying pVS18 and pVS54 was used. For CFP–FliM–CheY–YFP FRET studies of cheb cheb cells, JY37 carrying pVS18 and pVS54 was used. For the TIRF studies of single motors, JY40 carrying pPRW7 was used. For comparison of motors with wild-type flm and FliM–YFP, JY40 carrying pDFB72 and pKAF131, and JY40 carrying pRBW7 and pKAF131 were used. Cells were grown at 33 °C in 10 ml T broth (1% tryptone and 0.5% NaCl) supplemented with the appropriate antibiotics (ampicillin: 100 μg ml−1, kanamycin: 50 μg ml−1, chloramphenicol: 34 μg ml−1) and inducers (0.005% arabinose for the bead assay, 0.01% arabinose and 50 μM IPTG for the FRET studies, 100 μM IPTG for the TIRF studies) to an A590 of 0.45 to 0.50. Cells were centrifuged (10 min, at 13,000g), washed twice in 10 ml of motility medium (10 mM potassium phosphate/0.1 mM EDTA/1 μM methionine/10 mM lactic acid, pH 7.0), and resuspended in 10 ml of this medium. They were used immediately for experiments or stored at 4 °C for up to 2 h.

Bead assay and data analysis. Cells were sheared to truncate flagella by passing 1 ml of the washed-cell suspension 50 times between two syringes equipped with 23-gauge needles and connected by a 7-cm length of polyethylene tubing (0.58 mm internal diameter, catalogue no. 427411; Becton Dickinson). The sheared cell suspension was centrifuged and resuspended in 0.5 ml of motility medium. 50 μl of this suspension was plated on a glass coverslip coated with poly-t-lysine (0.01%, catalogue no. P4707; Sigma) and allowed to stand for 2 min, then 5 μl of 1.0-mm-diameter polystyrene latex beads (2.69%, catalogue no. 07310; Polysciences) was added, mixed by gentle pipetting, and allowed to stand for 2 min. The coverslip was installed as the top window of a flow chamber29 and rinsed with motility medium. The chamber was kept under a constant flow of buffer (400 μl min−1) by a syringe pump (Harvard Apparatus). Rotation of the bead was monitored with a laser dark-field setup described previously29. Outputs from the photomultiplier tubes were directly coupled to an eight-pole low-pass (325 nm). The fluorescent intensity centroid for each motor was calculated using ImageJ (National Institutes of Health)29. The number of exposures for each motor. The laser illumination was blocked between exposures. Images of the motor spots showed radially symmetric and approximately Gaussian intensity profiles. The width of these spots was about 5 pixels (325 nm). The fluorescent intensity centroid for each motor was calculated using a Gaussian maximum method described previously29,34. Specifically, an initial estimate was made based on the peak pixel intensity, a 9 × 9 pixel region of interest (ROI) was defined centring on the initial motor centroid, and the motor centroid was calculated as follows: first, a circular motor mask of diameter 300 nm was applied to the ROI centring on the current motor centroid. Second, pixel intensities within the motor mask were multiplied by a radially symmetric two-dimensional Gaussian mask of fixed half-width 170 nm, and a revised estimate for the motor centroid was calculated using a weighted average. Lastly, the previous two steps were iterated either 150 times or until the motor mask began clamping the side of the ROI. We also calculated the motor centroid with a two-dimensional Gaussian fitting, and both methods yielded comparable results. After the centroid was calculated, the background intensity was defined as the mean pixel intensity within the ROI but external to the final motor mask, and the motor intensity was calculated as the sum of all pixel intensities within the motor mask after subtraction of the background intensity from each pixel value.

The model assumes that CheY-P binding destabilizes FliM, so that when [Chey-P] suddenly decreases due to addition of attractant, koff (the off rate of each flm unit) decreases, while kon remains the same. When the number of flm units (N) in the C-ring reaches a new steady state, NKoff = (M − N)kon, where M is the maximum number of flm binding sites in a motor. The pre-stimulus N is assumed to be 34. During the response to the attractant step, the increment of N satisfies dN = ((M − (n + 34))kon − (n + 34)koff)dτ, while the increment of the normalized motor intensity satisfies dN/koff = dN/a − jfdt, where a is the normalization factor that converts the number of flm units to fluorescence intensity, and j is the fluorescence bleaching rate obtained by fitting the control curve (Fig. 4c). Solving these two differential equations with the initial conditions: n0(0) = 0, (N0/0) = 0 leads to:

\[
\frac{f}{(ka_0-koff)}M-34(koff+koff+t)}
\]

If the time of arrival of the attractant at the cell is t0 instead of 0, change t in the above equations to t − t0.

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