Response kinetics in the complex chemotaxis signalling pathway of *Rhodobacter sphaeroides*

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Chemotaxis is one of the best-characterized signalling systems in biology. It is the mechanism by which bacteria move towards optimal environments and is implicated in biofilm formation, pathogenesis and symbiosis. The properties of the bacterial chemosensory response have been described in detail for the single chemosensory pathway of *Escherichia coli*. We have characterized the properties of the chemosensory response of *Rhodobacter sphaeroides*, an α-proteobacterium with multiple chemotaxis pathways, under two growth conditions allowing the effects of protein expression levels and cell architecture to be investigated. Using tethered cell assays, we measured the responses of the system to step changes in concentration of the attractant propionate and show that, independently of the growth conditions, *R. sphaeroides* is chemotactic over at least five orders of magnitude and has a sensing profile following Weber’s Law. Mathematical modelling also shows that, as *E. coli*, *R. sphaeroides* is capable of showing fold-change detection (FCD). Our results indicate that general features of bacterial chemotaxis such as the range and sensitivity of detection, adaptation times, adherence to Weber’s Law and the presence of FCD may be integral features of chemotaxis systems in general, regardless of network complexity, protein expression levels and cellular architecture across different species.

1. Introduction

Chemotaxis allows swimming bacteria to move towards optimal environments for growth by performing temporal comparisons of chemoeffector concentrations [1,2]. For decades, the chemosensory system has been used as a paradigm for bacterial signalling systems and has been extensively studied both experimentally and via the use of mathematical modelling. Although chemotaxis is widespread among bacterial species and central to the establishment of symbioses [3–5], biofilm formation [6] and virulence [7,8], the processes and features of chemosensory signalling have been principally investigated in *Escherichia coli* [9]. This study investigates whether the underlying principles elucidated for *E. coli* hold for chemosensory systems in other species, especially those with more complex internal signalling networks.

Most bacteria swim by rotating semi-rigid, helical flagella and change direction every few seconds [10]. These changes are due to transient reversals in the direction of flagellar motor rotation causing tumbles during which bacteria are reoriented. When swimming resumes, it is generally in a new direction, resulting in the bacteria moving in a random walk [11]. In non-homogeneous environments, modulation of the tumbling frequency by the chemotaxis signalling pathway biases the overall bacterial movement in a favourable direction. *Escherichia coli* has a single chemotaxis signalling pathway (reviewed in...
Transmembrane receptors sense changes in chemoeffectors and the signal is relayed to the flagellar motor through a diffusible phosphorylatable protein, CheY. Cells adapt to the persistence of the signal by methylation/demethylation of the receptors, returning behaviour to its pre-stimulus pattern. The E. coli sensory system is extremely sensitive, detecting changes as low as 3 nM aspartate [15] and able to respond to changes in chemoeffect concentration over five to six orders in magnitude [10,16,17]. This system has been shown to sense relative changes in chemoeffect concentration and not absolute changes, behaviour thought to conform to Weber’s Law [16,18]. Recent work performed by Lazova et al. [19] has demonstrated not only that E. coli chemotaxis follows Weber’s Law, but that it also shows fold-change detection (FCD) as predicted by Shoval et al. [20]. FCD encompasses Weber’s Law but it is more exacting. For a sensory system displaying FCD, the entire shape of the response must be identical for fold changes in chemoeffect concentration, i.e. responses to identical fold changes show equal amplitudes and adaptation times, and a steady post-stimulus state identical to the pre-stimulus state, known as exact adaptation [19,20]. Our current study investigates whether these properties are specific to the single chemotaxis pathway in E. coli or are also observed in other, more complex, chemotaxis systems.

Rhodobacter sphaeroides is a purple non-sulphur α-proteobacterium able to grow using either aerobic or anaerobic respiration or photosynthesis and shows taxis to a wide range of stimuli, including sugars, light, oxygen and organic acids such as propionate [21–23]. Interestingly, this bacterium is able to tune its tactic responses to the environmental conditions with responses to certain stimuli such as oxygen and light depending on growth conditions [24–26]. It has a number of differences in its chemosensory pathway when compared with E. coli (see the electronic supplementary material, figure S1). A comparison of the response characteristics of the two species will suggest how universal the input : output function of chemosensory systems is across species. Unlike E. coli with a single chemosensory pathway, R. sphaeroides expresses two pathways under laboratory conditions [22,23]. The proteins of one chemosensory pathway localize with membrane-spanning receptors at the cell poles, sensing the extracellular environment, while those of a second pathway localize with soluble chemoreceptors in a cluster in the cytoplasm, sensing the intracellular environment. Signals from the two pathways, in the form of CheY-Ps, must balance to control the behaviour of a single flagellar motor, and this study is aimed at identifying whether the input : output relationship from this complex two chemosensory pathway system is similar to that from a more simple single chemosensory pathway.

Aerobically and photoheterotrophically grown R. sphaeroides cells also have different cell architecture, with the membrane invaginating extensively under photosynthetic conditions [27], possibly altering diffusion rates and interfering with chemosensory signalling. Under these different growth conditions, while both pathways are expressed, the expression levels of the two pathways alters [28,29]; and while both types of cell show chemotaxis, it is possible that their input : output characteristics might differ.

In this study, using tethered cell assays, we have measured the chemotactic responses of individual aerobically and photosynthetically grown cells to a range of step decreases in propionate concentration. Our results indicate that, independently of the growth conditions, R. sphaeroides is chemotactic over at least five orders of magnitude, can sense changes in concentration as low as 10 nM and has a sensing profile following Weber’s Law. We demonstrate that, irrespective of the growth conditions and despite variability between cells in terms of adaptation times and responsiveness to stimuli, our experimental data are consistent with R. sphaeroides showing FCD and mathematical modelling of this complex signalling pathway supports the fact that it is capable of demonstrating FCD. These data suggest that chemosensory stimuli sensed through two physically separate pathways, in cells with different cellular architectures, balance to produce an output response with the same characteristics as the output from the single well-characterized pathway of E. coli. In conjunction with other modelling data [30], this supports the notion that the underlying features of the chemosensory signalling system may be universal among diverse bacterial species.

2. Material and methods

2.1. Growth conditions

Rhodobacter sphaeroides WS8N [31] was grown in succinate medium [32] at 30°C either aerobically in the dark in 250 ml flasks containing 50 ml of medium shaken at 255 r.p.m. or photosynthetotrophic without shaking, in airtight 25 ml flasks illuminated with white light at low intensity (5 W m⁻²) to maximize membrane invagination. Cells were harvested at mid-exponential phase (OD₅₇₀ between 0.45 and 0.55) when cells are very motile. This also ensures limited self-shading in photosynthetic conditions and oxygen saturation for aerobic cultures.

2.2. Cell tethering and motion analysis

One millilitre of cells in mid-exponential phase was harvested, washed and resuspended in tethering buffer (10 mM Na-PiPES, containing chloramphenicol at 30 μg ml⁻¹ to stop protein synthesis). Cells were tethered by their flagella onto a coverslip by adding 10 μl of cell suspension with 2 μl of 10 000X diluted anti-flagella antibody which spontaneously adsorb to the coverslip glass. The coverslip was incubated for 20 min in a humidity chamber and then inverted onto a microscope flow chamber and tethering buffer or propionate solutions successively flowed through at a rate of 0.12 ml min⁻¹ during 5 min sequences. Cells were observed and recorded using 40× magnification under phase contrast (Nikon Optiphoto phase contrast Microscope). Tethered cells were recorded using a digital DALSA Genie-HM640 camera with an acquisition frame rate of 100 fps and an exposure time of 6 ms. Movies were analysed using software BRAS and rotation speeds of single-bacteria extracted and analysed using the graphical interface ‘click & mean’ [33]. For each type of experiment and each type of cells studied, at least three biological replicates were analysed.

2.3. Data analysis

Tethered cells were divided into two categories: unresponsive cells, showing no stop within 2 min of a reduction in propionate concentration; and responsive cells showing a stop within this time frame. The percentage of cells responding to a given stimulus was calculated from the total number of responsive and unresponsive cells obtained from experiments across at least three biological replicates. The general responses of single cells or populations and the variations within responses were determined from at least three measurements by calculating data.
were successively challenged with six different stimuli concentration ranging over five orders of magnitude. Cells
chemosensory system, we measured the
R. sphaeroides which
To determine the range of propionate concentrations over
3.1. Population responses conform to Weber’s Law
To determine the range of propionate concentrations over which R. sphaeroides is chemotactic and the sensitivity of R. sphaeroides chemosensory system, we measured the chemosensory responses of tethered cells to reductions in concentration ranging over five orders of magnitude. Cells were successively challenged with six different stimuli (10 nM to zero, 100 nM to zero, 1 µM to zero, 10 µM to zero, 100 µM to zero and 1 mM to zero) and their responses measured. As illustrated in figure 1a, a proportion of cells grown under either aerobic or photosynthetic conditions is sensitive to changes in propionate over at least five orders of magnitude. For both growth conditions, the majority of the cells (more than 50%) react to drops greater than 10 µM, with fewer cells responding to lower concentrations drops (figure 1a). The increase in the percentage of aerobically grown cells responding to a drop from 10 nM to zero is probably owing to intrinsic cell-to-cell variation (see §3.2) coupled with the low number of cells responding to these concentrations.

We then investigated whether R. sphaeroides reacts to absolute or relative changes in attractant concentration. As over 50 per cent of cells react to steps down in propionate concentration of between 1 mM and 10 µM to zero, we tested how an identical 10 µM drop in propionate concentration is sensed over different background concentrations ranging from 1 mM to 10 µM. We successively challenged cells with the following stimuli: 1010–1000 µM, 1000–110 µM, 110–100 µM, 100–20 µM, 20–10 µM and 10 µM to zero. This enabled us to alternate between 10 µM absolute drops in propionate concentration, corresponding to low relative changes in propionate concentration, and to high relative changes in propionate concentration. Figure 1b demonstrates that, in both aerobically and photosynthetically grown cells, 10 µM drops are poorly detected against high background concentrations while changes of fivefold or greater trigger responses in a significant percentage of cells. Rhodobacter sphaeroides, therefore, appears to sense relative and not absolute changes in attractant concentration. Interestingly, the percentages of responsive cells were generally higher for aerobically grown cells than for photosynthetically grown cells (figure 1b).

As our results showed that a significant percentage of cells react to a fivefold drop in concentration, we tested responses to fivefold drops in propionate concentration over background concentrations ranging from 12.5 mM to 4 µM. Our results indicate that a significant percentage of cells react to fivefold drops in propionate over a background range of 0.5 mM to 4 µM concentration (figure 1c). However, for background concentrations over 0.5 mM, the percentages of responsive cells decrease, dropping to less than 50 per cent for aerobic cells and almost zero for photosynthetically grown cells (figure 1c). Therefore, within a specific range of background concentrations and irrespective of the growth conditions, the cells react similarly to fivefold drops. This shows that R. sphaeroides reacts to relative changes in concentrations, and thus the responses follow Weber’s Law.

3.2. Single-cell analysis of adaptation times
An intriguing feature of E. coli chemotaxis is that the intracellular signalling pathway not only obeys Weber’s Law,
but also shows FCD [19]. As our results suggested that \textit{R. sphaeroides} sensing follows Weber’s Law, we investigated whether FCD could also be a feature of \textit{R. sphaeroides} chemosensing. In order to test this hypothesis, we analysed the adaptation times (the time taken to return to the pre-stimulus rotation pattern in the continued presence of the stimulus) of individual cells within the population to different fivefold drops over background concentration between 12.5 mM and 4 mM. In addition, we challenged bacteria with six identical fivefold drops (from 100 \textmu M propionate to 20 \textmu M) to determine the intrinsic variability in \textit{R. sphaeroides} chemosensory responses in terms of adaptation time.

Analysis of the responses of single cells reacting to six repeated challenges with identical fivefold drops (from 100 \textmu M propionate to 20 \textmu M) showed a degree of underlying intrinsic cell-to-cell variability in their response times (see figure 2a,b and electronic supplementary material, figure S2) consistent with that seen in other systems [34–36]. However, comparing these data to those obtained after challenging cells with different fivefold drops over background concentration between 12.5 mM and 4 \textmu M, the variations in response times for single cells challenged multiple times with different stimuli showed no significant difference to the intrinsic variations in adaptation times (see figure 2 and electronic supplementary material, figure S2; Kruskal–Wallis rank sum test, \( p \)-value > 0.05). Combining these single-cell responses to look at the median adaptation time for each population shows that, irrespective of the growth conditions, the variability observed in population adaptation times to different fivefold drops is similar to the basal response variability of \textit{R. sphaeroides} populations (see figure 3 and electronic supplementary material, figure S3). Interestingly, despite differences in protein copy number and cellular morphology, both photosynthetic and aerobic cell populations show similar adaptation times to fivefold drops in concentration (figure 3a,b). Thus, these data show that, independently of the growth conditions, \textit{R. sphaeroides} cells have similar chemotactic responses which are consistent with the requirements for FCD.

3.3. Can the \textit{Rhodobacter sphaeroides} chemosensory pathway show true FCD?

A fundamental feature of FCD is that the entire shape of the chemosensory response (amplitude, adaptation time and precision of return to pre-stimulus behaviour) must be identical for the same fold changes in input [20]. Owing to the large number of proteins within the complex chemotaxis system of \textit{R. sphaeroides}, it is not possible to experimentally measure the amplitudes and dynamics of all of the components of the system. Therefore, to determine whether the chemosensory pathway of \textit{R. sphaeroides} is capable of showing true FCD, we developed a nonlinear ODE model to characterize the changes in the intracellular signalling pathway components on different fivefold drops in attractant concentration. The mathematical model was formulated around earlier simpler models of the \textit{R. sphaeroides} signalling cascade [37], but

![Figure 2. Adaptation times of responsive single cells to fivefold drops in propionate concentrations. (a,c) Aerobic single-cell adaptation times. (b,d) Photosynthetic single-cell adaptation times. (a,b) Adaptation times to six successive drops from 100–20 \textmu M in propionate concentration. (c,d) Adaptation times to different fivefold drops in propionate concentration. Lozenges represent the median adaptation time of a single cell. The circles represent responses of a single cell to successive stimuli. Only cells responding to at least three stimuli are represented.](http://rsif.royalsocietypublishing.org/Downloadedonomic.png)
which did not include a model of adaptation. As such we inte-
grated a recent Monod–Wyman–Changeux (MWC) model of
*E. coli* receptor adaptation [38] with the underlying
*R. sphaeroides* signalling cascade in a similar manner to that of Clausznitzer
et al. [39] for *E. coli* chemotaxis. Full details of the mathematical
model and its parameter values are given in the electronic
supplementary material. This mathematical model is similar
in features to that of Tu et al. [40], which exhibited FCD under
specific conditions [20] and satisfies the conditions that
are shown by Hamadeh et al. [30] to be sufficient for the
*R. sphaeroides* chemotaxis system to show FCD.

The mathematical model was challenged with ligand step
changes which elicited the greatest cell response experimen-
tally, namely 100–20 µM and 20–4 µM. For FCD to occur,
each signalling protein must react with an identical response
curve to each of these fivefold changes, i.e. the CheY6-P
response curves must be identical to each other, as must
the CheY3-P curves, etc. As can be seen in figure 4, each
phosphorylated protein responds in an identical way to the
two fivefold drops as their response curves superimpose
exactly. These results, along with those discussed in
Hamadeh et al. [30], confirm that the *R. sphaeroides* signalling
cascade can exhibit FCD.

Interestingly, the shape of the adaptation curves for the
various proteins are different. Experimentally, CheY6-P has
been shown to be responsible for stopping the flagellar
motor [23], while CheY3-P and CheY4-P also bind the
motor but may modulate the effect of CheY6-P. We noted
that the levels of the motor stopping CheY6-P quickly
reached a saturating level during each fold change, while
all the remaining protein levels did not. This result can be
explained as follows. Because CheY6 is in high concentration
throughout the cell and its phosphorylation is dominated by
the transfer from CheA3-P rather than CheA2-P (see the
electronic supplementary material, table S1), any variation
in CheA3-P levels are mirrored by CheY6-P; the phospho-
transfer occurs over a much shorter timescale than that of
adaptation, and thus saturating levels of CheY6-P are
observed. In contrast, the concentration of CheY3, CheY4,
CheB1 and CheB2 is considerably less than that of CheY6.
Furthermore, with the exception of phosphotransfer between
CheA2-P and CheY4, and CheA2-P and CheB1, all the
remaining phosphotransfer rates are considerably slower
than that of CheA3-P to CheY6. Thus, we expect the time
taken for CheY3-P, CheY4-P, CheB1-P and CheB2-P to reach
saturation levels to be considerably slower such that it may
not occur during the adaptation time period. As CheY6-P is
the major output causing the motor to stop [23], having this
rapid transition between low and high concentrations may
provide for a rapid physiological output in the chemotaxis
response while other proteins follow a more graded tran-
sition, consistent with their having a role in the adaptation
process or a more subtle modulation of the motor output.

3.4. Population adaptation times to different stimuli

We also challenged the model with twofold and 10-fold
changes in attractant concentrations (figure 5), to identify
how the phosphorylation levels of the constituent proteins
was affected by the magnitude of the input. The time for
the model to adapt was measured as the time taken for the
major motor stopping CheY6-P levels to return to their
pre-stimulus levels. Interestingly, the twofold change
showed a response in the model with a short adaptation
time of ca 25 s, compared with ca 45 s for the fivefold
change and 55 s for the 10-fold change. When the model

![Figure 3. Adaptation times of responsive cell populations to fivefold drops in propionate concentrations. (a) Responses to six drops in concentration from 100–20 µM. (b) Responses to fivefold drops over different background concentrations. The number of cells analysed for each drop in concentration was between (a) 26 and 20 for photosynthetic (circles) cells, and 12 and 22 for aerobic (squares) cells, (b) 23 and 24 for photosynthetic cells, and six and 25 for aerobic cells. Only datasets comprising more than three cells are represented. Symbols represent data medians. The lower and the upper limits of the bars represent the first and the third quartiles of the data, respectively.](http://rsif.royalsocietypublishing.org/)
was tested against our experimental data, it was found that only a low percentage of cells show responses to a twofold change (figure 1b), probably because of the intrinsic cell-to-cell variations observed combined with the limit of time resolution for stop detection in our experimental set-up. However, cells challenged with 10-fold changes in propionate concentration for background concentrations between 1 mM and 10 mM showed adaptations times which were not longer for the 10-fold changes than for the fivefold changes (figure 6). This is consistent with the relatively small increase in adaptation times suggested by the model (figure 5) being similar to the underlying intrinsic variation shown by the R. sphaeroides cell populations (figure 3a). Thus, the adaptation times do not appear to increase linearly with the magnitude of the fold change.

4. Discussion

In this study, we have investigated whether the output of the bacterial chemosensory pathway is altered by increased network complexity, protein expression levels or cellular architecture compared with the simple well-studied pathway of E. coli. Chemosensory responses were studied for R. sphaeroides cells grown either photosynthetically or aerobically, i.e. in conditions in which cells have different cell architecture and chemosensory protein expression profiles [27–29].

Our results show that R. sphaeroides is sensitive to step changes in propionate concentration ranging over five to six orders of magnitude depending on the growth conditions (from 10 nM to about 10 mM). In this respect, R. sphaeroides is similar to E. coli which senses differences in attractant concentration over five to six orders of magnitude [10,16,17].
Therefore, despite significant differences in their chemotaxis pathways, the range and sensitivity over which R. sphaeroides is chemotactic is similar to that described for E. coli [10,16,17]. Interestingly, while all cells analysed behave similarly to a given stimulus regardless of growth conditions, photosynthetic populations of R. sphaeroides show a lower general percentage of cells responding to a given stimulus and the response of these cells saturate at high input concentrations earlier than cells in the aerobic populations. The reason for this is not currently known, but may reflect the lower average copy number of chemotaxis proteins expressed under photosynthetic growth conditions [28,29], and possibly the lower number of chemoreceptors [41] may become saturated by this is not currently known, but may reflect the lower average copy number of chemotaxis proteins expressed under photosynthetic growth conditions [28,29], and possibly the lower number of chemoreceptors [41] may become saturated by high levels of attractant.

In vivo FRET measurements, between the single CheY-P and its phosphatase CheZ, were used in E. coli to measure the entire shape of the chemosensory response (amplitude, adaptation time and precision) from a population of cells and to give a measure of the intracellular kinetics of the response and demonstrate FCD [19]. This is not possible in R. sphaeroides because it lacks a CheZ and it would be necessary to simultaneously measure the FRET interactions between three CheY proteins and two CheAs in this more complex signalling pathway, which is currently not technically possible. Therefore, we used a tethering assay to accurately quantify the whole input : output response as measured by the adaptation times of individual cells to different stimuli.

Figure 6. Adaptation times of responsive cell populations to fivefold changes (black) and 10-fold changes (white). Circles represent median responses of photosynthetic cell populations and squares median responses of aerobic cell populations. The number of cells analysed for each drop in concentration was between, for fivefold changes: 23 and 25 for photosynthetic cells and six and 25 for aerobic cells; for 10-fold changes: four and 18 for photosynthetic cells and three and 12 for aerobic cells. Only datasets comprising more than three cells are represented. The lower and the upper limits of the bars represent the first and the third quartiles of the data, respectively.

Our experimental results show that, irrespective of the growth conditions, individual R. sphaeroides cells display constant adaptation times to fold changes in concentration conforming to Weber’s Law and consistent with FCD for the entire signalling system. We then applied mathematical ODE-based modelling to characterize the concentrations of the major intracellular signalling proteins during responses to different fivefold drops in attractant concentration, as these are not directly measurable experimentally, and found that the model supported FCD in the R. sphaeroides chemotaxis pathway. Therefore, although we cannot experimentally measure response amplitudes, our data are consistent with the hypothesis that the R. sphaeroides chemosensory system shows FCD as recently demonstrated for the simpler pathway in E. coli [19]. Significantly, our results suggest that FCD occurs over similar background concentrations to the ones described in E. coli for methyl-aspartate [19], suggesting that neither the complexity of the chemosensory network in R. sphaeroides nor differences in cell architecture or protein expression profiles under different growth conditions influence FCD. This is also shown in the work of Hamadeh et al. [30] and supports the proposal that FCD could be a general feature of many biological sensory systems as hypothesized by Shoval et al. [20].

Analysis of the chemosensory responses of individual cells highlights that there is significant variability between cells in a genetically identical population, similar to observations/predictions made in Salmonella enterica S.v. Typhimurium [34] and E. coli [10,35,36]. Even though the majority of aerobic and photosynthetic cells show similar sensitivity and sensing features, a small proportion of cells can demonstrate higher sensitivity or different sensing properties by reacting to lower fold changes. In addition, for both aerobic and photosynthetic cells, adaptation times were shown to vary both from cell-to-cell and when an individual cell was repeatedly challenged with the same stimulus. Phenotypic variability in adaptation times and in responsiveness to chemoattractants could allow isogenic R. sphaeroides populations to optimize their search for nutrients and could represent a bet-hedging strategy to maximize survival.

The similarity in the chemotaxis responses of both aerobically and photoheterotrophically grown cells to each other, and to those previously determined for E. coli, suggests that general physiological features of chemotaxis may be a necessary consequence of their function. Our experimental data and modelling results, supported by the work of Hamadeh et al. [30], suggest that the range and sensitivity of chemoattractant detection, adaptation times, adherence to Weber’s Law and FCD may be integral features of many chemotaxis systems, regardless of differences in network complexity, chemosensory protein expression profiles and cell architecture in different bacterial species.

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