Data-driven quantification of the robustness and sensitivity of cell signaling networks

Sayak Mukherjee1,2, Sang-Cheol Seok1, Veronica J Vieland1,2,3 and Jayajit Das1,2,4,5,6

1 Battelle Center for Mathematical Medicine, The Research Institute at the Nationwide Children’s Hospital, The Ohio State University, 700 Children’s Drive, Columbus, OH 43205, USA
2 Department of Pediatrics, The Ohio State University, 700 Children’s Drive, Columbus, OH 43205, USA
3 Department of Statistics, The Ohio State University, 700 Children’s Drive, Columbus, OH 43205, USA
4 Department of Physics, The Ohio State University, 700 Children’s Drive, Columbus, OH 43205, USA
5 Biophysics Graduate Program, The Ohio State University, 700 Children’s Drive, Columbus, OH 43205, USA

E-mail: das.70@osu.edu

Received 20 May 2013
Accepted for publication 18 September 2013
Published 29 October 2013

Abstract
Robustness and sensitivity of responses generated by cell signaling networks has been associated with survival and evolvability of organisms. However, existing methods analyzing robustness and sensitivity of signaling networks ignore the experimentally observed cell-to-cell variations of protein abundances and cell functions or contain ad hoc assumptions. We propose and apply a data-driven maximum entropy based method to quantify robustness and sensitivity of Escherichia coli (E. coli) chemotaxis signaling network. Our analysis correctly rank orders different models of E. coli chemotaxis based on their robustness and suggests that parameters regulating cell signaling are evolutionary selected to vary in individual cells according to their abilities to perturb cell functions. Furthermore, predictions from our approach regarding distribution of protein abundances and properties of chemotactic responses in individual cells based on cell population averaged data are in excellent agreement with their experimental counterparts. Our approach is general and can be used to evaluate robustness as well as generate predictions of single cell properties based on population averaged experimental data in a wide range of cell signaling systems.

Introduction
Robustness of specific cell functions against intra- and extracellular perturbations is a salient feature of many biological systems [1–4]. For example, polarization of selected molecules across wide variations of protein abundances in yeast cells is necessary for mating and bud formation [2]; similarly, the ability of Escherichia coli (E. coli) cells to migrate to a nutrient rich region over a large range of nutrient concentration is related to their increased growth [4, 5]. Mutations increasing robustness of tumor cell survival and proliferation underscores the importance of robustness in diseases such as cancer [6]. Robustness in biological systems is often accompanied with processes that respond sensitively to perturbations [7, 8]. A striking example of this ‘robust yet fragile’ behavior is found in individual T cells, which can accommodate over tenfold variations in protein abundances [9], yet also mount binary all or none responses to pathogenic ligands based on the formation of non-covalent T cell receptor ligand complexes with lifetimes differing from each other by just few seconds [10–12]. These examples suggest that both robustness and sensitivity (lack of robustness) of specific cell functions are

6 Author to whom any correspondence should be addressed.
key to understanding the design principles underlying cell signaling and gene regulatory networks.

Measuring robustness, however, raises a number of challenges. First and foremost is the inability of standard methods to incorporate experimentally observed cell-to-cell variations of protein abundances and cell responses [5, 13, 14]. For example, ordinary differential equation (ODE) based models, used to describe deterministic signaling kinetics of concentrations of signaling molecules averaged over a cell population, ignore intrinsic stochastic fluctuations [13, 15, 16] in individual cells that occur due to thermal fluctuations in biochemical signaling reactions [1, 4, 17, 18]. Similarly, these models generally do not account for extrinsic noise fluctuations [16, 19] arising from cell-to-cell variations of steady state abundances of signaling proteins and the physical properties of the signaling environment (e.g., cell size, molecular crowding, number of spatial compartments, or spatial micro-domains of receptors). But the unaccounted for variation of copy numbers of the signaling molecules produced by these fluctuations can generate qualitatively different signaling outcomes in single cells compared to those predicted by the ODE models for a cell population [20–22]. Thus these models might not properly reflect the behavior of the signaling network in single cells or even the behavior of a cell population. Methods have been proposed for including intrinsic and extrinsic noise fluctuations in stochastic and spatially resolved in silico models [2]. However, these methods still do not incorporate cell-to-cell variations of cell responses and model selection requires imposition of ad hoc criteria. For example, in quantifying robustness of different models designed to produce spatial polarization of marker molecules, the models that produced a polarity score of greater than an arbitrary threshold value of 0.6 were considered to be able to produce polarization, whereas, the corresponding experiments with synthetic circuits showed a wide distribution of the polarity score ranging between 0 and values larger than 1 [2]. This can lead to erroneous conclusions regarding relative robustness of competing models, especially when the models show similar behavior. There are in addition computational challenges, whether working with deterministic or stochastic models, because of the large size of the parameter space, with many parameters required to describe strengths of interactions, total protein concentrations, and/or diffusion constants. Recent studies have proposed combining local Monte Carlo moves with principal component analysis (PCA) [18] or random walks in the parameter space [17] to address this issue.

Here we propose a novel data-driven approach based on maximum entropy (MaxEnt), a technique championed by Jaynes [23, 24], and maximum caliber (MaxCal) [25] to addresses these challenges. Our approach is entirely guided by available experimental data, measured in either a cell population or in individual cells, and it naturally combines intrinsic and extrinsic noise fluctuations in the cell signaling kinetics. MaxEnt has been widely used in diverse disciplines ranging from physics [26] to information theory [27] to biology [28–32] to estimate probability distributions of variables subject to constraints [24, 28, 30, 31]. In essence, the MaxEnt procedure yields the least structured, or least restricted, probability distribution for the underlying parameters, consistent with constraints imposed by available empirical data (such as average values). MaxCal [25, 33], also proposed by Jaynes, is an extension of MaxEnt to dynamical systems. These methods simultaneously allow us to directly incorporate stochastic properties of real networks, to avoid imposition of additional ad hoc modeling assumptions, and to bypass computational difficulties for some competing approaches. They also generate predictions regarding the distribution of specific attributes (e.g., adaptation time) of signaling kinetics in single cells, based on experimental observations that are only available at the cell population level. This addresses a common problem in inferring mechanisms underlying signaling kinetics in single cells in the absence of experimental observations in single cells.

But beyond simply addressing technical problems, we also propose that MaxEnt and MaxCal uniquely provide a direct measure of biological robustness. The existing methods analyze sensitivity and robustness of specific cell functions as model parameters are perturbed [1, 2, 4, 18, 34]. In contrast, we constrain cell population averaged measurements in our MaxEnt approach to infer how model parameters are varying in individual cells, and to test predictions using the inferred probability distribution with available single cell measurements. Therefore, a consistency between our approach and the existing methods will suggest evolutionary selection of model parameters depending on how their perturbations influence model robustness. If this is correct for a biological system, then when applied to compare multiple mechanistic models underlying robust systems, the most robust MaxEnt or MaxCal model should coincide with the correct model. The ability of our approach to predict distributions of single cell attributes in a cell population allows us to test against single cell experiments if the most robust model is indeed the correct model. In order both to illustrate the approach and also to establish a proof of principle, we apply our method to quantification of the robustness of *E. coli* chemotaxis.

The chemotactic behavior of *E. coli* cells is one of the best-characterized models of cell signaling to date [35, 36]. *E. coli* cells sense the presence of attractants (or repellants) in the medium with the help of transmembrane Tar receptors, and respond by swimming towards (or away) from the nutrient source. Upon binding to attractants, Tar receptors initiate a series of signaling events that lead to an increased bias towards anticlockwise rotations of the flagella motor causing directed movements in individual *E. coli* cells [35, 36]. Then as the bacterial cells arrive at the region of higher nutrient concentrations, the flagella motor movements are biased towards clockwise directions, and the cells return to their pre-stimulus state of random movements. This represents a perfect adaptive behavior, which has been demonstrated to be robust against variations of signaling protein concentrations, nutrient concentrations, rate constants, temperature changes, and strengths of interactions between proteins, both in experiments and in silico modeling [1, 5, 37, 38]. Recent experiments also provide data regarding cell-to-cell variations in chemotactic responses and protein abundances [5, 14, 39], in addition to the vast amount of cell population averaged
measurements carried out over many decades [40]. Therefore, 
E. coli chemotaxis is an ideal system in which to test and 
validate our approach.

We consider three different coarse grained or approximate 
models that were proposed to explain E. coli chemotaxis: 
(i) the fine-tuned (FT) model [41, 42], (ii) the Barkai–Leibler 
(BL) model [1], and (iii) the modified BL (MBL) model, 
which is a recently proposed modification of the BL model 
[5]. Both BL and MBL can describe the robustness of the 
neatly perfect adaptation behavior in E. coli to variations of 
the interaction strengths and nutrient concentrations. However, 
BL fails, whereas MBL succeeds in describing how the cells 
manage to restrain pre-stimulus steady state values of the 
CheY-P protein abundances within the working range of the 
flagella motors while being subject to wide variations in the 
abundances of the chemotactic proteins. The FT model fails to 
capture the robustness of E. coli chemotaxis and can reproduce 
the adaptive behavior only over a small range of parameters. 
We use stochastic simulations of the chemotaxis signaling 
network in individual E. coli cells, including intrinsic and 
extrinsic noise fluctuations, and use MaxEnt and MaxCal to 
quantify robustness of each of the three models, utilizing 
available cell population averaged and single cell data from 
the published literature as constraints. We also compare the 
models’ predictions regarding distributions of chemotactic 
responses and protein abundances in individual cells with 
available experimental data.

The remainder of the paper is organized as follows. 
In section 1 we describe our overall approach and study 
design. Section 2 presents results regarding the relative 
robustness of the three models. Section 3 compares predictions 
of distributions of single cell adaptation time and protein 
concentrations with results obtained from experiments. 
Section 4 is reserved for the discussion, while detailed methods 
are found in section 5 and supplementary material.

1. Approach and study design

Here we describe our approach in the context of chemotactic 
responses in individual E. coli cells responding to attractants 
added to the medium. The method can be generalized to any 
other cell signaling system where signaling kinetics data are 
available at the cell population and/or the single cell level.

1.1. E. coli chemotactic models

We consider three different coarse grained or approximate 
signaling models that were proposed to explain E. coli 
chemotaxis. Each model is composed of a set of biochemical 
reactions involving the chemotactic proteins, CheA, CheR, 
CheB, CheY, CheZ (only for MBL and FT), and the 
transmembrane Tar receptors. The models also differ from 
each other due to differences in molecular wiring between 
chemotactic proteins (figure 1). The kinetics of the chemotactic 
response for a given model is determined by the rate 
constants of different biochemical reactions, and also the 
total abundances of the associated proteins. The FT model 
(figure 1(A)) was among the first proposed models that could 
explain precise adaptation in E. coli within a narrow range 
of rate constants and protein concentrations [41, 42]. The 
BL model (figure 1(B)), proposed later, was able to capture 
the robustness of the nearly perfect nature of adaptation of 
E. coli chemotaxis to variations in rate constants or 
strengths of protein–protein interactions [1], as confirmed in 
cell population experiments [37]. However, the steady state 
concentration of phosphorylated CheY protein (or CheY-P) 
in the BL model is sensitive to large variations of protein 
concentrations in the model. Since the working range of the 
flagella motor is limited to small variations (~30%) 
from the optimal CheY-P concentration, the robustness of 
E. coli adaptation to large range of variation in the abundance 
of chemotactic proteins requires small variations of steady 
state CheY-P concentrations [5, 43]. Sourjik and colleagues 
proposed and experimentally tested a modified version of the 
BL model (MBL, figure 1(C)) that was able to restrain 
variations in CheY-P concentration to a small range [5].

All three models include biochemical reactions that 
describe the basic signaling events underlying E. coli 
chemotaxis (figure 1). Transmembrane Tar receptors bind 
to attractants (or repellants) in the medium and become 
deactive (or active). The kinase CheA, associated with activated 
Tar receptors, gets auto-phosphorylated and then transfers 
the phosphoryl group (PO4) to the cytosolic enzyme CheY 
rendering the latter active. Phosphorylated CheY controls 
rotational bias of the flagella motor, an increase in CheY-P 
concentrations leads to an enhanced clockwise bias of the 
flagella motors [43, 44], causing individual E. coli cells to 
tumble. Tar receptors also undergo methylation by an enzyme, 
CheR, and the enzyme CheB removes the methyl groups from 
the active form of the receptors. Methylated receptors induce 
CheY phosphorylation at an increased rate. When attractants 
are added to the medium, a decrease in the number of activated 
receptors leads to decrease in CheY phosphorylation, and the 
E. coli cells display a chemotactic response by executing 
a directed motion toward the nutrient source. However, the 
decrease in the number of active receptors results in an increase 
in the rate of methylation of the receptors due to the decreased 
rate of de-methylation by CheB. As methylation goes up, the 
activity of the receptors and the phosphorylation of CheY 
start to increase and eventually return to the pre-stimulus 
level, resulting in the restoration of random motion in E. coli. 
CheY-P concentrations return almost precisely to the pre-
stimulus level after a time interval, which generates the nearly 
perfect adaptive behavior.

The key difference in signaling between the FT and 
the BL model is that the enzyme CheB can demethylate 
only the active methylated receptors in BL, whereas, CheB 
demethylates both the active and the inactive receptors in 
the FT model (figure 1). This induces an integral feedback 
control process in the BL model [45], where the production 
of active methylated receptors, the key inducer of CheY-P, 
is regulated by the sum of the difference between the actual 
abundance of active methylated receptors with its steady state 
value over a time period. This produces a steady state of active 
methylated receptors independent of the ligand concentrations 
for a wide range of model parameters in the BL model. In
The fine-tuned model (FT), originally proposed by Knox et al [42] and Hauri and Ross [41], shows robust adaptation only for a small region in the parameter space. In this model the aspartate receptors (Tar) can shuttle between an active (denoted by the orange star) and an inactive conformation. The probability of the receptors to be in an active conformation depends on their state of methylation. Pair of enzymes namely CheR (R) and CheB-P (B) add and remove methyl groups from the receptors. CheB-P can demethylate receptors regardless of their state of methylation. The Tar receptors form complexes with a kinase CheA (A). CheA can phosphorylate itself with a rate proportional to the abundance of the active Tar. Phosphorylated CheA can transfer its phosphoryl group (PO4) either to the kinase CheB, rendering it capable of demethylation, or to another response regulatory protein called CheY (Y). Upon receiving the phosphoryl group from CheA, CheY renders itself active. Active form of CheY (CheY-P) diffuses across the cell and binds to the flagella motors causing them to tumble. The phosphatase CheZ (Z) de-activates the active form CheY. (B) Barkai–Leibler model (BL) was put forward to explain robust adaptation in the chemotactic network of bacterial E. coli. Unlike FT model, CheB, both the unphosphorylated and the phosphorylated form, demethylates only the active Tar receptors. This model also lacks the phosphatase CheZ and CheY-P undergoes auto-dephosphorylation. (C) MBL model proposed by Kollmann et al except for two differences, is similar to the BL model. The differences are: (i) Only the phosphorylated form of CheB, as opposed to CheB and CheB-P in the BL model, can de-methylate the active receptors, (ii) the dephosphorylation of the active CheY is done by the phosphatase called CheZ (Z). The presence of this phosphatase makes the steady state of CheY-P abundance relatively robust to the concerted over expression of the chemotactic proteins.

Figure 1. Three models for E. coli chemotaxis. (A) The fine-tuned model (FT), originally proposed by Knox et al [42] and Hauri and Ross [41], shows robust adaptation only for a small region in the parameter space. (B) Barkai–Leibler model (BL) was put forward to explain robust adaptation in the chemotactic network of bacterial E. coli. (C) MBL model proposed by Kollmann et al except for two differences, is similar to the BL model. The differences are: (i) Only the phosphorylated form of CheB, as opposed to CheB and CheB-P in the BL model, can de-methylate the active receptors, (ii) the dephosphorylation of the active CheY is done by the phosphatase called CheZ (Z). The presence of this phosphatase makes the steady state of CheY-P abundance relatively robust to the concerted over expression of the chemotactic proteins.
CheY-P, as well as, unmodified protein species such as CheY. The time interval \( \Delta \) is chosen to be smaller than or of the same order of the smallest reaction time scale (figure 2). We have \( N_F \gg N_T \), as a protein species can be modified during signaling, e.g., the signaling protein CheY-P is generated from the protein CheY. The properties of chemotactic responses in a single cell, such as the adaptation time or the precision of adaptation, depend on the stochastic trajectory \( \Gamma \) of that cell (figure 2(A)). Therefore, we represent the magnitude \( (f_a) \) of a specific property, \( a \) (e.g., adaptation time), of the chemotactic response displayed by a single cell by \( f_a \). Denoting the probability that a single cell follows a stochastic trajectory, \( \Gamma \), be \( P_T \), the distribution of \( f_a \) can be written as

\[
P(f_a) = \sum_{\Gamma} \delta_{f_a,f_a} P_T(\Gamma)
\]

where the summation is over a set of stochastic trajectories \( \{\Gamma\} \) observed in a population of \( E. \ coli \) cells. \( P_T \) can be calculated from the joint distribution \( P(n_j, t_0; n_q^{\text{total}}) \) of the signaling proteins at the pre-stimulus level and the total protein abundances first, and then by calculating the transition probabilities with which the initial state \( \{n_j\}, t_0 \) goes to the consecutive signaling states at different times. Therefore we have

\[
P_T = P(n_j, t_0; \{n_j\}, t_{n-1}; \ldots; \{n_j\}, t_1|\{n_j\}, t_0) \times P(n_j, t_0; n_q^{\text{total}}).
\]

The biochemical reactions in each model occur as Markov processes where the probability \( P(\{n_j(t_2)\}|\{n_j(t_1)\}) \) \( t_2 \geq t_1 \), that the state changes from \( \{n_j\}, t_1 \) to \( \{n_j\}, t_2 \) is given by the master equation [15],

\[
\frac{\partial P(n_j(t_2)|\{n_j(t_1)\})}{\partial t_2} = LP(n_j(t_2)|n_j(t_1)).
\]

In equation (3), \( L \) is a linear operator [15] dependent on the biochemical reaction rates, wiring of the signaling network, and the numbers of signaling proteins at time \( t_1 \).

The steady state conditional probability of the abundance of signaling molecules at the pre-stimulus state at \( t = t_0 \) given a fixed set of total protein abundances or \( P(n_j, t_0; n_q^{\text{total}}) \) can be calculated from equation (3) by setting the left hand side to zero and using the form for \( f_a \) without any attractant present in the medium. The joint probability distribution, \( P(n_j, t_0; \{n_q^{\text{total}}\}) \) then can be calculated using the relation \( P(n_j, t_0; \{n_q^{\text{total}}\}) = P(n_j, t_0; \{n_q^{\text{total}}\}) P(n_q^{\text{total}}) \), where, \( P(n_q^{\text{total}}) \) denotes the distribution of total protein abundances in single cells.

When experimental measurements are available for a population of cells, e.g. from western blot assays, data are available as average values of the properties of the chemotactic response,

\[
\frac{1}{\text{total \# of cells}} \sum_{\alpha=1}^{\text{total \# of cells}} f_a^\alpha = \sum_{\Gamma} f_a^\Gamma P_T = f_a^{\text{exp}}
\]

where \( f_a^{\text{exp}} \) denote the experimental average value for the \( a \)th property of the chemotactic response. Note that \( \sum_{\Gamma} f_a^\Gamma P_T \) refers to the expected value of the variable, \( f_a \), rather than the sample average. Therefore, the equality in equation (4) is strictly true only as the number of samples (i.e., total \# of
cells) becomes very large [47]. In finite samples, the average can deviate from the expected value, potentially leading to errors in the estimated distributions of the parameters (see the discussion section for further details). Note too that to make our notation clear, as previously noted, we will use \( f_a \) to denote properties of chemotactic responses (e.g., adaptation time or precision of adaptation), and \( n_q^{\text{total}} \) for protein abundances in individual cells. Single cell measurements can provide further details regarding how a specific property is distributed in a cell population, and they could allow for the calculation of variances or even higher order moments from the experimental data, i.e.,

\[
\frac{1}{\text{total # of cells}} \sum_{a=1}^{\text{total # of cells}} (f_a^n) = \sum_{a} f_{a|T} P_T = (\bar{f}_{a|\text{exp}})^n
\]

where the right hand side denotes average of the \( n \)th moment for the property \( f_a \) calculated from experimental data. In addition, average values of the total protein abundances from cell population level experiments, variance, and higher moments for the total protein numbers from single cell measurements, might be available, i.e.,

\[
\frac{1}{\text{total # of cells}} \sum_{a=1}^{\text{total # of cells}} (n_q^{\text{total}})_a = \sum_{[q_a]} n_q^{\text{total}} P([n_q^{\text{total}}])
\]

\[
\frac{1}{\text{total # of cells}} \sum_{a=1}^{\text{total # of cells}} (n_q^{\text{total}})_a^n = \sum_{[q_a]} (n_q^{\text{total}})_a^n P([n_q^{\text{total}}]) = (\bar{n}_q^{\text{exp}})^n
\]

where \( n_q^{\text{total}} \) denotes the total abundance of protein \( q \) in a single cell. The probability function \( P_T \) contains variations due to intrinsic and extrinsic noise fluctuations through \( P([n_j], \tau, [n_j], \tau_2, \ldots; [n_j], t_1, [n_j], t_0 | [n_q^{\text{total}}]) \) and \( P([n_q^{\text{total}}]) \).

Therefore, it is possible to choose different shapes of distributions of the total number of proteins or \( P([n_q^{\text{total}}]) \), which will satisfy the constraints imposed by the properties of the chemotactic response and/or the available cell population and the single cell data for total protein abundances. We seek to estimate the maximally varying, or the least structured distribution of the stochastic signaling trajectories, where the minimal structure in the distribution of the protein abundances arise solely due to the constraints imposed by the available experimental data. Such a distribution represents the maximal cell-to-cell variations the system can endure while reproducing the experimentally measured data at the single cell and the cell population level.

In order to estimate this distribution which incorporates the available experimental data, we maximize the Shannon entropy \( S \) constructed from \( P_T \) [33]

\[
S = -\sum_T P_T \ln P_T
\]

in the presence of the constraints imposed by equations (4)–(6). We refer to the resulting distribution, \( \hat{P}_T \), as the constrained MaxEnt distribution. We reiterate that while searching for a MaxEnt solution we considered different \( P_T \) arising from different distributions of the total protein concentrations and selected the \( \hat{P}_T \) that satisfied all the imposed constraints and produced the maximum value of \( S \). More details on the implementation can be found in the methods section and the supplementary material available at stacks.iop.org/PhysBio/10/066002/mmedia. In order to compare distribution of protein total abundances, \( \hat{P}(n_q^{\text{total}}) \) corresponding to \( \hat{P}_T \) with the unconstrained case, we construct a uniform distribution of the total protein concentration \( \hat{Q}(n_q^{\text{total}}) \). We chose the uniform distribution as it has the maximum uncertainty. Maximization of the path entropy, defined in equation (7), is also known as the MaxCal distribution, which we refer to as the MaxCal [25, 33], which can also be derived in either the constrained or the unconstrained form.

Because the unconstrained MaxEnt (or MaxCal) represents the greatest robustness available to a given model in the absence of any data, while the constrained MaxEnt (MaxCal) represents the greatest robustness available to the model while constraining its behavior to conform to experimental results, the difference between the two is a measure of the degree to which the model must deviate from the uniform distribution in order to accommodate the data, or in other words, how great a departure from the uniform distribution is required to bring the model into accordance with empirical observations on the behavior of the system. Thus the model exhibiting the maximum robustness is the model with the minimal relative entropy (MinRE),

\[
\text{MinRE} = \sum \hat{P}(n_q^{\text{total}}) \ln [\hat{P}(n_q^{\text{total}})]/\hat{Q}(n_q^{\text{total}})].
\]

Note that MinRE is a particular form of the Kullback–Leibler distance [48] in which the probability ratio reflects the constrained versus unconstrained MaxEnt distributions and the expected value of the ln probability ratio is taken with respect to the former. One advantage of using MinRE rather than Shannon’s entropy \( S \) (equation (7)) in quantifying robustness is that, unlike \( S \), the lower bound for MinRE is always zero, which makes MinRE a better metric than \( S \).

Under the constraints in equations (4)–(6), the maximization of \( S \) will lead to an estimate of \( \hat{P}(n_q^{\text{total}}) \),

\[
\hat{P}(n_q^{\text{total}}) = Z^{-1} \exp \left[ -\sum_{i=1}^{r} \lambda_i \left( \sum_{c} f_{a|c} P_c \right) 
\right. 
- \sum_{i=1}^{m} \kappa_{a|c} \left( \sum_{c} f_{a|c} \right)^n P_c 
\left. \right] 
\times \exp \left[ -\sum_{q=1}^{N_q} \eta_q n_q^{\text{total}} - \sum_{q=1}^{N_q} \mu_q n_q^{\text{total}} \right] 
\times \exp \left[ -\sum_{c} P_c \ln P_c \right],
\]

where, average values and the \( n \)th order moments of chemotactic properties, indexed as 1 to \( r \) and 1 to \( m \), respectively, and, average values and the \( n \)th order moments of total abundances of proteins indexed by 1 to \( N_q \) and 1 to \( N_q \), respectively, have been constrained. In the above expression,
\[ P_C = P\left\{ n_j \right\}, \quad \tau; \quad \left\{ n_j \right\}, \quad t_0; \quad \left\{ n_j \right\} \]  

is the conditional probability of generating a stochastic trajectory \( \Gamma_C \) represented by \( \left\{ n_j \right\}, \quad \tau; \quad \left\{ n_j \right\}, \quad t_0; \quad \left\{ n_j \right\} \) given a fixed set of total protein abundances, \( \left\{ n_j^{\text{total}} \right\} \). The sum over \( \Gamma_C \) essentially denotes averages over variations of stochastic trajectories due to intrinsic noise fluctuations. The Lagrange’s multipliers in the above equation, the Gillespie method \[49\]. Variations and the MBL models using a continuous time Monte Carlo stochastic biochemical signaling processes in the FT, BL, and MBL models when the cell population averaged value of \( \tau \) (or \( s \)) was constrained to the experimentally observed values (figures \(3\left( B \right) \) and \( C \)). Similar to figure \(3\left( A \right)\), the FT model was much less robust compared to the BL and MBL models when \( \tau \) was held fixed at the experimentally observed value of 245 s (figure \(3\left( B \right)\)). The adaptation module of the FT model lacks an integral feed forward mechanism, and consequently performs rather weakly in the face of variations in the protein concentrations. Thus, only a very narrow range of protein abundances can generate perfect adaptation. The MBL model again produced slightly higher values of MinRE compared to the BL model, due to the greater number of cells that did not adapt well compared to BL model. This occurs for the same reason mentioned above, as poor adaptation also leads to poor precision of adaptation. All three models produced comparable values of MinRE, with MBL and BL producing the smallest and the largest values, respectively, when the average \( p \) values were constrained (figure \(3\left( C \right)\)). This occurred due to the following reason. Since restricting the steady state abundances of CheY-P requires correlated variations of protein abundances in all the models \[5\], a relatively smaller subset of cells that were randomly assigned correlated variation of protein abundances when the protein abundances were initially drawn from uniform distributions were able to produce values of \( p \) closer to the experiments. This raised the values of MinRE in all the models compared to the cases when \( \tau \) or \( s \) were constrained. The MBL model, designed to produce smaller variations of \( p \) for correlated variations of protein concentrations, produced the smallest values of MinRE. The FT model produced lower values of MinRE compared to the BL model, as the presence of the phosphatase CheZ in the FT

2. Quantification of robustness of \emph{E. coli} chemotaxis

We first considered the robustness of the three models when the average value of precision of adaptation (or \( s \)) over a cell population was constrained (figure \(3\left( A \right)\)). As expected from the robustness analysis of the models reported in the literature, the BL and MBL models produced substantially smaller values of MinRE compared to that of the FT model. This increased robustness of the BL and MBL models is due to the presence of an integral feedback control \[45\]. The MBL model produced slightly higher values of MinRE compared to the BL model, especially at very small values of the average precision. The reason for this is that the speed of adaptation depends on the total abundance of CheB (for BL) and CheB-P (for MBL) (see supplementary material available at stacks.iop.org/PhysBio/10/066002/mmedia). The models fail to adapt properly when the copy numbers of the enzyme (total CheB for BL and CheB-P for MBL) demethylating the active receptors become very small. Since the copy number of the phosphorylated form, CheB-P can be much smaller than that of CheB, it is more likely for MBL to generate cases that do not adapt (see supplementary material and supplementary figure \(1\) available at stacks.iop.org/PhysBio/10/066002/mmedia).

Next we calculated the values of MinRE for the three models when the cell population averaged value of \( \tau \) (or \( \bar{\tau} \)) was constrained to the experimentally observed values (figures \(3\left( B \right) \) and \( C \)). Similar to figure \(3\left( A \right)\), the FT model was much less robust compared to the BL and MBL models when \( \bar{\tau} \) was held fixed at the experimentally observed value of 245 s (figure \(3\left( B \right)\)). The adaptation module of the FT model lacks an integral feed forward mechanism, and consequently performs rather weakly in the face of variations in the protein concentrations. Thus, only a very narrow range of protein abundances can generate perfect adaptation. The MBL model again produced slightly higher values of MinRE compared to the BL model, due to the greater number of cells that did not adapt well compared to BL model. This occurs for the same reason mentioned above, as poor adaptation also leads to poor precision of adaptation. All three models produced comparable values of MinRE, with MBL and BL producing the smallest and the largest values, respectively, when the average \( p \) values were constrained (figure \(3\left( C \right)\)). This occurred due to the following reason. Since restricting the steady state abundances of CheY-P requires correlated variations of protein abundances in all the models \[5\], a relatively smaller subset of cells that were randomly assigned correlated variation of protein abundances when the protein abundances were initially drawn from uniform distributions were able to produce values of \( p \) closer to the experiments. This raised the values of MinRE in all the models compared to the cases when \( \tau \) or \( s \) were constrained. The MBL model, designed to produce smaller variations of \( p \) for correlated variations of protein concentrations, produced the smallest values of MinRE. The FT model produced lower values of MinRE compared to the BL model, as the presence of the phosphatase CheZ in the FT
Figure 3. MinRE quantifies robustness of different models for E. coli chemotaxis. (A) Variation of MinRE with the average precision of adaptation $\bar{s}$ for three different models (BL in blue, MBL in green and FT in red). (B) MinRE for the three different models when the average time is constrained to $\bar{\tau} = 245$ s. The color scheme is same as (A). (C) Variation of MinRE with $\bar{p}$ for three different models. The same color scheme as (A) has been used. (D) Shows variation of MinRE with $\bar{s}$ and $\bar{p}$ when $\bar{\tau}$ is held fixed to the experimentally measured value $\bar{\tau}_{\text{expt}}$ ($= 245$ s). (E) Variation of MinRE with $\bar{s}$ and $\bar{p}$ when $\bar{\tau}$ and $\bar{\tau}^2$ are set equal to their experimentally measured values, $\bar{\tau}_{\text{expt}}$ ($= 245$ s) and $\bar{\tau}^2_{\text{expt}}$ ($= 62323.5$ s$^2$), respectively.

model ensures that the steady state concentration of CheY-P stays relatively resilient to the variations in the protein abundances [5].

Next we simultaneously constrained $\bar{s}$, $\bar{\tau}$, and $\bar{p}$ (figure 3(D)). The MBL model displayed smaller values MinRE or higher robustness behavior compared to the BL and the FT model. This is because the MBL model is designed to produce smaller values of $\bar{p}$, therefore, it can accommodate more variation in protein abundances than the other models while reproducing the average values of $\bar{s}$, $\bar{\tau}$ and $\bar{p}$. The constraint imposed by $\bar{p}$ has a greater role in regulating the MinRE values in the three-constraints case, since holding $\bar{p}$ fixed within small range requires substantially restricted variations in protein abundances. This is reflected in differences in MinRE of one or two orders of magnitudes in figure 3(C) compared to figures 3(A) and (B). The BL
model produced lower values of MinRE compared to the FT model. However, MinRE values were comparable for the BL and FT models. The large separation in MinRE values produced by the FT and the BL models when $\bar{s}$ or $\bar{r}$ was constrained is reduced in figure 3(D), since constraining $\bar{p}$ to a small range ($< 30\%$) in both the models requires substantial reductions in the variations of protein abundances (figures 3(C) and supplementary figure 2(D) available at stacks.iop.org/PhysBio/10/066002/mmedia). Therefore, when all three variables, $\bar{s}$, $\bar{r}$ and $\bar{p}$ were constrained both the FT and the BL model produced similar values of MinRE.

Finally, we constrained the cell population averaged value of $\tau^2$ (or $\bar{\tau}^2$) in addition to the average values of $\bar{s}$, $\bar{r}$ and $\bar{p}$ (figure 3(E)). We checked whether the value of $\bar{\tau}^2$ is independent of $\bar{r}$ by comparing experimental data for $\tau$ with an exponential or a Poisson distribution; a Gaussian distribution appears to be a better fit to $P(\tau)$ (details in supplementary table 4 in the supplementary material available at stacks.iop.org/PhysBio/10/066002/mmedia) suggesting that $\bar{\tau}^2$ does not depend on $\bar{r}$. In this case, the relative rank ordering between the three models remains unchanged from the three-constraints case shown in figure 3(D). However, the separation between the FT and the BL models increases, since in the FT model the cases that displayed poor adaptation also produced much larger values of $\tau$ compared to the BL model. Therefore, the FT model required a greater restriction in the protein abundances to reproduce the experimentally observed value of $\bar{\tau}^2$ compared to the BL model. The relative rank ordering of the models (MBL > BL > FT) based on the MinRE values remain unchanged (supplementary figure S6A available at stacks.iop.org/PhysBio/10/066002/mmedia) when variations of $\tau$ and $s$ were further constrained in addition to the above constraints ($\bar{s}$, $\bar{r}$, $\bar{p}$, and $\bar{\tau}^2$).

Thus overall, the MBL model was found to be consistently the most robust model when the chemotactic responses were constrained. In the following section, therefore, we restrict attention to this particular model.

3. Predictions of distributions of the individual cell attributes

We compared the predictions from equation (9) regarding distributions of properties of the chemotactic response and specific protein abundances in individual cells with experiments for the MBL model. The predicted distribution for $\tau$ when $\bar{s}$, $\bar{r}$ and $\bar{p}$ were constrained to their experimental counterparts showed a wider distribution for $\tau$ compared to that observed in experiments (supplementary figure 3(A) available at stacks.iop.org/PhysBio/10/066002/mmedia). When the variance of $r$ was constrained along with $\bar{s}$, $\bar{r}$ and $\bar{p}$, the predicted distribution agreed reasonably well with the experiments (figure 4(A)). However, we found that constraining $\bar{r}$ and $\bar{\tau}^2$ alone to their experimental values also produced a reasonable agreement between the predicted distribution of $\tau$ and the experiment (supplementary figure 3(B) available at stacks.iop.org/PhysBio/10/066002/mmedia). This behavior could arise if $\tau$ is not substantially correlated with $s$ and $p$ in individual cells for the set of output constraints investigated. To test this conjecture, we calculated the Pearson correlation co-efficients $[50]$ ($r_{\tau s}$, $r_{\tau p}$ and $r_{p p}$) between $\tau$, $s$ and $p$, respectively, under the joint distribution of $\tau$, $s$, $p$ and $\tau^2$ when the values of $\bar{s}$, $\bar{r}$, $\bar{p}$ and $\bar{\tau}^2$ were constrained to their experimental values. We found that $r_{\tau s}$ has the largest value ($r_{\tau s} = 0.0349 \ll 1$, $r_{\tau p} = 0.0087$ and $r_{p p} = 0.0254$), implying that $\tau$ is not strongly correlated with $s$ and $p$ in individual cells, resulting in distributions of $\tau$ that are primarily regulated by $s$ and $\tau^2$ in single cells.

Next, we compared the predicted distributions for protein abundances in single cells when $\bar{s}$, $\bar{r}$ and $\bar{\tau}^2$ were constrained to their experimental values from the single cell experiments reported in [5, 37, 39]. We calculated the first six moments from the predicted distribution of CheY abundance in single cells and compared them with their experimental counterparts [5]. Figure 4(B) shows that the predicted distribution produced much larger values for the moments to those observed in the experiments. When we further constrained the averages, variances and the co-variances of CheY and CheZ to the experimentally observed values, the predicted moments for CheY abundance again showed excellent agreement with experiments (figure 4(C)). The higher moments deviate slightly upwards from the $y = x$ curve, which can be indicative of the fact that the actual CheY distribution is a log normal distribution with a longer tail [5]. The predictions for CheZ abundances also showed similar behavior to that of CheY when $\bar{s}$, $\bar{r}$, $\bar{p}$ and $\bar{\tau}^2$ were constrained alone or in combination with average values and variances of CheY abundances in individual cells (figures 4(D) and (E)). We note that the uncertainties in the values of the higher moments of the variables can be largely due to the small size of the available data. Variances were estimated assuming normal distributions for those variables [51] and we tested our predictions within those variances ($\pm 1$ standard deviation). We also checked the independence of the variances and co-variances of the abundances of CheY and CheZ from the mean values (supplementary table 4 available at stacks.iop.org/PhysBio/10/066002/mmedia). The availability of additional data would improve our ability to further test these predictions.

4. Discussion

We showed that a data-driven MaxEnt based approach can successfully quantify robustness of signaling models of E. coli chemotaxis. The robustness is measured considering cell-to-cell variations of protein abundances and chemotactic responses in a population of E. coli cells that reproduce the experimentally observed chemotaxis in the wild type E. coli strain RP437. This approach is markedly different from a class of traditional experimental or in silico methods where parameters regulating the cell function(s) are perturbed one at a time or simultaneously to measure robustness of a signaling system. The robustness of a model in such perturbation studies is quantified by considering the range of variation in the model parameters that can be accommodated without changing the output responses. The larger the range of the perturbations that can be tolerated, the greater is
Figure 4. Comparison of single cell attributes as predicted from the MaxEnt approach with experiments. (A) Comparison of the distribution of the adaptation time $\tau$ calculated from the MaxEnt distribution with the four constraints ($\bar{\tau} = 245 s$, $\bar{\tau'} = 62323 s^2$, $\bar{s} = 0.02$ and $\bar{p} = 20\%$) for the MBL model of *E. coli* chemotaxis and with experiments (black stairs). (B) Comparison of the first six moments calculated from the distribution of CheY abundance obtained from the MaxEnt approach ($x$-axis, log scale) for the MBL model with the same moments obtained from experiments ($y$-axis, log scale) as reported in [5]. The MaxEnt calculation was performed using the same constraints as in (A). The black line shows the $y=x$ graph. (C) Similar comparison between the MaxEnt predictions and experiments as in (B) for the moments calculated for the distribution of CheY. The MaxEnt approach used three constraints ($\bar{\tau} = 245 s$, $\bar{s} = 0.02$ and $\bar{p} = 20\%$) along with the average values, variances and co-variances of abundances of CheY and CheZ obtained from experiments. The averages for the CheY abundance were obtained from Li *et al* [40] and the variances of CheY and CheZ abundances and co-variance between CheY and CheZ abundances were taken from Kollmann *et al* [5]. (D) Shows comparison between predictions ($x$-axis, log scale) generated from the MaxEnt approach and experiments ($y$-axis, log scale) for the first six moments calculated from the distribution of CheZ abundances in single *E. coli* cells. The distribution of CheZ abundances in the MaxEnt approach is calculated for the same constraints as in (A). The black line shows the $y=x$ graph as a guide. The same sources as in (B) are used to obtain the experimental data. (E) Same comparison as in (D). The distribution of the CheZ abundance in the MaxEnt approach is calculated using the same constraints as in (C). The same sources as in (C) are used to obtain the experimental data.

The robustness. Our MaxEnt based approach considers the range of variation of the parameters in individual cells such that the cell population is able to reproduce the experimentally observed population averaged cell signaling responses. In this approach, the larger the range of the cell-to-cell variation of model parameters, the greater is the
robustness. We show that the MaxEnt based quantification of robustness (figure 3 and supplementary figure 2 available at stacks.iop.org/PhysBio/10/066002/mmedia) is in agreement with the estimation of robustness using traditional perturbation studies. The MBL model turned out to be the most robust model against cell-to-cell variations occurring from intrinsic and extrinsic noise fluctuations, followed by the BL model, while reproducing experimentally observed chemotactic responses in individual cells. The FT model was found to be substantially less robust than either the MBL or the BL model. These results are consistent with the results from the previous studies investigating sensitivity of *E. coli* chemotaxis against variations of protein concentrations and kinetic rates using *in silico* modeling and overexpression experiments [1, 5, 37]. The agreement of the rank ordering of these models based on our MinRE measure with the existing robustness analysis for the three models we analyzed validates our approach. Furthermore, it shows that the relative robustness of these models remains unchanged when cell-to-cell variations of chemotactic responses as well and protein abundances are included in the quantification of robustness.

The agreement between the robustness of different models from the MaxEnt analysis and the traditional perturbation experiments and simulations points to an interesting issue in biology. Did robust cell signaling systems evolve to produce larger cell-to-cell variations in parameters that can accommodate large perturbations without changing cell responses? The agreement between the two methods points us to an affirmative answer. We further probed this question by comparing the single cell distributions predicted from our MaxEnt analysis with available single cell measurements in *E. coli* chemotaxis. We found that the single cell distributions of protein abundances of CheY and CheZ could produce the experimentally observed distributions when the average values, variances, and co-variances were constrained to the experimental data. The rank ordering of the robust models was consistent with that obtained from perturbation experiments when the above constraints were imposed. This provides further indication that evolutionary selection of abundances or magnitudes of parameters in a signaling network in individual cells is influenced by their ability to control the robustness and sensitivity of cell functions.

Our MaxEnt based approach is similar in spirit to some recent work in parameter estimation techniques for biochemical networks using Bayesian methods [52–54]. These methods infer distributions of parameters (e.g., rate constants in signaling networks) by evaluating the posterior distribution of the parameters given the available experimental data using Bayes’ rule; models producing larger Bayes factor are then considered to be more appropriate for explaining the measured data [53, 54]. These methods require assumption of specific distributional forms for both priors and likelihoods. By contrast, the MaxEnt approach is free from such assumptions, so that the inference is solely guided by the available experimental data. The connection between Bayesian and MaxEnt approaches has been explored elsewhere [55, 56]. However, the precise connection between the MinRE metric developed here and the Bayes Factor used in the Bayesian approaches remains as an interesting topic of future work.

Technical limitations often make it difficult to perform single cell experiments due to lack of appropriate antibodies or small concentrations of expressed proteins in individual cells. In such situations immunoassays (e.g., western blot) measuring cell population level abundances of signaling proteins are used to decipher underlying mechanisms. The MaxEnt based approach can be used in conjunction with these types of experimental assays to find the least structured distribution of protein abundances and attributes of cell responses. As our study with *E. coli* chemotaxis shows, these distributions, even after being constrained to reproduce the average values from the experimental data, may need additional constraints regarding variances and co-variances of the variables in order to reach agreement with the experimental information. However, the estimated distributions calculated using constraints on average values indicate the limits of the cell-to-cell variations that can be allowed to observe the experimental data.

As mentioned earlier, robustness and sensitivity of cell responses represent two sides of the same coin. Here we used MinRE to determine the robustness, or insensitivity of chemotactic responses in *E. coli* cells to parameter variation. However, the same method can be used to determine parameters that sensitively regulate cell functions; these parameters will vary in individual cells within a narrow range. The joint distribution of these parameters can be used to calculate co-variances between the parameters. The covariance matrix analyzed using PCA can determine the most sensitive parameters or linear combinations of a subset of parameters that represent directions of sensitive perturbations [57, 58]. Therefore, the MaxEnt based analysis could be used to identify sensitive and insensitive or ‘sloppy’ parameters [59].

In our investigations, we held the kinetic rates describing strengths of protein–protein interactions in the models fixed in individual cells. This represents a reasonable assumption in the signaling models we investigated, since kinetic rates are largely determined by the thermodynamics of protein–protein interactions. Therefore, for experiments done at a fixed temperature, very little cell-to-cell variation in the rates is expected. However, when signaling models approximate interactions between two proteins, which can be affected by molecular crowding [60, 61] and/or cell shapes, then cell-to-cell variations of such kinetic rates should be included in the calculations. Moreover, coarse-grained *in silico* models often approximate multiple steps in biochemical reactions by one-step reactions, e.g., through the Michaelis–Menten approximation, where kinetic rates of the reactions depend on protein abundances. These reaction rates could vary in *in silico* models describing signaling kinetics in individual cells due to cell-to-cell variations of protein abundances [62]. Such variations in kinetic rate constants can be easily incorporated under our approach.

In many situations, experimentally measured values of the kinetic rates of biochemical reactions and protein concentrations in cell signaling and gene regulatory systems are unavailable, especially in higher organisms. Moreover, data
such as strengths of protein–protein interactions, e.g., obtained in vitro using truncated protein domains, may incorrectly describe those interactions in vivo. Therefore, an increasingly popular systems biology approach is to subject in silico models to perturbations of parameters (rate constants and proteins concentrations), using as a measure of robustness the insensitivity of key model outputs to such perturbations [34, 63]. For computational reasons, models with small numbers of sensitive parameters are generally preferred [63]. Our approach can also be used for analysis of in silico models in such settings. In the absence of any experimental data, user provided criteria for accepting a parameter set, such as production of cell population averaged concentrations of specific signaling proteins within a particular range, can be used for estimating robustness of these models.

The proposed approach is not free from some limitations of MaxEnt as a technique for inferring parameters [47, 64, 65]. First, in order to set up the MaxEnt model, one needs to start with a set of constraints that are relevant for the system. But when little is known about the biology or the functional response of a signaling network, it may be difficult to determine which are the relevant constraints from an available set of variables describing qualitative aspects of signaling kinetics. Second, it is possible for the inferred distributions of parameters to be inconsistent with experiments. This can be an informative result in its own right, since it may imply that additional constraints are required. However, increasing the number of constraints also requires solving for an increasing number of Lagrange multipliers, which can pose a significant computational challenge, in some cases requiring the use of computationally intensive Monte Carlo algorithms [66]. Since signaling networks can easily contain a large number of reactions and molecular species [67, 68], MaxEnt calculations for such networks can become computationally challenging. Third, as previously noted, the constraints used in equations (4)–(6) refer to the expected values of variables rather than the sample averages, and in finite samples these two quantities can differ from one another, introducing error into the estimation of the Lagrange multipliers [47]. Since the Lagrange multipliers appear in the exponential functions in the estimation of \( \hat{P}(n^\text{total}) \), a small change in a Lagrange multiplier can potentially produce a large change in \( \hat{P}(n^\text{total}) \) blurring the differences between relative entropies or MinRE values for different models. However, we have evaluated the current results for robustness to address this issue, and our tests show (supplementary figure S6B available at stacks.iop.org/PhysBio/10/066002/mmedia) that even allowing for substantial deviations between the sample averages and expected values, the rank ordering of models remains unchanged. Finally, in small samples, underlying dependencies of higher moments on lower moments could be difficult to detect. If present but not accounted for, such dependencies could lead to poor estimation of MinRE (or robustness) and a MaxEnt model with lower predictive power.

The MaxEnt based approach is general and can be used for a wide range of cell signaling systems. The ability of the approach to predict distributions of single cell attributes in a cell population makes it particularly useful in selecting the correct model when one can construct competing mechanistic models consistent with existing experimental measurements. We have observed a surge in development of new technologies for measuring signaling kinetics in single cells in the recent years. We believe the systems biology community will find the MaxEnt based approach useful for deciphering new mechanisms using single cell and cell population averaged data.

5. Materials and methods

5.1. Data from E. coli experiments

The average values of the chemotactic protein abundances were taken from Li et al [40]. We considered the chemotactic response for 100 \( \mu M \) L-aspartate stimulation. The distribution of the adaptation time was obtained from Min et al [39] by digitizing figure 3(C) in that paper using an online web plot digitizer (http://arohatgi.info/WebPlotDigitizer/). The values of \( \tilde{\tau} \) and \( \tilde{\tau}^2 \) were calculated from the distribution thus obtained. The cell population level value for precision of adaptation for wildtype RP437 strain was obtained from Alon et al [37]. In their experiments, the measured cell population averaged ratio of the steady state tumbling frequency of the wild type E. coli cells in the absence of any nutrient to that of when 1 mM L-aspartate was added in the medium was equal to 0.98 ± 0.05. The perfect adaptation corresponds to a ratio of 1.0. We considered precision of adaptation in individual cells. For a stochastic trajectory \( \Gamma \) (figure 2(B)), we defined the precision of adaptation \( s^2 \) as the absolute value of the relative difference between the population averaged abundance of pre-stimulus CheY-P at the steady state (\( \bar{N}_{\text{CheY-P pre-stim}} \)) and the post stimulus steady state abundance of CheY-P (\( \bar{N}_{\text{CheY-P post-stim}} \)) in individual cells. We calculated \( \bar{N}_{\text{CheY-P post-stim}} \) by using the CheY-P abundance in single E. coli cells evaluated at \( t = 2000 \) s, which is about eight times larger than the average adaptation time. The precision of adaption in a single cell is given by, \( s^2 = |\bar{N}_{\text{CheY-P pre-stim}} - \bar{N}_{\text{CheY-P post-stim}}| \). When single E. coli cells adapt perfectly, \( s^2 \equiv 0 \). We calculate \( \bar{s} \) from \( s^2 \) using equation (4). We varied \( \bar{s} \) from 0.005 to 0.05 for figure 3 as we have considered a concentration of 100 \( \mu M \) L-aspartate in our simulations instead of 1 mM used in [37]. For figure 4, we have used a \( \bar{s} \) value of 1 – 0.98 = 0.02. In order to make the calculations computationally efficient we calculated the cell-population averaged quantity, \( \bar{N}_{\text{CheY-P pre-stim}} \), by solving the ODEs which ignored the intrinsic noise of each individual E. coli because the contribution of the intrinsic noise fluctuation to this average value was small (supplementary figure 4 available at stacks.iop.org/PhysBio/10/066002/mmedia).

The steady state abundance of CheY-P varies from cell to cell due to the variations of total protein abundances in individual E. coli cells. The variation of the steady state abundance of CheY-P needs to be within 30% from an optimal value for proper functioning of the flagellar motor [5]. We calculated the variation of steady state CheY-P abundance (\( p \)) in single E. coli cells using the equation

\[ p = \frac{\bar{N}_{\text{CheY-P pre-stim}}}{\bar{N}_{\text{CheY-P post-stim}}} \]
below, \( p = \frac{N_{\text{CheY-Pre-stim}} - N_{\text{CheY-optimal}}}{N_{\text{CheY-optimal}}} \). Optimal value of CheY-P, \( N_{\text{CheY-optimal}} \), is defined as the ODE based solution of the steady state value of the CheY-P when the total protein concentrations are set to the values quoted in Li et al [40]. \( \hat{p} \) was calculated from \( p \) using equation (4). For figure 3, \( \hat{p} \) has been varied from 10 to 30% whereas for figure 4 we have used an ad hoc value of 20%. We calculated \( N_{\text{CheY-P Pre-stim}} \) using the ODE solutions ignoring intrinsic noise fluctuations for the reasons mentioned above.

The single cell CheY distribution for the wildtype RP437 strain was taken from Kollman et al [5]. The plot was digitized using the web plot digitizer and the x-axis was rescaled to obtain the average number of the CheY-P quoted in [40]. The moments for abundances of CheY were calculated using this distribution. The moments for CheZ abundances were computed by digitizing the co-expression plot in [5].

5.2. Computational method

We use a rule based modeling software package BioNetGen [68] for the solution of the ODEs as well as to perform continuous time Monte Carlo. Using the BioNetGen software for simulating the models allows us to share our codes easily, in addition to varying the parameters efficiently in the simulations. The codes are available at http://planetx.nationwidechildrens.org/~jayajjit/. The biochemical networks for the BL and the MBL models are curated from [5]. The FT model is constructed by adding an extra module to the MBL model where CheB-P is allowed to de-phosphorylate the inactive receptors. All the simulations were initialized at \( t_{\text{initial}} = -800,000 \) s with protein abundances, \( \text{Tar} \), \( \text{CheRT} \), \( \text{CheB} \), \( \text{CheR} \), \( \text{CheY} \), and \( \text{CheZ} \) are set to zero. The superscript \( T \) refers to total abundances of the respective protein in a single cell. The non-zero protein abundances at \( t_{\text{initial}} \) are drawn from a uniform distribution \( U(0, U_{\text{F}}) \), where \( U_{\text{F}} \) is chosen to be roughly 10 times larger than the experimentally measured mean abundance of the corresponding chemotactic protein [40] (see supplementary table 2 available at stacks.iop.org/PhysBio/10/066002/mmedia for the mean values used in our MaxEnt calculations). Once a set of initial protein abundances is chosen, we solve the ODEs describing the chemotactic kinetics to obtain steady state values of the protein abundances. These steady state values for the protein abundances are used as initial conditions to simulate the chemotactic response of individual cells when 100 \( \mu \)M of L-aspartate is added to the medium. The time when the ligand is added is considered to be \( t = 0 \). The chemotactic response of the E. coli, given by the kinetic of the protein abundances, is calculated by solving the Master Equation in equation (3) exactly by the Gillespie method. The ligand receptor interactions are approximated by the rates at which the receptors can become active. The adaptation time \( \tau_{\text{G}} \) and the precision of adaptation \( \sigma_{\tau} \) are calculated using for each stochastic trajectory \( \Gamma \) representing the chemotactic response in an individual cell composed of abundances of signaling proteins recorded at regular time intervals for a time period of \( t_0 = 0 \) to \( t_0 = 2000 \) s; the time scale, \( t_0 = 2000 \) is much larger than the typical adaptation time for E. coli for a 100 \( \mu \)M L-aspartate stimulation. When the Che-Y-P abundance in an individual cell does not recover to the half of the pre-stimulus Che-Y-P level within 2000 s, we assign a very number \( (6 \times 10^8 \text{ s}) \) to \( \tau \) to mark the cell that did not adapt in a realistic time scale. In size of the sample (up to 70,000 single E. coli cells), each individual E. coli cell produced a unique stochastic trajectory. Therefore, in our simulations each stochastic trajectory could be identified with a single cell.

5.3. Calculation of MinRE

We seek for a solution of the form given in equation (9), when the average values and the \( n \)th moments of the chemotactic responses \( \{f_{\text{tar}}\} \) and the protein abundances \( \{n_{\text{pro}}\} \) are constrained to the value measured in the experiments. The Lagrange multipliers \( \{\lambda_n\} \), \( \{\kappa_n\} \), \( \{\mu_q\} \), and, \( \{\eta_n\} \) are obtained by solving the set of nonlinear simultaneous equations when equation (9) is substituted in equations (4)–(6) describing the constraints. We carry out the summation over the stochastic trajectories \( \Gamma_C \) then evaluate average values in the constraint equation in the following way. In the sample size we considered, each E. coli cell produces a unique chemotactic response composed of a stochastic trajectory describing time evolution of abundances of signaling proteins, therefore, we identified each trajectory by the single cell that generated it (figure 2). The single cells were assigned with identification numbers, such as cell #1, cell#2 and so on. The summation over the trajectories then is essentially the summation over the single cells used in the simulation. The uniqueness of the stochastic trajectories in our simulations also implies that for a particular stochastic trajectory, \( \Gamma_C = \{ \{n_j\}, t_0; \{n_j\}, t_{n-1}; \ldots; \{n_j\}, t_0 \} \) for a fixed set of total protein concentrations \( \{n_{\text{pro}}\} \), \( P_C = P(\{n_j\}, t_0; \{n_j\}, t_{n-1}; \ldots; \{n_j\}, t_0 \mid \{n_{\text{pro}}\}) \) is either equal to 1 (when \( \Gamma_C \) or the corresponding single cell is present in the samples we considered) or 0 (when \( \Gamma_C \) is absent in the samples). Thus, \( \sum_{\Gamma'} P_{C} \ln P_{C} = 0 \) and \( \sum_{\Gamma'} f_{\text{tar}} P_{C} = f_{\text{tar}} \Gamma_C \) for the trajectories we analyzed in the simulations. We used up to 70,000 single cells, the convergence of the results with the number of cells used is shown in the supplementary material (supplementary figure 5 available at stacks.iop.org/PhysBio/10/066002/mmedia). The Lagrange multipliers are then calculated using standard techniques used for solving nonlinear algebraic equations (web supplement). Then we calculate the minimum relative entropy MinRE given by equation (8). It is in principle possible that when a very large number of cells are present, two different single cells could produce the same stochastic trajectory and for such cases, \( \sum_{\Gamma} P_{C} \ln P_{C} \) will not vanish and \( \sum_{\Gamma} f_{\text{tar}} P_{C} \) will contain averages over multiple trajectories. However, occurrences of such events (e.g., the presence of pairs of identical stochastic trajectories) appear to be extremely rare for the rate constants and the ranges of the protein abundances we considered. We further tested this approximation by considering deterministic chemotactic...
signaling kinetics where the kinetics of signaling protein abundances only depend on the total protein abundances (as the kinetic rates are fixed for each cell), therefore, $P_C = 1$ when the deterministic kinetic trajectory of abundances of signaling proteins is present and $P_C = 0$, otherwise. When we used the same a priori uniform distribution for protein abundances as our stochastic simulations, the rank ordering of the FT, BL, and MBL models based on the MinRE values did not change compared to the stochastic simulations (supplementary figure S7 available at stacks.iop.org/PhysBio/10/066002/mmedia). The small differences in the values of the MinRE between the stochastic and the deterministic simulations show the dominance of extrinsic noise fluctuations over intrinsic noise fluctuations in determining robustness of the models (supplementary figure S7 available at stacks.iop.org/PhysBio/10/066002/mmedia). These results also demonstrate that associating a unique stochastic trajectory to a single cell is a good approximation for the calculation of MinRE in the stochastic simulations.

5.4. Prediction of single cell properties

Our MaxEnt approach allows us to predict distributions of single cell properties given by equation (9). The Lagrange multipliers in equation (9) are calculated using the procedure described above. Once the Lagrange multipliers are known, equation (9) essentially predicts the probability for an individual cell (indexed by the stochastic trajectory $\Gamma$) displaying chemotactic responses $\{r_{\Gamma}\}$ and the protein abundances $\{p_q^{\text{final}}\}$. We use this probability to calculate occurrence probabilities for the cells in the cell population that were initially assigned with protein abundances chosen from uniform distributions. The occurrence probability evaluated for an individual cell gives the maximally broad probability with which that particular cell should be present in the cell population so that the cell population is able to produce the population level and single cell measurements that were constrained in the MaxEnt calculation. For example, when $\tilde{\tau}$ is constrained to its experimentally measured value $\tilde{\tau}^{\exp}$, then the MaxEnt calculation produces a solution $P_{\tilde{\tau}} = Z_\tilde{\tau}^{-1} \exp[-\lambda_\tilde{\tau}, \tau_{\tilde{\tau}}]$, where, $\tau_{\tilde{\tau}}$, is the adaptation time for an individual cell executing a stochastic trajectory, $\Gamma$, and, $\lambda_\tilde{\tau}$ is the Lagrange multiplier. Once, $\lambda_\tilde{\tau}$ is evaluated using $\tilde{\tau}^{\exp}$, $P_{\tilde{\tau}}$ gives the probability that an individual cell, $\Gamma$, is selected in the cell population. The relationship between $P_{\tilde{\tau}}$ and $p(\tau)$ is given by equation (1). Following the same scheme, the other single cell properties, such as CheY abundance, or, CheZ abundance are evaluated. The moments of the distributions are then calculated using those distributions.

Acknowledgments

This work was supported by funding from the Research Institute at Nationwide Children’s Hospital and NIH grant AI090115 to JD, and NIH grant MH086117 to VJV. JD thanks C Jayaprabhash for discussions. We thank both the anonymous reviewers for making constructive suggestions.

JD, SM and VJV planned the research and analyzed the data; SM and SS performed simulations and calculations. JD, SM and VJV wrote the paper.

We declare no conflict of interest.

References

[1] Barkai N and Leibler S 1997 Robustness in simple biochemical networks Nature 387 913–7
[2] Chau A H, Walter J M, Gerardin J, Tang C and Lim W A 2012 Designing synthetic regulatory networks capable of self-organizing cell polarization Cell 151 320–32
[3] Eldar A, Dorfman R, Weiss D, Ashe H, Shilo B Z and Barkai N 2002 Robustness of the BMP morphogen gradient in drosophila embryonic patterning Nature 419 304–8
[4] Ma W, Trusina A, El-Samad H, Lim W A and Tang C 2009 Defining network topologies that can achieve biochemical adaptation Cell 138 760–73
[5] Kollmann M, Lodovik L, Bartholome K, Timmer J and Sourjik V 2005 Design principles of a bacterial signalling network Nature 438 504–7
[6] Kitano H 2004 Cancer as a robust system: implications for anticancer therapy Nature Rev. Cancer 4 227–35
[7] Carlson J M and Doyle J 2000 Highly optimized tolerance: robustness and design in complex systems Phys. Rev. Lett. 84 2529–32
[8] Kitano H 2004 Biological robustness Nature Rev. Genetics 5 826–37
[9] Feinerman O, Veiga J, Dorfman J R, Germain R N and Altan-Bonnet G 2008 Variability and robustness in T cell activation from regulated heterogeneity in protein levels Science 321 1081–4
[10] Daniels M A, Teixeiro E, Gill J, Hausmann B, Roubaty D, Holmberg K, Werlen G, Hollander G A, Gascoigne N R and Palmer E 2006 Thymic selection threshold defined by compartmentalization of Ras/MAPK signalling Nature 444 724–9
[11] Das J, Ho M, Zikherman J, Govern C, Yang M, Weiss A, Chakraborty A K and Roose J P 2009 Digital signaling and hysteresis characterize Ras activation in lymphoid cells Cell 136 337–51
[12] Prasad A, Zikherman J, Das J, Roose J P, Weiss A and Chakraborty A K 2009 Origin of the sharp boundary that discriminates positive and negative selection of thymocytes Proc. Natl Acad. Sci. USA 106 528–33
[13] Elowitz M B, Levine A J, Siggia E D and Swain P S 2002 Stochastic gene expression in a single cell Science 297 1183–6
[14] Korobkova E, Emonet T, Vilar J M, Shimizu T S and Cluzel P 2004 From molecular noise to behavioural variability in a single bacterium Nature 428 574–8
[15] Kampen N G V 1992 Stochastic Processes in Physics and Chemistry (Amsterdam: North-Holland)
[16] Swain P S, Elowitz M B and Siggia E D 2002 Intrinsically and extrinsically induced periodicities in gene expression Proc. Natl Acad. Sci. USA 99 12795–800
[17] Dayarian A, Chaves M, Sontag E D and Sengupta A M 2009 Shape, size, and robustness: feasible regions in the parameter space of biochemical networks PLoS Comput. Biol. 5 e1000256
[18] Hafner M, Koeppl H, Hasler M and Wagner A 2009 ‘Glocal’ robustness analysis and model discrimination for circadian oscillators PLoS Comput. Biol. 5 e1000534
[19] Bialek W 2012 Biophysics: Searching for Principles (Princeton, NJ: Princeton University Press)
[20] Artyomov M N, Das J, Kardar M and Chakraborty A K 2007 Purely stochastic binary decisions in cell signaling models without underlying deterministic bistabilities Proc. Natl Acad. Sci. USA 104 18958–63
