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Distributed classifier based on genetically engineered bacterial cell cultures

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Abstract

We describe a conceptual design of a distributed classifier formed by a population of genetically engineered microbial cells. The central idea is to create a complex

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classifier from a population of weak or simple classifiers. We create a master population of cells with randomized synthetic biosensor circuits that have a broad range of sensitivities towards chemical signals of interest that form the input vectors subject to classification. The randomized sensitivities are achieved by constructing a library of synthetic gene circuits with randomized control sequences (e.g. ribosome-binding sites) in the front element. The training procedure consists in re-shaping of the master population in such a way that it collectively responds to the “positive” patterns of input signals by producing above-threshold output (e.g. fluorescent signal), and below-threshold output in case of the “negative” patterns. The population re-shaping is achieved by presenting sequential examples and pruning the population using either graded selection/counterselection or by fluorescence-activated cell sorting (FACS). We demonstrate the feasibility of experimental implementation of such system computationally using a realistic model of the synthetic sensing gene circuits.

Keywords

Chemical pattern recognition, consensus classification, distributed sensing, machine learning, microbial population engineering, synthetic circuits

Introduction

Pattern recognition and classification is one of the most important statistical disciplines.¹ Its applications span across disciplines such as computer vision,² natural language processing,³ search engines,⁴ medical diagnosis,⁵ classification of DNA sequences,⁶ speech recognition,⁷ computational finance,⁸ fraud detection,⁹ and many others.¹⁰ In these con-

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7 texts usually a system that solves a pattern recognition problem learns from the “training”
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9 data presented to it. Using the “training” data such a system forms an internal model to
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11 classify new previously unseen data. Typically these models are built in regular comput-
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13 ers, although alternative approaches have been proposed.^{11,12}

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15 Many pattern recognition algorithms are biologically motivated. Biological organisms
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17 perform decision making based on classification of external environmental cues at all levels
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19 from intracellular (e.g. ref. 13) to organismal¹⁴ and even population-wide.¹⁵ The devel-
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21 opment of artificial neural network (ANN) classifiers was inspired by the brain’s natural
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23 ability to perform complex computational and classification tasks.¹⁶ The main principles
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25 of brain dynamics, its layered organization and ability to learn by adapting strengths
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27 of inter-neuron synaptic connections (plasticity) is mimicked in ANN by multilayered
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29 perceptrons and various learning algorithms.¹⁷

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31 A different learning approach is motivated by the adaptive immune system of jawed
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33 vertebrates which employs a population of lymphocytes (T and B cells) with a diverse
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35 genetically encoded library of recognition specificities in order to implement learning,
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37 memory, and pattern recognition capabilities.^{18,19} Lymphocytes with different receptor
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39 variants undergo essentially a supervised learning procedure in central lymphoid organs,
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41 being presented self-antigens as examples. Positive selection retains T cells capable of
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43 interacting with the major histocompatibility complex while negative selection eliminates
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45 self-reactive T and B cells. Subsequent exposure of mature lymphocytes to foreign anti-
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47 gens results in positive selection of the reactive clones. The adaptive immunity classifier
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49 is not a single complex multivariate system with parameters adjusted in course of learn-
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51 ing, Rather it is a distributed system that consists of a large number of relatively simple
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53 cellular classifiers and implements learning through deletion of outliers. When trained, it
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55 solves a consensus classification problem, reporting the absence of pathogens if and only
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7 if each cellular classifier does not respond to presented antigens. The main principles of
8 the natural adaptive immune system of jawed vertebrates have inspired the branch of
9 computer science known as Artificial Immune Systems (AIS).^{20–22}

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12 In this work we propose to use synthetic biology to adapt biological systems them-
13 selves for solving complex classification tasks. While the most straightforward solution
14 for this problem would be to design a single gene circuit that would produce output signal
15 sufficient for classification, in practice the uncertainty and wide dynamic range of possible
16 signals of interest would render this solution suboptimal. In such situations it appears
17 useful to borrow the principles underlying the distributed classification abilities of nat-
18 ural adaptive immune system and create a heterogeneous population of microorganisms
19 with different synthetic gene circuits capable of performing classification tasks based on
20 consensus strategy. The desired binary classifier should produce a positive response (for
21 example, above-threshold population-averaged fluorescence level) to positive inputs, and
22 negative response (below-threshold fluorescence) to negative ones. The learning algo-
23 rithm then should consist of shaping the population in such a way that the population
24 collectively “arrives” at a probably correct decision.

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27 The idea of aggregating many simple classifiers to yield a better classifier is a widely
28 used strategy in machine learning that capitalizes on the idea that using a set of classifiers
29 that produce barely better results than random guessing can achieve arbitrarily high
30 accuracy when combined appropriately.²³ It also can be cast as a function approximation
31 problem in which a complex target function is approximated by a weighted sum of multiple
32 simpler functions, such as radial basis functions²⁴ or wavelets.²⁵ Inspired by these ideas,
33 we propose to use genetically engineered cells with limited abilities to perform complex
34 classification tasks.

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37 Here we present a specific biological implementation of this general distributed pattern

recognition system concept using a model of synthetic gene regulatory circuits in engineered cell populations. The proposed implementation requires to build a cell library with genetically encoded randomized sensitivities to external chemical signals to be classified. Such libraries have been successfully constructed for optimization of synthetic biological circuits.²⁶⁻²⁸ One could in principle envision a system in which different strains were placed in different chambers and probed separately, and then the classification task would be done *in silico*. However, this approach would entail a complex multi-chamber, multi-channel system that would be difficult to operate in an open environment. Instead, we propose here to use a library to form a *master classifier* population that is pruned to learn how to solve a certain classification task. In the properly trained system, classification is based on a single population-averaged output, which simplifies the device design and operation considerably. The learning is done by examples: cells with erroneous outputs are gradually attenuated from the population, while the “correct” cells are amplified. As a result of multiple iterations of pruning/amplification, a distributed classifier trained for a specific task emerges from the master population. We envision that an arbitrary external input subject to recognition can be encoded by a combination of chemical inputs capable of generating the engineered cellular response. In this paper, however, we will consider the most straightforward case when the vector of chemical concentrations *is* the input signal subject to classification. In the following we demonstrate the general principle of this classification procedure and describe its implementation using a model of a synthetic genetic circuit based on the lambda phage P_{RM} promoter.¹³

Results and discussion

Learning by examples

In this section we describe the general idea behind the training of a distributed classifier by presenting a set of positive and negative examples. In the following, we denote the set of input variables to be classified as $\mathbf{x} \in \Re^M$. A classifier is the function $y = 2H(f(\mathbf{x}) - \theta) - 1$, such that if $f(\mathbf{x}) > \theta$, the answer is $y = 1$, and otherwise, $y = -1$. Here $H(\cdot)$ is the Heaviside function, θ is a scalar threshold, and $f(\mathbf{x})$ is the scalar function of the inputs. The heart of the pattern classifier is the function $f(\mathbf{x})$ that minimizes the classification errors for a given distribution of positive and negative inputs.

In general, the classifier function is not known *a priori* and has to be learned by training the classifier using examples (training data). The training of a classifier by examples consists in finding a function $f(\mathbf{x})$ that minimizes the error in mapping of a given set of N examples $\mathbf{x}_i$ to a set of binary outputs, $y_i = \{-1, +1\}$ which label the examples. In the following, we will say that if the output $y_i = +1$, the example i belongs to the “positive” class and if the output is $y_i = -1$ the example i belongs to the opposing “negative” class.

In our proposed population-based classifier, $\mathbf{x}$ will be a set of concentrations of chemical signals which cells are subjected to, and the cells are assumed to contain gene circuits that produce a fluorescent signal $z_i(\mathbf{x})$ in response to the input signal $\mathbf{x}$. The overall signal function, $f(\cdot)$, will be the normalized linear sum of the individual fluorescence signals from all N_c cells in the trained population:

$$f(\mathbf{x}) = \frac{1}{N_c} \sum_{i=1}^{N_c} z_i(\mathbf{x}). \quad (1)$$

The key to the “trainability” of the distributed classifier is to prepare a master population of cells with broadly varied functions $z_i(\mathbf{x})$, so this population can be appropriately shaped to perform the needed classification task. This can be achieved using synthetic gene regulatory circuits with randomized control elements such as promoter regions, ribosome binding sites, or other sequences as described in detail below.

We assume that the master population contains cells that individually provide correct answers to subsets of the data to be classified, but that in general no single cell provides correct answers to all data (weak classifiers). The goal of training is therefore to shape the master population to create a distributed consensus classifier that performs better than any cell individually. Such training must amplify the cells providing correct answers as frequently as possible and conversely suppress the cells with poor overall performance. Thus, our proposed learning procedure consists in modifying the composition of the cell population based on the examples with known outcomes. The details of our training algorithm for the specific implementation of a genetic sensing circuit are outlined below.

Distributed classification with randomized synthetic gene sensors

Here we outline an implementation of the proposed distributed classifier with a diverse population of bacterial cells containing synthetic sensory genetic circuits with randomized parameters. For simplicity we focus on a scalar input with only one chemical signal affecting the gene circuit, however the same approach can be straightforwardly generalized to the multi-dimensional vector input. In our two-gene design (Figure 1) the sensing and the reporting functionalities are split between the two genetic modules. The sensing module is monotonically induced by the external chemical signal X and drives the synthesis of a transcription factor U . The second promoter is regulated by U and drives the expression of a reporter protein, for example, green fluorescent protein (GFP). The re-

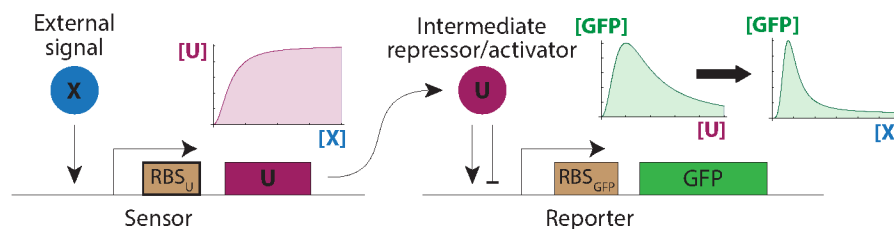
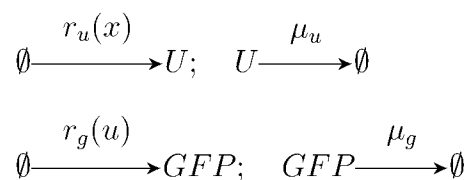


Figure 1: A modular genetic circuit proposed for implementing a distributed genetic classifier. Sensing and response functionalities are split into separate modules. In the first module (sensor), an inducible promoter drives the expression of the transcription factor U in response to the applied signaling molecule X . The response function of the promoter is chosen to be monotonic (see inset). In the second module (reporter), another inducible promoter drives the expression of a reporter (GFP) in response to induction by U . The promoter is activated by intermediate concentrations of U and inhibited by high concentrations of U . Thus the resulting response function of the entire two-promoter circuit to the concentration of signaling molecule is bell-shaped for the relevant values of the circuit parameters as shown on the inset.

porter promoter is activated by U at intermediate concentrations and inhibited at higher concentrations, thus being active only within a finite range of concentrations of U . The classic well-characterized example of such promoter is the promoter P_{RM} of phage lambda which is activated by intermediate concentrations and is repressed by high concentrations of the lambda repressor protein CI.^{13,29}

This two-gene circuit can be modeled by the following set of biochemical reactions



where x and u are the concentrations of X and U , $r_u(x)$ and $r_g(u)$ are the effective production rates of U and GFP , respectively, and μ_u and μ_g are the degradation rates of U and GFP . The rates of gene expression in this system can be described by standard

Hill functions:

$$r_u(x; m_u) = m_u \cdot \frac{\alpha A_u^{p_u} + x^{p_u}}{A_u^{p_u} + x^{p_u}} \quad (2)$$

$$r_g(u; m_g) = m_g \cdot \frac{A_g^{p_g} u^{p_g}}{(A_g^{p_g} + u^{p_g})^2} \quad (3)$$

where α describes the basal expression from the sensor promoter in the absence of the signaling molecule X , A_u is the dissociation constant of X with the sensor promoter, the Hill coefficient p_u characterizes the cooperativity of activation of the sensor promoter, p_g characterizes the cooperativity of activation and repression of the reporter promoter by the transcription factor U , A_g is the dissociation constant for activation and repression of the reporter promoter by U (we assume the activation and repression cooperativities and dissociation constants to be the same), m_u and m_g describe the overall strength of production of U and GFP respectively. Noteworthy, such Hill functions based model provides a simple yet adequate approximation of more complex response functions required to describe real promoters²⁹ (see Supporting Information, section 3).

In mass-action approximation the dynamics of GFP production in this system is described by the following system of ordinary differential equations:

$$\frac{du}{dt} = r_u(x; m_u) - \mu_u u \quad (4)$$

$$\frac{dz}{dt} = r_g(u; m_g) - \mu_g z \quad (5)$$

where z is the concentration of GFP. The steady state concentration of GFP as a function of the concentration of the external chemical signal X can be found from the equations (4) and (5) as

$$z^*(x; m_u, m_g) = \frac{r_g(r_u(x; m_u)/\mu_u; m_g)}{\mu_g} \quad (6)$$

The function $z^*(x)$ is bell-shaped in a broad range of $m_u/\mu_u \in (A_g, A_g/\alpha)$ (Figure 2). Varying m_u/μ_u allows to create a library of circuits which act as low-pass filters ($m_u/\mu_u \leq A_g$), high-pass filters ($m_u/\mu_u \geq A_g/\alpha$), or tunable bandpass filters for the intermediate values of m_u/μ_u . As described in the following sections, such a library can be used to train a cell population-based distributed classifier. Since common sensor promoters can have a regulatory range of over 10^3 ($\alpha = 10^{-3}$),^{30,31} to create a library that contains low-pass, bandpass, and high-pass filters, the m_u/μ_u ratio has to be varied at least $1/\alpha = 10^3$ fold within $[A_g, A_g/\alpha]$ range (see Figure 2). Such libraries have been widely constructed experimentally: m_u can be varied over more than 10^5 fold range by varying the DNA sequence within and near the ribosome binding site of the gene of interest;^{32,33} this range can be further expanded by modulating the sensor promoter strength;³⁴ m_u/μ_u ratio can also be modulated by varying the stability of the U coding mRNA as well as the stability of the protein U itself.^{26,35–37}

The modular architecture of the classifier circuit proposed here allows us to independently select and optimize the sensor and the reporter functionalities in order to maximize the classifier performance. Well-characterized sensor promoters, which can be induced by a variety of chemical signals with the appropriate monotonic response, are common^{30,31,38–40} and can be readily combined with the reporter promoter such as the well characterized lambda phage promoter P_{RM} .^{13,29} For these reasons the proposed two-gene sensory circuit appears to be well-suited for experimental implementation of a distributed cell population based classifier.

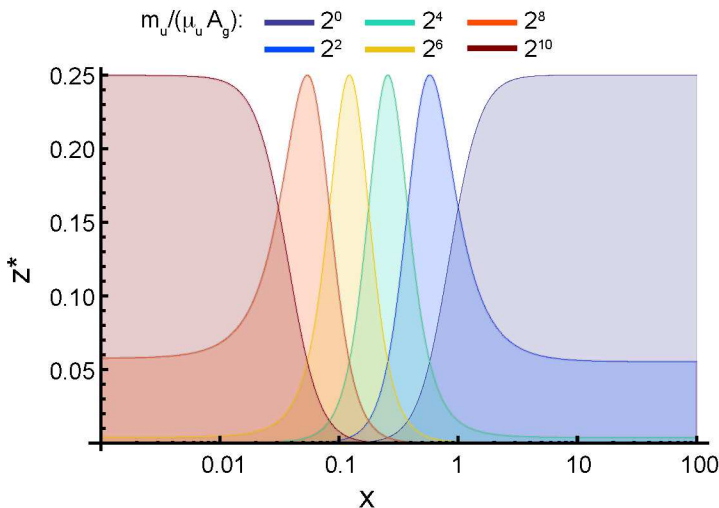


Figure 2: Steady state GFP concentration (z^*) as a function of the concentration of the external chemical signal X for the modular classifier circuit shown for a range of m_u values representing a range of the relative strengths of the sensor promoter (Figure 1). Non-dimensional circuit parameters are $\mu_u = \mu_g = m_g = A_u = 1$, $A_g = 20$, $p_g = p_u = 2$, $\alpha = 10^{-3}$.

Classifier training algorithm

In order to train a classifier we need to be able to sort individual cells based on their response to a set of known examples. A hard-decision algorithm implies that if a given example x_i belongs to a positive class, $y_i = +1$, and the GFP level in j -th cell is above a threshold $z_j^*(x_i) > \theta$, then that particular cell should be retained in the population because the cell provides the correct answer. Meanwhile, the cells not reaching the threshold expression level after presenting a positive example should be removed from the population. On the other hand, if a negative example is presented ($y_i = -1$), the cells generating above-threshold fluorescence should be eliminated and the cells which are below threshold should be retained. By this selection mechanism we ensure that only the cells generating correct answers survive.

However, this hard-decision training algorithm in most practical situations leads to

poor performance. As mentioned above, in general each cell is a weak classifier and so it can not provide the complete classification solution. If positive and/or negative categories encompass a broad range or several distinct ranges of inputs, a sub-population in which all cells generate above-threshold output for positive examples and below-threshold output for negative examples would be empty. Thus, an outright elimination of all cells producing “incorrect answer” to any of many training examples would eventually lead to elimination of all cells. To avoid this undesirable outcome, a “soft-training” algorithm has to be employed in which even the “erroneous” cells have a chance to remain in the population, and so the resultant population-based classifier will produce the correct answer only by the population average, not the unanimous decision of all cells.

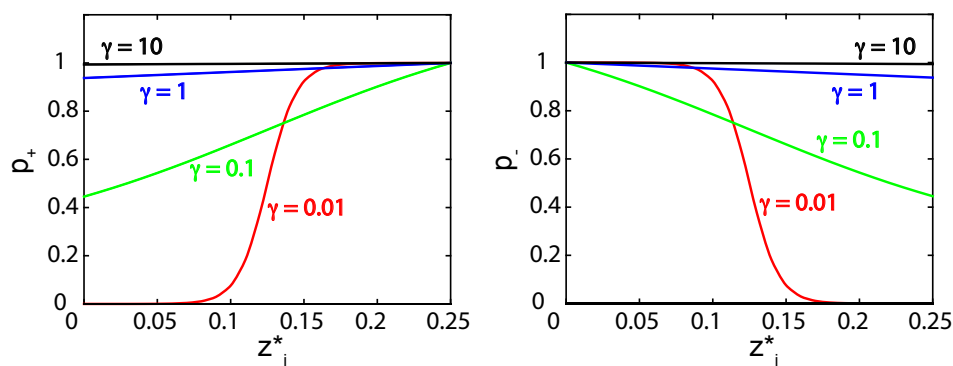


Figure 3: Cell survival probabilities during training upon presentation of a positive ($p_+(z_j^*; \gamma)$) or a negative example ($p_-(z_j^*; \gamma)$).

We postulate that the elimination of cells from the population is governed by two sigmoidal cell survival probability functions $p_+(z_j^*) = (1 + \xi)^{-1} + (1 + \xi \exp(-z_j^*/\gamma))^{-1}$ and $p_-(z_j^*) = 1 + (1 + \xi)^{-1} - (1 + \xi \exp(-z_j^*/\gamma))^{-1}$ for positive and negative examples correspondingly, where $\xi = \exp((8\gamma)^{-1})$ and $0 \leq z_j^* \leq 1/4$ are chosen to encompass the entire range of possible values of z_j (eq.(6)) (Figure 3). These functions are chosen such that the cells that respond perfectly to the presented example ($z_j^* = 1/4$ or $z_j^* = 0$ for a positive or a negative example correspondingly) are retained in the population

($p_{\pm}(z_j^*) = 1$). Otherwise the cells are eliminated from the population with a probability that depends on the cell fluorescence z_j^* and the “rigidity” of training γ . For very small γ , cells not responding to the currently presented input are eliminated with high probability. This can potentially eliminate many cells required to recognize other positive examples x_i from the pattern being taught leading to the poor overall performance. For too large γ (weak elimination), the learning rate slows down significantly requiring unreasonably large number of training iterations in order to achieve high performance. Therefore an optimal value of the parameter γ can be chosen to balance performance with the learning speed. In general this value will be different for each individual problem. Experimentally these selection functions can be implemented via fluorescence activated cell sorting (FACS) or well established genetically encoded positive/negative selection systems.⁴¹ Conveniently, in the latter case positive or negative example is indicated by applying the corresponding selective compound to the cells and its concentration can be used to adjust the rigidity of training (γ).

As usual in the design of classifiers, the output element of the classifier must be a threshold element. In the distributed classifier described here, the mean population fluorescence $f(x)$ has to be compared with a suitably chosen threshold θ such that the above-threshold fluorescence ($f(x) > \theta$) corresponds to the positive class, and below-threshold fluorescence ($f(x) < \theta$) to the negative class. The optimal threshold θ can be found by presenting the training set of examples and minimizing the percentage of incorrect answers. The training procedure described above can be formalized as Algorithm 1.

In the following sections we use this algorithm to train a computational model of the distributed gene expression classifier and analyze its performance using sets of simulated data. We generate these sets using the model of the synthetic gene circuit (6) presented above. In order to simulate the combined effects of biological and instrumental noise on

Algorithm 1 Algorithm for training the gene expression classifier.

Input: The data set x_i with N examples. The class type y_i for each example. The total number of cells N_c . The number of iterations of the algorithm, N_{iter} .

Output: The trained set consisting of N_c selected cells; classification threshold θ .

for iteration $k \leftarrow 1$ to N_{iter} **do**

 Choose a random example from x_i in the range $i = 1, \dots, N$ {Apply the chosen chemical input to the cells.}

for $j \leftarrow 1$ to N_c **do**

 Eliminate the cell with probability $1 - p_{\pm}(z_j^*(x_k))$ (for $y_i = \pm 1$) with random cell replacement from the previous iteration or from the master population for $k = 1$.

end for

end for

for training example $i \leftarrow 1$ to N **do**

 Use selected cells to measure the mean population fluorescence:

$$f(x_i) \leftarrow N_c^{-1} \sum_{j=1}^{N_c} z_j(x_i)$$

end for

Find threshold θ for which $\sum_{k=1}^N y_k [2H(f(x_k) - \theta) - 1]$ is maximal

the performance of the classifier we will assume that the resultant mean fluorescence of the population contains both additive and multiplicative noise:

$$f(x) = \frac{1}{N_c} \sum_{i=1}^{N_c} z_i(x)(1 + \varepsilon) + \zeta. \quad (7)$$

where ε and ζ are independent normal random variables with the respective means of 0 and $\sigma/4$ and standard deviations σ and $\sigma/4$. As usual in performance evaluation, each data set is divided into two parts, one for training and another for testing, and the overall performance of the classifier is measured as the percentage of correct answers on the test sets. The classifier performance is calculated using Algorithm 2.

Classification problems

Before we present our numerical results demonstrating the performance of the distributed classifier, let us discuss the motivation for using non-separable data sets as representative

Algorithm 2 Algorithm for testing the gene expression classifier.

Input: The testing data set x_i with N examples. The class type y_i for each example. The total number of cells N_c .

Output: The performance of the classifier.

Use all the cells selected during training.

correct $\leftarrow$ 0

for $i \leftarrow 1$ to N **do**

 Measure average population fluorescence $f(x_i)$

if $f(x_i) > \theta$ **and** $y_i = 1$ **then**

 correct=correct+1

end if

if $f(x_i) < \theta$ **and** $y_i = -1$ **then**

 correct=correct+1

end if

end for

performance=correct/ N

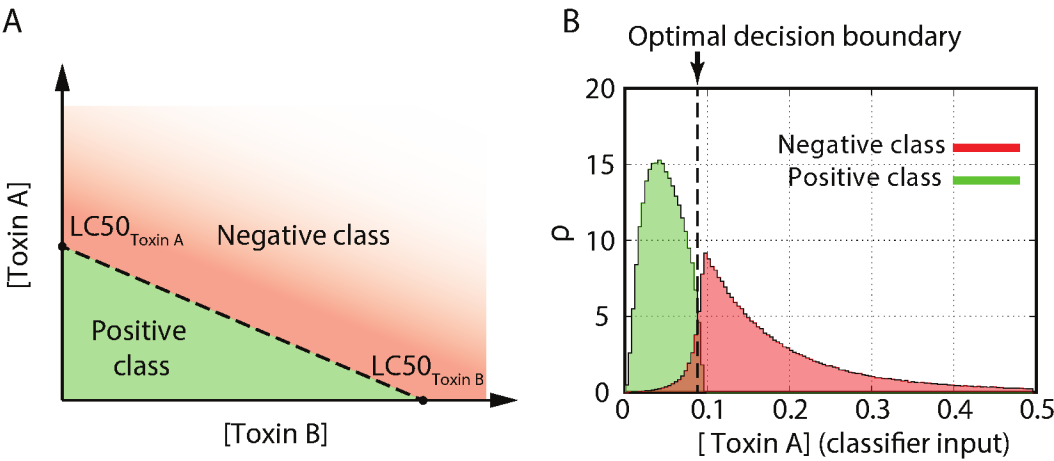


Figure 4: (A) Definition of positive and negative classes for two additively interacting toxins. (B) Positive and negative classes separable in two dimensions become non-separable in one dimension, i.e. when only the concentration of one of the toxins can be measured directly and the other becomes a hidden random variable. For illustrative purposes we assumed log-normal distributions of toxin concentrations. A histogram of 10^6 examples is shown. Optimal decision boundary for a naive Bayes classifier is shown with a dashed line.

examples of real world classification problems. If the classes were completely separated, then a properly trained classifier would be able to identify the boundary (or a manifold in a multi-dimensional input space) between the classes and perform classification with 100% accuracy. However in reality different classes often overlap, and therefore 100% accuracy of classification cannot be attained. Real-world biological and medical data frequently consists of overlapping (non-separable) classes.^{42,43} To illustrate how non-separable classes can emerge even in a simple biochemical system we consider an example of two toxins that additively contribute to the overall toxic effect on the cells. In such case, in a two-dimensional plane of two toxin concentrations there exist a straight line where the overall toxicity is equal to a certain threshold^{44,45}, thus the positive and negative classes are separable (Figure 4(A)). For specificity, we define a positive class as a domain of concentrations of Toxin A and Toxin B which cause less than 50% population mortality after a specified test duration; all other combinations of concentrations by definition are assigned to the negative class. Now, let us assume that the concentrations of both toxins are distributed according to some random distribution (e.g. log-normal), but our sensor can only measure concentration of only one of the toxins (e.g. Toxin A). In this case the two-dimensional positive and negative classes are projected onto Toxin A axis and become non-separable (Figure 4(B)). The concentration of the unmeasured Toxin B becomes a hidden variable that makes positive and negative classes non-separable. More generally, all the unknown or unmeasured environmental conditions can render two sets of measured conditions non-separable. In such cases the classification task consists of learning the optimal discrimination boundary along Toxin A axis that maximizes the discriminatory power of the classifier at the level below 100% accuracy. In the absence of *a priori* information about the known and unknown toxin distributions and their effect on cell viability, the optimal decision boundary will have to be inferred solely from random

examples presented to the classifier, thus the performance of a real classifier will be even lower.

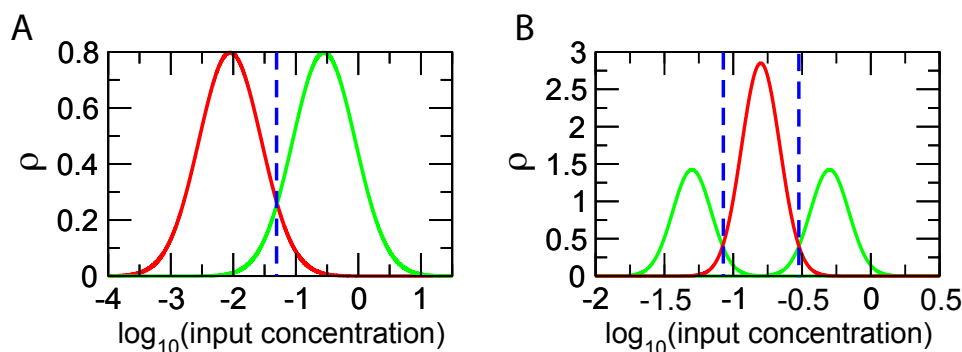


Figure 5: The data is generated from two (left) or three (right) log-normal distributions generating the positive class (green) and the negative class (red) examples. The optimal thresholds for discriminating between the classes are represented by the dashed vertical lines. The maximum performance achievable by a classifier trained on infinite amount of examples from the two distributions is 93.3%, from the three distributions is 94.8%. The minimum in both cases is 50% which is equivalent to a random answer selection.

In order to clearly demonstrate the salient features of the classifier and study the limits of its performance we will continue with two “idealized” classification problems. First, we consider the case of discriminating data generated by two log-normal distributions of input chemical concentration with some overlap between the two classes (Figure 5A). This example is qualitatively similar to the two-toxin example discussed above, except that there is a nonzero probability of having points far from the decision boundaries (it is a harder problem). After that, we test the distributed classifier on a more challenging problem of discriminating the data generated by complex distributions (one unimodal and another bimodal), when the negative class is surrounded by the positive class on both sides (Figure 5B).

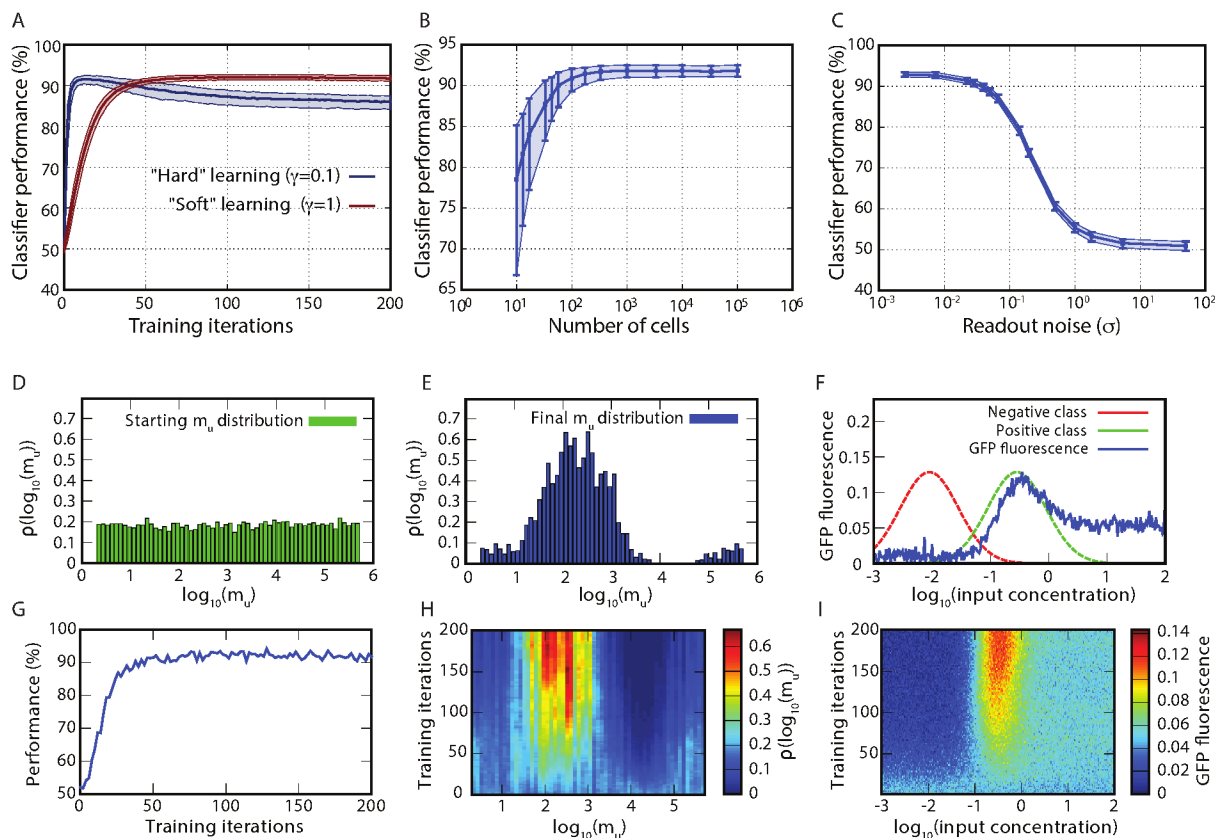


Figure 6: Classification results for the data set drawn from two unimodal classes, Figure 5A. (A) Evolution of the classifier performance for $\gamma = 0.1$ (“hard” learning; blue) and $\gamma = 1$ (“soft” learning; red), population size $N_c = 10^4$ cells. The classifier performance versus cell population size N_c (B) or GFP fluorescence readout noise σ (C); $\gamma = 1$ in (B) and (C), $N_c = 10^4$ in (C). The median and interquartile range of the distribution of the classifier performance calculated from 10^3 different stochastic realizations are shown in (A)–(C), readout noise $\sigma = 1/35$ in (A) and (B). (D)–(I) Evolution of the parameters of the classifier before and after training – an example trajectory. The parameters used are $\gamma = 1$, $N_c = 10^4$, $\sigma = 1/35$. It illustrates the shift in the distribution of parameters due to the training process of elimination of cells. The distribution of RBS/promoter strengths m_u before training (D) and after 200 training iterations (E). (F) Normalized GFP fluorescence of the ensemble of cells $f(x)$ (blue) after 200 training iterations, log-normal distributions generating positive (green) and negative (red) class examples. (G) Evolution of the classifier performance in this realization. Evolution of m_u distribution (H) and normalized cumulative GFP fluorescence $f(x)$ (I).

Discrimination of two unimodal classes

As the first example we used the data drawn from two log-normal distributions centered at $\log(x)=-0.55$ (class +1) and $\log(x)=-2.05$ (class -1) with standard deviation of 0.5 (Figure 5A). The data has been generated on a log-scale since it is the natural scale of the response of the genetic circuit used in the classifier (Figure 2). Figure 5A gives an illustration of the two generating distributions. The optimal theoretical performance is determined by choosing the threshold that separates the positive-class distribution from the negative one in a manner that minimizes discrimination errors. In this particular case, the optimal threshold value that leads to the minimal number of errors and therefore to the maximum performance is located exactly in the middle between the peaks of the two distributions which is indicated by the blue dashed vertical line ($\log(x) = -1.3$). Due to the nonseparability of the two classes it is impossible to solve this classification problem with 100% success rate. The optimal theoretical performance for this problem is 93.3%. An ideal classifier could approach this value for infinite amount of training and evaluation data.

Using a “hard” learning strategy (small $\gamma = 0.1$), the population-based classifier achieves high performance of 91.6% in just 12 iterations, however with further training the performance deteriorates achieving 86.1% after 200 iterations (Figure 6A). These results are obtained using $N_c = 10^4$ cells (all other circuit parameters are as described in Figure 2). Such deterioration of performance is a well known phenomenon in machine learning, where early stopping is often applied in similar situations.^{46,47}

In contrast, as discussed above, a “soft” learning strategy ($\gamma = 1.0$) allows one to achieve the maximum performance (92.0%) with high robustness over a wide range of training iterations (Figure 6A). Relatively small number of cells is sufficient to achieve such performance (Figure 6B). The classifier is also robust to readout noise (Figure 6C).

The evolution of the parameters of the classifier during “soft” training is demonstrated in Figure 6D–I. The training process leads to a robust selection of the cells with those values of the genetic diversity parameter m_u that allow the normalized population-wide GFP response $f(x)$ to roughly track the positive class data distribution (Figure 6E–F, H–I). Correspondingly it is possible to reliably select such a threshold θ (Algorithm 2) that allows the classifier to achieve performance close to the theoretical maximum for this problem after roughly 50 training iterations (Figure 6G).

We also analyzed the performance of the distributed classifier on the more realistic unimodal example of dual-toxin cell viability classification with one toxin measured and another unobserved which was introduced in the beginning of this Section (see Supporting Information section 2). The classifier achieves high performance of about 95% after learning just 15 examples using “hard” ($\gamma = 0.1$) training strategy (Figure S3 of the Supporting Information).

Discrimination of a bimodal and a unimodal class

This is a more challenging one-dimensional classification problem where a negative class is “sandwiched” by the positive class on both sides (Figure 5B). The negative class data is generated by a log-normal distribution centered at $\log(x)=-0.8$. The positive class data is generated from two equivalent log-normal distributions centered at $\log(x)=-1.3$ and -0.3 . The standard deviation of all three distributions is 0.14. The distributions are normalized such that on average the equal number of examples is drawn from the negative and positive classes. The maximum theoretical performance on this problem is 94.8%, determined as in the example above. We use the same genetic circuit parameters as in the previous example (see Figure 2).

This classification problem exemplifies the differences between the “hard” and “soft”

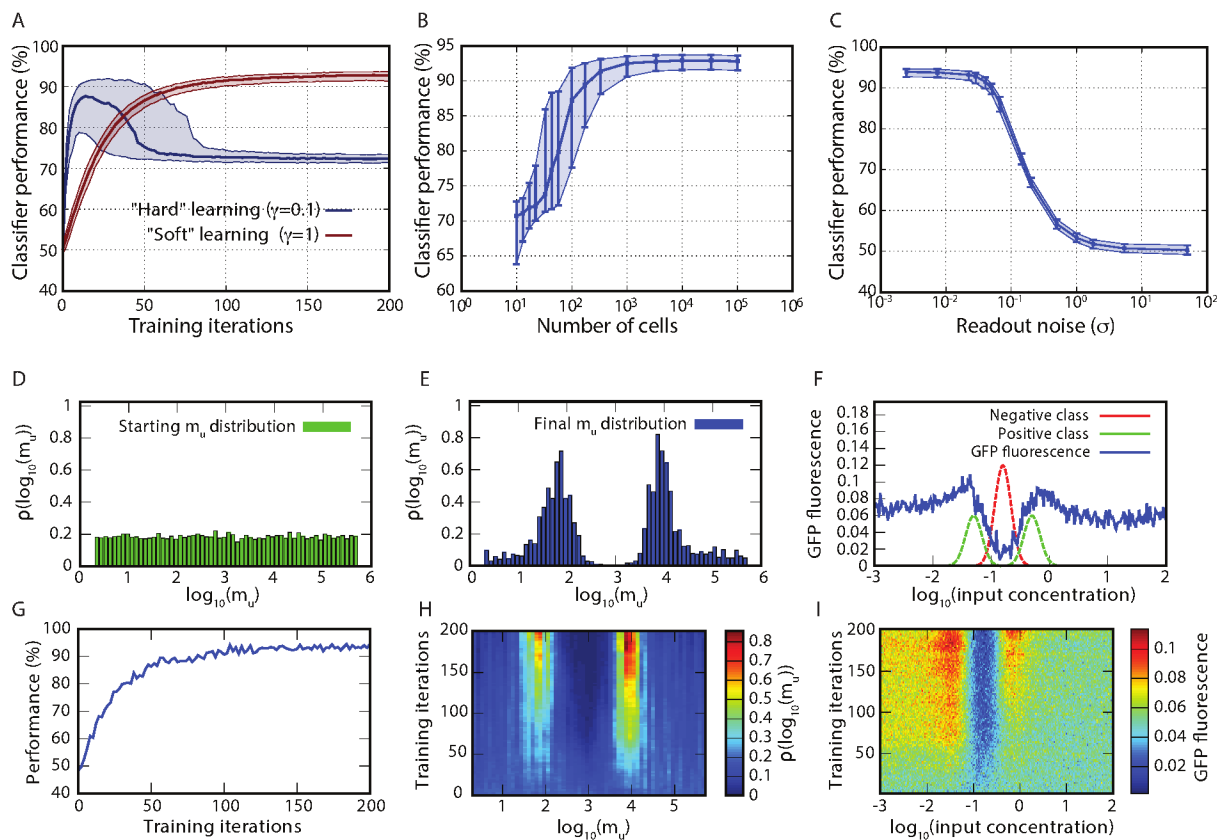


Figure 7: Classification results for the data set drawn from one bimodal and one unimodal classes, Figure 5B. (A) Evolution of the classifier performance for $\gamma = 0.1$ (“hard” learning; blue) and $\gamma = 1$ (“soft” learning; red), population size $N_c = 10^4$ cells. The classifier performance versus cell population size N_c (B) or GFP fluorescence readout noise σ (C); $\gamma = 1$ in (B) and (C), $N_c = 10^4$ in (C). The median and interquartile range of the distribution of the classifier performance calculated from 10^3 different stochastic realizations are shown in (A)–(C), readout noise $\sigma = 1/35$ in (A) and (B). (D)–(I) Evolution of the parameters of the ensemble of cells before and after training – an example trajectory. The parameters used are $\gamma = 1$, $N_c = 10^4$, $\sigma = 1/35$. It illustrates the shift in the distribution of parameters due to the training process of elimination of cells. The distribution of RBS/promoter strengths m_u before training (D) and after 200 training iterations (E). (F) Normalized GFP fluorescence of the ensemble of cells $f(x)$ (blue) after 200 training iterations, log-normal distribution generating positive (green) and negative (red) class examples. (G) Evolution of the classifier performance in this realization. Evolution of m_u distribution (H) and normalized cumulative GFP fluorescence $f(x)$ (I).

learning strategies (Figure 7A). Similarly to the previous example the “hard” learning strategy ($\gamma = 0.1$) initially leads to rapid improvement of classifier performance, reaching the maximum performance of 87.7% in 14 iterations. However unlike the previous example this increase is not robust and is characterized by high stochastic variability (compare Figure 6A and Figure 7A). With further training the performance quickly degrades due to stochastic extinction of the cells fitting one of the two positive class distribution peaks (see the details below). These problems can be ameliorated by employing “soft” training strategy ($\gamma = 1$) (Figure 7A). In this case the maximum performance is higher (92.9%) and is achieved much more reliably albeit in noticeably higher number of training iterations. Similarly to the previous example, relatively low number of cells is sufficient to achieve this performance (Figure 7B). The performance is robust with respect to the readout noise (Figure 7C). An example of the evolution of the parameters of the classifier during “soft” training is shown on Figure 7D–I.

Analytical description of soft-learning performance

We have developed an analytical theory that describes the training and the resulting performance of the cell-based classifier in the limit of “soft” learning (see Supporting Information, section 1). This analytical description is exact in the case of infinitely “soft” learning and infinite number of cells. It is based on the assumption that at each training iteration the distribution of cells is changed by infinitely small amount, correspondingly requiring an infinite number of training iterations to achieve finite changes in cell distribution. Thus the discrete iteration step can be replaced by continuous “time” t , and the evolution of the number of cells n_i with a given value m_i of genetic diversity parameter

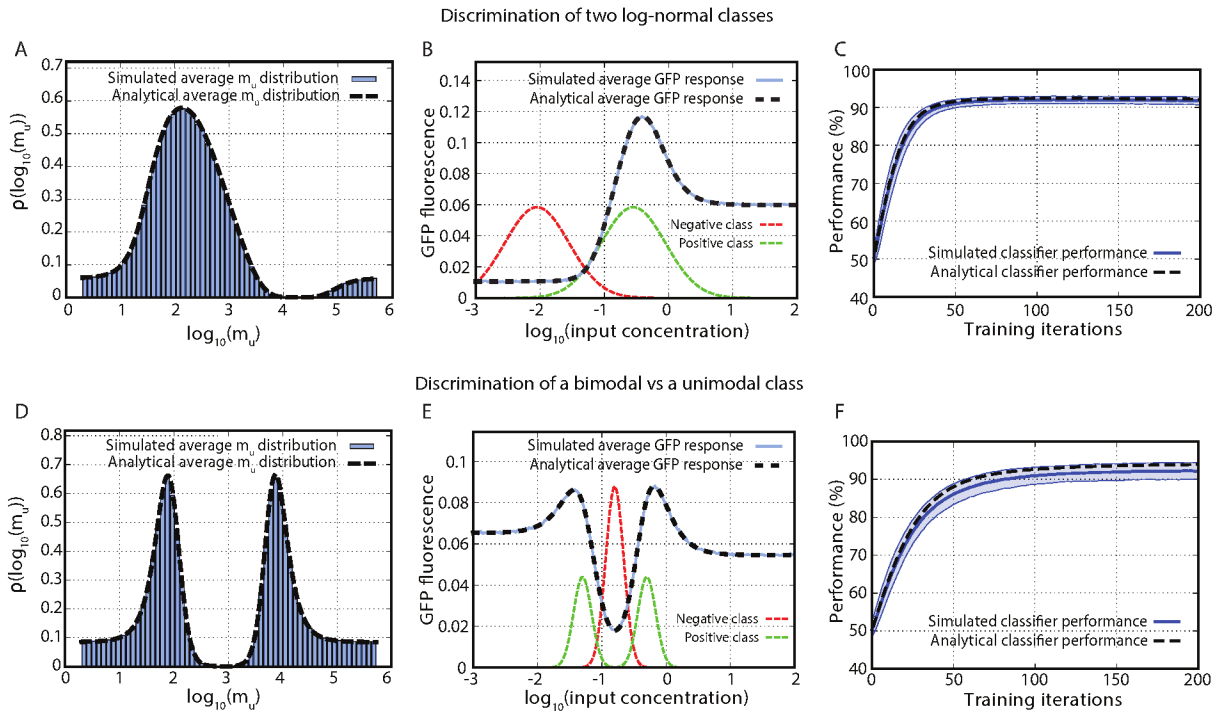


Figure 8: Comparison of the analytical theory with the simulation results for the case of soft-learning ($\gamma = 1$). (A)–(C) Discrimination of two log-normal classes. (D)–(F) Discrimination of a bimodal vs a unimodal class. (A), (D) Ensemble average distribution of the genetic diversity parameter m_u calculated after 200 training iterations from 10^3 independent simulations (blue bars) vs the analytical prediction (black dashed line). (B), (E) Ensemble average population-wide GFP responses calculated after 200 training iterations from 10^3 independent simulations (blue line) vs the analytical prediction (black dashed line). (C), (F) Ensemble average classifier performance calculated from 10^3 independent simulations vs the analytical prediction; the simulation results are shown as mean $\pm$ standard deviation. All other simulation parameters are the same as in Figure 6D–I, 7D–I

m_u , in the course of training can be described by a differential equation

$$\frac{dn_i}{dt} = \lambda_i n_i - \frac{n_i}{N_c} \sum_k \lambda_k n_k$$

where N_c is the total number of cells (fixed) and λ_i is a “shaping factor” which depends on the distributions of the positive and negative training examples $w_{\pm}(x)$, m_i -dependent sin-

gle cell GFP response functions $z(x; m_i)$ and the corresponding cell survival probabilities $p_{\pm}(z)$ (see Supporting Information, section 1). The solution of this equation approximates an ensemble average solution in the case of finite “softness”, number of training steps (N), and finite number of cells by formally setting $t = N/2$. Figure 8 demonstrates the good agreement between the analytical solutions for the ensemble- averaged m_u distribution, population average GFP response, and the ensemble average classifier performance and the simulation results for “soft” learning with $\gamma = 1$. Note that the performance figures calculated analytically are consistently higher than the ensemble average performance figures calculated from the simulations (Figure 8C, F). This is largely due to the fact that the analytical approximation is based on asymptotic analytical solution which ignores the stochastic effects due to the finite number of iterations.

This analytical theory allows to quickly estimate the parameters for optimal training of the distributed cell population based classifier for a given classification problem. In comparison the stochastic simulations necessary to estimate the same training parameters are significantly more computationally expensive. If necessary the results of the analytical approximation can be confirmed by stochastic simulations.

Another important result that follows from the analytical approximation is that early stopping must be always used in order to maximize the classifier performance, even in the case of infinitely “soft” learning, since after large number of training iterations ($N \rightarrow \infty$) only the cells with the maximum λ_i survive. In most practical cases it means that the cells with only one particular value of m_i survive thus generally leading to poor classifier performance.

Summary and Discussion

In this paper we introduced a conceptual design of a distributed classifier in the form of a population of microorganisms containing a strategically constructed library of sensory gene circuits. We described an algorithm of training such a classifier by pruning the master population after iterative presentation of known positive and negative input-output examples. We characterized both numerically and analytically the performance of the proposed classifier based on a particular sensory gene circuit that is induced by the external chemical input within a certain range of concentrations and repressed outside of it. A library containing a broad distribution of such circuits with different sensitivities to the chemical input in individual cells within the master population can be constructed by randomizing the control sequences within the input genetic element of the sensing circuit. We demonstrated that after appropriate training the distributed classifier can achieve nearly-optimal performance in solving the task of discriminating nonseparable input data sets.

In this theoretical study we have not addressed in detail several important issues that are likely to arise in the experimental implementation of the distributed classifier. One such issue is the specific procedure to retain the “good” cells and discard the “bad” ones. The most straightforward way to do this is to use a fluorescence-activated cell sorting (FACS). However, typical commercial FACS software allows one only to select cells deterministically with a measured fluorescent value (or a combination of multiple values) within a certain range - the process known as “gating” - rather than to sort cells with a certain probability based on their fluorescence levels. We envisage two ways to circumvent this problem. First, any smooth probability function can be approximated to a required precision by a step function by dividing the entire range of cell fluorescence on a sufficiently high number of bins. Cells with the fluorescence values falling within each bin can

be independently sorted using standard FACS systems and later combined in proportions according to the corresponding probabilities of the smooth probability function. Alternatively, a probabilistic selection algorithm could implement directly by straightforward adaption of the FACS control software if the hardware programming interface is available. A potentially more interesting approach allowing for autonomous and adaptable training could be to engineer an additional synthetic gene circuit controlling cell growth based on the output signal, using for example one of well-characterized positive/negative selection systems.⁴¹

In this study we assumed that the promoter strength parameters are initially distributed uniformly over a broad range, however experimentally the distribution would likely be non-uniform. This non-uniformity may affect the ability to train the classifier to an arbitrary classification task. In order to estimate how sensitive the performance of the proposed classification system may be to non-uniformity of the initial distribution of parameters we have calculated the performance of our distributed classifier starting with model distributions of varying degrees of non-uniformity (Supporting Information, section 4). Based on these calculations we estimate that depending on the classification problem a moderate degree of non-uniformity can be tolerated. However, the more uniform the initial distribution is, the better the final performance of the trained classifier will be, so the design of a uniform library is one of the important considerations for the future experimental implementation of the classifier. Such a library may have to be designed and synthesized, preselected from a random library, or both (see the discussion in Supporting Information, section 4). Finally, we did not take into consideration the growth and division of cells that can affect the distribution of cells within the trained population if cells with different sensory circuit parameters differ in their doubling rates, for example due to respective differences in metabolic burden exerted by these circuits.

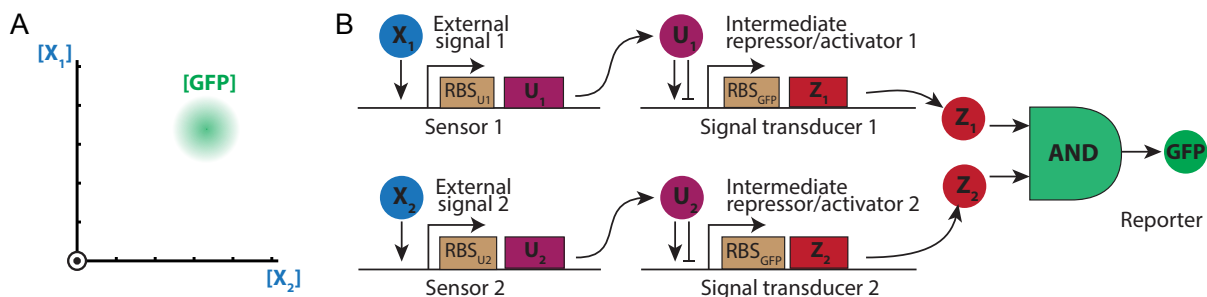


Figure 9: (A) The desired bell-shaped response of the multidimensional classification circuit. (B) The genetic circuit proposed for use in a distributed genetic classifier with multiple inputs per cell. Independent sensing and response functionalities are combined using an appropriate biological AND gate. The resulting response function of the entire two-promoter circuit is bell-shaped with respect to the inputs X_1 and X_2 for the relevant choice of parameters.

The particular design of a distributed classifier presented here is suitable for classification of a scalar input (single chemical inducer X) only. Many real-world classification problems involve multidimensional inputs. The one-dimensional classifier described in this paper can be generalized to solve multidimensional problems. In the simplest case of multidimensional classification where each dimension can be classified independently, the problem can be trivially solved by a cell population classifier consisting of a mixture of cells capable of responding individually to just one input using the circuit described in Figure 1. However, such approach would fail for more complex classification problems when a multidimensional distribution of positive or negative outcomes is not the direct product of corresponding one-dimensional distributions. These more complex problems can be solved using a classifier built with the cells endowed with circuits that are sensitive to multiple inputs. An example of such circuit design for a two-dimensional input is shown in Figure 9. In this design the input signals are sensed separately by the corresponding two-stage modules similar to described above. The outputs of these two modules are then combined by a genetic AND gate. A number of circuits performing logical operations

including AND have been developed and characterized recently.^{48–50} Similarly to the one-dimensional circuit the output of this circuit is a two-dimensional bell-shaped function of the two inputs (Figure 9A). The parameters of both sensory circuits can be randomized as before. These multidimensional classifiers can be trained and the performance can be characterized in the same way as in the case of the one-dimensional classifier.

From synthetic biology perspective, our modeling study opens an intriguing possibility to engineer genetically diversified microbial populations to solve classification tasks which are difficult or impossible to solve within a single microbial cell. Such approach is known in machine learning theory where it was proven that consensus classifiers made of appropriately combined weak classifiers can achieve high performance.^{51,52} We anticipate that beyond conceptual academic interest, autonomous cell population-based classifiers can have biotechnological and medical applications. The proposed circuit and its training algorithm could potentially be used to facilitate biosensor design. An autonomous classifier could directly produce a biologically relevant response, for example a drug or a signaling molecule can be synthesized *in situ* when the measured input concentration falls within or outside an optimal range. Such autonomous systems capable of responding to a complex input can be used e.g. to control bioreactors or as prosthetic control systems for use in living organisms. On a more general level, our findings suggest that perhaps the genetic and phenotypic diversity found in many natural populations,^{53,54} along with other potential survival benefits⁵⁵ can play an important role in forming a decentralized “social intelligence”^{15,56} capable of solving complex computational tasks in a noisy and unpredictable environment.

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Graphical TOC Entry

