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Design and Evolution of Engineered Biological Systems

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(http://openwetware.org/wiki/Jason_Kelly)

Abstract

To date, engineered biological systems have been constructed via a variety of ad hoc approaches. The resulting systems should be thought of as pieces of at. We are interested in exploring how existing forward engineering approaches might be combined with directed evolution to make routine the construction of engineered biological systems. We have specified a procedure for construction of biological systems via screening of subcomponent libraries and rational re-assembly. We have begun development of tools to enable this approach, including a FACS-based screening system to rapidly measure the input/output function of a genetic circuit. Additionally, we have begun designed a microfluid cystem that enables more sophisticated screening and selection functions. Specifically, a microfluidic chemostat integrated with a cell sorter (i.e., a sortostat). This microscope-based system will enable us to evaluate whether or not more complicated screens and selectical use in service of evolving engineered biological systems.

Motivation

In our framework, engineered biological systems are made from devices. Devices, in turn, are made from parts. Parts are units of basic biological function (e.g., a ribosome binding site). Devices are combinations of parts that perform some simple operation (e.g., a Boolean NOT function). We have to solve several problems to reliably combine devices into functional systems.

1. Common Signal Carrier

A carrier signal which is independent of the specifics of the device is essential to composing devices into systems.



2. Signal Level Matching

The levels of the common signal must match between devices. For example, in analogy to digital systems, both devices must have the same definition of a 1 or 0 if they are to communicate.



3. Resistance to internal noise

Our devices must work inside living cells. The intracellular environment is noisy, small, dense, and uncertain. Devices should be designed to function robustly in the presence of expected fluctuations in their local environment. (Or local environment should be made more receptive to hosting devices)

4. Stability of the information encoding the system

Engineered genetic systems are subject to mutation and natural election. As a result, we must develop methods to control the stability of the genetic information encoding our devices.

Library-Based Construction

A strategy of constructing engineered biological systems via directed evolution from standard parts libraries offers several advantages over a purely rational approach.

- 1. Some characterization of devices becomes implicit in their construction due to screening for I/O function of device.
- Signal matching between devices is more easily accomplished[1]
 Library diversity yields functional subcomponents with diverse characteristics not specifically selected for (latency, etc.) improving
- likelihood of achieving a functional final system. 4. Complexity of biological substrate makes effective modeling of system
- performance in response to rational changes very challenging. 1. Rationally design system from standard devices

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2. Identify devices with screenable I/O functions



For instance, oscillatory behavior would not be easily screenable – however, some of the subcomponents of an oscillating system might be screenable. In this simple system, subcomponents as well as the final system can be screened for function.

3. Generation of sub-libraries from a base device





Since no screening was done for characteristics such as the switching speed in response to inducer, we expect that bistable switches with a variety of speed swill be isolated from the final system library.



The plasmic will be based on pSiExX3 [Registry of standard biological Parts], pSiExX3 is derived from the variable copy plasmid system, pScANS[2]. The plasmid contains the F replication origin (copy number <10) and also the P1 lytic origin (copy number >100). Replication at the lytic origin (copy number should be JPTG. This will facilitate screening of devices at low copy number (expected operating conditions for our systems) while allowing for induction to high copy number to increase DNA preparation yield for subsequent construction steps.



The latest version of the screening plasmid contains RNase E sites to create independence between the mRNA stability of the device being screened and the mRNA stability of the fluorescent proteins. In particular, we suspect mRFP1 to contain internal RNaseE cut sites and have added a hairpin 5' of the coding region to slow degradation by RNase E. [4] An earlier version of the screening plasmid was tested with a tetR-based inverter. The curves look qualitatively correct, however expression levels of the fluorescent proteins were rather low.

Sortostat Design & Motivation



Total Reactor Volume = 16nL
 Sorting chamber = 1/50th of total reactor volume
 Modification and extension of design by Balagadde et al. [5]

A microfluidic chemostal integrated with a cell sorter, which we call a "sort-ostat", will enable more complicated selections to be applied to a population of cells in continuous culture. In particular, time varying selective pressures as well as very specific selection strengths can be applied. We will evaluate whether or not these more sophisticated selective pressures will be of practical use in service of evolving engineered biological systems. Selection can be based on any characteristic that can be reliably measured via microscopy. Lastly, since this is a physical selection (rather than a chemical one), it may be more difficult for cells to find unexpected methods to evade selection.

Sortostat Preliminary Results



Sortostat was run with no selective pressure after being inoculated with cells growing in log phase from a batch culture. Analysis of steady state region (>6hrs) suggests that the %CFP cells found in the sorting chamber is binomially distributed (0.01 significance level)

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Selective Pressure Turned ON



Sortostat was run with selective pressure, sorting against cells expressing YFP. Based on the rate of sorting events (1/3 min⁻¹) and initial cell counts, the mathematical model prediction effect of sorting on the population relatively well. Graph depicts the performance limits of the device based on a mathematical model at maximum screening rate for populations 50 10de cells / reactor. Smaller populations 51 vider distribution and thus will face a great selective pressure

Future Work

Further characterization and specification of device performance

- Tuning of oscillation frequency by selective pressure
 Selection for reduction in noise in gene expression across population
- Selection for a specific expression level of a fluorescent protein.
 Other ideas?

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