Informatics Tools for Designing Oligos and Assemblies

There are many steps involved in the DNA synthesis and assembly. We have implemented a suite of informatics tools to assist the design of oligos, the assembly of larger constructs, and novel enzymology solutions to significantly improve the scale and reduce the cost.

DNA Synthesis Workflows at JGI

DNA synthesis allows digital sequence information to be translated into biology by generating DNA in a template-independent manner. Sequencing databases are used to design DNA parts using design software developed at JBEI. The parts are then assembled to larger size constructs in a similar process.

Adding Value to Sequence Information

Sequence information specially from metagenomes and single cell genomes is essentially digital information that cannot be easily accessed for biological hypothesis testing. This represents a limiting factor in the development of sequence based applications. DNA synthesis allows digital sequence information to be translated into biology by generating DNA in a template-independent manner.

Informatics Tools for Designing Oligos and Assemblies

These informatics tools are publicly available. Listed here are some examples of what these tools can assist in the design of DNA parts and assembly. Users have the options to use any of these tools to design their own constructs. J5 is an automated DNA assembly software developed at JBEI (j5.jbei.org). It provides a web-based graphical user interface for external users of the JGI's DNA Synthesis Program.

Laboratory Processes

Typically 48-96 basic parts can be synthesized in a week by a single FTE. Several robotic workstations are applied to handle the labor intensive steps shown in red. The basic parts are then assembled to larger size constructs in a similar process.

Synthesis Design and Status Reporting Platform

We plan to integrate the DIVA (Design, Implementation, Validation Automation) platform developed at JBEI with the DNA synthesis pipeline. This will provide a web-based graphical user interface for external users of the JGI's DNA Synthesis Program.

Systematic Survey of GH1 Enzyme Diversity

Our initial demonstration project was to functionally characterize the natural diversity of GH1 enzymes. GH1s participate in the last step of biomass breakdown (cellobiose to glucose). We identified GH1 candidate genes from NR, CAZy and JGI metagenomics databases, performed multiple sequence alignments, constructed phylogenetic trees and selected 200 representatives that cover the maximum sequence space (selected GH1 genes are represented by red dots on tree outer circle).

We have synthesized and over-expressed our 200 GH1 candidate genes. Initial biochemical characterization at different temperatures and pHs has revealed a wide range of functional properties, validating our phylogenomics approach.

Synthetic Biology Vision at JGI

Synthetic biology approaches combined with large scale genomics information, will result in novel sequence driven applications. This will require substantial output and cost improvements as well as increasingly complex regulation and circuitry tools.

Examples of Proposed Genes

- Glycosyl Hydrolases
- Polyketide Synthases
- Plant Transcription Factors
- Lignin Peroxidases
- Cyanobacterial Carboxysomes
- Mevalonate Pathway
- Nitrogenases

The FY13 synthesis capacity will be used primarily to support Bioenergy Research Centers (BRCs) and our Community Sequencing Programs (CSPs). Additional capacity will be used to develop proof-of-principle projects.

The work conducted by the U.S. Department of Energy Joint Genome Institute is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231.