The Constructor: a web application optimizing cloning strategies based on modules from the registry of standard biological parts

Matthijn C Hesselman1†, Jasper J Koehorst1†, Thijs Slijkhuis1, Dorett I Odoni1, Floor Hugenholtz1,2 and Mark W J van Passel1,3*

Abstract

Synthetic biology is an emerging field that combines molecular biology with engineering principles, which requires abstraction levels applied to a modular biological componentry. The Registry of Standard Biological Parts harbours such a repository of standardized parts, and thereby facilitates the combination of complex molecular modules to novel genetic circuits and devices. However, since finding the best parts for a pre-determined genetic design can be time consuming, we devised the Constructor, a web tool that recommends the smallest number of cloning steps for pre-designed circuits, and implements user-defined quality checks.

We present the Constructor (www.systemsbiology.nl/the_constructor) as a constructive web tool that simplifies the in silico assembly of pre-designed gene circuitries from standard parts, reducing both planning and subsequent cloning time.

Keywords: Synthetic Biology, BioBrick parts, Transcription units, Genetic circuit

Biological engineering

In recent years striking biological circuits have been fabricated, often resembling devices from electrical engineering. Examples include a genetic toggle switch [1], oscillators [2], a rewritable memory unit [3], but also more complex features such as a DNA-guided assembly line [4]. These sorts of devices provide a starting ground for further engineering. However, next to the development of stable and streamlined microbial chassis [5-7], biological engineering requires accessible, modular, standard parts with reliable characteristics. Even though several repositories have been constructed [8], the most well-known library is the Registry of Standard Biological Parts (http://partsregistry.org). It is widely used in the international Genetically Engineered Machine (iGEM) competition [9] aimed at teaching synthetic biology to undergraduate students [10,11], but also outside of this competition [12]. Users can design new devices from so-called BioBrick parts, the standard genetic parts, and are encouraged to submit new parts and devices to this Registry, which has grown to approximately 20,000 entries in 2012.

However, the vast number of entries in this Registry as well as the numerous parts with an unconfirmed status can frustrate the straightforward design of new genetic circuits. Furthermore, often several variants of parts and devices are available in the databases, which makes it difficult to find the shortest and most reliable strategy to clone new gene circuits. We found a need to automate querying the Registry in order to find the most straightforward cloning strategy for any pre-designed genetic circuit. This would simultaneously reduce time spent on finding appropriate parts in the Registry. We therefore set out to code the Constructor, a web-based application that recommends the smallest number of biological parts for a user-defined gene circuit, reducing both dry-lab and wet-lab time.

* Correspondence: Mark.vanPassel@wur.nl
† Equal contributors
1 iGEM Wageningen UR, Wageningen University, Wageningen, The Netherlands
2 Laboratory of Systems and Synthetic Biology, Wageningen University, Wageningen, The Netherlands
Full list of author information is available at the end of the article

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The Constructor

The web interface of the Constructor accepts user-defined genetic circuits built from individual transcription units (TUs). A TU consists of a promoter, a ribosomal binding site (RBS), a coding sequence (CDS) and a terminator (see also Figure 1). Arrangements with multiple RBSs and CDSs between a promoter and terminator, i.e., an operon, are allowed. Next, users can give a complex genetic circuit consisting of a number of TUs as input. To our knowledge, there is no evidence that the physical location of TUs on a plasmid has any relevance for the function of the genetic circuit. Therefore the Constructor permutes all possible arrangements of the different TUs making up a circuit. For example, if the genetic circuit consists of three separate TUs (in the order A-B-C, where each letter signifies a TU), the Constructor uses all six possible arrangements in its query (A-B-C, B-A-C, B-C-A, ...).

For each circuit arrangement, an SQL database containing information of ~20,000 parts (assembly standard 10 compatible) is queried. Since each circuit arrangement has a different sequence of genetic elements, each arrangement could collect a few arrangement-specific BioBrick parts. Through the combination of these specific BioBrick parts and the unique arrangement of genetic elements, some circuit arrangements could be constructed with fewer parts than others. An extra filter can be set if required on the availability, validity and quality of selected BioBrick parts by The Constructor.

Of all these possible combinations, the top three arrangements that could be assembled with the smallest number of BioBrick parts are shown as a recommendation to the user (Figure 1).

Validation

We tested several proposed cloning strategies from five iGEM 2011 team projects, and compared these to the cloning strategies recommended by the Constructor. We find that the number of cloning steps predicted by the Constructor with appropriate parts criteria is generally smaller than those that were undertaken by the teams (Table 1), indicating the usefulness of the Constructor in

| iGEM 2011 team | iGEM project title | # of TUs in the designed circuit | # of used Biobrick Parts | Processing time (min) |
|----------------|--------------------|---------------------------------|--------------------------|-----------------------|
| KAIST-Korea1   | E. casso           | 2                               | 11                       | 7                     | < 1 min              |
| UANL_Mty-Mexico2 | S.C.I.E.N.C.E.: Simple Code Interpretation Enabling Circuit in E. coli | 7 | 26 | 25 | 674 min |
| XMU-China3     | i-ccdB: intelligent Control of Cell Density in Bacteria | 3 | 6 | 6 | < 1 min |
| NTNU-Trondheim4 | Red Fluorescent Stress Sensor | 2 | 5 | 4 | < 1 min |
| Wageningen UR5 | The Synchronocillator: a Synchronized Oscillatory System | 4 | 7 | 6 | < 2 min |

1 http://2011.igem.org/Team:KAIST-Korea.
2 http://2011.igem.org/Team:UANL_Mty-Mexico.
3 http://2011.igem.org/Team:XMU-China.
4 http://2011.igem.org/Team:NTNU_Trondheim.
5 http://2011.igem.org/Team:Wageningen UR.

In these examples, the gene circuits contained between 2 and 7 Transcription Units (TUs).
optimizing cloning strategies. Specifically, in four out of five cases, the Constructor recommends cloning strategies that involve fewer BioBrick parts, and therefore fewer cloning steps. Also, in most cases, the processing time of the Constructor is less than two minutes. The most complex genetic circuit, from the iGEM team UANL_Mty-Mexico, consisting of seven transcriptional units, took approximately ten hours to run. Still, this is expected to be less than manually querying the Registry for the optimal cloning strategy.

Potential extensions
The Constructor facilitates the cloning strategy of complex pre-designed genetic circuits from elements of the Registry of Standard Biological Parts. Although the Constructor specifically focuses on this Registry, the same search algorithms for assembly optimization recommendations can be applied to other extensive and well-defined parts libraries. Furthermore, the Constructor uses a straightforward transcriptional unit concept, which could be expanded by including different parts such as splicing signals. Finally, the web tool could be further optimised by suggesting alternatives for certain parts, like available reporter genes with another fluorescent ability, or different inducible promoters.

Availability
The Constructor is available at www.systemsbiology.nl/the_Constructor. Help functions and a tutorial are provided with test cases of user defined gene circuits. All scripts are available from the authors upon request.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
MCH conceived the initial design of the software, programmed the algorithms, and helped draft the manuscript. JK participated in the design of the study, further developed the algorithms, added functionalities and helped to draft the manuscript. TS participated in the design and coordination of the study, added functionalities, and helped to draft the manuscript. MWJvP participated in the design and coordination of the study, further developed the algorithms, and helped draft the manuscript. JJK participated in the design of the application. FH participated in the design and coordination of the study, and helped to draft the manuscript. MWJvP is Systems Biology (NCSB), which is part of the Netherlands Genomics Initiative reading the manuscript. FH is funded by the Netherlands Consortium for Wageningen UR iGEM student teams, specifically Brendan Ryback for critically suggesting alternatives for certain parts, like available reporter genes with another fluorescent ability, or different inducible promoters.

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