Towards predictable 5'-UTRs in Saccharomyces cerevisiae

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Introduction

Need for tools to fine tune pathways in yeast

The ability to precisely fine tune novel biosynthetic pathways in living cells is a fundamental requirement for the fast development of new microbial cell factories. Typically, protein levels are adjusted through engineering of the transcriptional and translational level. Besides efforts in yeast on the transcriptional level like the creation of synthetic promoter and terminator libraries, the development of synthetic transcription factors and the modification of transcription factor binding sites, less progress has been made in modifying the translation rate of genes for the predictive regulation of protein levels.

Varying gene translation rates

We developed a framework in S. cerevisiae for the de novo design of A'2 untranslated regions (AU2UTRs) with a predictive outcome on translation initiation rate. Therefore, we developed a partial least square (PLS) regression model based on a dataset of AU2UTR sequences of Dvir et al. (2013) [1]. Next, this model was employed for the creation of completely new AU2UTRs leading to protein abundances (PA) within a specified target range. Finally, in silico generated AU2UTRs with different combinations of promoters, 5'-UTRs and coding sequences were evaluated in vivo.

Methods

PLS regression model [2]

\[
Y_j = \beta_0 + \beta_1 X_{j,1} + \ldots + \beta_k X_{j,k} + \epsilon_j
\]

(k varies from 1 to 13)

Y: protein abundance
X: 5'-UTR feature

For the model construction, a dataset of 2041 RPL8A 5'-UTR sequences [1] was used. This dataset was split in a training set to build the model and a test set for model validation.

Results

in silico 5'-UTR design

The design of novel 5'-UTR sequences with a predictive outcome of protein expression levels is performed by an iterative process.

- Evaluation of the library scatter
- Degenerations, mutations & recombinations
- UTR feature calculation & prediction PA

As output, 2 libraries of eight 5'-UTRs covering the wanted protein abundance range is given.

Example of the scatter of an in silico generated library.

An R^2 of 0.72 was obtained for the test set, indicating that our PLS model has the potential to successfully predict protein abundance.

in vivo validation of new 5'-UTRs

In both cases, an R^2 of 0.70 or higher was obtained, indicating the predictive power of our 'UTR calculator'. In future work, the generality of our model will be tested with other promoters and UTRs.

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References