Gene Expression

mRNAStab - A web application for mRNA stability analysis
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ABSTRACT
Eukaryotic gene expression is regulated both at the transcription and the mRNA degradation levels. The implementation of functional genomics methods that allow the simultaneous measurement of transcription (TR) and degradation (DR) rates for thousands of mRNAs is a huge improvement in this field. One of the best established methods for mRNA stability determination is Genomic Run-On (GRO). It allows the measurement of DR, TR and mRNA levels during cell dynamic responses. Here we offer a software package that provides improved algorithms for determination of mRNA stability during dynamic GRO experiments.

Availability: The program mRNAStab is freely accessible at http://mRNAStab.uv.es/
Implementation: mRNAStab is written in C, PHP and R
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1 INTRODUCTION
mRNA is the messenger molecule transferring information from genes to proteins. It is by its own nature and function quite unstable (see Parker 2012 and Pérez-Ortín et al., 2007 for discussion). There is a variable turnover that depends on synthesis (transcription rate, TR) conducted by RNA polymerase II and degradation rate (DR) conducted by a complex system of RNases and other proteins or RNAs (Parker, 2012). The importance of TR in determining the levels of mRNAs is widely documented, however, the parallel importance of DR in it is only starting to be recognized (Pérez-Ortín et al., 2007).

There are several methods currently available for DR determination. Some of them are able to measure mRNA stabilities genomewide (see Chen et al, 2008; Schwald B, et al., 2012 and Supplementary data for a critical description and comparison). The most popular although prone to mistakes and biases (Wang et al., 2002; Marín-Navarro et al., 2011) is the transcription shut-off and subsequent determination of mRNA decay: assuming first-order kinetics, the degradation constant (k\textsubscript{d}) and the mRNA half-life (HL = ln2/ k\textsubscript{d}) can be calculated. The TR and mRNA concentrations can be determined genomewide by using a genomic run-on protocol (GRO) which allows to estimate the HLs of mRNAs even under non steady-state conditions (García-Martínez et al., 2004; Fan et al. 2012). This possibility is extremely interesting because it allows to determine TR, mRNA levels and HLs in a single experiment during the fast transcriptional response to an imposed environmental change (Castells-Roca et al., 2011 and references therein). This is the simplest protocol currently available for monitoring mRNA turnover in dynamic situations. However, it involves complex numerical analysis and costly computing. Now we describe here the mRNAStab package which offers two alternative algorithms (programs “StepK” and “RampK”) for determining mRNA HLs from data generated by GRO experiments. Additionally, it also includes a program (“Shutoff”) to calculate k\textsubscript{d} from simple decay experiments (e.g. Wang et al., 2002). All three programs are devised to cope with genome-wide data (TR and/or mRNA time sequences for thousands of genes).

2 WHAT THE PROGRAMS DO
Typically, the GRO experiment delivers values of TR (TR\textsubscript{0}, TR\textsubscript{1}, TR\textsubscript{2}...) and mRNA concentration ([mRNA]\textsubscript{0}, [mRNA]\textsubscript{1}, [mRNA]\textsubscript{2}...) simultaneously determined at discrete times (t\textsubscript{0}, t\textsubscript{1}, t\textsubscript{2}...). By assuming a linear variation of TR for a particular gene between consecutive pointwise determinations (e.g., values of TR and [mRNA] at t\textsubscript{0} and t\textsubscript{1}) with slope p:

\[ p = (TR_1 - TR_0)/(t_1 - t_0) \] 

Eq. 1

a mean value for the first-order degradation constant (k\textsubscript{d}) throughout the considered time interval can be determined (Pérez-Ortín et al., 2007; see Suppl. data). This calculation is sequentially translated to subsequent intervals and afterwards to all genes. A program following this procedure has been already available for spreadsheets (Marín-Navarro et al., 2011) but a faster version (in C) is now included in mRNAStab with the name of StepK.

An inconvenient feature of StepK is that mean values for k\textsubscript{d} jump stepwise from one interval to the next. A more realistic change would be to suppose that k\textsubscript{d} varies also linearly in between sample points as TR is assumed to do. Under these conditions, the rate of mRNA change for a gene would be:

\[ \frac{d[mRNA]}{dt} = TR_0 + p \cdot (t - t_0) - [k_d(t) + q \cdot (t - t_0)] \cdot [mRNA] \] 

Eq. 2

where q is the (unknown) rate of k\textsubscript{d} change with time, assumed constant for the time interval under consideration. For the first interval (between t\textsubscript{0} and t\textsubscript{1}) the initial value of k\textsubscript{d} [i.e., (k\textsubscript{d}(t_0))] is calculated supposing an initial steady state (since time 0 is usually

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taken before the onset of the environmental change triggering the transcriptional response) as $TR_d$ divided by $[mRNA]_0$. The RampK program integrates numerically Eq. 2 for the considered time interval by Euler’s method taking a random initial $q$ value. This value is contrasted by checking the difference between the calculated and experimental value of mRNA concentration at the end of the interval, and iteratively refined through a bisection algorithm. Once the value of $q$ has been determined to a chosen precision, the final value of $(k_d)_1$ (i.e., the value of $k_d$ at $t_1$) is calculated as:

$$(k_d)_1 = (k_d)_0 + q \cdot (t_1 - t_0) \quad \text{Eq. 3}$$

and this is used as initial value of $k_d$ for integrating the next interval (between $t_1$ and $t_2$). Similarly, the procedure is extended to subsequent intervals and further to all genes.

Therefore, under slightly different assumptions, both programs (StepK and RampK) deliver a time course for $k_d$ from experimental determinations of TR and mRNA concentration sampled at a sequence of time points. StepK estimates a mean value of $k_d$ for each interval while RampK gives the $k_d$ values at the interval extremes assuming a linear time course between them. Shutoff is an additional program included in this website to treat data of mRNA decay after transcriptional arrest (Wang et al., 2002). Thus, data will be mRNA concentrations at different time points. Shutoff calculates $k_d$ fitting these data to an exponential decay by minimizing square deviations.

3 STRUCTURE OF THE WEBSITE

We offer a novel implementation of a web site holding the algorithms described above. The user has the chance to work with a set of well known tools, all gathered in a user friendly web application. The algorithms run on our server. It introduces the notion of Experiment which contains one or more Operations, including Input, Output and Settings as child notions for the latter.

The input file must have a TR and/or a RA set of columns, each one having the time in the header. If one of the types is missing, neither StepK nor RampK can run. Furthermore, the first column always contains the unique names of the genes analyzed. The application consists of 3 layers: the UI, the algorithms and the storage, built using the Yii framework. It can be viewed as 3 tier architecture (see Figure 1), employing different languages at each level. The first one uses PHP with JavaScript/HTML/CSS. It controls the algorithms execution written in R (UPGMA, Sota, Kmeans – not implemented by us – and Shutoff), C (StepK and RampK) and PHP (Filtering). The last layer consists of the file system (storing of the inputs/outputs of each algorithm) and the database (store the successfully executed operations for a registered user).

4 USAGE AND APPLICATION

One can create an account or to use the application anonymously. Both cases offer the same features when it comes to capabilities. The difference between them lies in the possibility to save the experiments with their history, inputs and results when an account is used. A user must upload his/her file and then he/she can use the algorithms to process it. The results (visual or text) can be downloaded onto the user’s computer. There is also a FAQ section where more details about the algorithms and the workflow can be found. This website was tested under Mozilla Firefox, Google Chrome and Microsoft Internet Explorer.

5 CONCLUSIONS

We offer a web application to analyze the stability of the genes from microarray data. It contains two new applications capable of estimating the degradation constant ($k_d$) and, therefore, the mRNA stability for a list of time points of TR and mRNA.

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