A python module to normalize microarray data by the quantile adjustment method

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Microarray technology is widely used for gene expression research targeting the development of new drug treatments. In the case of a two-color microarray, the process starts with labeling DNA samples with fluorescent markers (cyanine 635 or Cy5 and cyanine 532 or Cy3), then mixing and hybridizing them on a chemically treated glass printed with probes, or fragments of genes. The level of hybridization between a strand of labeled DNA and a probe present on the array is measured by scanning the fluorescence of spots in order to quantify the expression based on the quality and number of pixels for each spot. The intensity data generated from these scans are subject to errors due to differences in fluorescence efficiency between Cy5 and Cy3, as well as variation in human handling and quality of the sample. Consequently, data have to be normalized to correct for variations which are not related to the biological phenomena under investigation. Among many existing normalization procedures, we have implemented the quantile adjustment method using the python computer language, and produced a module which can be run via an HTML dynamic form. This module is composed of different functions for data files reading, intensity and ratio computations and visualization. The current version of the HTML form allows the user to visualize the data before and after normalization. It also gives the option to subtract background noise before normalizing the data. The output results of this module are in agreement with the results of other normalization tools.

1. Introduction

The two-color microarray technique involves the immobilization of thousands of DNA fragments (probes) on a substrate, generally a ~6 cm² glass cover slip. Sample RNAs extracted from test and control tissue sources are converted to cDNA and labelled with two fluorescent markers, a green Cy3 and a red Cy5 cyanine. Then the labelled CDNAs are hybridized with the probes on the microarray (Schena et al., 1995). Gene expression levels are quantified by a scanner which estimates the fluorescence intensity in two wavelengths for each spot. The scanner computes the signal and background values for each spot via a dedicated algorithm (Yang et al., 2002). The reliability of these values can be affected by technical factors. The aim of data normalization is to adjust variations due to these factors in order to obtain reliable results that reflect real gene expression (Smyth and Speed, 2003).

There are two types of normalization methods: linear and nonlinear. Each of these methods may be applied at local or global level. In the linear method, the normalization consists of finding a proportional constant between data from the Cy5 and Cy3 dyes. This can be done using a set of probes for known or housekeeping genes on the array or by using data from all probes. Nonlinear approaches are more general and considered to be more powerful. LOWESS, based on local regression (Cleveland, 1979), and quantile adjustment methods are the
most widely used statistical methods for microarray data normalization (Fujita et al., 2006; Saviozzi et al., 2006; Dabney and Storey, 2007; Oshlack et al., 2007). However, the quantile method has the advantage to be quicker and simpler than the LOWESS. The algorithm of the quantile adjustment method was described by Bolstad et al. (2003).

Our module implements the quantile algorithm (read the manual for more details). It involves sorting intensity values before doing the calculation of their mean for each rank. After that, means are redistributed according the ranks of the initial list of values. An important advantage of the quantile method is that it does not require parameter tuning like more complex methods such as LOWESS.

There are several existing tools for microarray analysis, some of which are commercial. Most of these do not have available the source codes for modification and customization for special situations. Due to this limitation, it is important for biologists to contribute to the development of more flexible tools. In this spirit, we decided to develop a normalization module with source code freely available to anyone who wants to modify it in order to fit his/her own needs. Specifically, we have implemented a Python (Guido van Rossum, 2005) module for microarray data normalization using the quantile adjustment method which can be run via a web interface. As far as we know, there is no module for quantile adjustment normalization available in the biopython library; our attempt tries to fill this lack. We chose Python due to its clean syntax, easy typing and many available libraries which are well adapted to the problem we are addressing.

2. Method

The module is composed of several functions which are processing all calculations for data normalization and for graphical view. There is also a CGI (Common Gateway Interface) script which provides access to the normalization calculation via a web server where the module is installed. An HTML form that allows the user to submit data and the server sends back the result to the user is also included. In summary, the implemented module is capable of:

- read a data file with .GPR extension → input
- compute normalization according to quantile method
- view results as a graphically and as text → output
- present web interface for option choice

2.1. Input file

Currently accepted input file of our implementation is the .GPR (GenePix Results) (in Molecular Devices, 2010). This kind of file has a header comment which includes experiment date, description of the scanner parameters and the type of experiment. Our program analyzes only the data of signal and background.

2.2. Web interface form

Fig. 1 shows the form which allows a user to submit a data file (.gpr extension, GenePix) to the program. A user can select the signal of his preference (Median or Mean) before he or she proceeds with the normalization by checking a button in the second column. The box corresponding to subtract background can be checked if the user decides to subtract the background value from the signal value (Fig. 1). A CGI script is included for interaction between the module source code and the web interface.

From this form, the input file is uploaded from the field File Name, and then a user can select median or mean data. He can check one of these boxes (before or after normalization) to view data plots.

2.3. Output file

The output data of the program has four columns. It can contain from 20 to about 50,000 lines. A graphical review of the data is produced by plotting ratios against intensities.

In the output file, the first and second columns are normalized signal values (Cy5 and Cy3). Those data have been used to get ratio and intensity values. The prefix (Norm_) of columns names means that Ratio and intensity values (log 2) have been computed after data normalization. In the case where the program is executed with normalization option checked, in the output file, the prefix preceding intensity and ratio.

2.4. Plot view of ratio against intensity

Fig. 2 shows the plot of ratio and intensity before and after normalization.

Fig. 2(a) is graphical view of data (median values) before normalization. The ratio values are not symmetric to the zero ratio line while this is the case for the second plot (Fig. 2(b)). From Fig. 2(b): (i) if the ratio log value is greater than a given threshold,
e.g. 1, corresponding genes are up-regulated. (ii) If ratio log value is less than a given threshold, e.g. \( \frac{1}{C_0} \), corresponding genes are down-regulated. (iii) If the log ratio is equal to 0, the expression level of genes did not change. All genes between the upper and lower threshold (around the \( M = 0 \) line) are not changed ones.

Data generated by the module are similar to those obtained with standard software named Elea (Microarray & Sequencing Platform, 2009). For error quantification between values obtained from the two methods (our module versus Elea), we have calculated Mean Square Error (MSE) of couple values. The results are respectively 1.9985E-5 and 8.2341E-6 for ratios and mean intensities (Fig. 3). These error values are quasi null, validating the results of our module.

Fig. 3(right) shows that most of the ratio data points are along the diagonal of the scatterplot, again illustrating that the results obtained from both two tools are in good agreement. All intensity data points are likewise along the diagonal of the scatterplot (Fig. 3(left)).

3. Discussion

Normalizing microarray data is required to adjust for variations which are not related to the biological phenomena under study (Knudsen, 2004). There are many tools for microarray data normalization based on the two principal methods, the quantile method and LOWESS. One good example comes from a web site named MIDAS (microarray data analysis system) accessible via its URL (TM4-MIDAS, 2005), where there are efficient tools for microarray data analysis, including normalization, clustering and functional database search modules.

In spite of the availability of those tools which include graphical interfaces, and executables run on command line such as the

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**Fig. 2.** Graphical view of genes repartition according to ratio values.

**Fig. 3.** Scatterplot of ratio \( M \) made with results of our module and with Elea quantile method result (left). Scatterplot of intensity \( A \) made with results of our module and with Elea quantile method result (right).

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limma library in Bioconductor (Gentleman et al., 2004), there is a real need for software which may be adapted to users' options. Tools which can be run via Internet are largely used nowadays (Oliveres, 2008). For these reasons we developed this normalization module with an HTML interface module.

The main limit of online normalization tools is network bandwidth. Uploading microarray data files, which are often large, requires significant bandwidth to be time-efficient.

In future versions of our module a MySQL database will be included to improve its handling of submitted data management. In this case, if the user sends consecutively the same data file to a server, the file name will be saved, and the module will be able send back to the user the result of his first submission.

Authors' contributions

IB implemented the module and wrote the manuscript. DD has designed the project and reviewed the manuscript. JPT participated in implementing the quantile module and in reviewing the manuscript. NCM participated in web dynamic form design and in editing the manuscript. DS, SD, SFT, MSM participated to the manuscript. Knudsen, S., 2004. Guide to Analysis of DNA Microarray Data, 2nd edition. Wiley Liss, ISBN: 184 pp.

Availability and requirements

The code source of this module may be obtained from: http://taylor0.biology.ucla.edu/~manoukis/Quantile.tar.gz.

The installation of this module requires having python and several libraries (Numpy, matplotlib) available, in addition to a web server such as Karrigell and Apache (more details are provided in the program download).

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.meegid.2010.10.008.

References


Microarray & Sequencing Platform: «IGBMC, Strasbourg, France. IGBMC Microarray and Sequencing Platform».

Molecular Devices, Inc.©2010. Produced in the US. <http://www.moleculardevi-
ces.com/pages/software/gn_genepix_file_formats.html#gpr> (10.05.10).


TM4-MIDAS 2005, Dana-Farber Cancer Institute, 44 Binney St., Boston, MA, USA. <http://www.tm4.org/midas.html> (17.05.10).