FDR made easy in differential feature discovery and correlation analyses

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ABSTRACT

Summary: Rapid progress in technology, particularly in high-throughput biology, allows the analysis of thousands of genes or proteins simultaneously, where the multiple comparison problems occur. Global false discovery rate (gFDR) analysis statistically controls this error, computing the ratio of the number of false positives over the total number of rejections. Local FDR (lFDR) method can associate the corrected significance measure with each hypothesis testing for its feature-by-feature interpretation. Given the large feature number and sample size in any genomics or proteomics analysis, FDR computation, albeit critical, is both beyond the regular biologists’ specialty and computationally expensive, easily exceeding the capacity of desktop computers. To overcome this digital divide, a web portal has been developed that provides bench-side biologists easy access to the server-side computing capabilities to analyze for FDR, differential expressed genes or proteins, and for the correlation between molecular data and clinical measurements.

Availability: http://translationalmedicine.stanford.edu/Mass-Conductor/FDR.html

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION

The ability to simultaneously analyze vast number of genes or proteins in genomics, proteomics and imaging offers both unprecedented scientific opportunities and significant statistical challenges. The concurrent statistical test of thousands of null hypotheses leads to the multiple-testing problem, demanding that the expected proportion of false discoveries among all discoveries. This can be achieved either by the overly conservative Bonferroni correction or the analysis of the global false discovery rate (gFDR) (Benjamini and Hochberg, 1995). After determining the gFDR test threshold for significance, the IFDR (lFDR) analysis can compute and assign significance measures to all features. The IFDR analysis (Efron et al., 2001) addresses one drawback of the gFDR, statistically distinguishing features that are close to the threshold and therefore more likely to be falsely positive from those that are not.

Although important, multiple testing corrections do not always enter into practitioners’ considerations. The FDR computation usually requires some programming with either commercial packages (e.g. SAS, S-Plus) or programs (e.g. C, FORTRAN or R). In reality, few biologists or clinicians are proficient in software coding, scripting or are well informed of packages’ computational performance (Westfall et al., 1999) and potential memory allocation problems. Due to the large feature number and sample size, the FDR analysis can also be computationally expensive and most likely exceeds the capacity of desktop or notebook computers. To overcome these barriers, a web portal has been developed for the bench-side biologists such that they can easily access the server-side computing tools and generate graphic output for meaningful interpretation. Permutation-based gFDR analysis was implemented as previously described (Tusher et al., 2001) and the IFDR analysis integrated previous implementations of the FDR estimators (Aubert et al., 2004). The analysis results are summarized in plots and excel tables online, and are also e-mailed to the user from the server.

2 ANALYSIS GUIDELINES

There are two applications deployed at the web site: (i) differential analysis by Student’s t-test or the correlation tests lower than a threshold deemed statistically significant and are defined as total discoveries. Given the same threshold, permutations of the molecular measurements construct false positive feature sets of varied sizes. To assess the overall quality of the discoveries, thresholds ranging from the minimum single test P-value up to 1 are surveyed comprehensively. The box-whisker graphs, shown in Supplementary Figure 1 (P-value threshold range, left panel: minimum to 1.0; right panel: minimum to 0.05), illustrate the spread of the distribution of the sizes of the false discovery (FD) sets, using...
Another utility of lFDR is to assist the exploration of the biological significance of any given functional class or regulatory network. As shown in Figure 1A, the mean, median and 95th percentile of the feature-specific test $P$-value (Supplementary Fig. 2) are surveyed comprehensively. At each threshold, discoveries in both the input dataset (total discoveries) and the permutations of the input dataset (false discoveries, distribution described by a box-whisker graph) are counted to calculate gFDR. TD: total discoveries. (B) IFDR analysis. Each feature-specific test $P$-value is plotted against its corresponding lFDR.

To use 5% FDR as guideline to control overall false positive during multiple hypothesis testing is arbitrary. Therefore, the IFDR, which measures the significance that can be attached to each feature, is more appealing to the experimentalists because it can directly estimate the probability for the feature to be a false positive. Another utility of IFDR is to assist the exploration of the biological mechanism, as previously suggested (Aubert et al., 2004). The significance of any given functional class or regulatory network can be computed by summing the IFDRs of all component features. The graphic plots contrast the feature-specific $P$-value thresholds, balancing the need to maximize the chance of genuine discoveries while minimizing that of the false ones, the generally accepted 5% FDR is commonly used to guide the selection of the suitable feature-specific $P$-value threshold.

A 'box' (25–75%) and 'whiskers' to break down data by percentile. As shown in Figure 1A, the mean, median and 95th percentile of the sizes of these FD sets when divided by the total discoveries are used to estimate the mean, median and 95th percentile of the FDR, respectively. We found that the largest 5th percentile of the sizes of the FD sets are usually quite different from the remaining 95th percentile, exaggerating the totals of potential FDs. Therefore, 95 percentile of the FD distribution can serve as a good empirical upper bound for the FDR. To choose a good feature-specific $P$-value threshold, the need to maximize the chance of genuine discoveries while minimizing that of the false ones, the generally accepted 5% FDR is commonly used to guide the selection of the suitable feature-specific $P$-value threshold.

In addition to the graphic outputs, the results of feature-specific tests and FDRs are also summarized in excel files. For the correlation analysis between molecular data and a clinical variable, results are described in two tables: one summarizes all the feature-specific correlation $P$-values and the correlation coefficient estimates; and the other lists the feature-correlation $P$-values and the corresponding IFDRs.

3 OUTLOOK

Our FDR analysis portal is mainly designed for biologists or clinicians to analyze FDR in the hope of discovering genuinely differential features such as genes or proteins with statistical significance or correlations between molecular and clinical datasets to explore translational mechanisms. The analysis begins with a raw data upload and ends with a set of data sheets and plots for easy data drilling and visualization. With all the functions established in the Stanford server no informatics knowledge is required for the end user and computational capacity is guaranteed. This FDR portal should be of general interest to those working in the field of high-throughput biology to address the multiplicity of the statistical tests.

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REFERENCES


