

Multiple Testing Procedures with Applications to Genomics

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- [Katherine S. Pollard](#), Genome Center and Department of Statistics, UC Davis.

References

This presentation is based on the forthcoming book

[S. Dudoit and M. J. van der Laan \(2007\). *Multiple Testing Procedures with Applications to Genomics*, Springer Series in Statistics.](#)

Related articles, lecture notes, and software may be downloaded from Sandrine Dudoit's website

www.stat.berkeley.edu/~sandrine

and Mark van der Laan's website

www.stat.berkeley.edu/~laan.

Outline

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- Motivation: Multiple Hypothesis Testing Problems in Genomics.
- Overview of Main Contributions: Multiple Hypothesis Testing Framework and Procedures.
- Application: Multiple Tests of Association with Biological Annotation Metadata.
- Software Implementation: Bioconductor R Package `multtest`.

[Mark van der Laan.](#)

- Test Statistics Null Distribution.
- Resampling-Based Empirical Bayes Multiple Testing Procedures.

[Houston Gilbert.](#) FDR-Controlling Resampling-Based Empirical Bayes Multiple Testing Procedures: Simulation Study and Application to Microarray-Based Genetic Mapping of Gene Expression in *S. cerevisiae*.

Multiple Hypothesis Testing Problems in Genomics

- **High-throughput microarray gene expression analysis.**
 - Identification of **differentially expressed** (DE) genes. Testing for associations between gene expression measures and possibly censored biological and clinical covariates and outcomes.
 - Identification of **co-expressed** (CE) genes. Testing for associations in the expression measures of sets of genes across biological conditions.
- **Biological annotation metadata analysis.** Testing for associations between gene expression measures and biological annotation metadata.
E.g. Gene Ontology (GO) terms; PubMed abstracts.

Multiple Hypothesis Testing Problems in Genomics

- **Protein sequence analysis.** Testing for associations between phenotypes and codon/amino acid mutations.
E.g. Association between viral replication capacity and HIV-1 sequence variation.
- **Genetic mapping of complex traits.** Testing for associations between (sets of) phenotypes and genotypes.
E.g. Phenotypes: Affectedness status; transcript (i.e., mRNA) levels.
Genotypes: Single nucleotide polymorphisms (SNP); SNP haplotypes; microsatellite marker genotypes.
- **Mass-spectroscopy.** Testing for associations between phenotypes and protein mass-spectroscopy measures.
E.g. Association between leukemia class (ALL vs. AML) and mass-to-charge ratios.

Multiple Hypothesis Testing Problems in Genomics

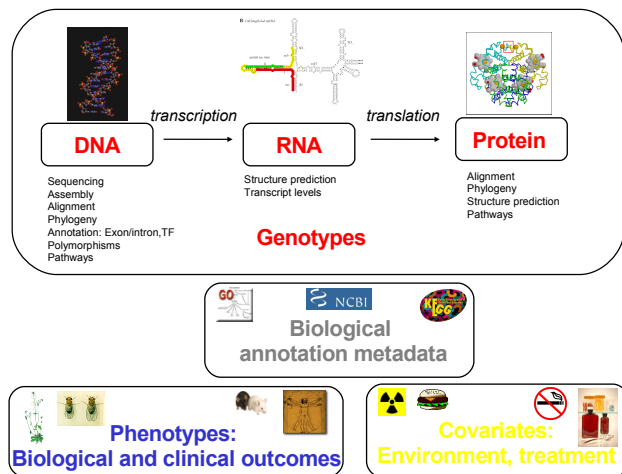


Figure 1: Biomedical and genomic data.

Multiple Hypothesis Testing Problems in Genomics

- Inference for **high-dimensional multivariate distributions**, with **complex and unknown dependence structures** among variables.
- **Broad range of parameters** of interest.
E.g. Regression coefficients in non-linear models relating patient survival data to genome-wide transcript levels, DNA copy numbers, or SNP genotypes;
measures of association between GO annotation and parameters of the distribution of microarray expression measures;
pairwise correlation coefficients between transcript levels.
- **Many null hypotheses**, in the thousands or even millions.
- **Complex and unknown dependence structures among test statistics.**
E.g. Directed acyclic graph (DAG) structure of GO terms;
Galois lattice for multilocus composite SNP genotypes.

Main Contributions: General and Unified Framework

Motivation.

- Large-scale multiple testing problems, e.g., genomics.
- Limitations of existing multiple testing methods, in terms of scope, Type I error rates, marginal nature, distributional assumptions, etc.

Main contributions.

- Foundations of a **general and unified methodology** for multiple hypothesis testing.
- **Resampling-based joint multiple testing procedures** (MTP) for controlling a broad class of Type I error rates, such as **generalized tail probabilities** and **generalized expected values** for arbitrary functions of the numbers of Type I errors and rejected hypotheses.
- **Software** implementation: Bioconductor R package `multtest`.

Main Contributions: General and Unified Framework

General and unified framework for multiple hypothesis testing.

- General definition of **null and alternative hypotheses** in terms of **submodels** for the data generating distribution,

$$H_0(m) \equiv I(P \in \mathcal{M}(m)) \quad \text{vs.} \quad H_1(m) \equiv I(P \notin \mathcal{M}(m)). \quad (1)$$

- General definition of **test statistics** and **rejection regions**.
- General definition of **Type I error rates** (and power) as **arbitrary parameters** $\Theta(F_{V_n, R_n})$ of the joint distribution of the numbers of Type I errors V_n and rejected hypotheses R_n .
- General definition of **adjusted p-values** and parameter **confidence regions** for **arbitrary Type I error rates**.

➡ Dudoit and van der Laan (2007, Chapter 1), Dudoit et al. (2004b), Pollard and van der Laan (2004).

Multiple Hypothesis Testing Framework: Error Rates

Table 1: *Type I and Type II errors in multiple hypothesis testing.*

		Null hypotheses			
		Non-rejected, \mathcal{R}_n^c	Rejected, \mathcal{R}_n		
True, \mathcal{H}_0	$W_n = \mathcal{R}_n^c \cap \mathcal{H}_0 $	$V_n = \mathcal{R}_n \cap \mathcal{H}_0 $	h_0		
False, \mathcal{H}_1	$U_n = \mathcal{R}_n^c \cap \mathcal{H}_1 $	$S_n = \mathcal{R}_n \cap \mathcal{H}_1 $	h_1		
	$M - R_n$	R_n	M		

Type I errors: $\mathcal{R}_n \cap \mathcal{H}_0$ Type II errors: $\mathcal{R}_n^c \cap \mathcal{H}_1$

Multiple Hypothesis Testing Framework: Error Rates

Type I error rates. Define a **Type I error rate** as a **parameter** $\theta_n = \Theta(F_{V_n, R_n})$ of the joint distribution F_{V_n, R_n} of the numbers of Type I errors $V_n = |\mathcal{R}_n \cap \mathcal{H}_0|$ and rejected hypotheses $R_n = |\mathcal{R}_n|$.

Specifically, consider **generalized tail probability** (gTP) error rates,

$$gTP(q, g) \equiv \Pr(g(V_n, R_n) > q), \quad (2)$$

and **generalized expected value** (gEV) error rates,

$$gEV(g) \equiv E[g(V_n, R_n)], \quad (3)$$

for arbitrary functions $g(V_n, R_n)$ of the numbers of Type I errors V_n and rejected hypotheses R_n .

Multiple Hypothesis Testing Framework: Error Rates

Number of false positives, $g(v, r) = v$.

Generalized family-wise error rate (gFWER):

$$gFWER(k) = \Pr(V_n > k).$$

Per-family error rate (PFER):

$$PFER = E[V_n].$$

Proportion of false positives among the rejected hypotheses, $g(v, r) = v/r$.

Tail probability for the proportion of false positives (TPFPF):

$$TPFPF(q) = \Pr(V_n/R_n > q).$$

False discovery rate (FDR):

$$FDR = E[V_n/R_n].$$

Type I error rates based on the **proportion of false positives among the rejected hypotheses** are particularly appealing for large-scale testing problems, as they do not increase exponentially with the number of tested hypotheses.

Multiple Hypothesis Testing Framework: Adjusted p -Values

As in the case of single hypothesis testing, the **results of a multiple testing procedure** $\mathcal{R}_n(\alpha)$ are reported in terms of the following quantities.

- **Rejection regions** for the test statistics.
- **Confidence regions** for the parameters of interest.
- **Adjusted p -values**. The **adjusted p -value** $\tilde{P}_{0n}(m)$, for null hypothesis $H_0(m)$, is the **smallest nominal Type I error level** α of the **multiple hypothesis testing procedure** $\mathcal{R}_n(\alpha)$ at which one would reject $H_0(m)$, given the data. That is,

$$\begin{aligned} \tilde{P}_{0n}(m) &\equiv \inf \{ \alpha \in [0, 1] : \text{Reject } H_0(m) \text{ at nominal MTP level } \alpha \} \\ &= \inf \{ \alpha \in [0, 1] : m \in \mathcal{R}_n(\alpha) \}, \quad m = 1, \dots, M. \end{aligned}$$

Multiple Hypothesis Testing Framework: Adjusted p -Values

Reporting the results of a MTP in terms of **adjusted p -values**, as opposed to only rejection or not of the null hypotheses, offers several **advantages**.

- Adjusted p -values can be defined for **any Type I error rate** (e.g., gFWER, TPFPF, or FDR).

E.g. FWER-controlling single-step Bonferroni (1936) MTP:

$$\tilde{P}_{0n}(m) = \min \{ M P_{0n}(m), 1 \}.$$

FDR-controlling step-up Benjamini and Hochberg (1995) MTP:

$$\tilde{P}_{0n}(O_n(m)) = \min_{h=m, \dots, M} \left\{ \min \left\{ \frac{M}{h} P_{0n}(O_n(h)), 1 \right\} \right\}.$$

- The **smaller** the adjusted p -value $\tilde{P}_{0n}(m)$, the **stronger the evidence** against the corresponding null hypothesis $H_0(m)$. Thus, one **rejects** $H_0(m)$ for **small adjusted p -values** $\tilde{P}_{0n}(m)$.

Multiple Hypothesis Testing Framework: Adjusted p -Values

- They reflect the strength of the evidence against each null hypothesis in terms of the **Type I error rate for the entire MTP**.
- They are **flexible summaries** of a MTP, in the sense that results are supplied for **all Type I error levels** α , i.e., the level α need not be chosen ahead of time.
- They provide convenient **benchmarks to compare different MTPs**, whereby smaller adjusted p -values indicate a less conservative procedure.
- **Plots of sorted adjusted p -values** allow investigators to examine sets of rejected hypotheses associated with various Type I error rates (e.g., gFWER, TPFPF, or FDR) and nominal levels α . Such plots provide tools to decide on an appropriate combination of the number of rejected hypotheses and tolerable false positive rate for a particular experiment and available resources.

Main Contributions: Test Statistics Null Distribution

Test statistics null distribution.

- **General characterization** of a proper null distribution in terms of **null domination conditions** for the test statistics for the true null hypotheses.
- **Explicit construction** of two main types of test statistics null distributions.
 - **Null shift and scale-transformed test statistics null distribution**, based on user-supplied upper bounds for the means and variances of the test statistics for the true null hypotheses.
 - **Null quantile-transformed test statistics null distribution**, based on user-supplied marginal test statistics null distributions.
- **Resampling procedures** (e.g., non-parametric and model-based bootstrap) for **consistent estimation** of the null distribution and of the corresponding test statistic cut-offs, parameter confidence regions, and adjusted p -values.

Main Contributions: Test Statistics Null Distribution

- Only concerned with controlling the Type I error rate under the **true data generating distribution**.
The concepts of weak and strong control of a Type I error rate are therefore irrelevant.
 - Directly consider a **null distribution for the test statistics** rather than a data generating null distribution.
The latter approach does not necessarily provide proper Type I error control under the true distribution.
- ⇒ Dudoit and van der Laan (2007, Chapter 2), Dudoit et al. (2004b), van der Laan and Hubbard (2006), Pollard and van der Laan (2004).

⇒ ... more in Mark van der Laan's presentation.

Main Contributions: Multiple Testing Procedures

Joint multiple testing procedures.

- **Joint single-step common-cut-off** and **common-quantile procedures** for controlling **general Type I error rates** $\Theta(F_{V_n})$, defined as arbitrary parameters of the distribution of the number of Type I errors V_n .
E.g. Generalized family-wise error rate (gFWER),
 $gFWER(k) = 1 - F_{V_n}(k) = \Pr(V_n > k)$.
⇒ Dudoit and van der Laan (2007, Chapter 4), Dudoit et al. (2004b), Pollard and van der Laan (2004).
- **Joint step-down common-cut-off (maxT)** and **common-quantile (minP) procedures** for controlling the **family-wise error rate** (FWER),
 $FWER = gFWER(0) = 1 - F_{V_n}(0) = \Pr(V_n > 0)$.
⇒ Dudoit and van der Laan (2007, Chapter 5), van der Laan et al. (2004a).

Main Contributions: Multiple Testing Procedures

- (Marginal/joint single-step/stepwise common-cut-off/common-quantile) **augmentation multiple testing procedures** (AMTP) for controlling **generalized tail probability** (gTP) error rates, $gTP(q, g) = \Pr(g(V_n, R_n) > q)$, based on an initial gFWER-controlling procedure.
E.g. gFWER: $g(v, r) = v$; TPPFP: $g(v, r) = v/r$.
⇒ Dudoit and van der Laan (2007, Chapter 6), Dudoit et al. (2004a), van der Laan et al. (2004b).
- **Joint resampling-based empirical Bayes procedures** for controlling **generalized tail probability** and **generalized expected value** error rates.
⇒ Dudoit and van der Laan (2007, Chapter 7), van der Laan et al. (2005).

⇒ ... more in Mark van der Laan's and Houston Gilbert's presentations.

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

Three-step road map. We have proposed a road map that leads to

- the general characterization and explicit construction of a proper **test statistics null distribution** Q_0 ;
- **joint single-step procedures** for controlling Type I error rates defined as arbitrary parameters $\Theta(F_{V_n})$ of the distribution of the number of Type I errors V_n (e.g., gFWER).

The main idea is to substitute control of the **unknown parameter** $\Theta(F_{V_n})$, for the **true distribution** F_{V_n} of the **number of Type I errors**, by control of the corresponding **known parameter** $\Theta(F_{R_0})$, for the **null distribution** F_{R_0} of the **number of rejected hypotheses**.

➡ Dudoit and van der Laan (2007, Chapter 4), Dudoit et al. (2004b), Pollard and van der Laan (2004).

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

Procedure 1 [Three-step road map for controlling Type I error rates $\Theta(F_{V_n})$]

1. **Null domination conditions for the Type I error rates $\Theta(F_{V_n})$ and $\Theta(F_{V_0})$.**
Select a test statistics null distribution Q_0 such that

$$(\limsup_{n \rightarrow \infty}) \quad \Theta(F_{V_n}) \leq \Theta(F_{V_0}). \quad (\text{ND}\Theta)$$

2. **Monotonicity of the Type I error rate mapping Θ .**

$$\Theta(F_{V_0}) \leq \Theta(F_{R_0}). \quad (5)$$

3. **Control of $\Theta(F_{R_0})$.** Select rejection regions so that

$$\Theta(F_{R_0}) \leq \alpha. \quad (6)$$

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

Specifically, consider single-step procedures with one-sided rejection regions, so that

$$\mathcal{R}_n = \{m : T_n(m) > c(m)\}.$$

Among the family of MTPs that satisfy the Type I error constraint

$$\Theta(F_{R_0}) \leq \alpha,$$

for $R_0 = R(c|Q_0) = \sum_{m=1}^M \mathbf{I}(Z(m) > c(m))$ and $Z \sim Q_0$, we have explicitly derived two types of procedures:

- Procedure 2, based on a **common cut-off** for all test statistics;
- Procedure 3, with **common-quantile cut-offs** for the test statistics.

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

Procedure 2 [$\Theta(F_{V_n})$ -controlling single-step common-cut-off procedure]

For controlling the Type I error rate $\Theta(F_{V_n})$ at level α , the set of rejected null hypotheses is of the form $\mathcal{R}_n = \{m : T_n(m) > c_0\}$, where the **common cut-off** c_0 is the **smallest** (i.e., least conservative) value for which $\Theta(F_{R_0}) \leq \alpha$. Adjusted p -values are given by

$$\tilde{P}_{0n}(m) = \Theta(F_{R(T_n(m)^{(M)}|Q_0)}), \quad m = 1, \dots, M, \quad (7)$$

where $T_n(m)^{(M)}$ denotes an M -vector of common cut-offs equal to $T_n(m)$.

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

- **gFWER** control, i.e., $\Theta(F_{V_n}) = 1 - F_{V_n}(k)$: **single-step $T(k+1)$** procedure, based on the **$(k+1)$ st largest test statistic**,

$$\tilde{p}_{0n}(m) = \Pr_{Q_0}(Z^{\circ}(k+1) \geq t_n(m)), \quad m = 1, \dots, M. \quad (8)$$

- **FWER** control, i.e., $\Theta(F_{V_n}) = 1 - F_{V_n}(0)$: **single-step maxT** procedure, based on the **maximum test statistic**,

$$\tilde{p}_{0n}(m) = \Pr_{Q_0} \left(\max_{m=1, \dots, M} Z(m) \geq t_n(m) \right), \quad m = 1, \dots, M. \quad (9)$$

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

Procedure 3 [$\Theta(F_{V_n})$ -controlling single-step common-quantile procedure]

For controlling the Type I error rate $\Theta(F_{V_n})$ at level α , the set of rejected null hypotheses is of the form $\mathcal{R}_n = \{m : T_n(m) > c_0(m)\}$, where $c_0(m) = Q_{0,m}^{-1}(\delta_0) = \inf \{z \in \mathbb{R} : Q_{0,m}(z) \geq \delta_0\}$ is the δ_0 -quantile of the marginal null distribution $Q_{0,m}$. The **common quantile probability** δ_0 is chosen as the **smallest** (i.e., least conservative) value for which $\Theta(F_{R_0}) \leq \alpha$. Adjusted p -values are given by

$$\tilde{P}_{0n}(m) = \Theta(F_{R(q_0^{-1}(1-P_{0n}(m)))|Q_0}), \quad m = 1, \dots, M, \quad (10)$$

where $P_{0n}(m)$ is the unadjusted p -value for null hypothesis $H_0(m)$,

$$P_{0n}(m) = \bar{Q}_{0,m}(T_n(m)) = 1 - Q_{0,m}(T_n(m)), \quad (11)$$

and $q_0^{-1}(\delta) = (Q_{0,m}^{-1}(\delta) : m = 1, \dots, M)$ denotes an M -vector of δ -quantiles for the marginal null distributions $Q_{0,m}$.

$\Theta(F_{V_n})$ -Controlling Single-Step Procedures

- **gFWER** control, i.e., $\Theta(F_{V_n}) = 1 - F_{V_n}(k)$: **single-step $P(k+1)$** procedure, based on the **$(k+1)$ st smallest unadjusted p -value**,

$$\tilde{p}_{0n}(m) = \Pr_{Q_0}(P_0^{\circ}(k+1) \leq p_{0n}(m)), \quad m = 1, \dots, M. \quad (12)$$

- **FWER** control, i.e., $\Theta(F_{V_n}) = 1 - F_{V_n}(0)$: **single-step minP** procedure, based on the **minimum unadjusted p -value**,

$$\tilde{p}_{0n}(m) = \Pr_{Q_0} \left(\min_{m=1, \dots, M} P_0(m) \leq p_{0n}(m) \right), \quad m = 1, \dots, M. \quad (13)$$

gTP-Controlling Augmentation Procedures

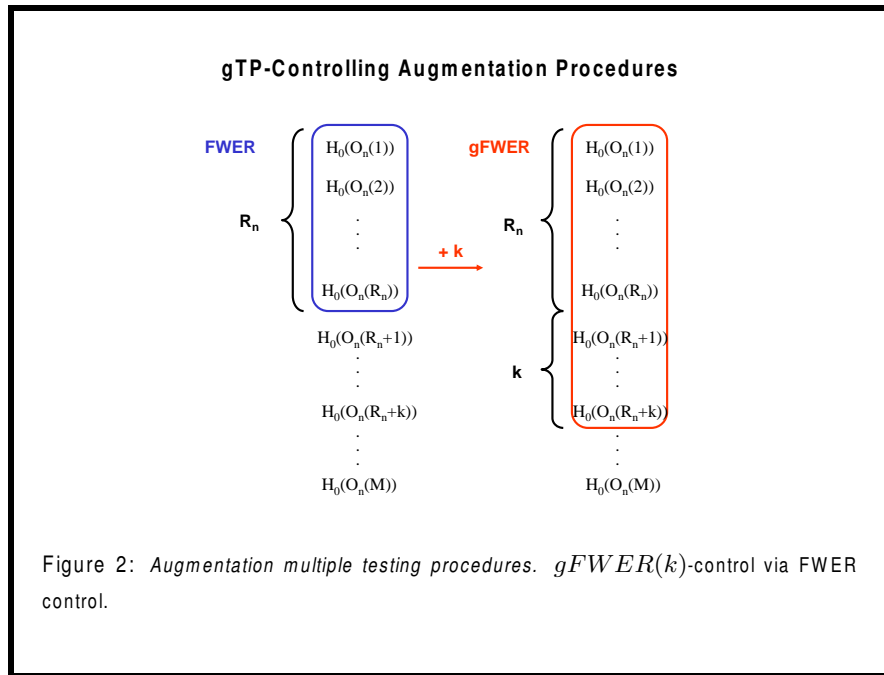
In order to control a new target Type I error rate, an **augmentation multiple testing procedure** (AMTP) adds suitably chosen null hypotheses to the set of hypotheses already rejected by an initial MTP.

Given any **initial gFWER-controlling MTP**, we have derived AMTPs for controlling **generalized tail probability** (gTP) error rates,

$gTP(q, g) = \Pr(g(V_n, R_n) > q)$, for arbitrary functions $g(V_n, R_n)$ of the numbers of Type I errors V_n and rejected hypotheses R_n .

$$\begin{array}{ccc} \text{gFWER} & \text{AMTP} & \text{gTP} \\ \mathcal{R}_n(\alpha) & \implies & \mathcal{R}_n^+(\alpha) = \mathcal{R}_n(\alpha) \cup \mathcal{A}_n(\alpha) \\ \Pr(V_n > k_0) \leq \alpha & & \Pr(g(V_n^+, R_n^+) > q) \leq \alpha \end{array}$$

➡ Dudoit and van der Laan (2007, Chapter 6), Dudoit et al. (2004a), van der Laan et al. (2004b).



gTP-Controlling Augmentation Procedures

Procedure 4 [gTP-controlling augmentation multiple testing procedure]

Consider any $gFWER(k_0)$ -controlling procedure $\mathcal{R}_n(\alpha)$, with adjusted p -values $\tilde{P}_{0n}(m)$ and indices $O_n(m)$ so that $\tilde{P}_{0n}(O_n(1)) \leq \dots \leq \tilde{P}_{0n}(O_n(M))$. This initial $gFWER$ -controlling procedure rejects the following $R_n(\alpha) = |\mathcal{R}_n(\alpha)|$ null hypotheses,

$$\mathcal{R}_n(\alpha) = \{m : \tilde{P}_{0n}(m) \leq \alpha\} = \{O_n(m) : m = 1, \dots, R_n(\alpha)\}.$$

For controlling $gTP(q, g)$ at level α , the augmentation multiple testing procedure rejects the $R_n(\alpha)$ null hypotheses specified by the initial $gFWER$ -controlling MTP, as well as the next $A_n(\alpha)$ most significant null hypotheses,

where

$$A_n(\alpha) \equiv \max \{m \in \{0, \dots, M - R_n(\alpha)\} : g(k_0 + m, R_n(\alpha) + m) \leq q\} \quad (14)$$

The set of rejected null hypotheses for the gTP -controlling AMTP is

$$\mathcal{R}_n^+(\alpha) \equiv \{O_n(m) : m = 1, \dots, R_n(\alpha) + A_n(\alpha)\} \quad (15)$$

and the adjusted p -values satisfy

$$\tilde{P}_{0n}(O_n(m)) = \tilde{P}_{0n}^+(O_n(S_n(m))), \quad (16)$$

where $S_n : \{1, \dots, M\} \rightarrow \{1, \dots, M\}$ is an integer shift function defined by

$$S_n(m) \equiv R_n^+(\tilde{P}_{0n}(O_n(m))) = m + A_n(\tilde{P}_{0n}(O_n(m))). \quad (17)$$

gTP-Controlling Augmentation Procedures

Intuitively, a gTP -controlling AMTP keeps rejecting null hypotheses until $g(k_0 + m, R_n + m)$ reaches the bound q for false positives.

The adjusted p -values for the AMTP are shifted versions of the adjusted p -values of the initial $gFWER$ -controlling MTP.

$gFWER$ -controlling AMTP. ($g(v, r) = v, k_0 = 0$)

$$\tilde{P}_{0n}^+(O_n(m)) = \begin{cases} 0, & \text{if } m \leq k \\ \tilde{P}_{0n}(O_n(m - k)), & \text{if } m > k \end{cases}. \quad (18)$$

TPFPF-controlling AMTP. ($g(v, r) = v/r, k_0 = 0$)

$$\tilde{P}_{0n}^+(O_n(m)) = \tilde{P}_{0n}(O_n(\lceil (1 - q)m \rceil)), \quad m = 1, \dots, M. \quad (19)$$

gTP-Controlling Augmentation Procedures

- Any **gFWER-controlling procedure** (marginal/joint single-step/stepwise common-cut-off/common-quantile) provides immediately and trivially MTPs controlling a **broad class of Type I error rates**, e.g., gFWER and TPPFP.
- One can **build on the large pool of available FWER-controlling MTPs**, such as the single-step and step-down maxT and minP procedures.
- **Adjusted p -values** for an AMTP are simply **shifted** versions of the ordered adjusted p -values for the initial MTP.
- **$gFWER(k)$ -controlling AMTP** guarantees **at least k rejected null hypotheses**.
- AMTPs augment the set of null hypotheses rejected by an initial MTP **conservatively**, in the sense that every additional rejected hypothesis is counted as a false positive.
- Unlike many procedures controlling the proportion of false positives, which assume either independence or specific dependence structures for the joint distribution of the test statistics, AMTPs provide gTP control for **general data generating distributions**, i.e., **arbitrary joint distributions for the test statistics**.

Resampling-Based Empirical Bayes Procedures


Many commonly-used MTPs share the following **two conservative features**.

- **Test statistics joint distribution**. The unknown test statistics joint distribution is replaced by a null distribution that satisfies **null domination** conditions.
E.g. $\Theta(F_{V_n}) \leq \Theta(F_{V_0})$ as in Step 1 of the road map of Procedure 1 and $\Theta(F_{V_n})$ -controlling single-step Procedures 2 and 3.
- **Set of true null hypotheses**. The unknown set of true null hypotheses \mathcal{H}_0 is replaced by the **complete set of null hypotheses** $\{1, \dots, M\}$.
E.g. $\Theta(F_{V_0}) \leq \Theta(F_{R_0})$ as in Step 3 of the road map of Procedure 1 and $\Theta(F_{V_n})$ -controlling single-step Procedures 2 and 3;
counting every additional rejected hypothesis as a Type I error in Equation (14) of gTP-controlling augmentation Procedure 4;
controlling FDR at level $(h_0/M)\alpha \leq \alpha$ as in step-up Benjamini and Hochberg (1995) procedure.

Resampling-Based Empirical Bayes Procedures

In order to achieve more power regarding the second point, one can adopt an empirical Bayes approach and generate **random guessed sets of true null hypotheses** \mathcal{H}_{0n} under a suitable distribution $Q_{0n}^{\mathcal{H}}$.

We have provided a general characterization and explicit constructions for **resampling-based empirical Bayes** procedures that control **generalized tail probability** and **generalized expected value** error rates, e.g., FDR.

 Dudoit and van der Laan (2007, Chapter 7), van der Laan et al. (2005).

 ... more in Mark van der Laan's and Houston Gilbert's presentations.

Main Contributions: General and Unified Framework

N. B. Compared to previously-proposed approaches, our multiple testing procedures based on a null-transformed test statistics null distribution offer the following advantages.

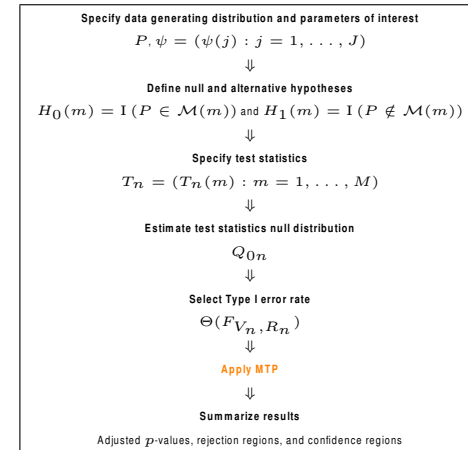
- **General and unified framework** for multiple hypothesis testing.
- **Proper Type I error control** for **general**
 - **data generating distributions**, with **arbitrary dependence structures** among variables;
 - **null hypotheses**, defined in terms of **submodels** for the data generating distribution;
 - **test statistics**, e.g., t -statistics, χ^2 -statistics, F -statistics;
 - Type I error rates, such as, **generalized tail probabilities** and **generalized expected values** for arbitrary functions of the numbers of Type I errors and rejected hypotheses.

Main Contributions: General and Unified Framework

- Do not rely on restrictive and questionable assumptions on the joint distribution of the test statistics, such as, independence, Simes' Inequality, subset pivotality.
- Account for the joint distribution of the test statistics \implies more power than procedures based solely on marginal distributions, i.e., unadjusted p -values.
- Report results using adjusted p -values.

Main Contributions: General and Unified Framework

Table 2: Multiple hypothesis testing flowchart.



Main Contributions: General and Unified Framework

Apply MTP

FWER	$\Pr(V_n > 0)$	Single-step common-cut-off maxT Single-step common-quantile minP Step-down common-cut-off maxT Step-down common-quantile minP Resampling-based empirical Bayes
gFWER	$\Pr(V_n > k)$	Single-step common-cut-off $T(k + 1)$ Single-step common-quantile $P(k + 1)$ Augmentation Resampling-based empirical Bayes
General	$\Theta(F_{V_n})$	Single-step common-cut-off Single-step common-quantile Resampling-based empirical Bayes
TPFP	$\Pr(V_n / R_n > q)$	Augmentation Resampling-based empirical Bayes
gTP	$\Pr(g(V_n, R_n) > q)$	Augmentation Resampling-based empirical Bayes
FDR	$E[V_n / R_n]$	TPFP-based Resampling-based empirical Bayes
gEV	$E[g(V_n, R_n)]$	gTP-based Resampling-based empirical Bayes
General	$\Theta(F_{g(V_n, R_n)})$	Resampling-based empirical Bayes

Main Contributions: General and Unified Framework

- Data generating distribution: $P \in \mathcal{M}$.
- Parameters: $\psi = (\psi(j) : j = 1, \dots, J)$, where $\psi(j) = \Psi(P)(j)$.
- Null and alternative hypotheses: $H_0(m) = I(P \in \mathcal{M}(m))$ and $H_1(m) = I(P \notin \mathcal{M}(m))$, where $\mathcal{M}(m) \subseteq \mathcal{M}$, $m = 1, \dots, M$.
- Data and empirical distribution: $\mathcal{X}_n = \{X_i : i = 1, \dots, n\} \stackrel{IID}{\sim} P, P_n$.
- Test statistics: $T_n = (T_n(m) : m = 1, \dots, M)$, where $T_n(m) = T(m; \mathcal{X}_n) = T(m; P_n)$.
- Test statistics null distribution: Q_0 (or estimator thereof, Q_{0n}).
- Multiple testing procedure and rejection regions: $\mathcal{R}_n = \mathcal{R}(T_n, Q_{0n}, \alpha) = \{m : T_n(m) \in \mathcal{C}_n(m)\} = \{m : H_0(m) \text{ is rejected}\}$.
- Type I error rate: $\theta_n = \Theta(F_{V_n, R_n})$, where $V_n = |\mathcal{R}_n \cap \mathcal{H}_0| = \#$ Type I errors and $R_n = |\mathcal{R}_n| = \#$ rejected hypotheses.
- Type II error rate/power: $\vartheta_n = \Theta(F_{U_n, R_n})$, where $U_n = |\mathcal{R}_n^c \cap \mathcal{H}_1| = \#$ Type II errors.
- Summaries of results: Adjusted p -values, test statistic rejection regions, parameter confidence regions.


Tests of Association with Biological Annotation Metadata

Experimental data, such as microarray gene expression measures, gain much in relevance from their association with **biological annotation metadata**, i.e., **data on data**.

E.g. GenBank sequences, GO terms, KEGG pathways, PubMed abstracts.

A challenging and fascinating area of research for statisticians concerns the development of methods for **relating experimental data to the wealth of metadata** available publicly on the WWW.

Tasks include **accessing and pre-processing** the data, **making inference** from these data, and **summarizing and interpreting** the results.

 Dudoit and van der Laan (2007, Chapter 10), Dudoit et al. (2007).

Tests of Association with Biological Annotation Metadata

In this context, an important class of statistical problems involves **testing for associations** between

- **gene-annotation profiles**, i.e., **known fixed features** of a genome,
- **gene-parameter profiles**, i.e., **unknown parameters of the distribution of variable features** of this genome in a population of interest.

Here, features of a genome are said to be **fixed**, if they remain constant among population units. In contrast, **variable** features are allowed to differ among population units.

The parameter of interest then corresponds to **measures of association between known gene-annotation profiles and unknown gene-parameter profiles**.

Tests of Association with Biological Annotation Metadata

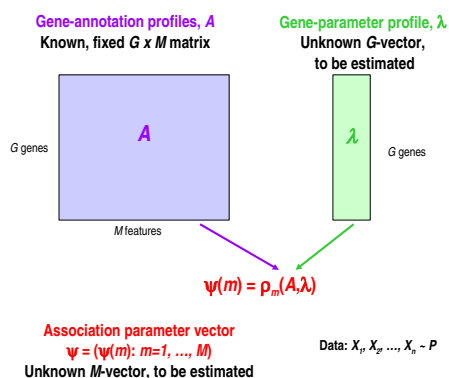


Figure 3: *Parameters for tests of association with biological annotation metadata.*

Tests of Association with Biological Annotation Metadata

Gene-annotation profiles. Gene-annotation profiles refer to features of a genome that are assumed to be **known** and **constant** among population units.

Fixed features of interest typically consist of **gene annotation metadata**, that reflect current knowledge on **gene properties**, such as, nucleotide and protein sequences, regulation, and function.

E.g. Gene Ontology (GO, www.geneontology.org) annotation.

Gene pathway membership (e.g., Kyoto Encyclopedia of Genes and Genomes, KEGG, www.genome.ad.jp/kegg).

Gene regulation by particular transcription factors, presence or absence of certain motifs in gene control region (e.g., Transcription Factor DataBase, TRANSFAC, www.gene-regulation.com).

Exon/intron counts/lengths/nucleotide distributions.

N. B. Features are **fixed in time** only for a given version/release of the corresponding database(s).

Tests of Association with Biological Annotation Metadata

Let $A = (A(g, m) : g = 1, \dots, G; m = 1, \dots, M)$ denote a $G \times M$ **gene-annotation matrix**, providing data on M features for G genes in an organism of interest.

Row $A(g, \cdot) = (A(g, m) : m = 1, \dots, M)$ is an M -dimensional gene-specific feature vector for the g th gene.

Column $A(\cdot, m) = (A(g, m) : g = 1, \dots, G)$ is a G -dimensional **gene-annotation profile** for the m th feature.

Tests of Association with Biological Annotation Metadata

Gene-parameter profiles. Gene-parameter profiles concern the **distribution of variable features** of a genome in a well-defined population.

Gene-specific variables of interest reflect **cellular type/state/activity** under particular conditions and include microarray measures of transcript levels and comparative genomic hybridization (CGH) measures of DNA copy numbers.

E.g. Vector of genome-wide mean transcript levels in a population of heat-shocked yeast cells.

Vector of regression coefficients relating survival to genome-wide transcript levels or DNA copy numbers in a population of cancer patients.

Tests of Association with Biological Annotation Metadata

Let $X = (X(j) : j = 1, \dots, J) \sim P \in \mathcal{M}$ denote a J -dimensional random vector, with data generating distribution P belonging to a (possibly non-parametric) model \mathcal{M} .

Let the parameter mapping $\Lambda : \mathcal{M} \rightarrow \mathbb{R}^G$ define a G -dimensional **gene-parameter profile**, $\Lambda(P) = \lambda = (\lambda(g) : g = 1, \dots, G) \in \mathbb{R}^G$.

While gene-annotation profiles are known and fixed, gene-parameter profiles are typically **unknown** and need to be **estimated**, e.g., from a microarray experiment involving a sample of population units.

Tests of Association with Biological Annotation Metadata

The **association parameter** of interest is an M -vector

$$\psi = (\psi(m) : m = 1, \dots, M) = \rho(A, \lambda), \quad (20)$$

of association measures between the **gene-annotation profiles** A and a **gene-parameter profile** λ .

In the simplest case, one could define the M association parameters univariately, i.e., let

$$\psi(m) = \rho_m(A(\cdot, m), \lambda), \quad (21)$$

where $\rho_m(\cdot, \cdot)$ provides a measure of association between two G -vectors (e.g., t -statistic, χ^2 -statistic, Pearson correlation coefficient).

Tests of Association with Biological Annotation Metadata

Our approach to multiple tests of association with biological annotation metadata **differs in a number of important ways** from current approaches, such as those developed for inference with GO.

General gene-annotation profiles.

Existing approaches typically consider binary gene-annotation profiles, e.g., vectors of indicators of GO term annotation.

Our general definition of gene-annotation profiles allows consideration of arbitrary qualitative and quantitative fixed features of a genome, e.g., membership of genes to any number of pathways or clusters, exon/intron counts/lengths/nucleotide distributions, mean transcript levels.

Tests of Association with Biological Annotation Metadata

General gene-parameter profiles.

Existing approaches typically consider binary gene-parameter profiles, e.g., vectors of indicators of differential expression.

Our general definition of gene-parameter profiles allows consideration of a much broader class of testing problems, concerning arbitrary qualitative and quantitative parameters, such as, differences in mean expression levels or regression coefficients relating expression levels to clinical outcomes.

Estimated gene-parameter profiles.

Existing approaches typically assume known gene-parameter profiles. For example, the list of DE genes from a microarray experiment is usually treated as known and fixed in subsequent analyses with GO, while in fact it corresponds to an unknown and estimated parameter.

Distinguishing between the definition of a parameter and inference concerning this parameter provides a more rigorous and general formulation of the statistical question.

Tests of Association with Biological Annotation Metadata

General tests of association.

Common approaches to tests of association with GO annotation are typically limited to tests of independence in 2×2 contingency tables (e.g., based on the hypergeometric distribution, Fisher's exact test). Rows correspond to gene annotation with a given GO term (fixed binary gene-annotation profile) and columns to a gene property of interest, such as differential expression (treated as a fixed binary gene-parameter profile).

Our approach allows consideration of a broader class of biological testing problems, while properly accounting for the fact that gene-parameter profiles are usually unknown and replaced by a random (i.e., data-driven) estimator.

Tests of Association between GO Annotation and DE in ALL

Our proposed approach to tests of association with biological annotation metadata is illustrated using the **acute lymphoblastic leukemia (ALL) microarray** dataset of Chiaretti et al. (2004), with the aim of **relating Gene Ontology (GO) annotation to differential expression (DE)** among ALL samples.

The **BCR/ABL fusion** is the molecular analogue of the **Philadelphia chromosome**, one of the most frequent cytogenetic abnormalities in human leukemias. A number of recent articles have investigated the **prognostic relevance** of the BCR/ABL fusion in adult ALL of the B-cell lineage.

Consider the following two related questions.

- Identifying **differentially expressed genes** between B-cell ALL with the BCR/ABL fusion and cytogenetically normal NEG B-cell ALL.
- Identifying **GO terms associated with differential expression**.

Tests of Association between GO Annotation and DE in ALL

Acute lymphoblastic leukemia dataset. The Chiaretti et al. (2004) ALL dataset comprises, for each of 128 ALL cell samples,

- 12,625 **microarray expression measures** (Affymetrix chip series HG-U95Av2);
- 21 **phenotypes** (i.e., covariates and outcomes).

Focus on

- $n = 79$ **B-cell ALL** cell samples of the **BCR/ABL** and **NEG** molecular types;
- $G = 2,071$ **filtered genes** with unique Entrez Gene IDs.

Tests of Association between GO Annotation and DE in ALL

The Gene Ontology. The **Gene Ontology** (GO) Consortium (www.geneontology.org) provides **ontologies**, i.e., structured and controlled vocabularies, to describe gene products in terms of their associated **biological processes** (BP), **cellular components** (CC), and **molecular functions** (MF).

For each of the three ontologies, GO terms are organized in a **directed acyclic graph** (DAG), i.e., a **directed graph** (one-way edges) containing **no cycles** (no path starts and ends at the same vertex).

The GO Consortium and other organizations provide **mappings** between GO terms and genes in various organisms.

Tests of Association between GO Annotation and DE in ALL

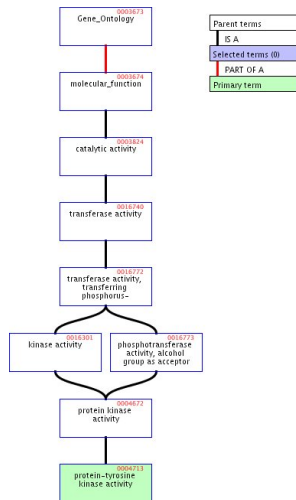


Figure 4: *Gene Ontology*. Portion of the MF DAG for the GO term *protein-tyrosine kinase activity* (GO:0004713). EBI QuickGO browser (www.ebi.ac.uk/ego).

Tests of Association between GO Annotation and DE in ALL

GO gene-annotation profiles. For each of the three gene ontologies, assemble a $G \times M$ **binary gene-annotation matrix** A , indicating for each gene g whether it is annotated with each GO term m ,

$$A(g, m) = \begin{cases} 1, & \text{if gene } g \text{ is annotated with GO term } m \\ 0, & \text{otherwise} \end{cases}$$

Consider only the M GO terms annotating at least 10 of the $G = 2,071$ filtered genes.

Ontology	Number of terms, M
Biological Process	367
Cellular Component	81
Molecular Function	185

Tests of Association between GO Annotation and DE in ALL

Null hypotheses. Test the M null hypotheses of **no association** between GO gene-annotation profiles $A(\cdot, m)$ and a DE gene-parameter profile λ .

	Testing scenario		
	Continuous		Binary
	MT[t, t]	MT[d, t]	MT[\neq, χ]
DE gene-parameter, λ	Standardized difference of means	Unstandardized	Indicator
GO gene-annotation, A		Indicator	Indicator
DE and GO association, ρ		t -statistic	χ^2 -statistic
Null hypotheses, $H_0(m)$	$\rho(A(\cdot, m), \lambda) = 0$		$\rho(A(\cdot, m), \lambda) \leq 1$

Tests of Association between GO Annotation and DE in ALL

Testing scenarios MT[t, t] and MT[d, t].

- DE gene-parameter profile, λ : **Continuous**
Standardized/Unstandardized differences of means in BCR/ABL vs. NEG B-cell ALL
Estimator: Two-sample Welch t -statistics/differences of empirical means
- GO gene-annotation profiles, A : **Binary**
- DE and GO association measure, ρ : Two-sample Welch t -statistic

Testing scenario MT[\neq, χ]. (The usual approach.)

- DE gene-parameter profile, λ : **Binary**
Indicators of DE between BCR/ABL and NEG B-cell ALL
Estimator: Based on adjusted p -values for FWER-controlling permutation-based single-step maxT procedure, with a pre-specified proportion of DE genes (MT[$\neq, \chi : \gamma G$]) or significance level (MT[$\neq, \chi : \alpha$])
- GO gene-annotation profiles, A : **Binary**
- DE and GO association measure, ρ : χ^2 -statistic

Tests of Association between GO Annotation and DE in ALL

Test statistics. Unstandardized difference statistics:

$$T_n(m) = \sqrt{n}(\psi_n(m) - \psi_0(m)).$$

Test statistics null distribution. Non-parametric bootstrap estimator of the null shift-transformed test statistics null distribution, $B = 5,000$.

Multiple testing procedure. FWER-controlling single-step maxT procedure.

Tests of Association between GO Annotation and DE in ALL

Results: DE between BCR/ABL and NEG B-cell ALL. Two-sided tests using two-sample Welch t -statistics and FWER-controlling bootstrap-based single-step maxT MTP.

- 16 DE genes** at nominal FWER level $\alpha = 0.05$.
- DE genes tend to be **over-expressed** in cell samples with the BCR/ABL fusion (14/16 positive t -statistics).
- The **ABL1 gene** shows the most over-expression in BCR/ABL cell samples.
- DE genes appear to be related to **apoptosis** or **oncogenesis**.

Tests of Association between GO Annotation and DE in ALL

Results: Association between GO annotation and DE in ALL.

- Adjusted p -values tend to be quite large, with only a handful of GO terms identified as being significantly associated with BCR/ABL vs. NEG DE.
- Little overlap between the binary and continuous testing scenarios.
- Testing scenarios based on binary DE gene-parameter profiles tend to be more conservative than scenarios based on continuous profiles and lack robustness with respect to the somewhat arbitrary DE/non-DE gene dichotomization (i.e., the number of DE genes).
- Testing scenarios based on standardized and unstandardized continuous DE gene-parameter profiles lead to very similar results.
- GO terms associated with BCR/ABL vs. NEG DE tend to concentrate in certain branches of the DAGs.
- Some of the genes annotated with the identified GO terms have been linked to the BCR/ABL proto-oncogene and have been suggested as potential targets for molecular therapies of leukemia.

Tests of Association between GO Annotation and DE in ALL

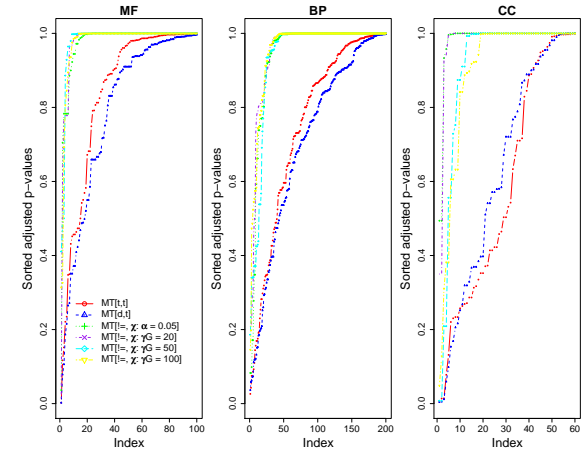


Figure 5: GO terms associated with BCR/ABL vs. NEG DE, adjusted p -values.

Tests of Association between GO Annotation and DE in ALL

Table 3: GO terms associated with BCR/ABL vs. NEG DE.

	Nominal FWER level, α								
	0.05	0.10	0.20	0.05	0.10	0.20	0.05	0.10	0.20
MT[t, t]	2	6	14	3	4	5	1	1	3
MT[d, t]	1	5	16	3	5	7	1	2	4
MT[≠, χ : α = 0.05]	0	3	5	0	0	0	1	1	1
MT[≠, χ : γG = 20]	0	0	0	0	0	0	1	1	1
MT[≠, χ : γG = 50]	0	0	1	2	2	2	0	0	0
MT[≠, χ : γG = 100]	0	0	2	1	1	2	0	0	0

BP

CC

MF

Tests of Association between GO Annotation and DE in ALL

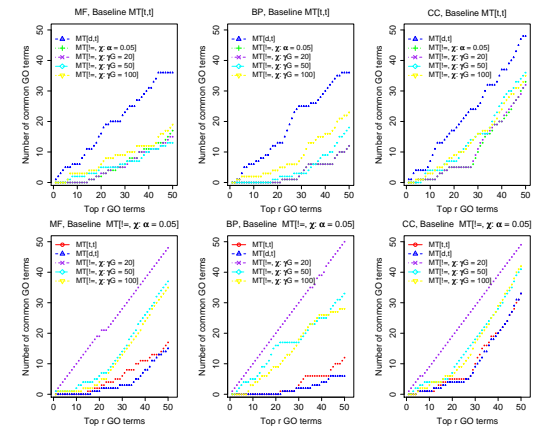


Figure 6: GO terms associated with BCR/ABL vs. NEG DE, common terms between testing scenarios.

Tests of Association between GO Annotation and DE in ALL

Table 4: GO terms associated with BCR/ABL vs. NEG DE, top 20 BP GO terms.

BP, Scenario MT[t , t]			
GO term ID	GO term	$A_1(m)$	$\hat{P}_{0n}(m)$
GO:008152	metabolism	1076	2.6e-02
GO:044237	cellular metabolism	1045	4.3e-02
GO:009058	biosynthesis	187	7.5e-02
GO:044238	primary metabolism	1002	7.5e-02
GO:044249	cellular biosynthesis	169	8.6e-02
GO:006091	generation of precursor metabolites and energy	98	9.3e-02
GO:019882	antigen presentation	15	1.1e-01
GO:030333	antigen processing	14	1.4e-01
GO:006916	anti-apoptosis	21	1.6e-01
GO:043066	negative regulation of apoptosis	26	1.7e-01
GO:043069	negative regulation of programmed cell death	26	1.7e-01
GO:007154	cell communication	390	1.8e-01
GO:006457	protein folding	52	1.9e-01
GO:007165	signal transduction	351	1.9e-01
GO:000226	microtubule cytoskeleton organization and biogenesis	14	2.3e-01
GO:006082	organic acid metabolism	65	2.5e-01
GO:006163	purine nucleotide metabolism	29	2.8e-01
GO:007155	cell adhesion	59	2.8e-01
GO:007028	cytoplasm organization and biogenesis	10	3.0e-01
GO:019752	carboxylic acid metabolism	63	3.1e-01

Tests of Association between GO Annotation and DE in ALL

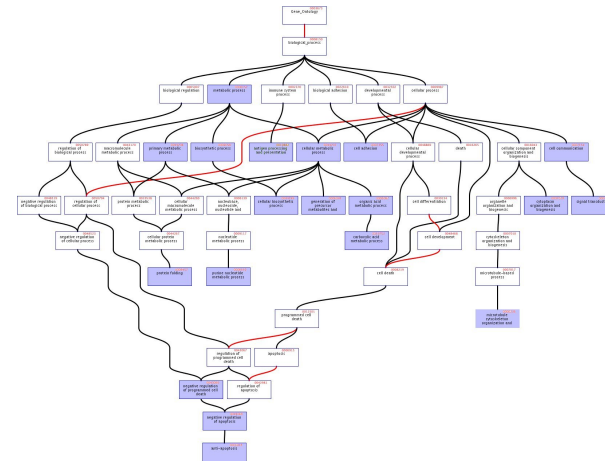


Figure 7: GO terms associated with BCR/ABL vs. NEG DE, DAG for top 20 BP GO terms.

Tests of Association between GO Annotation and DE in ALL

Table 5: GO terms associated with BCR/ABL vs. NEG DE, top 20 CC GO terms.

CC, Scenario MT[t , t]			
GO term ID	GO term	$A_1(m)$	$\hat{P}_{0n}(m)$
GO:0005840	ribosome	25	5.6e-03
GO:0030529	ribonucleoprotein complex	77	1.4e-02
GO:0005830	cytosolic ribosome (sensu Eukaryota)	11	1.4e-02
GO:0043234	protein complex	334	7.8e-02
GO:0005886	plasma membrane	200	1.3e-01
GO:0005829	cytosol	78	2.2e-01
GO:0005737	cytoplasm	578	2.3e-01
GO:0005887	integral to plasma membrane	125	2.3e-01
GO:0031226	intrinsic to plasma membrane	125	2.3e-01
GO:0019866	inner membrane	37	2.6e-01
GO:0005743	mitochondrial inner membrane	28	2.6e-01
GO:0005746	mitochondrial electron transport chain	11	2.7e-01
GO:000502	proteasome complex (sensu Eukaryota)	26	2.7e-01
GO:0000323	lytic vacuole	28	2.9e-01
GO:0005764	lysosome	28	2.9e-01
GO:0005576	extracellular region	54	3.1e-01
GO:0005773	vacuole	29	3.2e-01
GO:0005622	intracellular	1152	3.4e-01
GO:0043228	non-membrane-bound organelle	218	3.5e-01
GO:0043232	intracellular non-membrane-bound organelle	218	3.5e-01

Tests of Association between GO Annotation and DE in ALL

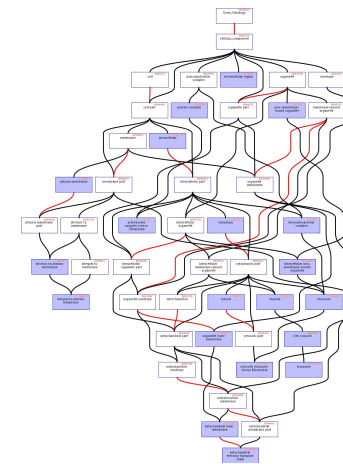


Figure 8: GO terms associated with BCR/ABL vs. NEG DE, DAG for top 20 CC GO terms.

Tests of Association between GO Annotation and DE in ALL

Pre-processing and filtering.

- Three-step **robust multichip average** (RMA) pre-processing for all 128 ALL samples (Bolstad et al., 2005).
- **Base 2 logarithmic transformation**.
- **Intensity-based filtering** (von Heydebreck et al., 2004). Retain only probes with: (i) fluorescence intensities greater than 100 (absolute scale) for at least 25% of the 79 cell samples and (ii) interquartile range (IQR) of the fluorescence intensities for the 79 cell samples greater than 0.5 (log base 2 scale).
- Average the expression measures of **multiple probes mapping to the same gene** (i.e., same Entrez Gene ID).

Tests of Association between GO Annotation and DE in ALL

R and Bioconductor packages. (R Release 2.2.1; Bioconductor Release 1.7)

- **multtest** (Version 1.8.0): Resampling-based multiple testing procedures.
- **ALL** (Version 1.0.2): Microarray expression measures and phenotypes for Chiaretti et al. (2004) ALL study.
- **annotate** (Version 1.8.0): General annotation software package.
- **annaffy** (Version 1.2.0): Annotating and generating HTML reports for Affymetrix chip data.
- **hgu95av2** (Version 1.10.0): Affymetrix chip-specific metadata package.
- **GO** (Version 1.10.0): GO-specific metadata package.

Software Implementation: Bioconductor R Package **multtest**


The multiple testing procedures developed in Dudoit and van der Laan (2007) and related articles are implemented in the **R package multtest**, released as part of the **Bioconductor Project**, an open-source software project for the analysis of biomedical and genomic data.

Please consult the package documentation (e.g., helpfiles, manuals) and book chapters for details.

Bioconductor R package: **multtest**.

Authors: Katherine S. Pollard, Yongchao Ge, and Sandrine Dudoit.

URL: www.bioconductor.org.

 Dudoit and van der Laan (2007, Section 13.1), Pollard et al. (2005).

Software Implementation: Bioconductor R Package **multtest**

Test statistics. *t*-statistics for tests of regression coefficients in linear models and Cox proportional hazards survival models;

F-statistics for tests of equality of means in one-way and two-way designs.

Weighted and robust rank-based versions of the above test statistics are implemented.

Test statistics null distribution. Bootstrap null shift and scale-transformed; permutation.

Software Implementation: Bioconductor R Package `multtest`

Multiple testing procedures.

- **FWER control.**
 - Marginal single-step Bonferroni (1936), step-down Holm (1979), and step-up Hochberg (1988).
 - Joint single-step maxT and minP (Ch. 4 in Dudoit and van der Laan, 2007; Dudoit et al., 2004b; Pollard and van der Laan, 2004).
 - Joint step-down maxT and minP (Ch. 5 in Dudoit and van der Laan, 2007; van der Laan et al., 2004a).
- **gFWER and TPPFP control.** Augmentation multiple testing procedures (Ch. 6 in Dudoit and van der Laan, 2007; van der Laan et al., 2004b).
- **FDR control.**
 - Marginal step-up Benjamini and Hochberg (1995) and Benjamini and Yekutieli (2001).
 - TPPFP-based (Ch. 6 in Dudoit and van der Laan, 2007; van der Laan et al., 2004b).

Software Implementation: Bioconductor R Package `multtest`

- **Numerical summaries.** Parameter estimates; test statistics; unadjusted and adjusted p -values; test statistic cut-offs; parameter confidence regions; estimated test statistics null distribution.
- **Graphical summaries.** Type I error rates vs. # rejections; # rejections vs. adjusted p -values; adjusted p -values vs. test statistics (“volcano” plots).
- **Software design.**
 - **Function closure.** Allow uniform data input for all MTPs; facilitate the extension of the package’s functionality, by implementing, for example, new types of test statistics.
 - **Class/method object-oriented programming.** Represent and operate on the results of multiple testing procedures.


Software Implementation: SAS Macros

SAS macros are available to compute the following components of a MTP:

- t -statistics;
- non-parametric bootstrap estimates of the null shift and scale-transformed test statistics null distribution;
- adjusted p -values for the FWER-controlling single-step maxT procedure;
- adjusted p -values for the gFWER- and TPPFP-controlling augmentation procedures.

Author: M. D. Birkner.

URL: www.stat.berkeley.edu/~sandrine/MTBook.

 Dudoit and van der Laan (2007, Section 13.2), Birkner et al. (2005).

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