

# **ComparativeMarkerSelection Documentation**

Module name: ComparativeMarkerSelection

**Description:** Computes significance values for features using several metrics,

including FDR(BH), Q Value, FWER, Feature-Specific P-Value,

and Bonferroni.

Authors: Joshua Gould, Gad Getz, Stefano Monti, Alper Uzun (Broad

Institute) gp-help@broad.mit.edu

The ComparativeMarkerSelection module includes several approaches to determine the features that are most closely correlated with a class template and the significance of that correlation. The module outputs a file containing the following columns:

- 1. **Rank** The rank of the feature within the dataset based on the value of the test statistic. If a two-sided p-value is computed, the rank is with respect to the absolute value of the statistic.
- 2. **Feature** The feature name.
- 3. **Description** The description of the feature.
- 4. **Score** The value of the test statistic.
- 5. **Feature P** The feature-specific p-value based on permutation testing.
- 6. *Feature P Low* The estimated lower bound for the feature p-value.
- 7. **Feature P High** The estimated upper bound for the feature p-value.
- FDR (BH) An estimate of the false discovery rate by the Benjamini and Hochberg procedure (1). The FDR is the expected proportion of erroneous rejections among all rejections.
- 9. **Q Value** An estimate of the FDR using the procedure developed by Storey and Tibshirani (6).
- 10. **Bonferroni** The value of the Bonferroni correction applied to the feature specific p-value.
- 11. **maxT** The adjusted *p*-values for the maxT multiple testing procedure described in Westfall (7), which provides strong control of the FWER.
- 12. **FWER** (Family Wise Error Rate) The probability of at least one null hypothesis/feature having a score better than or equal to the observed one. This measure is not feature-specific.
- 13. **Fold Change** The class zero mean divided by the class one mean.
- 14. Class Zero Mean The class zero mean.
- 15. Class Zero Standard Deviation The class zero standard deviation.
- 16. Class One Mean The class one mean.
- 17. Class One Standard Deviation The class one standard deviation.
- 18. **k** If performing a two-sided test or a one-sided test for markers of class zero, the number of permuted scores greater than or equal to the observed score. If testing for markers of class one, then the number of permuted scores less than or equal to the observed score.

The results from the ComparativeMarkerSelection algorithm can be viewed with the ComparativeMarkerSelectionViewer.



## Parameters:

Name	Description		
input filename	The input fileres, .gct		
cls filename	The class filecls		
confound variable cls	The class file containing the confounding variablecls		
filename	If you specify a confounding variable class file, permutations shuffle		
	the phenotype labels only within the subsets defined by that class		
	file. For example, in Lu, Getz, et. al. (2005), to select features that		
	best distinguish tumors from normal samples on all tissue types,		
	tissue type is treated as the confounding variable. In this case, the		
	confounding variable class file lists each tissue type as a phenotype		
	and associates each sample with its tissue type. Consequently,		
	•	Selection performs permutations, it	
		labels only among samples with the same	
	tissue type.		
test direction	The test to perform (up-regulated for class 0, up-regulated for class		
	· ·	ComparativeMarkerSelection performs the	
	two-sided test.		
test statistic The statistic to use:			
	t-test		
	i-lest	$\mu_A - \mu_B$	
		$  \sigma_A^2 \sigma_B^2  $	
		\ +	
		V nA nB where	
		μ is the average	
		$\sigma$ is the standard deviation	
		n is the number of samples	
	t-test (median)	same as t-test, but uses median	
		rather than average	
	t-test (min std)	same as t-test, but enforces a	
		minimum value for σ (minimal	
		standard deviation)	
	t-test (median, min std)	same as t-test, but uses median	
		rather than average and enforces a	
		minimum value for σ (minimal	
		standard deviation)	
	SNR	$\mu_A - \mu_B$	
	(signal-to-noise ratio)	$\frac{1}{\sigma_{A} + \sigma_{B}}$	
		where	
		μ is the average	
		$\sigma$ is the standard deviation	
	SNR (median)	same as SNR, but uses median	
		rather than average	
	SNR (min std)	same as SNR, but enforces a	
		minimum value for σ (minimal	
		standard deviation)	
	SNR (median, min std)	same as SNR, but uses median	



		rather than average and enforces a minimum value for σ (minimal	
		standard deviation)	
	Paired t-test	computes a paired, 2-sample t-	
		statistic	
min std	Used only if <i>test statistic</i> includes the min std option. If $\sigma$ is less than		
	min std, σ is set to min std.		
number of	The number of permutations to perform (use 0 to calculate		
permutations	asymptotic p-values). The number of permutations you specify		
	depends on the number of hypotheses being tested and the		
	significance level that you want to achieve (3). The greater the		
	number of permutations, the more accurate the p value. Asymptotic		
	p values are calculated using the p value obtained from the standard		
	independent two-sample t-test.		
complete	Whether to perform all possible permutations. By default, complete		
	is set to no and <i>number of permutations</i> determines the number of		
	permutations performed. If you have a <b>small</b> number of samples,		
	you might want to perform all possible permutations.		
balanced	Whether to perform balanced permutations.		
random seed	The seed for the random number generator.		
smooth p values	Whether to smooth p-values by using the Laplace's Rule of		
	Succession. By default, sr	mooth p values is set to yes, which means	
	p-values are always less t	han 1.0 and greater than 0.0.	
phenotype test	phenotype test Tests to perform when cls file has more than two classe		
	versus-all, all pairs.		
	Note: The p-values obtain	ed from the one-versus-all comparison are	
	not fully corrected for mult	iple hypothesis testing.	
output file	The name of the output file		

### **Output Files:**

An odf file of type ComparativeMarkerSelection

#### References:

- 1. Benjamini, Y. and Hochberg, Y. (1995) Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. *Journal of the Royal Statistical Society. Series B (Methodological)*. **57**(1): p. 289-300.
- 2. Golub, T., Slonim, D. et al. (1999) Molecular Classification of Cancer: Class Discovery and Class Prediction by Gene Expression. *Science* **286**, 531-537.
- 3. Good, P. (2000) Permutation Tests: A Practical Guide for Testing Hypotheses, 2<sup>nd</sup> Ed., New York: Springer-Verlag
- 4. Gould J., Getz G., Monti S., Reich M., and Mesirov J.P. (2006) Comparative gene marker selection suite. *Bioinformatics* **22**, 1924-1925; doi:10.1093/bioinformatics/btl196.
- 5. Lu, J., Getz, G., Miska, E., et al. (2005) MicroRNA Expression Profiles Classify Human Cancers. *Nature* **435**, 834-838
- 6. Storey, J.D. and R. Tibshirani (2003) Statistical significance for genomewide studies. *PNAS*, **100**(16): p. 9440-9445.



7. Westfall, P.H. and S. S. Young (1993) Resampling-Based Multiple Testing: Examples and Methods for p-Value Adjustment. *Wiley Series in Probability and Statistics*. New York: Wiley.

## Platform dependencies:

Module type: Gene List Selection

CPU type: any
OS: any
Java JVM level: 1.5
Language: Java, R