# jtGWAS

Efficient Jonckheere-Terpstra Test Statistics

2017-08-14

## Outline

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Session Information

#### Introduction

- This document provides an example for using the jtGWAS package to calculate the Jonckheere-Terpstra test statistics for large data sets (multiple markers and genome-wide SNPs) commonly encountered in GWAS.
- ▶ The calculation of the standardized test statistic employs the null variance equation as defined by Hollander and Wolfe (1999, eq. 6.19) to account for ties in the data.
- ▶ The major algorithm in this package is written in C++, which is ported to R by Rcpp, to facilitate fast computation.
- Features of this package include:
  - 1 OpenMP supported parallelization
  - 2 Customized output of top *m* significant SNPs for each marker
  - 3  $O(N \times \log(N))$  computational complexity (where N is the number of the samples)

## itGWAS

```
res <- jtGWAS(X, G, outTopN=15, numThreads=1, standardized=TRUE)</pre>
```

#### Function aguments:

- X: Matrix of marker levels, with sample IDs as row names and marker IDs as column names.
- G: Matrix of genotypes, with sample IDs as row names and SNP IDs as column names.
- outTopN: Number of top statistics to return (i.e., the largest *n* standardized statistics). The default value is 15. If outTopN is set to NA, all results will be returned.
- numThreads: Number of threads to use for parallel computation. The default value is 1 (sequential computation).
- standardized: A boolean to specify whether to return standardized statistics or non-standardized statistics. The default value is TRUE, returning standardized statistics.

Users may wish to consider the dplyr::recode() function for converting non-numeric group indices into ordinal values for argument G.

#### Returned Values

#### Function returns:

- J: A matrix of standardized/non-standardized Jonckheere Terpstra test statistics, depending on option standardized, with column names from input X. If outTopN is not NA, results are sorted within each column.
- gSnipID: If outTopN is not NA, this is a matrix of column names from G associated with top standardized Jonckheere Terpstra test statistics from J. Otherwise this is an unsorted vector of column names from input G.

### Simulate Data

1 Define the number of markers, patients, and SNPs:

```
    num_patient
    <- 100</td>

    num_marker
    <- 4</td>

    num_SNP
    <- 50</td>
```

- 2 Create two matrices containing marker levels and genotype information.
  - X\_pat\_mak contains the patients' marker levels.
  - b. G\_pat\_SNP contains the patients' genotypes.

# Load Package

Load jtGWAS (after installing its dependent packages):

library(jtGWAS)

## **Example Execution**

```
JTStat <- jtGWAS(X_pat_mak, G_pat_SNP, outTopN=10)
summary(JTStat, marker2Print=1:4, SNP2Print=1:5)</pre>
```

```
##
##
                    Johckheere-Terpstra Test for Large Matrices
                       P-values for Top Standardized Statistics
##
                Mrk:1|
                                    Mrk:21
                                                        Mrk·31
                                                                            Mrk·41
        SNPID P-valuel
                            SNPID P-valuel
                                                SNPID P-valuel
                                                                    SNPTD P-value!
##
       SNP:35 1.7e-021
                          SNP:35 2.0e-02|
                                               SNP:20 1.9e-021
                                                                   SNP:46 1.2e-021
       SNP:17 1.7e-02
                          SNP:7 5.7e-02|
                                               SNP:49 3.5e-02|
                                                                   SNP:34 1.7e-02|
##
       SNP:27 7.0e-02| SNP:46 9.1e-02|
                                               SNP:26 3.8e-021
                                                                 SNP:47 3.7e-021
       SNP:28 7.0e-021
                         SNP:40 9.4e-021
                                               SNP:47 5.6e-021
                                                                   SNP:23 4.5e-021
##
##
       SNP:14 8.8e-021
                          SNP:29 1.5e-01|
                                               SNP:30 9.7e-021
                                                                   SNP:16 5.1e-021
```

# Example Execution: Statistics in the Summary

## summary(JTStat, marker2Print=1:4, SNP2Print=1:5, printP=FALSE)

##								
##								
##	Johckheere-Terpstra Test for Large Matrices							
##	* Top Standardized Statistics							
##								
##								
##		Mrk:1		Mrk:2		Mrk:3		Mrk:4
##								
##		J*	SNPID	J*	SNPID	J*	SNPID	J*
##								
##	SNP:35	-2.390	SNP:35	-2.331	SNP:20	2.350	SNP:46	2.505
##	SNP:17	-2.388	SNP:7	1.905	SNP:49	-2.106	SNP:34	2.396
##	SNP:27	-1.813	SNP:46	1.693	SNP:26	-2.072	SNP:47	-2.089
##	SNP:28	1.813	SNP:40	1.676	SNP:47	1.914	SNP:23	2.000
##	SNP:14	1.706	SNP:29	1.432	SNP:30	-1.662	SNP:16	1.955

# Example Execution: Sorting in the Summary

```
JTAll <- jtGWAS(X_pat_mak, G_pat_SNP, outTopN=NA)
summary(JTAll, marker2Print=1:4, SNP2Print=1:3)
summary(JTAll, marker2Print=1:4, outTopN=3)</pre>
```

```
##
         Johckheere-Terpstra Test
##
##
       P-values Based on Standardized Statistics
##
             Mrk:1
                       Mrk:2
                                 Mrk:3
##
                                           Mrk:4
## SNP:1 0.2931953 0.7711424 0.5877522 0.1610595
## SNP:2 0.9120503 0.6085816 0.8017917 0.3169385
## SNP:3 0.5081104 0.3280014 0.5546797 0.2823776
##
##
                     Johckheere-Terpstra Test for Large Matrices
                       P-values for Top Standardized Statistics
                 Mrk·11
         SNPID P-valuel
                             SNPID P-value
                                                 SNPID P-valuel
                                                                     SNPID P-valuel
        SNP:35 1.7e-02|
                            SNP:35 2.0e-02|
                                                SNP:20 1.9e-02|
                                                                    SNP:46 1.2e-02
##
        SNP:17 1.7e-02|
                           SNP:7 5.7e-02|
                                                SNP:49 3.5e-021
                                                                    SNP:34 1.7e-02
        SNP:27 7.0e-02|
                          SNP:46 9.1e-02|
                                                SNP:26 3.8e-021
                                                                    SNP:47 3.7e-02
```

## References

Hollander, M. and Wolfe, D. A., *Nonparametric Statistical Methods*. New York, Wiley, 2nd edition, 1999.

## Session Information

- ► R version 3.3.3 (2017-03-06), x86\_64-pc-linux-gnu
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- ▶ Other packages: jtGWAS 1.5.1, knitr 1.13
- ▶ Loaded via a namespace (and not attached): Rcpp 0.12.4, evaluate 0.8, formatR 1.2.1, highr 0.5.1, magrittr 1.5, stringi 1.0-1, stringr 1.0.0, tools 3.3.3

```
## [1] "Start Time Mon Aug 14 13:53:03 2017"
## [1] "End Time Mon Aug 14 13:53:03 2017"
```