Pleiotropy for quantitive traits: pleio

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Example Data

The pleio package contains a pre-made simulated dataset with multiple quantitative traits simulated from a multivariate normal distribution with common correlation structure, correlation of 0.5, and genotypes simulated based on minor allele frequency of 0.2, and assumes that the regression coefficients (beta's) for traits 2 and 3 have non-zero values, while all other traits are not associated with dose of minor allele.

Here, we load the simulated dataset and show

```
## load package and dataset
require(pleio)
## Loading required package: pleio
## Loading required package: Matrix
data(pleio.qsim)
## preview simulated data
head(y)
                t1
                            t2
                                       t3
                                                  t4
                                                              t5
                                                                         t6
## [1,] -0.6795023
                   1.07928655 -0.3447868
                                           0.7757427
                                                      0.2441987
                                                                 0.2849724
## [2,] -0.5310806  0.06419289 -0.8963973
                                           0.2715258 -0.4579625 -1.2279082
## [3,]
        1.0244553 1.18181268
                               0.1977245
                                           0.3652163
                                                      0.3497826
## [4,] -0.3054566 -0.49250649
                                1.0348276 -0.1683192
                                                      1.8157372
## [5,] -0.4246874 -1.57093941 -0.3505441
                                           0.7817763 -2.4317261 -1.0474812
## [6,]
        2.0213843 1.45198074 0.2430903
                                           2.2323818 1.2128110
table(geno)
## geno
## 312 170
            18
```

Sequential Pleiotropy Tests

The pleio.q.sequential function is a high-level way to perform sequential tests of the number of traits (and which traits) are associated with a genotype. The algorithm starts with testing the usual multivariate null hypthothesis that all beta's are zero. If this test rejects, because the p-value is less than a user-spiecifed threshold, then allow one beta to be non-zero in order to test whether the remaining beta's = 0. If the test of one non-zero beta rejects, then allow two non-zero beta's, and continue this sequential testing until the p-value for a test is greater than the specified threshold. When the sequential testing stops, one can conclude

that the final model contains the non-zero beta's, while all other beta's are inferred to be zero. Below we run two functions, *pleio.q.fit*, which performs pre-calculations on the models to be tested, and *pleio.q.sequential*, which performs the sequential pleiotropy tests.

The final result lists the indices of the non-zero beta's (the indices of the traits associated with a genotype), and the p-value that tests the fit of the final model. A p-value greater than the threshold is expected for the final model, showing that the final model fits the data well. For this example, the sequential approach stopped at 2 traits because the p-value is greater than the *pval.threshold* argument given of 0.05.

```
fit <- pleio.q.fit(y, geno)

test.seq <- pleio.q.sequential(fit, pval.threshold=.05)
test.seq

## $pval
## [1] 0.2744734
##
## $index.beta
## [1] 2 3</pre>
```

Equivalent Steps to Sequential Fit

The sequential steps above can be performed with more user control using pleio.q.test, with count.nonzero.beta as the number of non-zero betas for the null hypothesis. The result of pleio.q.test contains the global test statistic, degrees of freedom (df), p-value for testing the model, the index of the non-zero beta's in the model, and a data.frame called "tests" that contains the tests performed for the null hypothesis models (i.e., the indices of the non-zero beta's and the corresponding statistic, tk, for each model). For m traits, and k = count.nonzero.beta, there are m-choose-k models in the set that are considered in the null hypothesis, and the minimum tk test statistic over the set provides the global test statistic reported.

```
test0 <- pleio.q.test(fit, count.nonzero.beta = 0)
test0</pre>
```

```
## $stat
   [1] 37.08576
##
##
## $pval
## [1] 1.694389e-06
##
## $df
## [1] 6
##
## $index.nonzero.beta
## [1] 0
##
## $tests
##
     index.1
                     t.k
            0 37.08576
## 1
test1 <- pleio.q.test(fit, count.nonzero.beta = 1)</pre>
test1
```

```
## $stat
## [1] 23.53879
##
## $pval
## [1] 0.000266202
##
## $df
## [1] 5
##
## $index.nonzero.beta
## [1] 3
##
## $tests
## index.1
## 1
         1 32.44700
          2 24.26597
## 2
## 3
        3 23.53879
## 4
         4 28.02627
## 5
         5 37.01172
## 6
          6 36.32446
test2 <- pleio.q.test(fit, count.nonzero.beta = 2)</pre>
test2
## $stat
## [1] 5.1274
##
## $pval
## [1] 0.2744734
##
## $df
## [1] 4
##
## $index.nonzero.beta
## [1] 2 3
## $tests
##
     index.1 index.2 tk
## 1
     1 2 21.43943
## 2
          1
                 3 21.50900
                 4 20.53175
## 3
           1
                 5 32.01780
## 4
           1
## 5
         1
                 6 31.14303
## 6
           2
                 3 5.12740
          2
                 4 17.74535
## 7
## 8
          2
                 5 24.14999
## 9
          2
                 6 24.26479
## 10
         3
                 4 17.37530
## 11
          3
                 5 23.31140
## 12
          3
                6 23.24654
## 13
          4
                 5 27.59217
           4
                  6 25.76981
## 14
## 15
           5
                  6 36.13547
```