

ASSET(Association analysis for SubSETs) Package

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Introduction

The ASSET package consists of two main functions: (1) h.traits and (2) h.types. The function h.traits is suitable for conducting meta-analysis of possibly different traits when summary level data are available from individual studies. The function allows for correlation among different studies/traits, which, for example, may arise due to shared subjects across studies. This function can also be used to conduct "meta-analysis" across multiple correlated traits on the same individuals by appropriately specifying the correlation matrix for the multivariate trait. Input arguments to this function are vectors/matrices of the estimated log-odds ratios, standard errors and number of cases and controls for each SNP and study. The function h.types is suitable for analysis of case-control studies when cases consist of distinct disease subtypes. This function assumes individual level data are available. The main input argument for h.types is a data frame containing the SNP variables, response variable and covariates for all subjects.

```
> library(ASSET)
```

Examples of h.traits

Get the path to the data.

```
> datafile <- system.file("sampleData", "vdata.rda", package="ASSET")
```

Load the data frames. There are 4 data frames, data1 - data4 for the 4 independent studies. Each study has the SNPs SNP1-SNP3 genotyped, and information on each subject's age and case-control status. Each SNP is coded as the number of copies of the minor allele or NA for missing genotypes.

```
> load(datafile)
> data1[1:5, ]
```

	CC	AGE	SNP1	SNP2	SNP3
456	1	70	1	0	0
457	1	55	1	0	0
458	1	48	0	0	1
459	1	72	1	0	2
460	1	74	2	0	0

```

> SNPs    <- paste("SNP", 1:3, sep="")
> nSNP   <- length(SNPs)
> studies <- paste("STUDY", 1:4, sep="")
> nStudy  <- length(studies)

```

Let us determine the number of non-missing cases and controls for each SNP and study.

```

> case    <- matrix(data=NA, nrow=nSNP, ncol=nStudy)
> control <- matrix(data=NA, nrow=nSNP, ncol=nStudy)
> for (i in 1:nStudy) {
+   data <- eval(parse(text=paste("data", i, sep="")))
+   caseVec <- data[, "CC"] == 1
+   controlVec <- !caseVec
+   for (j in 1:nSNP) {
+     temp <- !is.na(data[, SNPs[j]])
+     case[j, i] <- sum(caseVec & temp, na.rm=TRUE)
+     control[j, i] <- sum(controlVec & temp, na.rm=TRUE)
+   }
+ }
> case
      [,1] [,2] [,3] [,4]
[1,] 1897 1363 1714 686
[2,] 1909 1369 1726 691
[3,] 1875 1341 1732 696

> control
      [,1] [,2] [,3] [,4]
[1,] 1955 1802 1262 667
[2,] 1955 1773 1268 670
[3,] 1925 749 1269 674

```

Run a logistic regression for each SNP and study

```

> beta  <- matrix(data=NA, nrow=nSNP, ncol=nStudy)
> sigma <- matrix(data=NA, nrow=nSNP, ncol=nStudy)
> for (i in 1:nStudy) {
+   data <- eval(parse(text=paste("data", i, sep="")))
+   for (j in 1:nSNP) {
+     data[, "SNP"] <- data[, SNPs[j]]
+     fit <- glm(CC ~ AGE + SNP, data=data, family=binomial())
+     coef <- summary(fit)$coefficients
+     beta[j, i] <- coef["SNP", 1]
+     sigma[j, i] <- coef["SNP", 2]
+   }
+ }
> beta
      [,1]      [,2]      [,3]      [,4]
[1,] 0.30837615 0.09041508 0.1799979 0.13116360
[2,] 0.09311754 0.20472698 0.1465665 0.05729745
[3,] -0.08212701 0.08909210 -0.0621090 0.01181724

```

```

> sigma
      [,1]      [,2]      [,3]      [,4]
[1,] 0.04637132 0.05410822 0.05931264 0.07970842
[2,] 0.10214703 0.08211686 0.09299885 0.11889584
[3,] 0.04954003 0.07202424 0.05736282 0.08468467

```

>

Call the h.traits function. Since the studies are independent, we do not need to specify the cor option.

```
> res <- h.traits(SNPs, studies, beta, sigma, case, control, meta=TRUE)
```

Compute a summary table. Notice that in the Subset.2sided results, the first 2 SNPs have missing values for OR.2, CI.low.2, and CI.high.2 since the estimated betas were all positive for these SNPs.

```
> h.summary(res)
```

```
$Meta
  SNP      Pvalue      OR CI.low CI.high
1 SNP1 3.268265e-12 1.218  1.216   1.220
2 SNP2 3.666743e-03 1.150  1.145   1.156
3 SNP3 2.994911e-01 0.968  0.967   0.970

$Subset.1sided
  SNP      Pvalue      OR CI.low CI.high          Pheno
1 SNP1 2.202379e-11 1.268  1.183   1.359 STUDY1,STUDY3,STUDY4
2 SNP2 3.389452e-02 1.196  1.014   1.412     STUDY2,STUDY3
3 SNP3 3.190294e-01 0.929  0.804   1.074     STUDY1,STUDY3

$Subset.2sided
  SNP      Pvalue      Pvalue.1    Pvalue.2      OR.1 CI.low.1 CI.high.1      OR.2
1 SNP1 5.154931e-11 5.154931e-11 1.0000000 1.268   1.181   1.361   NA
2 SNP2 5.527301e-02 5.527301e-02 1.0000000 1.196   0.996   1.437   NA
3 SNP3 1.582707e-01 3.604528e-01 0.1020498 1.093   0.903   1.323  0.929
  CI.low.2 CI.high.2          Pheno.1      Pheno.2
1       NA        NA STUDY1,STUDY3,STUDY4
2       NA        NA     STUDY2,STUDY3
3  0.851    1.015     STUDY2 STUDY1,STUDY3
```

Instead of searching over all possible subsets, let us define our own subset function to determine which nsubsets to search over. We will only consider subsets where the first m traits are in the subset ($m = 1, 2, \dots$). The DLM p-value will also be computed using only these subsets.

```
> sub.def <- function(logicalVec) {
+   sum <- sum(logicalVec)
+   ret <- all(logicalVec[1:sum])
+   ret
+ }
```

Call the h.traits function with the zmax.args pval.args options defined

```
> res <- h.traits(SNPs, studies, beta, sigma, case, control, meta=TRUE,
+                   zmax.args=list(sub.def=sub.def), pval.args=list(sub.def=sub.def))

> h.summary(res)

$Meta
  SNP      Pvalue      OR CI.low CI.high
1 SNP1 3.268265e-12 1.218  1.216   1.220
2 SNP2 3.666743e-03 1.150  1.145   1.156
3 SNP3 2.994911e-01 0.968  0.967   0.970

$Subset.1sided
  SNP      Pvalue      OR CI.low CI.high          Pheno
1 SNP1 2.096558e-11 1.218  1.150   1.290 STUDY1,STUDY2,STUDY3,STUDY4
2 SNP2 1.430355e-02 1.169  1.032   1.325     STUDY1,STUDY2,STUDY3
3 SNP3 2.477190e-01 0.921  0.801   1.059     STUDY1

$Subset.2sided
  SNP      Pvalue      Pvalue.1    Pvalue.2    OR.1 CI.low.1 CI.high.1  OR.2
1 SNP1 1.713498e-11 1.713498e-11 1.00000000 1.218   1.150   1.290   NA
2 SNP2 9.590357e-03 9.590357e-03 1.00000000 1.169   1.039   1.316   NA
3 SNP3 6.352379e-02 2.085630e-01 0.05586139 1.093   0.951   1.256  0.929
  CI.low.2 CI.high.2          Pheno.1    Pheno.2
1       NA           NA STUDY1,STUDY2,STUDY3,STUDY4
2       NA           NA     STUDY1,STUDY2,STUDY3
3  0.862   1.002           NA           STUDY2 STUDY1,STUDY3
```

Session Information

```
> sessionInfo()

R version 3.0.2 (2013-09-25)
Platform: x86_64-unknown-linux-gnu (64-bit)

locale:
[1] LC_CTYPE=en_US.UTF-8        LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8         LC_COLLATE=C
[5] LC_MONETARY=en_US.UTF-8     LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8        LC_NAME=C
[9] LC_ADDRESS=C                LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:
[1] grid      stats     graphics  grDevices utils     datasets  methods 
[8] base

other attached packages:
[1] ASSET_1.0.0 rmeta_2.16  msm_1.2    MASS_7.3-29
```

```
loaded via a namespace (and not attached):  
[1] mvtnorm_0.9-9996 splines_3.0.2    survival_2.37-4  tools_3.0.2
```