

Package ‘qfratio’

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Type Package

Title Moments of Ratios of Quadratic Forms Using Recursion

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Description Evaluates moments of ratios (and products) of quadratic forms in normal variables, specifically using recursive algorithms developed by Bao and Kan (2013) <[doi:10.1016/j.jmva.2013.03.002](https://doi.org/10.1016/j.jmva.2013.03.002)> and Hillier et al. (2014) <[doi:10.1017/S0266466613000364](https://doi.org/10.1017/S0266466613000364)>. Originally developed as a supplement to Watanabe (2022) <[doi:10.1101/2022.11.02.514929](https://doi.org/10.1101/2022.11.02.514929)> for evaluating average evolvability measures in evolutionary quantitative genetics, but can be used for a broader class of moments. Generating functions for these moments are also closely related to the top-order zonal and invariant polynomials of matrix arguments.

License GPL (>= 3)

URL <https://github.com/watanabe-j/qfratio>

BugReports <https://github.com/watanabe-j/qfratio/issues>

Imports Rcpp, MASS

LinkingTo Rcpp, RcppEigen

Suggests gsl, mvtnorm, graphics, stats, testthat (>= 3.0.0), rlang (>= 0.4.7), knitr, rmarkdown

Encoding UTF-8

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Config/testthat/edition 3

VignetteBuilder knitr, rmarkdown

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Author Junya Watanabe [aut, cre, cph]
(<<https://orcid.org/0000-0002-9810-5286>>)

Maintainer Junya Watanabe <jw2098@cam.ac.uk>

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qfratio-package	<i>qfratio: Moments of Ratios of Quadratic Forms Using Recursion</i>
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Description

This package is for evaluating moments of ratios (and products) of quadratic forms in normal variables, specifically using recursive algorithms developed by Bao et al. (2013) and Hillier et al. (2014) (see also Smith, 1989, 1993; Hillier et al., 2009). It was originally developed as a supplement to Watanabe (2022) for evaluating average evolvability measures in evolutionary quantitative genetics, but can be used for a broader class of moments.

Details

The primary front-end functions of this package are `qfrm()` and `qfmrn()` for evaluating moments of ratios of quadratic forms. These pass arguments to one of the several “internal” (though exported) functions which do actual calculations, depending on the argument matrices and exponents. In addition, there are a few functions to calculate moments of products of quadratic forms (integer exponents only; `qfpm`).

There are many internal functions for calculating coefficients in power-series expansion of generating functions for these moments (`d1_i`, `d2_ij`, `d3_ijk`, `dtil2_pq`) using “super-short” recursions (Bao and Kan, 2013; Hillier et al. 2014). Some of these coefficients are related to the top-order zonal and invariant polynomials of matrix arguments.

See package vignette (`vignette("qfratio")`) for more details.

The DESCRIPTION file:

```
Package:          qfratio
Type:             Package
Title:            Moments of Ratios of Quadratic Forms Using Recursion
Version:          1.0.1
Date:             2023-03-31
Authors@R:       c(person("Junya", "Watanabe", email = "jw2098@cam.ac.uk", role = c("aut", "cre", "cph"), comment = "I am the author, creator, and maintainer of this package.")
Description:      Evaluates moments of ratios (and products) of quadratic forms in normal variables, specifically using the method of moments.
License:          GPL (>= 3)
URL:              https://github.com/watanabe-j/qfratio
BugReports:       https://github.com/watanabe-j/qfratio/issues
Imports:          Rcpp, MASS
LinkingTo:        Rcpp, RcppEigen
Suggests:         gsl, mvtnorm, graphics, stats, testthat (>= 3.0.0), rlang (>= 0.4.7), knitr, rmarkdown
Encoding:         UTF-8
Roxygen:          list(markdown = TRUE)
RoxygenNote:     7.2.3
Config/testthat/edition: 3
VignetteBuilder: knitr, rmarkdown
Author:           Junya Watanabe [aut, cre, cph] (<https://orcid.org/0000-0002-9810-5286>)
Maintainer:      Junya Watanabe <jw2098@cam.ac.uk>
```

Index of help topics:

Ap_int_cmE	Internal C++ functions
KiK	Matrix square root and generalized inverse
S_fromUL	Make covariance matrix from eigenstructure
a1_pk	Recursion for a_p,k
d1_i	Coefficients in polynomial expansion of generating function-single matrix
d2_ij	Coefficients in polynomial expansion of generating function-for ratios with two matrices
d3_ijk	Coefficients in polynomial expansion of generating function-for ratios with three matrices
dtil2_pq	Coefficients in polynomial expansion of generating function-for products
hgs	Calculate hypergeometric series
is_diagonal	Is this matrix diagonal?
iseq	Are these vectors equal?
new_qfrm	Construct qfrm object
print.qfrm	Methods for qfrm and qfpm objects
qfmrn	Moment of multiple ratio of quadratic forms in normal variables

qfpm	Moment of (product of) quadratic forms in normal variables
qfratio-package	qfratio: Moments of Ratios of Quadratic Forms Using Recursion
qfrm	Moment of ratio of quadratic forms in normal variables
rqfr	Monte Carlo sampling of ratio/product of quadratic forms
sum_counterdiag	Summing up counter-diagonal elements
tr	Matrix trace function

Author/Maintainer

Junya Watanabe jw2098@cam.ac.uk

References

- Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:[10.1016/j.jmva.2013.03.002](https://doi.org/10.1016/j.jmva.2013.03.002).
- Hillier, G., Kan, R. and Wang, X. (2009) Computationally efficient recursions for top-order invariant polynomials with applications. *Econometric Theory*, **25**, 211–242. doi:[10.1017/S0266466608090075](https://doi.org/10.1017/S0266466608090075).
- Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:[10.1017/S0266466613000364](https://doi.org/10.1017/S0266466613000364).
- Smith, M. D. (1989) On the expectation of a ratio of quadratic forms in normal variables. *Journal of Multivariate Analysis*, **31**, 244–257. doi:[10.1016/0047259X\(89\)900651](https://doi.org/10.1016/0047259X(89)900651).
- Smith, M. D. (1993) Expectations of ratios of quadratic forms in normal variables: evaluating some top-order invariant polynomials. *Australian Journal of Statistics*, **35**, 271–282. doi:[10.1111/j.1467-842X.1993.tb01335.x](https://doi.org/10.1111/j.1467-842X.1993.tb01335.x).
- Watanabe, J. (2022) Exact expressions and numerical evaluation of average evolvability measures for characterizing and comparing **G** matrices. *bioRxiv* preprint, 2022.11.02.514929. doi:[10.1101/2022.11.02.514929](https://doi.org/10.1101/2022.11.02.514929).

See Also

- [qfrm](#): Moment of simple ratio of quadratic forms
- [qfmr](#): Moment of multiple ratio of quadratic forms
- [qfpm](#): Moment of product of quadratic forms
- [rqfr](#): Monte Carlo sampling of ratio/product of quadratic forms

Examples

```
## Symmetric matrices
nv <- 4
A <- diag(nv:1)
B <- diag(sqrt(1:nv))
D <- diag((1:nv)^2 / nv)
```

```

mu <- nv:1 / nv
Sigma <- matrix(0.5, nv, nv)
diag(Sigma) <- 1

## Expectation of (x^T A x)^2 / (x^T x)^2 where x ~ N(0, I)
qfrm(A, p = 2)
## And a Monte Carlo mean of the same
mean(rqfr(1000, A = A, p = 2))

## Expectation of (x^T A x)^1/2 / (x^T x)^1/2 where x ~ N(0, I)
(res1 <- qfrm(A, p = 1/2))
plot(res1)
## A Monte Carlo mean
mean(rqfr(1000, A = A, p = 1/2))

## (x^T A x)^2 / (x^T B x)^3 where x ~ N(mu, Sigma)
(res2 <- qfrm(A, B, p = 2, q = 3, mu = mu, Sigma = Sigma))
plot(res2)
## A Monte Carlo mean
mean(rqfr(1000, A = A, B = B, p = 2, q = 3, mu = mu, Sigma = Sigma))

## Expectation of (x^T A x)^2 / (x^T B x) (x^T x) where x ~ N(0, I)
(res3 <- qfmr(A, B, p = 2, q = 1, r = 1))
plot(res3)
## A Monte Carlo mean
mean(rqfmr(1000, A = A, B = B, p = 2, q = 1, r = 1))

## Expectation of (x^T A x)^2 where x ~ N(0, I)
qfm_Ap_int(A, 2)
## A Monte Carlo mean
mean(rqfp(1000, A = A, p = 2, q = 0, r = 0))

## Expectation of (x^T A x) (x^T B x) (x^T D x) where x ~ N(mu, I)
qfpm_ABDpqr_int(A, B, D, 1, 1, 1, mu = mu)
## A Monte Carlo mean
mean(rqfp(1000, A = A, B = B, D = D, p = 1, q = 1, r = 1, mu = mu))

```

a1_pk

Recursion for a_{p,k}

Description

a1_pk() is an internal function to calculate $a_{p,k}$ ($a_{r,l}$ in Hillier et al. 2014; eq. 24), which is used in the calculation of the moment of such a ratio of quadratic forms in normal variables where the denominator matrix is identity.

Usage

```
a1_pk(L, mu = rep.int(0, n), m = 10L)
```

Arguments

L	Eigenvalues of the argument matrix; vector of λ_i
mu	Mean vector μ for \mathbf{x}
m	Scalar to specify the desired order

Details

This function implements the super-short recursion described in Hillier et al. (2014 eqs. 31–32). Note that $w_{r,i}$ there should be understood as $w_{r,l,i}$ with the index l fixed for each $a_{r,l}$.

See Also

[qfrm_ApIq_int\(\)](#), in which this function is used (for noncentral cases only)

Ap_int_cmE

Internal C++ functions

Description

These are internal C++ functions called from corresponding R functions when `use_cpp = TRUE`. Direct access by the user is **not** assumed. All parameters are assumed to be appropriately structured.

Usage

```
Ap_int_cmE(A, p = 1, thr_margin = 100)
Ap_int_nmE(A, mu, p = 1, thr_margin = 100)
ABpq_int_cvE(LA, LB, p = 1, q = 1, thr_margin = 100)
ABpq_int_cmE(A, LB, p = 1, q = 1, thr_margin = 100)
ABpq_int_nvE(LA, LB, mu, p = 1, q = 1)
ABpq_int_nmE(A, LB, mu, p = 1, q = 1)
ABDpqr_int_cvE(LA, LB, LD, p = 1, q = 1, r = 1, thr_margin = 100)
ABDpqr_int_cmE(A, LB, D, p = 1, q = 1, r = 1, thr_margin = 100)
ABDpqr_int_nvE(LA, LB, LD, mu, p = 1, q = 1, r = 1)
ABDpqr_int_nmE(A, LB, D, mu, p = 1, q = 1, r = 1)
ApIq_int_cmE(A, p = 1, q = 1, thr_margin = 100)
```

```
ApIq_int_nmE(A, mu, p = 1, q = 1, thr_margin = 100)
```

```
ApIq_npi_cvE(  
  LA,  
  bA,  
  p = 1,  
  q = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)
```

```
ApIq_npi_nvE(  
  LA,  
  UA,  
  bA,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBq_int_cvE(  
  LA,  
  LB,  
  bB,  
  p = 1,  
  q = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)
```

```
ApBq_int_cmE(  
  A,  
  LA,  
  UA,  
  LB,  
  bB,  
  p = 1,  
  q = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)
```

```
ApBq_int_nvE(  
  LA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)
```

```
ApBq_int_nmE(  
  A,  
  LA,  
  UA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)
```

```
ApBq_npi_cvE(  
  LA,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBq_npi_cmE(  
  A,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L
```



```
)  
  
ApBq_npi_nvE(  
  LA,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)  
  
ApBq_npi_nmE(  
  A,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)  
  
ApBIqr_int_cvE(  
  LA,  
  LB,  
  bB,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100  
)  
  
ApBIqr_int_cmE(  
  A,  
  LA,  
  UA,  
  LB,  
  bB,  
  p = 1,  
  q = 1,
```

```
    r = 1,  
    m = 100L,  
    error_bound = TRUE,  
    thr_margin = 100  
)
```

```
ApBIqr_int_nvE(  
  LA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBIqr_int_nmE(  
  A,  
  LA,  
  UA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
ApBIqr_npi_cvE(  
  LA,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBIqr_npi_cmE(  
  A,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
ApBIqr_npi_nvE(  
  LA,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBIqr_npi_nmE(  
  A,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
IpBDqr_gen_cvE(  
  LB,  
  LD,  
  bB,  
  bD,  
  p = 1,
```

```
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
IpBDqr_gen_cmE(  
  LB,  
  D,  
  bB,  
  bD,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
IpBDqr_gen_nvE(  
  LB,  
  LD,  
  bB,  
  bD,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
IpBDqr_gen_nmE(  
  LB,  
  D,  
  bB,  
  bD,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
ApBDqr_int_cvE(  
  LA,  
  LB,  
  LD,  
  bB,  
  bD,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBDqr_int_cmE(  
  A,  
  LB,  
  D,  
  bB,  
  bD,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
ApBDqr_int_nvE(  
  LA,  
  LB,  
  LD,  
  bB,  
  bD,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBDqr_int_nmE(  
  A,  
  LB,  
  D,  
  bB,
```

```
    bD,  
    mu,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 0L  
)
```

```
ApBDqr_npi_cvE(  
    LA,  
    LB,  
    LD,  
    bA,  
    bB,  
    bD,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 0L  
)
```

```
ApBDqr_npi_cmE(  
    A,  
    LB,  
    D,  
    bA,  
    bB,  
    bD,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 0L  
)
```

```
ApBDqr_npi_nvE(  
    LA,  
    LB,  
    LD,  
    bA,  
    bB,  
    bD,  
    mu,
```

```
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
ApBDqr_npi_nmE(  
A,  
LB,  
D,  
bA,  
bB,  
bD,  
mu,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 0L  
)
```

```
ApIq_npi_nvEc(  
LA,  
UA,  
bA,  
mu,  
p = 1,  
q = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
ApBq_npi_cvEc(  
LA,  
LB,  
bA,  
bB,  
p = 1,  
q = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
ApBq_npi_cmEc(  
  A,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)  
  
ApBq_npi_nvEc(  
  LA,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)  
  
ApBq_npi_nmEc(  
  A,  
  LB,  
  bA,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)  
  
ApBIqr_int_nvEc(  
  LA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  error_bound = TRUE,
```



```
    thr_margin = 100,  
    nthreads = 1L  
)  
  
ApBIqr_int_nmEc(  
  A,  
  LA,  
  UA,  
  LB,  
  bB,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  error_bound = TRUE,  
  thr_margin = 100,  
  nthreads = 0L  
)  
  
ApBIqr_npi_cvEc(  
  LA,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)  
  
ApBIqr_npi_cmEc(  
  A,  
  LB,  
  bA,  
  bB,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)  
  
ApBIqr_npi_nvEc(  
  LA,
```

```
LB,  
bA,  
bB,  
mu,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
ApBIqr_npi_nmEc(  
A,  
LB,  
bA,  
bB,  
mu,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 0L  
)
```

```
IpBDqr_gen_cvEc(  
LB,  
LD,  
bB,  
bD,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
IpBDqr_gen_cmEc(  
LB,  
D,  
bB,  
bD,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,
```

```
    thr_margin = 100,  
    nthreads = 0L  
)
```

```
IpBDqr_gen_nvEc(  
  LB,  
  LD,  
  bB,  
  bD,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
IpBDqr_gen_nmEc(  
  LB,  
  D,  
  bB,  
  bD,  
  mu,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 0L  
)
```

```
ApBDqr_int_cvEc(  
  LA,  
  LB,  
  LD,  
  bB,  
  bD,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  thr_margin = 100,  
  nthreads = 1L  
)
```

```
ApBDqr_int_cmEc(  
  A,
```

```
LB,  
D,  
bB,  
bD,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 0L  
)
```

```
ApBDqr_int_nvEc(  
LA,  
LB,  
LD,  
bB,  
bD,  
mu,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 1L  
)
```

```
ApBDqr_int_nmEc(  
A,  
LB,  
D,  
bB,  
bD,  
mu,  
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 0L  
)
```

```
ApBDqr_npi_cvEc(  
LA,  
LB,  
LD,  
bA,  
bB,
```

```
    bD,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 1L  
)
```

```
ApBDqr_npi_cmEc(  
    A,  
    LB,  
    D,  
    bA,  
    bB,  
    bD,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 0L  
)
```

```
ApBDqr_npi_nvEc(  
    LA,  
    LB,  
    LD,  
    bA,  
    bB,  
    bD,  
    mu,  
    p = 1,  
    q = 1,  
    r = 1,  
    m = 100L,  
    thr_margin = 100,  
    nthreads = 1L  
)
```

```
ApBDqr_npi_nmEc(  
    A,  
    LB,  
    D,  
    bA,  
    bB,  
    bD,  
    mu,
```

```
p = 1,  
q = 1,  
r = 1,  
m = 100L,  
thr_margin = 100,  
nthreads = 0L  
)
```

```
ApIq_npi_nvEl(  
  LA,  
  UA,  
  bA,  
  mu,  
  p = 1L,  
  q = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 1L  
)
```

```
ApBq_npi_cvEl(  
  LA,  
  LB,  
  bA,  
  bB,  
  p = 1L,  
  q = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 1L  
)
```

```
ApBq_npi_cmEl(  
  A,  
  LB,  
  bA,  
  bB,  
  p = 1L,  
  q = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 0L  
)
```

```
ApBq_npi_nvEl(  
  LA,  
  LB,  
  bA,
```

```
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBq_npi_nmEl(  
    A,  
    LB,  
    bA,  
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
ApBIqr_int_nvEl(  
    LA,  
    LB,  
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    error_bound = TRUE,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBIqr_int_nmEl(  
    A,  
    LA,  
    UA,  
    LB,  
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    error_bound = TRUE,  
)
```

```
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
ApBIqr_npi_cvEl(  
    LA,  
    LB,  
    bA,  
    bB,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBIqr_npi_cmEl(  
    A,  
    LB,  
    bA,  
    bB,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
ApBIqr_npi_nvEl(  
    LA,  
    LB,  
    bA,  
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBIqr_npi_nmEl(  
    A,  
    LB,  
    bA,
```



```
    bB,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
IpBDqr_gen_cvEl(  
    LB,  
    LD,  
    bB,  
    bD,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
IpBDqr_gen_cmEl(  
    LB,  
    D,  
    bB,  
    bD,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
IpBDqr_gen_nvEl(  
    LB,  
    LD,  
    bB,  
    bD,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
)  
  
IpBDqr_gen_nmEl(  
  LB,  
  D,  
  bB,  
  bD,  
  mu,  
  p = 1L,  
  q = 1L,  
  r = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 0L  
)  
  
ApBDqr_int_cvEl(  
  LA,  
  LB,  
  LD,  
  bB,  
  bD,  
  p = 1L,  
  q = 1L,  
  r = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 1L  
)  
  
ApBDqr_int_cmEl(  
  A,  
  LB,  
  D,  
  bB,  
  bD,  
  p = 1L,  
  q = 1L,  
  r = 1L,  
  m = 100L,  
  thr_margin = 100L,  
  nthreads = 0L  
)  
  
ApBDqr_int_nvEl(  
  LA,  
  LB,  
  LD,
```

```
    bB,  
    bD,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBDqr_int_nmEl(  
    A,  
    LB,  
    D,  
    bB,  
    bD,  
    mu,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 0L  
)
```

```
ApBDqr_npi_cvEl(  
    LA,  
    LB,  
    LD,  
    bA,  
    bB,  
    bD,  
    p = 1L,  
    q = 1L,  
    r = 1L,  
    m = 100L,  
    thr_margin = 100L,  
    nthreads = 1L  
)
```

```
ApBDqr_npi_cmEl(  
    A,  
    LB,  
    D,  
    bA,  
    bB,  
    bD,
```

```

    p = 1L,
    q = 1L,
    r = 1L,
    m = 100L,
    thr_margin = 100L,
    nthreads = 0L
)

ApBDqr_npi_nvEl(
  LA,
  LB,
  LD,
  bA,
  bB,
  bD,
  mu,
  p = 1L,
  q = 1L,
  r = 1L,
  m = 100L,
  thr_margin = 100L,
  nthreads = 1L
)

ApBDqr_npi_nmEl(
  A,
  LB,
  D,
  bA,
  bB,
  bD,
  mu,
  p = 1L,
  q = 1L,
  r = 1L,
  m = 100L,
  thr_margin = 100L,
  nthreads = 0L
)

rqfpE(nit, A, B, D, p, q, r, mu, Sigma)

```

Arguments

A, B, D	Argument matrices passed as Eigen::Matrix. Symmetry is assumed.
p, q, r	Exponents for A, B, and D. Passed as double or long double.
thr_margin	Optional argument to adjust the threshold for scaling. See “Scaling” in d1_i .
mu	Mean vector μ for x passed as Eigen::Array

LA, LB, LD	Eigenvalues of the argument matrices passed as Eigen::Array
bA, bB, bD	Scaling coefficients for A , B , and D . Passed as double or long double.
m	Integer to specify the order of polynomials at which the series expression is truncated. Passed as Eigen::Index (aka std::ptrdiff_t or long long int)
error_bound	bool to specify whether the error bound is returned
UA	Matrix whose columns are eigenvectors of A corresponding to LA. Passed as Eigen::Matrix.
nthreads	int to specify the number of threads in OpenMP-enabled functions. See “Multithreading” in qfrm .
nit	int to specify the number of iteration or sample size
Sigma	Covariance matrix Σ for x . Passed as Eigen::Matrix.

Details

At present, ApIq_int_nmE() calls the R function `gsl::hyperg_1F1()`, so will not be much faster than the R equivalent. Ideally, the C++ library GSL (or the like) should be used with **RcppGSL**, but this is not done to ensure portability.

rqpE uses `Rcpp::rnorm()`, which may not be particularly efficient.

Value

All return a list via `Rcpp::List` of the following (as appropriate):

- `$ans`: Exact moment, from double or long double
- `$ansseq`: Series for the moment, from Eigen::Array
- `$errseq`: Series of errors, from Eigen::Array
- `$twosided`: Logical, from bool
- `$diminished`: Logical, from bool

Functions

- `Ap_int_cmE()`: `qfm_Ap_int()`, central
- `Ap_int_nmE()`: `qfm_Ap_int()`, noncentral
- `ABpq_int_cvE()`: `qfpm_ABpq_int()`, central and vector
- `ABpq_int_cmE()`: `qfpm_ABpq_int()`, central and matrix
- `ABpq_int_nvE()`: `qfpm_ABpq_int()`, noncentral and vector
- `ABpq_int_nmE()`: `qfpm_ABpq_int()`, noncentral and matrix
- `ABDpqr_int_cvE()`: `qfpm_ABDpqr_int()`, central and vector
- `ABDpqr_int_cmE()`: `qfpm_ABDpqr_int()`, central and matrix
- `ABDpqr_int_nvE()`: `qfpm_ABDpqr_int()`, noncentral and vector
- `ABDpqr_int_nmE()`: `qfpm_ABDpqr_int()`, central and matrix
- `ApIq_int_cmE()`: `qfrm_ApIq_int()`, central

- ApIq_int_nmE(): qfrm_ApIq_int(), noncentral
- ApIq_npi_cvE(): qfrm_ApIq_npi(), central
- ApIq_npi_nvE(): qfrm_ApIq_npi(), noncentral
- ApBq_int_cvE(): qfrm_ApBq_int(), central and vector
- ApBq_int_cmE(): qfrm_ApBq_int(), central and matrix
- ApBq_int_nvE(): qfrm_ApBq_int(), noncentral and vector
- ApBq_int_nmE(): qfrm_ApBq_int(), noncentral and matrix
- ApBq_npi_cvE(): qfrm_ApBq_npi(), central and vector
- ApBq_npi_cmE(): qfrm_ApBq_npi(), central and matrix
- ApBq_npi_nvE(): qfrm_ApBq_npi(), noncentral and vector
- ApBq_npi_nmE(): qfrm_ApBq_npi(), noncentral and matrix
- ApBIqr_int_cvE(): qfmr_ApBIqr_int(), central and vector
- ApBIqr_int_cmE(): qfmr_ApBIqr_int(), central and matrix
- ApBIqr_int_nvE(): qfmr_ApBIqr_int(), noncentral and vector
- ApBIqr_int_nmE(): qfmr_ApBIqr_int(), noncentral and matrix
- ApBIqr_npi_cvE(): qfmr_ApBIqr_npi(), central and vector
- ApBIqr_npi_cmE(): qfmr_ApBIqr_npi(), central and matrix
- ApBIqr_npi_nvE(): qfmr_ApBIqr_npi(), noncentral and vector
- ApBIqr_npi_nmE(): qfmr_ApBIqr_npi(), noncentral and matrix
- IpBDqr_gen_cvE(): qfmr_IpBDqr_gen(), central and vector
- IpBDqr_gen_cmE(): qfmr_IpBDqr_gen(), central and matrix
- IpBDqr_gen_nvE(): qfmr_IpBDqr_gen(), noncentral and vector
- IpBDqr_gen_nmE(): qfmr_IpBDqr_gen(), noncentral and matrix
- ApBDqr_int_cvE(): qfmr_ApBDqr_int(), central and vector
- ApBDqr_int_cmE(): qfmr_ApBDqr_int(), central and matrix
- ApBDqr_int_nvE(): qfmr_ApBDqr_int(), noncentral and vector
- ApBDqr_int_nmE(): qfmr_ApBDqr_int(), noncentral and matrix
- ApBDqr_npi_cvE(): qfmr_ApBDqr_npi(), central and vector
- ApBDqr_npi_cmE(): qfmr_ApBDqr_npi(), central and matrix
- ApBDqr_npi_nvE(): qfmr_ApBDqr_npi(), noncentral and vector
- ApBDqr_npi_nmE(): qfmr_ApBDqr_npi(), noncentral and matrix
- ApIq_npi_nvEc(): qfrm_ApIq_npi(), noncentral, coefficient-wise scaling
- ApBq_npi_cvEc(): qfrm_ApBq_npi(), central and vector, coefficient-wise scaling
- ApBq_npi_cmEc(): qfrm_ApBq_npi(), central and matrix, coefficient-wise scaling
- ApBq_npi_nvEc(): qfrm_ApBq_npi(), noncentral and vector, coefficient-wise scaling
- ApBq_npi_nmEc(): qfrm_ApBq_npi(), noncentral and matrix, coefficient-wise scaling
- ApBIqr_int_nvEc(): qfmr_ApBIqr_int(), noncentral and vector, coefficient-wise scaling

- ApBIqr_int_nmEc(): qfmrn_ApBIqr_int(), noncentral and matrix, coefficient-wise scaling
- ApBIqr_npi_cvEc(): qfmrn_ApBIqr_npi(), central and vector, coefficient-wise scaling
- ApBIqr_npi_cmEc(): qfmrn_ApBIqr_npi(), central and matrix, coefficient-wise scaling
- ApBIqr_npi_nvEc(): qfmrn_ApBIqr_npi(), noncentral and vector, coefficient-wise scaling
- ApBIqr_npi_nmEc(): qfmrn_ApBIqr_npi(), noncentral and matrix, coefficient-wise scaling
- IpBDqr_gen_cvEc(): qfmrn_IpBDqr_gen(), central and vector, coefficient-wise scaling
- IpBDqr_gen_cmEc(): qfmrn_IpBDqr_gen(), central and matrix, coefficient-wise scaling
- IpBDqr_gen_nvEc(): qfmrn_IpBDqr_gen(), noncentral and vector, coefficient-wise scaling
- IpBDqr_gen_nmEc(): qfmrn_IpBDqr_gen(), noncentral and matrix, coefficient-wise scaling
- ApBDqr_int_cvEc(): qfmrn_ApBDqr_int(), central and vector, coefficient-wise scaling
- ApBDqr_int_cmEc(): qfmrn_ApBDqr_int(), central and matrix, coefficient-wise scaling
- ApBDqr_int_nvEc(): qfmrn_ApBDqr_int(), noncentral and vector, coefficient-wise scaling
- ApBDqr_int_nmEc(): qfmrn_ApBDqr_int(), noncentral and matrix, coefficient-wise scaling
- ApBDqr_npi_cvEc(): qfmrn_ApBDqr_npi(), central and vector, coefficient-wise scaling
- ApBDqr_npi_cmEc(): qfmrn_ApBDqr_npi(), central and matrix, coefficient-wise scaling
- ApBDqr_npi_nvEc(): qfmrn_ApBDqr_npi(), noncentral and vector, coefficient-wise scaling
- ApBDqr_npi_nmEc(): qfmrn_ApBDqr_npi(), noncentral and matrix, coefficient-wise scaling
- ApIq_npi_nvEl(): qfrm_ApIq_npi(), noncentral
- ApBq_npi_cvEl(): qfrm_ApBq_npi(), central and vector, long double
- ApBq_npi_cmEl(): qfrm_ApBq_npi(), central and matrix, long double
- ApBq_npi_nvEl(): qfrm_ApBq_npi(), noncentral and vector, long double
- ApBq_npi_nmEl(): qfrm_ApBq_npi(), noncentral and matrix, long double
- ApBIqr_int_nvEl(): qfmrn_ApBIqr_int(), noncentral and vector, long double
- ApBIqr_int_nmEl(): qfmrn_ApBIqr_int(), noncentral and matrix, long double
- ApBIqr_npi_cvEl(): qfmrn_ApBIqr_npi(), central and vector, long double
- ApBIqr_npi_cmEl(): qfmrn_ApBIqr_npi(), central and matrix, long double
- ApBIqr_npi_nvEl(): qfmrn_ApBIqr_npi(), noncentral and vector, long double
- ApBIqr_npi_nmEl(): qfmrn_ApBIqr_npi(), noncentral and matrix, long double
- IpBDqr_gen_cvEl(): qfmrn_IpBDqr_gen(), central and vector, long double
- IpBDqr_gen_cmEl(): qfmrn_IpBDqr_gen(), central and matrix, long double
- IpBDqr_gen_nvEl(): qfmrn_IpBDqr_gen(), noncentral and vector, long double
- IpBDqr_gen_nmEl(): qfmrn_IpBDqr_gen(), noncentral and matrix, long double
- ApBDqr_int_cvEl(): qfmrn_ApBDqr_int(), central and vector, long double
- ApBDqr_int_cmEl(): qfmrn_ApBDqr_int(), central and matrix, long double
- ApBDqr_int_nvEl(): qfmrn_ApBDqr_int(), noncentral and vector, long double
- ApBDqr_int_nmEl(): qfmrn_ApBDqr_int(), noncentral and matrix, long double
- ApBDqr_npi_cvEl(): qfmrn_ApBDqr_npi(), central and vector, long double
- ApBDqr_npi_cmEl(): qfmrn_ApBDqr_npi(), central and matrix, long double
- ApBDqr_npi_nvEl(): qfmrn_ApBDqr_npi(), noncentral and vector, long double
- ApBDqr_npi_nmEl(): qfmrn_ApBDqr_npi(), noncentral and matrix, long double
- rqfpE(): rqfp()

default value empirically seems to work well in most conditions, but use a large `thr_margin` (e.g., `1e5`) if you encounter numerical overflow. (The C++ functions use an equivalent expression, `std::numeric_limits<Scalar>::max() / thr_margin / Scalar(n)`, with `Scalar` being `double` or `long double`.)

In these R functions, the scaling happens order-wise; i.e., it influences all the coefficients of the same order in multidimensional coefficients (in `d2_ij` and `d3_ijk`) and the coefficients of the subsequent orders.

These scaling factors are recorded in the attribute `"logscale"` of the return value, which is a vector/matrix/array whose size is identical to the return value, so that `value / exp(attr(value, "logscale"))` equals the original quantities to be obtained (if there were no overflow).

The `qfrm` and `qfmr` functions handle return values of these functions by first multiplying them with hypergeometric coefficients (which are typically $\ll 1$) and then scaling the products back to the original scale using the recorded scaling factors. (To be precise, this typically happens within `hgs` functions.) The C++ functions handle the problem similarly (but by using separate objects rather than attributes).

However, this procedure does not always mitigate the problem in multiple series; when there are very large and very small coefficients in the same order, smaller ones can diminish/underflow to the numerical 0 after repeated scaling. (The `qfrm` and `qfmr` functions try to detect and warn against this by examining whether any of the highest-order terms is 0 .) The present version of this package has implemented two methods to mitigate this problem, but only through C++ functions. One is to use the `long double` variable type, and the other is to use coefficient-wise scaling (see `qfrm` and `qfmr`).

Value

Vector of length $m + 1$, corresponding to the 0th, 1st, ..., and m th order terms. Hence, the $[m + 1]$ -th element should be extracted when the coefficient for the m th order term is required.

Has the attribute `"logscale"` as described in "Scaling" above.

References

- Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:10.1016/j.jmva.2013.03.002.
- Hillier, G., Kan, R. and Wang, X. (2009) Computationally efficient recursions for top-order invariant polynomials with applications. *Econometric Theory*, **25**, 211–242. doi:10.1017/S0266466608090075.
- Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:10.1017/S0266466613000364.

See Also

`qfpm`, `qfrm`, and `qfmr` are major front-end functions that utilize these functions

`dti12_pq` for \tilde{d} used for moments of a product of quadratic forms

`d2_ij` and `d3_ijk` for d , h , \tilde{h} , and \hat{h} used for moments of ratios of quadratic forms

d2_ij	<i>Coefficients in polynomial expansion of generating function—for ratios with two matrices</i>
-------	---

Description

These are internal functions to calculate the coefficients in polynomial expansion of joint generating functions for two quadratic forms in potentially noncentral multivariate normal variables, $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \mathbf{I}_n)$. They are primarily used in calculations around moments of a ratio involving two or three quadratic forms.

Usage

```
d2_ij_m(
  A1,
  A2,
  m = 100L,
  p = m,
  q = m,
  thr_margin = 100,
  fill_all = !missing(p) || !missing(q)
)

d2_ij_v(
  L1,
  L2,
  m = 100L,
  p = m,
  q = m,
  thr_margin = 100,
  fill_all = !missing(p) || !missing(q)
)

d2_pj_m(A1, A2, m = 100L, p = 1L, thr_margin = 100)

d2_1j_m(A1, A2, m = 100L, thr_margin = 100)

d2_pj_v(L1, L2, m = 100L, p = 1L, thr_margin = 100)

d2_1j_v(L1, L2, m = 100L, thr_margin = 100)

h2_ij_m(
  A1,
  A2,
  mu = rep.int(0, n),
  m = 100L,
  p = m,
```

```

    q = m,
    thr_margin = 100,
    fill_all = !missing(p) || !missing(q)
  )

h2_ij_v(
  L1,
  L2,
  mu = rep.int(0, n),
  m = 100L,
  p = m,
  q = m,
  thr_margin = 100,
  fill_all = !missing(p) || !missing(q)
)

htil2_pj_m(A1, A2, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)
htil2_1j_m(A1, A2, mu = rep.int(0, n), m = 100L, thr_margin = 100)
htil2_pj_v(L1, L2, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)
htil2_1j_v(L1, L2, mu = rep.int(0, n), m = 100L, thr_margin = 100)
hhat2_pj_m(A1, A2, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)
hhat2_1j_m(A1, A2, mu = rep.int(0, n), m = 100L, thr_margin = 100)
hhat2_pj_v(L1, L2, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)
hhat2_1j_v(L1, L2, mu = rep.int(0, n), m = 100L, thr_margin = 100)

```

Arguments

A1, A2	Argument matrices. Assumed to be symmetric and of the same order.
m	Integer-alike to specify the desired order along A2/L2
p, q	Integer-alikes to specify the desired orders along A1/L1 and A2/L2, respectively.
thr_margin	Optional argument to adjust the threshold for scaling (see “Scaling” in <code>d1_i</code>)
fill_all	Logical to specify whether all the output matrix should be filled. See “Details”.
L1, L2	Eigenvalues of the argument matrices
mu	Mean vector μ for \mathbf{x}

Details

`d2_**_*`(`)` functions calculate $d_{i,j}(\mathbf{A}_1, \mathbf{A}_2)$ in Hillier et al. (2009, 2014) and Bao and Kan (2013). These are also related to the top-order invariant polynomials $C_{[k_1],[k_2]}(\mathbf{A}_1, \mathbf{A}_2)$ in the following

way: $d_{i,j}(\mathbf{A}_1, \mathbf{A}_2) = \frac{1}{k_1!k_2!} \left(\frac{1}{2}\right)_{k_1+k_2} C_{[k_1],[k_2]}(\mathbf{A}_1, \mathbf{A}_2)$, where $(x)_k = x(x+1)\dots(x+k-1)$ (Chikuse 1987; Hillier et al. 2009).

`h2_ij_*`() and `htil2_pj_*`() functions calculate $h_{i,j}(\mathbf{A}_1, \mathbf{A}_2)$ and $\tilde{h}_{i,j}(\mathbf{A}_1; \mathbf{A}_2)$, respectively, in Bao and Kan (2013). Note that the latter is denoted by the symbol $\hat{h}_{i,j}$ in Hillier et al. (2014). `hhat2_pj_*`() functions are for $\hat{h}_{i,j}(\mathbf{A}_1; \mathbf{A}_2)$ in Hillier et al. (2014), used to calculate an error bound for truncated sum for moments of a ratio of quadratic forms. The mean vector $\boldsymbol{\mu}$ is a parameter in all these.

There are two different situations in which these coefficients are used in calculation of moments of ratios of quadratic forms: **1**) within an infinite series for one of the subscripts, with the other subscript fixed (when the exponent p of the numerator is integer); **2**) within a double infinite series for both subscripts (when p is non-integer) (see Bao and Kan 2013). In this package, the situation **1** is handled by the `*_pj_*` (and `*_1j_*`) functions, and **2** is by the `*_ij_*` functions.

In particular, the `*_pj_*` functions always return a $(p+1) \times (m+1)$ matrix where all elements are filled with the relevant coefficients (e.g., $d_{i,j}$, $\tilde{h}_{i,j}$), from which, typically, the $[p+1,]$ -th row is used for subsequent calculations. (Those with `*_1q_*` are simply fast versions for the commonly used case where $p=1$.) On the other hand, the `*_ij_*` functions by default return a $(m+1) \times (m+1)$ matrix whose upper-left triangular part (including the diagonals) is filled with the coefficients ($d_{i,j}$ or $h_{i,j}$), the rest being 0, and all the coefficients are used in subsequent calculations.

(At present, the `*_ij_*` functions also have the functionality to fill all coefficients of a potentially non-square output matrix, but this is less efficient than `*_pj_*` functions so may be omitted in the future development.)

Those ending with `_m` take matrices as arguments, whereas those with `_v` take eigenvalues. The latter can be used only when the argument matrices share the same eigenvectors, to which the eigenvalues correspond in the orders given, but is substantially faster.

This package also involves C++ equivalents for most of these functions (which are suffixed by `E` for `Eigen`), but these are exclusively for internal use and not exposed to the user.

Value

$(p+1) \times (m+1)$ matrix for the `*_pj_*` functions.

$(m+1) \times (m+1)$ matrix for the `*_ij_*` functions.

The rows and columns correspond to increasing orders for \mathbf{A}_1 and \mathbf{A}_2 , respectively. And the 1st row/column of each dimension corresponds to the 0th order (hence $[p+1, q+1]$ for the (p, q) -th order).

Has the attribute "logscale" as described in the "Scaling" section in `d1_i`. This is a matrix of the same size as the return itself.

References

- Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:10.1016/j.jmva.2013.03.002.
- Chikuse, Y. (1987) Methods for constructing top order invariant polynomials. *Econometric Theory*, **3**, 195–207. doi:10.1017/S026646660001029X.
- Hillier, G., Kan, R. and Wang, X. (2009) Computationally efficient recursions for top-order invariant polynomials with applications. *Econometric Theory*, **25**, 211–242. doi:10.1017/S0266466608090075.

Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:10.1017/S0266466613000364.

See Also

[qfrm](#) and [qfmrn](#) are major front-end functions that utilize these functions
[dti12_pq](#) for \tilde{d} used for moments of a product of quadratic forms
[d3_ijk](#) for equivalents for three matrices

d3_ijk	<i>Coefficients in polynomial expansion of generating function—for ratios with three matrices</i>
--------	---

Description

These are internal functions to calculate the coefficients in polynomial expansion of joint generating functions for three quadratic forms in potentially noncentral multivariate normal variables, $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \mathbf{I}_n)$. They are primarily used in calculations around moments of a ratio involving three quadratic forms.

Usage

```
d3_ijk_m(
  A1,
  A2,
  A3,
  m = 100L,
  p = m,
  q = m,
  r = m,
  thr_margin = 100,
  fill_across = c(!missing(p), !missing(q), !missing(r))
)

d3_ijk_v(
  L1,
  L2,
  L3,
  m = 100L,
  p = m,
  q = m,
  r = m,
  thr_margin = 100,
  fill_across = c(!missing(p), !missing(q), !missing(r))
)
```

```

d3_pjk_m(A1, A2, A3, m = 100L, p = 1L, thr_margin = 100)

d3_pjk_v(L1, L2, L3, m = 100L, p = 1L, thr_margin = 100)

h3_ijk_m(
  A1,
  A2,
  A3,
  mu = rep.int(0, n),
  m = 100L,
  p = m,
  q = m,
  r = m,
  thr_margin = 100,
  fill_across = c(!missing(p), !missing(q), !missing(r))
)

h3_ijk_v(
  L1,
  L2,
  L3,
  mu = rep.int(0, n),
  m = 100L,
  p = m,
  q = m,
  r = m,
  thr_margin = 100,
  fill_across = c(!missing(p), !missing(q), !missing(r))
)

htil3_pjk_m(A1, A2, A3, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)

htil3_pjk_v(L1, L2, L3, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)

hhat3_pjk_m(A1, A2, A3, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)

hhat3_pjk_v(L1, L2, L3, mu = rep.int(0, n), m = 100L, p = 1L, thr_margin = 100)

```

Arguments

A1, A2, A3	Argument matrices. Assumed to be symmetric and of the same order.
m	Integer-alike to specify the desired order along A2/L2 and A3/L3
p, q, r	Integer-alikes to specify the desired orders along A1/L1, A2/L2, and A3/L3, respectively.
thr_margin	Optional argument to adjust the threshold for scaling (see “Scaling” in d1_i)
fill_across	Logical vector of length 3, to specify whether each dimension of the output matrix should be filled.

L1, L2, L3	Eigenvalues of the argument matrices
mu	Mean vector μ for \mathbf{x}

Details

All these functions have equivalents for two-matrix cases ([d2_ij](#)), to which the user is referred for documentations. The primary difference of these functions from the latter is the addition of arguments for the third matrix \mathbf{A}_3/L_3 .

`d3_*jk_*` functions calculate $d_{i,j,k}(\mathbf{A}_1, \mathbf{A}_2, \mathbf{A}_3)$ in Hillier et al. (2009, 2014) and Bao and Kan (2013). These are also related to the top-order invariant polynomials as described in [d2_ij](#).

`h3_ijk_*`, `htil3_pjk_*`, and `hhat3_pjk_*` functions calculate $h_{i,j,k}(\mathbf{A}_1, \mathbf{A}_2, \mathbf{A}_3)$, $\tilde{h}_{i,j,k}(\mathbf{A}_1; \mathbf{A}_2, \mathbf{A}_3)$, and $\hat{h}_{i,j,k}(\mathbf{A}_1; \mathbf{A}_2, \mathbf{A}_3)$, respectively, as described in the package vignette. These are equivalent to similar coefficients described in Bao and Kan (2013) and Hillier et al. (2014).

The difference between the `*_pjk_*` and `*_ijk_*` functions is as described for `*_pj_*` and `*_ij_*` (see “Details” in [d2_ij](#)). The only difference is that these functions return a 3D array. In the `*_pjk_*` functions, all the slices along the first dimension (i.e., `[i, ,]`) are an upper-left triangular matrix like what the `*_ij_*` functions return in the 2D case; in other words, the return has the coefficients for the terms that satisfy $j + k \leq m$ for all $i = 0, 1, \dots, p$. Typically, the `[p + 1, ,]`-th slice is used for subsequent calculations. In the return of the `*_ijk_*` functions, only the triangular prism close to the `[1, 1, 1]` is filled with coefficients, which correspond to the terms satisfying $i + j + k \leq m$.

Value

$(p + 1) * (m + 1) * (m + 1)$ array for the `*_pjk_*` functions

$(m + 1) * (m + 1) * (m + 1)$ array for the `*_ijk_*` functions (by default; see “Details”).

The 1st, 2nd, and 3rd dimensions correspond to increasing orders for \mathbf{A}_1 , \mathbf{A}_2 , and \mathbf{A}_3 , respectively. And the 1st row/column of each dimension corresponds to the 0th order (hence `[p + 1, q + 1, r + 1]` for the (p, q, r) -th order).

Has the attribute “logscale” as described in the “Scaling” section in [d1_i](#). This is an array of the same size as the return itself.

References

Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:10.1016/j.jmva.2013.03.002.

Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:10.1017/S0266466613000364.

See Also

[qfmrn](#) is a major front-end function that utilizes these functions

[dti12_pq](#) for \tilde{d} used for moments of a product of quadratic forms

[d2_ij](#) for equivalents for two matrices

dtl2_pq	<i>Coefficients in polynomial expansion of generating function—for products</i>
---------	---

Description

These are internal functions to calculate the coefficients in polynomial expansion of joint generating functions for two or three quadratic forms in potentially noncentral multivariate normal variables, $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \mathbf{I}_n)$. They are primarily used to calculate moments of a product of two or three quadratic forms.

Usage

```
dtl2_pq_m(A1, A2, mu = rep.int(0, n), p = 1L, q = 1L)
```

```
dtl2_1q_m(A1, A2, mu = rep.int(0, n), q = 1L)
```

```
dtl2_pq_v(L1, L2, mu = rep.int(0, n), p = 1L, q = 1L)
```

```
dtl2_1q_v(L1, L2, mu = rep.int(0, n), q = 1L)
```

```
dtl3_pqr_m(A1, A2, A3, mu = rep.int(0, n), p = 1L, q = 1L, r = 1L)
```

```
dtl3_pqr_v(L1, L2, L3, mu = rep.int(0, n), p = 1L, q = 1L, r = 1L)
```

Arguments

A1, A2, A3	Argument matrices. Assumed to be symmetric and of the same order.
mu	Mean vector $\boldsymbol{\mu}$ for \mathbf{x}
p, q, r	Integer-alikes to specify the order along the three argument matrices
L1, L2, L3	Eigenvalues of the argument matrices

Details

`dtl2_pq_m()` and `dtl2_pq_v()` calculate $\tilde{d}_{p,q}(\mathbf{A}_1, \mathbf{A}_2)$ in Hillier et al. (2014). `dtl2_1q_m()` and `dtl2_1q_v()` are fast versions for the commonly used case where $p = 1$. Similarly, `dtl3_pqr_m()` and `dtl3_pqr_v()` are for $\tilde{d}_{p,q,r}(\mathbf{A}_1, \mathbf{A}_2, \mathbf{A}_3)$.

Those ending with `_m` take matrices as arguments, whereas those with `_v` take eigenvalues. The latter can be used only when the argument matrices share the same eigenvectors, to which the eigenvalues correspond in the orders given, but is substantially faster.

These functions calculate the coefficients based on the super-short recursion algorithm described in Hillier et al. (2014: sec. 4.2).

Value

A $(p + 1) * (q + 1)$ matrix for the 2D functions, or a $(p + 1) * (q + 1) * (r + 1)$ array for the 3D functions.

The 1st, 2nd, and 3rd dimensions correspond to increasing orders for \mathbf{A}_1 , \mathbf{A}_2 , and \mathbf{A}_3 , respectively. And the 1st row/column of each dimension corresponds to the 0th order (hence $[p + 1, q + 1]$ for the (p, q) -th moment).

References

Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:10.1017/S0266466613000364.

See Also

`qfpm` is a front-end functions that utilizes these functions

`d1_i` for a single-matrix equivalent of \tilde{d}

 hgs

Calculate hypergeometric series

Description

These internal functions calculate (summands of) hypergeometric series.

`hgs_1d()` calculates the hypergeometric series $c \frac{(a_1)_i}{(b)_i} d_i$

`hgs_2d()` calculates the hypergeometric series $c \frac{(a_1)_i (a_2)_j}{(b)_{i+j}} d_{i,j}$

`hgs_3d()` calculates the hypergeometric series $c \frac{(a_1)_i (a_2)_j (a_3)_k}{(b)_{i+j+k}} d_{i,j,k}$

Usage

`hgs_1d(dks, a1, b, lconst = 0)`

`hgs_2d(dks, a1, a2, b, lconst = 0)`

`hgs_3d(dks, a1, a2, a3, b, lconst = 0)`

Arguments

`dks` $(m + 1)$ vector for d_i , $(m + 1) * (m + 1)$ square matrix for $d_{i,j}$, or $(m + 1) * (m + 1) * (m + 1)$ array for $d_{i,j,k}$ ($i, j, k = 0, 1, \dots, m$)

`a1, a2, a3` Numerator parameters

`b` Denominator parameter

`lconst` Scalar $\log c$

Details

The denominator parameter `b` is assumed positive, whereas the numerator parameters can be positive or negative. The signs of the latter will be reflected in the result.

Value

Numeric with the same dimension with `dks`

<code>iseq</code>	<i>Are these vectors equal?</i>
-------------------	---------------------------------

Description

This internal function is used to determine whether two vectors/matrices have the same elements (or, a vector/matrix is all equal to 0) using `all.equal()`. Attributes and dimensions are ignored as they are passed as vectors using `c()`.

Usage

```
iseq(x, y = rep.int(0, length(x)), tol = .Machine$double.eps * 100)
```

Arguments

<code>x</code>	Main target vector/matrix in <code>all.equal()</code>
<code>y</code>	current in <code>all.equal()</code> . Default zero vector.
<code>tol</code>	Numeric to specify tolerance in <code>all.equal()</code>

See Also

[all.equal](#)

<code>is_diagonal</code>	<i>Is this matrix diagonal?</i>
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Description

This internal function is used to determine whether a square matrix is diagonal (within a specified tolerance). Returns TRUE when the absolute values of all off-diagonal elements are below `tol`, using `all.equal()`.

Usage

```
is_diagonal(A, tol = .Machine$double.eps * 100, symmetric = FALSE)
```

Arguments

A	Square matrix. No check is done.
tol	Numeric to specify tolerance in <code>all.equal()</code>
symmetric	If FALSE (default), sum of absolute values of the corresponding lower and upper triangular elements are examined with a doubled <code>tol</code> . If TRUE, only the lower triangular elements are examined assuming symmetry.

See Also

[all.equal](#)

 KiK

Matrix square root and generalized inverse

Description

This internal function calculates the decomposition $\mathbf{S} = \mathbf{K}\mathbf{K}^T$ for an $n \times n$ covariance matrix \mathbf{S} , so that \mathbf{K} is an $n \times m$ matrix with m being the rank of \mathbf{S} . Returns this \mathbf{K} and its generalized inverse, \mathbf{K}^- , in a list.

Usage

```
KiK(S, tol = .Machine$double.eps * 100)
```

Arguments

S	Covariance matrix. Symmetry and positive (semi-)definiteness are checked.
tol	Tolerance to determine the rank of \mathbf{S} . Eigenvalues smaller than this value are considered zero.

Details

At present, this utilizes `svd()`, although there may be better alternatives.

Value

List with \mathbf{K} and $i\mathbf{K}$, with the latter being \mathbf{K}^-

new_qfrm	<i>Construct qfrm object</i>
----------	------------------------------

Description

These are internal “constructor” functions used to make qfrm and qfpm objects, which are used as a return value from the [qfrm](#), [qfprm](#), and [qfpm](#) functions.

Usage

```
new_qfrm(
  statistic,
  error_bound = NULL,
  terms = statistic,
  seq_error = NULL,
  exact = FALSE,
  twosided = FALSE,
  alphaout = FALSE,
  singular_arg = FALSE,
  diminished = FALSE,
  ...,
  class = character()
)

new_qfpm(statistic, exact = TRUE, ..., class = character())
```

Arguments

statistic	Terminal value (partial sum) for the moment. When missing, obtained as <code>sum(terms)</code> .
error_bound	Terminal error bound. When missing, obtained as <code>seq_error[length(seq_error)]</code> .
terms	Terms in series expression for the moment along varying polynomial degrees
seq_error	Vector of error bounds corresponding to <code>cumsum(terms)</code>
exact, twosided, alphaout, singular_arg	Logicals used to append attributes to the resultant error bound (see “Value”)
diminished	Logical used to append attribute to the resultant statistic and terms (see “Value”)
...	Additional arguments for accommodating subclasses
class	Character vector to (pre-)append classes to the return value

Value

`new_qfrm()` and `new_qfpm()` return a list of class qfrm and `c(qfpm, qfrm)`, respectively. These classes are defined for the print and plot methods.

The return object is a list of 4 elements which are intended to be:

- `$statistic`: evaluation result (`sum(terms)`)

- `$terms`: vector of 0th to m th order terms
- `$error_bound`: error bound of statistic
- `$seq_error`: vector of error bounds corresponding to partial sums (`cumsum(terms)`)

When the result is exact, `$terms` can be of length 1 and equal to `$statistic`. This is always the case for the `qfpm` class.

When the relevant flags are provided in the constructor, `$error_bound` and `$seq_error` have the following attributes which control behaviors of the `print` and `plot` methods:

- `"exact"`: indicates whether the moment is exact
- `"twosided"`: indicates whether the error bounds are two-sided
- `"alphaout"`: indicates whether any of the scaling factors (`alphaA`, `alphaB`, `alphaD`) is outside $(0, 1]$, when error bound does not strictly hold
- `"singular"`: indicates whether the relevant argument matrix is (numerically) singular, in which case the error bound is invalid

Similarly, when `diminished = TRUE`, `$statistic` and `$terms` have the attribute `"diminished"` being `TRUE`, which indicates that numerical underflow/diminishing happened during scaling (see `"Scaling"` in `d1_i`).

See Also

[qfrm](#), [qfmr](#), [qfpm](#): functions that return objects of these classes

[methods.qfrm](#): the `print` and `plot` methods

print.qfrm

Methods for qfrm and qfpm objects

Description

Straightforward `print` and `plot` methods are defined for `qfrm` and `qfpm` objects which result from the `qfrm`, `qfmr`, and `qfpm` functions.

Usage

```
## S3 method for class 'qfrm'
print(
  x,
  digits = getOption("digits"),
  show_range = !is.null(x$error_bound),
  ...
)

## S3 method for class 'qfpm'
print(x, digits = getOption("digits"), ...)
```

```
## S3 method for class 'qfrm'
plot(
  x,
  add_error = length(x$seq_error) > 0,
  add_legend = add_error,
  ylim = x$statistic * ylim_f,
  ylim_f = c(0.9, 1.1),
  xlab = "Order of evaluation",
  ylab = "Moment of ratio",
  col_m = "royalblue4",
  col_e = "tomato",
  lwd_m = 1,
  lwd_e = 1,
  lty_m = 1,
  lty_e = 2,
  pos_leg = "topright",
  ...
)
```

Arguments

x	qfrm or qfpm object
digits	Number of significant digits to be printed.
show_range	Logical to specify whether the possible range for the moment is printed (when available). Default TRUE when available.
...	In the plot methods, passed to plot.default . In the print methods, ignored (retained for the compatibility with the generic method).
add_error	Logical to specify whether the sequence of error bounds is plotted (when available). Default TRUE when available.
add_legend	Logical to specify whether a legend is added. Turned on by default when add_error = TRUE.
ylim, ylim_f	ylim is passed to plot.default . By default, this is automatically set to ylim_f times the terminal value of the series expression (x\$statistic). ylim_f is by default c(0.9, 1.1).
xlab, ylab	Passed to plot.default
col_m, col_e, lwd_m, lwd_e, lty_m, lty_e	col, lwd, and lty to plot the sequences of the moment (***_m) and its error bound (***_e)
pos_leg	Position of the legend, e.g., "topright", "bottomright", passed as the first argument for legend

Details

The print methods simply display the moment `x$statistic` (typically a partial sum), its error bound `x$error_bound` (when available), and the possible range of the moment (`x$statistic` to `x$statistic + x$error_bound` in case of one-sided error bound; `x$statistic - x$error_bound` to `x$statistic + x$error_bound` in case of two-sided).

The `plot` method is designed for quick inspection of the profile of the partial sum of the series along varying orders `cumsum(x$terms)`. When the object has a sequence for error bounds `x$seq_error`, this is also shown with a broken line (by default). When the object has an exact moment (i.e., resulting from `qfrm_ApIq_int()` or the `qfpm` functions), a message is thrown to tell inspection of the plot will not be required in this case.

Value

The `print` method invisibly returns the input.

The `plot` method is used for the side effect (and invisibly returns `NULL`).

See Also

[new_qfrm](#): descriptions of the classes and their “constructors”

Examples

```
nv <- 4
A <- diag(nv:1)
B <- diag(1:nv)
mu <- rep.int(1, nv)

res1 <- qfrm(A, B, p = 3, mu = mu)
print(res1)
print(res1, digits = 5)
print(res1, digits = 10)

## Default plot: ylim too narrow to see the error bound at this m
plot(res1)

## With extended ylim
plot(res1, ylim_f = c(0.8, 1.2), pos_leg = "topleft")

## In this case, it is easy to increase m
(res2 <- qfrm(A, B, p = 3, mu = mu, m = 200))
plot(res2)
```

qfmr

Moment of multiple ratio of quadratic forms in normal variables

Description

`qfmr()` is a front-end function to obtain the (compound) moment of a multiple ratio of quadratic forms in normal variables in the following special form: $E\left(\frac{(\mathbf{x}^T \mathbf{A} \mathbf{x})^p}{(\mathbf{x}^T \mathbf{B} \mathbf{x})^q (\mathbf{x}^T \mathbf{D} \mathbf{x})^r}\right)$, where $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \boldsymbol{\Sigma})$. Like `qfrm()`, this function calls one of the following “internal” functions for actual calculation, as appropriate.

`qfmr_ApBIqr_int()`: For $\mathbf{D} = \mathbf{I}_n$ and positive-integral p

`qfmm_ApBIqr_npi()`: For $\mathbf{D} = \mathbf{I}_n$ and non-integral p
`qfmm_IpBDqr_gen()`: For $\mathbf{A} = \mathbf{I}_n$
`qfmm_ApBDqr_int()`: For general \mathbf{A} , \mathbf{B} , and \mathbf{D} , and positive-integral p
`qfmm_ApBDqr_npi()`: For general \mathbf{A} , \mathbf{B} , and \mathbf{D} , and non-integral p

Usage

```

qfmm(
  A,
  B,
  D,
  p = 1,
  q = p/2,
  r = q,
  m = 100L,
  mu = rep.int(0, n),
  Sigma = diag(n),
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100,
  ...
)

qfmm_ApBIqr_int(
  A,
  B,
  p = 1,
  q = 1,
  r = 1,
  m = 100L,
  mu = rep.int(0, n),
  error_bound = TRUE,
  check_convergence = c("relative", "strict_relative", "absolute", "none"),
  use_cpp = TRUE,
  cpp_method = c("double", "long_double", "coef_wise"),
  nthreads = 0,
  alphaB = 1,
  tol_conv = .Machine$double.eps^(1/4),
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100,
  thr_margin = 100
)

qfmm_ApBIqr_npi(
  A,
  B,
  p = 1,
  q = 1,
  r = 1,

```



```
m = 100L,  
mu = rep.int(0, n),  
check_convergence = c("relative", "strict_relative", "absolute", "none"),  
use_cpp = TRUE,  
cpp_method = c("double", "long_double", "coef_wise"),  
nthreads = 0,  
alphaA = 1,  
alphaB = 1,  
tol_conv = .Machine$double.eps^(1/4),  
tol_zero = .Machine$double.eps * 100,  
tol_sing = .Machine$double.eps * 100,  
thr_margin = 100  
)  
  
qfmm_IpBDqr_gen(  
  B,  
  D,  
  p = 1,  
  q = 1,  
  r = 1,  
  mu = rep.int(0, n),  
  m = 100L,  
  check_convergence = c("relative", "strict_relative", "absolute", "none"),  
  use_cpp = TRUE,  
  cpp_method = c("double", "long_double", "coef_wise"),  
  nthreads = 0,  
  alphaB = 1,  
  alphaD = 1,  
  tol_conv = .Machine$double.eps^(1/4),  
  tol_zero = .Machine$double.eps * 100,  
  tol_sing = .Machine$double.eps * 100,  
  thr_margin = 100  
)  
  
qfmm_ApBDqr_int(  
  A,  
  B,  
  D,  
  p = 1,  
  q = 1,  
  r = 1,  
  m = 100L,  
  mu = rep.int(0, n),  
  check_convergence = c("relative", "strict_relative", "absolute", "none"),  
  use_cpp = TRUE,  
  cpp_method = c("double", "long_double", "coef_wise"),  
  nthreads = 0,  
  alphaB = 1,
```

```

alphaD = 1,
tol_conv = .Machine$double.eps^(1/4),
tol_zero = .Machine$double.eps * 100,
tol_sing = .Machine$double.eps * 100,
thr_margin = 100
)

qfmm_ApBDqr_npi(
  A,
  B,
  D,
  p = 1,
  q = 1,
  r = 1,
  m = 100L,
  mu = rep.int(0, n),
  check_convergence = c("relative", "strict_relative", "absolute", "none"),
  use_cpp = TRUE,
  cpp_method = c("double", "long_double", "coef_wise"),
  nthreads = 0,
  alphaA = 1,
  alphaB = 1,
  alphaD = 1,
  tol_conv = .Machine$double.eps^(1/4),
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100,
  thr_margin = 100
)

```

Arguments

A, B, D	Argument matrices. Should be square. Will be automatically symmetrized.
p, q, r	Exponents for A , B , and D , respectively. By default, q equals p/2 and r equals q. If unsure, specify all explicitly.
m	Order of polynomials at which the series expression is truncated. <i>M</i> in Hillier et al. (2009, 2014).
mu	Mean vector μ for x
Sigma	Covariance matrix Σ for x . Accommodated only by the front-end qfmm(). See “Details” in qfrm .
tol_zero	Tolerance against which numerical zero is determined. Used to determine, e.g., whether mu is a zero vector, A or B equals the identity matrix, etc.
tol_sing	Tolerance against which matrix singularity and rank are determined. The eigenvalues smaller than this are considered zero.
...	Additional arguments in the front-end qfmm() will be passed to the appropriate “internal” function.
error_bound	Logical to specify whether an error bound is returned (if available).

check_convergence	Specifies how numerical convergence is checked (see “Details”). Options: <ul style="list-style-type: none"> • “relative”: default; magnitude of the last term of the series relative to the sum is compared with <code>tol_conv</code> • “strict_relative” or TRUE: same, but stricter than default by setting <code>tol_conv = .Machine\$double.eps</code> (unless a smaller value is specified by the user) • “absolute”: absolute magnitude of the last term is compared with <code>tol_conv</code> • “none” or FALSE: skips convergence check
use_cpp	Logical to specify whether the calculation is done with C++ functions via Rcpp. TRUE by default.
cpp_method	Method used in C++ calculations to avoid numerical overflow/underflow (see “Details”). Options: <ul style="list-style-type: none"> • “double”: default; fastest but prone to underflow in some conditions • “long_double”: same algorithm but using the long double variable type; robust but slow and memory-inefficient • “coef_wise”: coefficient-wise scaling algorithm; most robust but variably slow
nthreads	Number of threads used in OpenMP-enabled C++ functions. See “Multithreading” in qfrm .
tol_conv	Tolerance against which numerical convergence of series is checked. Used with <code>check_convergence</code> .
thr_margin	Optional argument to adjust the threshold for scaling (see “Scaling” in d1_i). Passed to internal functions (d1_i , d2_ij , d3_ijk) or their C++ equivalents.
alphaA, alphaB, alphaD	Factors for the scaling constants for A , B , and D , respectively. See “Details” in qfrm .

Details

The usage of these functions is similar to [qfrm](#), to which the user is referred for documentation. It is assumed that $\mathbf{B} \neq \mathbf{D}$ (otherwise, the problem reduces to a simple ratio).

When **B** is identity or missing, this and its exponent *q* will be swapped with **D** and *r*, respectively, before `qfmr_ApBIqr_***()` is called.

The existence conditions for the moments of this multiple ratio can be reduced to those for a simple ratio, provided that one of the null spaces of **B** and **D** is a subspace of the other (including the case they are null). The conditions of Bao and Kan (2013: proposition 1) can then be applied by replacing *q* and *m* there by *q + r* and $\min(\text{rank}(\mathbf{B}), \text{rank}(\mathbf{D}))$, respectively (see also Smith 1989: p. 258 for nonsingular **B**, **D**). An error is thrown if these conditions are not met in this case. Otherwise (i.e., **B** and **D** are both singular and neither of their null spaces is a subspace of the other), it seems difficult to define general moment existence conditions. A sufficient condition can be obtained by applying the same proposition with a new denominator matrix whose null space is union of those of **B** and **D** (Watanabe, 2022). A warning is thrown if that condition is not met in this case.

Most of these functions, excepting `qfmm_ApBIqr_int()` with zero μ , involve evaluation of multiple series, which can suffer from numerical overflow and underflow (see “Scaling” in `d1_i` and “Details” in `qfrm`). To avoid this, `cpp_method = "long_double"` or `"coef_wise"` options can be used (see “Details” in `qfrm`).

Value

A `qfrm` object, as in `qfrm()` functions.

References

Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:10.1016/j.jmva.2013.03.002.

Smith, M. D. (1989). On the expectation of a ratio of quadratic forms in normal variables. *Journal of Multivariate Analysis*, **31**, 244–257. doi:10.1016/0047259X(89)900651.

Watanabe, J. (2022). Exact expressions and numerical evaluation of average evolvability measures for characterizing and comparing **G** matrices. *bioRxiv* preprint, 2022.11.02.514929. doi:10.1101/2022.11.02.514929.

See Also

`qfrm` for simple ratio

Examples

```
## Some symmetric matrices and parameters
nv <- 4
A <- diag(nv:1)
B <- diag(sqrt(1:nv))
D <- diag((1:nv)^2 / nv)
mu <- nv:1 / nv
Sigma <- matrix(0.5, nv, nv)
diag(Sigma) <- 1

## Expectation of  $(x^T A x)^2 / (x^T B x) (x^T x)$  where  $x \sim N(0, I)$ 
(res1 <- qfmm(A, B, p = 2, q = 1, r = 1))
plot(res1)

# The above internally calls the following:
qfmm_ApBIqr_int(A, B, p = 2, q = 1, r = 1) ## The same

# Similar result with different expression
# This is a suboptimal option and throws a warning
qfmm_ApBIqr_npi(A, B, p = 2, q = 1, r = 1)

## Expectation of  $(x^T A x) / (x^T B x)^{(1/2)} (x^T D x)^{(1/2)}$  where  $x \sim N(0, I)$ 
(res2 <- qfmm(A, B, D, p = 1, q = 1/2, r = 1/2))
plot(res2)

# The above internally calls the following:
qfmm_ApBDqr_int(A, B, D, p = 1, q = 1/2, r = 1/2) ## The same
```

```

## Average response correlation between A and B
(res3 <- qfmrn(crossprod(A, B), crossprod(A), crossprod(B),
              p = 1, q = 1/2, r = 1/2))
plot(res3)

## Same, but with  $x \sim N(\mu, \Sigma)$ 
(res4 <- qfmrn(crossprod(A, B), crossprod(A), crossprod(B),
              p = 1, q = 1/2, r = 1/2, mu = mu, Sigma = Sigma))
plot(res4)

## Average autonomy of D
(res5 <- qfmrn(B = D, D = solve(D), p = 2, q = 1, r = 1))
plot(res5)

```

qfpm

Moment of (product of) quadratic forms in normal variables

Description

Functions to obtain (compound) moments of a product of quadratic forms in normal variables, i.e., $E((\mathbf{x}^T \mathbf{A} \mathbf{x})^p (\mathbf{x}^T \mathbf{B} \mathbf{x})^q (\mathbf{x}^T \mathbf{D} \mathbf{x})^r)$, where $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \boldsymbol{\Sigma})$.

qfm_Ap_int() is for $q = r = 0$ (simple moment)

qfpm_ABpq_int() is for $r = 0$

qfpm_ABDpqr_int() is for the product of all three powers

Usage

```

qfm_Ap_int(
  A,
  p = 1,
  mu = rep.int(0, n),
  Sigma = diag(n),
  use_cpp = TRUE,
  cpp_method = "double",
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100
)

```

```

qfpm_ABpq_int(
  A,
  B,
  p = 1,
  q = 1,
  mu = rep.int(0, n),
  Sigma = diag(n),

```

```

use_cpp = TRUE,
cpp_method = "double",
tol_zero = .Machine$double.eps * 100,
tol_sing = .Machine$double.eps * 100
)

qfpm_ABDpqr_int(
  A,
  B,
  D,
  p = 1,
  q = 1,
  r = 1,
  mu = rep.int(0, n),
  Sigma = diag(n),
  use_cpp = TRUE,
  cpp_method = "double",
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100
)

```

Arguments

A, B, D	Argument matrices. Should be square. Will be automatically symmetrized.
p, q, r	Exponents for A , B , and D , respectively. By default, these are set to the same value. If unsure, specify all explicitly.
mu	Mean vector μ for x
Sigma	Covariance matrix Σ for x
use_cpp	Logical to specify whether the calculation is done with C++ functions via Rcpp. TRUE by default.
cpp_method	Variable type used in C++ calculations. In these functions this is ignored.
tol_zero	Tolerance against which numerical zero is determined. Used to determine, e.g., whether mu is a zero vector, A or B equals the identity matrix, etc.
tol_sing	Tolerance against which matrix singularity and rank are determined. The eigenvalues smaller than this are considered zero.

Details

These functions implement the super-short recursion algorithms described in Hillier et al. (2014: sec. 3.1–3.2 and 4). At present, only positive integers are accepted as exponents (negative exponents yield ratios, of course). All these yield exact results.

Value

A `qfpm` object which has the same elements as those returned by the `qfrm` functions. Use `$statistic` to access the value of the moment.

See Also

[qfrm](#) and [qfmrn](#) for moments of ratios

Examples

```
## Some symmetric matrices and parameters
nv <- 4
A <- diag(nv:1)
B <- diag(sqrt(1:nv))
D <- diag((1:nv)^2 / nv)
mu <- nv:1 / nv
Sigma <- matrix(0.5, nv, nv)
diag(Sigma) <- 1

## Expectation of (x^T A x)^2 where x ~ N(0, I)
qfm_Ap_int(A, 2)

## This is the same but obviously less efficient
qfpm_ABpq_int(A, p = 2, q = 0)

## Expectation of (x^T A x) (x^T B x) (x^T D x) where x ~ N(0, I)
qfpm_ABDpqr_int(A, B, D, 1, 1, 1)

## Expectation of (x^T A x) (x^T B x) (x^T D x) where x ~ N(mu, Sigma)
qfpm_ABDpqr_int(A, B, D, 1, 1, 1, mu = mu, Sigma = Sigma)

## Expectations of (x^T x)^2 where x ~ N(0, I) and x ~ N(mu, I)
## i.e., roundabout way to obtain moments of
## central and noncentral chi-square variables
qfm_Ap_int(diag(nv), 2)
qfm_Ap_int(diag(nv), 2, mu = mu)
```

qfrm

Moment of ratio of quadratic forms in normal variables

Description

`qfrm()` is a front-end function to obtain the (compound) moment of a ratio of quadratic forms in normal variables, i.e., $E\left(\frac{(\mathbf{x}^T \mathbf{A} \mathbf{x})^p}{(\mathbf{x}^T \mathbf{B} \mathbf{x})^q}\right)$, where $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \boldsymbol{\Sigma})$. Internally, `qfrm()` calls one of the following functions which does the actual calculation, depending on \mathbf{A} , \mathbf{B} , and p . Usually the best one is automatically selected.

`qfrm_ApIq_int()`: For $\mathbf{B} = \mathbf{I}_n$ and positive-integral p .

`qfrm_ApIq_npi()`: For $\mathbf{B} = \mathbf{I}_n$ and non-positive-integral p (fraction or negative).

`qfrm_ApBq_int()`: For general \mathbf{B} and positive-integral p .

`qfrm_ApBq_npi()`: For general \mathbf{B} and non-integral p .

Usage

```
qfrm(  
  A,  
  B,  
  p = 1,  
  q = p,  
  m = 100L,  
  mu = rep.int(0, n),  
  Sigma = diag(n),  
  tol_zero = .Machine$double.eps * 100,  
  tol_sing = .Machine$double.eps * 100,  
  ...  
)  
  
qfrm_ApIq_int(  
  A,  
  p = 1,  
  q = p,  
  m = 100L,  
  mu = rep.int(0, n),  
  use_cpp = TRUE,  
  cpp_method = "double",  
  nthreads = 1,  
  tol_zero = .Machine$double.eps * 100,  
  thr_margin = 100  
)  
  
qfrm_ApIq_npi(  
  A,  
  p = 1,  
  q = p,  
  m = 100L,  
  mu = rep.int(0, n),  
  error_bound = TRUE,  
  check_convergence = c("relative", "strict_relative", "absolute", "none"),  
  use_cpp = TRUE,  
  cpp_method = c("double", "long_double", "coef_wise"),  
  nthreads = 1,  
  alphaA = 1,  
  tol_conv = .Machine$double.eps^(1/4),  
  tol_zero = .Machine$double.eps * 100,  
  tol_sing = .Machine$double.eps * 100,  
  thr_margin = 100  
)  
  
qfrm_ApBq_int(  
  A,  
  B,
```



```

p = 1,
q = p,
m = 100L,
mu = rep.int(0, n),
error_bound = TRUE,
check_convergence = c("relative", "strict_relative", "absolute", "none"),
use_cpp = TRUE,
cpp_method = "double",
nthreads = 1,
alphaB = 1,
tol_conv = .Machine$double.eps^(1/4),
tol_zero = .Machine$double.eps * 100,
tol_sing = .Machine$double.eps * 100,
thr_margin = 100
)

qfrm_ApBq_npi(
  A,
  B,
  p = 1,
  q = p,
  m = 100L,
  mu = rep.int(0, n),
  check_convergence = c("relative", "strict_relative", "absolute", "none"),
  use_cpp = TRUE,
  cpp_method = c("double", "long_double", "coef_wise"),
  nthreads = 0,
  alphaA = 1,
  alphaB = 1,
  tol_conv = .Machine$double.eps^(1/4),
  tol_zero = .Machine$double.eps * 100,
  tol_sing = .Machine$double.eps * 100,
  thr_margin = 100
)

```

Arguments

A, B	Argument matrices. Should be square. Will be automatically symmetrized.
p, q	Exponents corresponding to A and B , respectively. When only one is provided, the other is set to the same value. Should be length-one numeric (see “Details” for further conditions).
m	Order of polynomials at which the series expression is truncated. <i>M</i> in Hillier et al. (2009, 2014).
mu	Mean vector $\boldsymbol{\mu}$ for x
Sigma	Covariance matrix $\boldsymbol{\Sigma}$ for x . Accommodated only by the front-end qfrm(). See “Details”.
tol_zero	Tolerance against which numerical zero is determined. Used to determine, e.g., whether mu is a zero vector, A or B equals the identity matrix, etc.

tol_sing	Tolerance against which matrix singularity and rank are determined. The eigenvalues smaller than this are considered zero.
...	Additional arguments in the front-end qfrm() will be passed to the appropriate “internal” function.
use_cpp	Logical to specify whether the calculation is done with C++ functions via Rcpp. TRUE by default.
cpp_method	Method used in C++ calculations to avoid numerical overflow/underflow (see “Details”). Options: <ul style="list-style-type: none"> • “double”: default; fastest but prone to underflow in some conditions • “long_double”: same algorithm but using the long double variable type; robust but slow and memory-inefficient • “coef_wise”: coefficient-wise scaling algorithm; most robust but variably slow
nthreads	Number of threads used in OpenMP-enabled C++ functions. 0 or any negative value is special and means one-half of the number of processors detected. See “Multithreading” in “Details”.
thr_margin	Optional argument to adjust the threshold for scaling (see “Scaling” in d1_i). Passed to internal functions (d1_i, d2_ij, d3_ijk) or their C++ equivalents.
error_bound	Logical to specify whether an error bound is returned (if available).
check_convergence	Specifies how numerical convergence is checked (see “Details”). Options: <ul style="list-style-type: none"> • “relative”: default; magnitude of the last term of the series relative to the sum is compared with tol_conv • “strict_relative” or TRUE: same, but stricter than default by setting tol_conv = .Machine\$double.eps (unless a smaller value is specified by the user) • “absolute”: absolute magnitude of the last term is compared with tol_conv • “none” or FALSE: skips convergence check
alphaA, alphaB	Factors for the scaling constants for A and B , respectively. See “Details”.
tol_conv	Tolerance against which numerical convergence of series is checked. Used with check_convergence.

Details

These functions use infinite series expressions based on the joint moment-generating function (with the top-order zonal/invariant polynomials) (see Smith 1989, Hillier et al. 2009, 2014; Bao and Kan 2013), and the results are typically partial (truncated) sums from these infinite series, which necessarily involve truncation errors. (An exception is when $\mathbf{B} = \mathbf{I}_n$ and p is a positive integer, the case handled by qfrm_ApIq_int().)

The returned value is a list consisting of the truncated sequence up to the order specified by m , its sum, and error bounds corresponding to these (see “Values”). The print method only displays the terminal partial sum and its error bound (when available). Use plot() for visual inspection, or the ordinary list element access as required.

In most cases, p and q should be nonnegative (in addition, p should be an integer in qfrm_ApIq_int() and qfrm_ApBq_int() when used directly), and an error is thrown otherwise. The only exception

is `qfrm_ApIq_npi()` which accepts negative exponents to accommodate $\frac{(\mathbf{x}^T \mathbf{x})^q}{(\mathbf{x}^T \mathbf{A} \mathbf{x})^p}$. Even in the latter case, the exponents should have the same sign. (Technically, not all of these conditions are necessary for the mathematical results to hold, but they are enforced for simplicity).

When `error_bound = TRUE` (default), `qfrm_ApBq_int()` evaluates a truncation error bound following Hillier et al. (2009: theorem 6) or Hillier et al. (2014: theorem 7) (for zero and nonzero means, respectively). `qfrm_ApIq_npi()` implements similar error bounds. No error bound is known for `qfrm_ApBq_npi()` to the author's knowledge.

For situations when the error bound is unavailable, a *very rough* check of numerical convergence is also conducted; a warning is thrown if the magnitude of the last term does not look small enough. By default, its relative magnitude to the sum is compared with the tolerance controlled by `tol_conv`, whose default is `.Machine$double.eps^(1/4)` ($\approx 1.2e-04$) (see `check_convergence`).

When `Sigma` is provided, the quadratic forms are transformed into a canonical form; that is, using the decomposition $\Sigma = \mathbf{K}\mathbf{K}^T$, where the number of columns m of \mathbf{K} equals the rank of Σ , $\mathbf{A}_{\text{new}} = \mathbf{K}^T \mathbf{A} \mathbf{K}$, $\mathbf{B}_{\text{new}} = \mathbf{K}^T \mathbf{B} \mathbf{K}$, and $\mathbf{x}_{\text{new}} = \mathbf{K}^{-1} \mathbf{x} \sim N_m(\mathbf{K}^{-1} \boldsymbol{\mu}, \mathbf{I}_m)$. `qfrm()` handles this by transforming `A`, `B`, and `mu` and calling itself recursively with these new arguments. Note that the "internal" functions do not accommodate `Sigma` (the error for unused arguments will happen). For singular Σ , one of the following conditions should be met for the above transformation to be valid: **1)** $\boldsymbol{\mu}$ is in the range of Σ ; **2)** \mathbf{A} and \mathbf{B} are in the range of Σ ; or **3)** $\mathbf{A}\boldsymbol{\mu} = \mathbf{B}\boldsymbol{\mu} = \mathbf{0}_n$. An error is thrown if none is met with a singular `Sigma`.

The existence of the moment is assessed by the eigenstructures of \mathbf{A} and \mathbf{B} , p , and q , according to Bao and Kan (2013: proposition 1). An error will result if the conditions are not met.

Straightforward implementation of the original recursive algorithms can suffer from numerical overflow when the problem is large. Internal functions (`d1_i`, `d2_ij`, `d3_ijk`) are designed to avoid overflow by order-wise scaling. However, when evaluation of multiple series is required (`qfrm_ApIq_npi()` with nonzero `mu` and `qfrm_ApBq_npi()`), the scaling occasionally yields underflow/diminishing of some terms to numerical \emptyset , causing inaccuracy. A warning is thrown in this case. (See also "Scaling" in `d1_i`.) To avoid this problem, the C++ versions of these functions have two workarounds, as controlled by `cpp_method`. **1)** The "long_double" option uses the long double variable type instead of the regular double. This is generally slow and most memory-inefficient. **2)** The "coef_wise" option uses a coefficient-wise scaling algorithm with the double variable type. This is generally robust against underflow issues. Computational time varies a lot with conditions; generally only modestly slower than the "double" option, but can be the slowest in some extreme conditions.

For the sake of completeness (only), the scaling parameters β (see the package vignette) can be modified via the arguments `alphaA` and `alphaB`. These are the factors for the inverses of the largest eigenvalues of \mathbf{A} and \mathbf{B} , respectively, and should be between 0 and 2. The default is 1, which should suffice for most purposes. Values larger than 1 often yield faster convergence, but are *not* recommended as the error bound will not strictly hold (see Hillier et al. 2009, 2014).

Multithreading:

All these functions use C++ versions to speed up computation by default. Furthermore, some of the C++ functions, in particular those using more than one matrix arguments, are parallelized with OpenMP (when available). Use the argument `nthreads` to control the number of OpenMP threads. By default (`nthreads = 0`), one-half of the processors detected with `omp_get_num_procs()` are used. This is except when all the argument matrices share the same eigenvectors and hence the calculation only involves element-wise operations of eigenvalues. In that case, the calculation is typically fast without parallelization, so `nthreads` is automatically set to 1 unless explicitly

specified otherwise; the user can still specify a larger value or θ for (typically marginal) speed gains in large problems.

Dependency note:

An exact expression of the moment is available when p is integer and $\mathbf{B} = \mathbf{I}_n$ (handled by `qfrm_ApIq_int()`), but this requires evaluation of a confluent hypergeometric function when μ is nonzero (Hillier et al. 2014: theorem 4). This is done via `gsl::hyperg_1F1()` if the package `gsl` is installed (which this package Suggests). Otherwise, the function uses the ordinary infinite series expression (Hillier et al. 2009), which is less accurate and slow, and throws a message. (In this case, the calculation is handled without C++ codes.) It is recommended to install that package if an accurate estimate is desired for that case.

Value

A `qfrm` object consisting of the following:

- `$statistic`: evaluation result (`sum(terms)`)
- `$terms`: vector of 0th to m th order terms
- `$error_bound`: error bound of statistic
- `$seq_error`: vector of error bounds corresponding to partial sums (`cumsum(terms)`)

References

- Bao, Y. and Kan, R. (2013) On the moments of ratios of quadratic forms in normal random variables. *Journal of Multivariate Analysis*, **117**, 229–245. doi:10.1016/j.jmva.2013.03.002.
- Hillier, G., Kan, R. and Wang, X. (2009) Computationally efficient recursions for top-order invariant polynomials with applications. *Econometric Theory*, **25**, 211–242. doi:10.1017/S0266466608090075.
- Hillier, G., Kan, R. and Wang, X. (2014) Generating functions and short recursions, with applications to the moments of quadratic forms in noncentral normal vectors. *Econometric Theory*, **30**, 436–473. doi:10.1017/S0266466613000364.
- Smith, M. D. (1989) On the expectation of a ratio of quadratic forms in normal variables. *Journal of Multivariate Analysis*, **31**, 244–257. doi:10.1016/0047259X(89)900651.
- Smith, M. D. (1993) Expectations of ratios of quadratic forms in normal variables: evaluating some top-order invariant polynomials. *Australian Journal of Statistics*, **35**, 271–282. doi:10.1111/j.1467-842X.1993.tb01335.x.

See Also

[qfmr](#) for multiple ratio

Examples

```
## Some symmetric matrices and parameters
nv <- 4
A <- diag(nv:1)
B <- diag(sqrt(1:nv))
mu <- nv:1 / nv
Sigma <- matrix(0.5, nv, nv)
```

```

diag(Sigma) <- 1

## Expectation of  $(x^T A x)^2 / (x^T x)^2$  where  $x \sim N(0, I)$ 
## An exact expression is available
(res1 <- qfrm(A, p = 2))

# The above internally calls the following:
qfrm_ApIq_int(A, p = 2) ## The same

# Similar result with different expression
# This is a suboptimal option and throws a warning
qfrm_ApIq_npi(A, p = 2)

## Expectation of  $(x^T A x)^{1/2} / (x^T x)^{1/2}$  where  $x \sim N(0, I)$ 
## Note how quickly the series converges in this case
(res2 <- qfrm(A, p = 1/2))
plot(res2)

# The above calls:
qfrm_ApIq_npi(A, p = 0.5)

# This is not allowed (throws an error):
try(qfrm_ApIq_int(A, p = 0.5))

##  $(x^T A x)^2 / (x^T B x)^3$  where  $x \sim N(0, I)$ 
(res3 <- qfrm(A, B, 2, 3))
plot(res3)

##  $(x^T A x)^2 / (x^T B x)^2$  where  $x \sim N(\mu, I)$ 
## Note the two-sided error bound
(res4 <- qfrm(A, B, 2, 2, mu = mu))
plot(res4)

##  $(x^T A x)^2 / (x^T B x)^2$  where  $x \sim N(\mu, \Sigma)$ 
(res5 <- qfrm(A, B, p = 2, q = 2, mu = mu, Sigma = Sigma))
plot(res5)

# Sigma is not allowed in the "internal" functions:
try(qfrm_ApBq_int(A, B, p = 2, q = 2, Sigma = Sigma))

# In res5 above, the error bound didn't converge
# Use larger m to evaluate higher-order terms
plot(print(qfrm(A, B, p = 2, q = 2, mu = mu, Sigma = Sigma, m = 300)))

```

Description

`rqfr()`, `rqfmr()`, and `rqfp()` calculate a random sample of a simple ratio, multiple ratio (of special

form), and product, respectively, of quadratic forms in normal variables of specified mean and covariance (standard multivariate normal by default). These functions are primarily for empirical verification of the analytic results provided in this package.

Usage

```
rqfr(nit = 1000L, A, B, p = 1, q = p, mu, Sigma, use_cpp = TRUE)
```

```
rqfmr(nit = 1000L, A, B, D, p = 1, q = p/2, r = q, mu, Sigma, use_cpp = TRUE)
```

```
rqfp(nit = 1000L, A, B, D, p = 1, q = 1, r = 1, mu, Sigma, use_cpp = TRUE)
```

Arguments

nit	Number of iteration or sample size. Should be an integer-alike of length 1.
A, B, D	Argument matrices (see “Details”). Assumed to be square matrices of the same order. When missing, set to the identity matrix. At least one of these should be specified.
p, q, r	Exponents for A, B, D, respectively (see “Details”). Assumed to be numeric of length 1 each. See “Details” for default values.
mu	Mean vector $\boldsymbol{\mu}$ for \mathbf{x} . Default zero vector.
Sigma	Covariance matrix $\boldsymbol{\Sigma}$ for \mathbf{x} . Default identity matrix. mu and Sigma are assumed to be of the same order as the argument matrices.
use_cpp	Logical to specify whether an C++ version is called or not. TRUE by default.

Details

These functions generate a random sample of $\frac{(\mathbf{x}^T \mathbf{A} \mathbf{x})^p}{(\mathbf{x}^T \mathbf{B} \mathbf{x})^q}$ (rqfr()), $\frac{(\mathbf{x}^T \mathbf{A} \mathbf{x})^p}{(\mathbf{x}^T \mathbf{B} \mathbf{x})^q (\mathbf{x}^T \mathbf{D} \mathbf{x})^r}$ (rqfmr()), and $(\mathbf{x}^T \mathbf{A} \mathbf{x})^p (\mathbf{x}^T \mathbf{B} \mathbf{x})^q (\mathbf{x}^T \mathbf{D} \mathbf{x})^r$ (rqfp()), where $\mathbf{x} \sim N_n(\boldsymbol{\mu}, \boldsymbol{\Sigma})$. (Internally, rqfr() and rqfmr() just call rqfp() with negative exponents.)

When only one of p and q is provided in rqfr(), the other (missing) one is set to the same value.

In rqfmr(), q and r are set to p/2 when both missing, and set to the same value when only one is missing. When p is missing, this is set to be q + r. If unsure, specify all these explicitly.

In rqfp(), p, q and r are 1 by default, provided that the corresponding argument matrices are given. If both an argument matrix and its exponent (e.g., D and r) are missing, the exponent is set to 0 so that the factor be unity.

Value

Numeric vector of length nit.

See Also

[qfrm](#) and [qfpm](#) for analytic moments

Examples

```

p <- 4
A <- diag(1:p)
B <- diag(p:1)
D <- diag(sqrt(1:p))

## By default B = I, p = q = 1;
## i.e., (x^T A x) / (x^T x), x ~ N(0, I)
rqfr(5, A)

## (x^T A x) / ((x^T B x)(x^T D x))^(1/2), x ~ N(0, I)
rqfmr(5, A, B, D, 1, 1/2, 1/2)

## (x^T A x), x ~ N(0, I)
rqfp(5, A)

## (x^T A x) (x^T B x), x ~ N(0, I)
rqfp(5, A, B)

## (x^T A x) (x^T B x) (x^T D x), x ~ N(0, I)
rqfp(5, A, B, D)

## Example with non-standard normal
mu <- 1:p / p
Sigma <- matrix(0.5, p, p)
diag(Sigma) <- 1
rqfr(5, A, mu = 1:p / p, Sigma = Sigma)

## Compare Monte Carlo sample and analytic expression
set.seed(3)
mcres <- rqfr(1000, A, p = 2)
mean(mcres)
(anres <- qfrm(A, p = 2))
stats::t.test(mcres, mu = anres$statistic)

```

sum_counterdiag

Summing up counter-diagonal elements

Description

sum_counterdiag() sums up counter-diagonal elements of a square matrix from the upper-left; i.e., $c(X[1, 1], X[1, 2] + X[2, 1], X[1, 3] + X[2, 2] + X[3, 1], \dots)$ (or a sequence of $\sum_{i=1}^k x_{i,k-i+1}$ for $k = 1, 2, \dots$). sum_counterdiag3D() does a comparable in a 3D cubic array. No check is done on the structure of X.

Usage

```
sum_counterdiag(X)
```

sum_counterdiag3D(X)

Arguments

X Square matrix or cubic array

S_fromUL *Make covariance matrix from eigenstructure*

Description

This is an internal utility function to make covariance matrix from eigenvectors and eigenvalues. Symmetry is assumed for the original matrix.

Usage

S_fromUL(evec, evalues)

Arguments

evec Matrix whose columns are eigenvectors
 evalues Vector of eigenvalues

tr *Matrix trace function*

Description

This is an internal function. No check is done on the structure of X.

Usage

tr(X)

Arguments

X Square matrix whose trace is to be calculated

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