# Package 'FnR' 

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cients between individuals us-
ing the method of Van Vleck (2007) [https://pubmed.ncbi.nlm.nih.gov/18050089/](https://pubmed.ncbi.nlm.nih.gov/18050089/).
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## $R$ topics documented:

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\text { calcR . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . } 2
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```
calcR
    Compute numerator relationship coefficients between two distinct
    groups of individuals
```


## Description

Compute numerator relationship coefficients between two distinct groups of individuals

## Usage

calcR(ped, set1, set2, type = "notdam-notsire", f = c(), d = c())

## Arguments

ped : A data frame with integer columns corresponding to ID, SIRE, and DAM. IDs should be sequential, starting from 1. Missing parents (SIRE and DAM) are denoted as 0 .
set1 : A set of individual IDs.
set2 : A set of individual IDs, distinct from set1.
type : "notdam-notsire" (default), "sire-sire", or "dam-dam" relationships
"notdam-notsire" requires set1 and set2 individuals not to be members of ped\$DAM and ped\$SIRE, respectively.
"sire-sire" requires set1 and set2 individuals to be members of ped\$SIRE. "dam-dam" requires set1 and set2 individuals to be members of ped\$DAM.
f : (Optional) If available, the vector of inbreeding coefficients for the whole pedigree (without dummy progeny) or from the previous calculation of inbreeding coefficients with less number of animals in the pedigree.
d
: (Optional) If available, the vector of the diagonal elements of the diagonal matrix $\mathbf{D}$ in $\mathbf{A}=\mathbf{T D T}^{\prime}$ where $\mathbf{A}$ is the numerator relationship matrix, for the whole pedigree (without dummy progeny) or from the previous calculation of inbreeding coefficients with less number of animals in the pedigree.

## Value

: Numerator relationship coefficients between set1 and set2 individuals in the form of a matrix (a partition of the numerator relationship matrix $\mathbf{A}$ ).

## Examples

```
# A sample pedigree data frame:
ped <- data.frame(
    ID = 1:12,
    SIRE = c(0, 0, 0, 2, 2, 0, 4, 6, 0, 6, 10, 10),
    DAM = c(0, 0, 0, 1, 1, 0, 3, 5, 7, 8, 9, 0)
)
```

```
# Example 1: Calculate relationship coefficients between two groups of animals,
# one's members not among dams, and the members of the other not among sires.
calcR(ped, set1 = c(12, 6), set2 = c(11, 8), type = "notdam-notsire")
# Since `"notdam-notsire"` is the default type, `type = "notdam-notsire"` might be omitted.
# Example 2: Calculate relationship coefficients between dam 7 and dams 8 and 9.
calcR(ped, set1 = 7, set2 = 8:9, type = "dam-dam")
# Example 3: Calculate relationship coefficients between sires 2 & 6 and sires 4 & 10.
calcR(ped, set1 = c(2, 6), set2 = c(4, 10), type = "sire-sire")
# Example 5: Repeat example 2 with inbreeding coefficients provided.
f <- rep(0, 12)
f[10] <- 0.25
f[11] <- 0.015625
calcR(ped, set1 = 7, set2 = 8:9, type = "dam-dam", f = f)
# Example 6: Repeat example 3 with inbreeding and d coefficients provided.
d <- c(1, 1, 1, 0.5, 0.5, 1, 0.5, 0.5, 0.75, 0.5, 0.4375, 0.6875)
calcR(ped, set1 = c(2, 6), set2 = c(4, 10), type = "sire-sire", f = f, d = d)
```

resume_inbreed Calculate inbreeding coefficients from scratch or resume for new indi-
viduals in the pedigree

## Description

Calculate inbreeding coefficients from scratch or resume for new individuals in the pedigree

## Usage

resume_inbreed(ped, $f=c(), d=c()$, export_d = FALSE)

## Arguments

ped : A data frame with integer columns corresponding to ID, SIRE, and DAM. IDs should be sequential, starting from 1. Missing parents (SIRE and DAM) are denoted as 0 .
f
: (Optional) If available, the vector of inbreeding coefficients from the previous calculation of inbreeding coefficients with less number of animals in the pedigree.
d
: (Optional) If available, the vector of the diagonal elements of the diagonal matrix $\mathbf{D}$ in $\mathbf{A}=\mathbf{T D T}^{\prime}$ from the previous calculation of inbreeding coefficients with less number of animals in the pedigree, where $\mathbf{A}$ is the numerator relationship matrix.
export_d : FALSE (default) or TRUE. If TRUE, vector $d$ is retuned for future use.

## Value

: Vector of inbreeding coefficients if export_d == FALSE, or a list containing the vector of inbreeding coefficients and the vector of $d$ coefficients if export_d == TRUE.

## Examples

```
# A sample pedigree data frame:
ped <- data.frame(
    ID = 1:12,
    SIRE = c(0, 0, 0, 2, 2, 0, 4, 6, 0, 6, 10, 10),
    DAM = c(0, 0, 0, 1, 1, 0, 3, 5, 7, 8, 9, 0)
)
oldped <- ped[1:9, ]
(oldrun <- resume_inbreed(oldped, export_d = TRUE))
resume_inbreed(ped)
resume_inbreed(ped, f = oldrun$f)
resume_inbreed(ped, f = oldrun$f, d = oldrun$d)
```


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