

# Package ‘AlgebraicHaploPackage’

October 31, 2015

**Type** Package

**Title** Haplotype Two Snips Out of a Paired Group of Patients

**Version** 1.2

**Date** 2015-10-26

**Author** Jan Wolfertz

**Maintainer** Jan Wolfertz <Jan.wolfertz@uni-duesseldorf.de>

**Depends** R (>= 3.1.3)

**Suggests** compiler

**Description** Two unordered pairs of data of two different snips positions is haplotyped by resolving a small number of closed equations.

**LazyLoad** yes

**License** GPL-2

**NeedsCompilation** no

**Repository** CRAN

**Date/Publication** 2015-10-31 08:46:53

## R topics documented:

AlgebraicHaploPackage-package . . . . .	2
callhaplotype . . . . .	3
cubic . . . . .	4
findoptimal . . . . .	6
haplotypeit . . . . .	7
optimalfrequency . . . . .	8
<b>Index</b>	<b>10</b>

---

AlgebraicHaploPackage-package

*AlgebraicHaploPackage— The package haplotype a contingency of two pairs of snips of a sample.*

---

## Description

Assume a group of patients of an unordered pair of data per snips. The contingency table of 2 snips on different positions are calculated. A haplotype 2x2 most likely contingency table is guessed.

## Details

Package: AlgebraicHaploPackage  
 Type: Package  
 Version: 1.2  
 Date: 2015-10-26  
 License: GPL2.0

## Author(s)

Jan Wolfertz.

Maintainer: jan.wolfertz@uni-duesseldorf.de

## References

- [1] David Clayton. "An r package for analysis of whole-genome association studies." *Human Heredity*, 64(1):45 - 51, 2007. doi: doi:10.1001/archgenpsychiatry.2010.25. <http://archpsyc.jamanetwork.com/article.aspx?articleid=117111>
- [2] Nathan Jacobson. *Basic Algebra I: Second Edition* (Dover Books on Mathematics). Dover Publication, 2009. [3] Montgomery Slatkin Laurent Excofie. Maximum-likelihood estimation of molecular haplotype frequencies in a diploid population. *Molecular biology and evolution*, 12(5):921 - 927, 1995. URL <http://mbe.oxfordjournals.org/content/12/5/921.full.pdf>. [4] Tianhua Niu. Algorithms for inferring haplotypes. *Genetic Epidemiology*, 27:334347, 2004. doi: DOI: 10.1002/gepi.20024. URL <http://biostat.gru.edu/Journal> [5] Werner A. Stahel. *Statistische Datenanalyse Eine*. Vieweg Verlag, 2002.

## Examples

```
print("The second example: \n")
dd=matrix(c(1212, 2, 0, 679, 0,0,75,0,0), byrow=TRUE, nrow=3)
colnames(dd)=c("CC", "CT", "TT")
rownames(dd)=c("CC", "CT", "TT")
callhaplotype(dd)
### Check the result of the cubic equation of the second example
print("#####")
print("Check the result of the cubic equation of the second example: \n")
```

```

temp2haplo =as.numeric(t(dd));
t2h=temp2haplo
haplotypeit(t2h[1],t2h[2],t2h[3],t2h[4],t2h[5],t2h[6],t2h[7],t2h[8],t2h[9]);
rm(temp2haplo)
rm(t2h)
### Third example
print("#####")
print("Third example : \n")
dd3=matrix(c(1030,678,123,1,1,0,0,0,0),ncol=3,byrow=TRUE)
colnames(dd3)=c("AA","AG","GG")
rownames(dd3)=c("CC","CT","TT")
callhaplotype(dd3)
### Check for alternative solutions
print("#####")
print("Check for alternative solutions: \n")
temp2haplo =as.numeric(t(dd3));
t2h=temp2haplo;
haplotypeit(t2h[1],t2h[2],t2h[3],t2h[4],t2h[5],t2h[6],t2h[7],t2h[8],t2h[9]);
rm(temp2haplo)
rm(t2h)
print("#####")
print("#####")
print("This tests the result of the first example of the article \n")
dd2=matrix(c(4,0,0,0,30,0,0,0,23),ncol=3,byrow=TRUE)
callhaplotype(dd2)
callhaplotype(dd2)/(2*57)
print("#####")

```

---

callhaplotype

*calculates the cotigency table of the haplotypes*


---

### Description

It starts with a contingency table of pairs of haplotypes and ends up with the haplotypes one. The heterocygote cases are the middle of the column and rows.

### Usage

```
callhaplotype(dd)
```

### Arguments

dd                    This is a contingency table. Rows and columns are in the order are AA, AB, BB.

### Details

A 2x2 contingency table of haplotypes is calculated. The most likely solution had been choosen.

**Value**

The haplotype contingency table is returned. All entries are numeric.

**Note**

The differences are the coice of the solution of the cubic equations. About 4 percent differences and about 7 assuming 1 per thousand. For data export or import you can use a different package.

**Author(s)**

Jan Wolfertz

**References**

David Clayton. "An r package for analysis of whole-genome association studies." *Human Heredity*, 64(1):45 - 51, 2007. doi: doi:10.1001/archgenpsychiatry.2010.25. URL <http://archpsyc.jamanetwork.com/article.aspx?articleid=117111>  
Jan Wolfertz(in press.):""

**Examples**

```
print("#####")
dd2=matrix(c(4,0,0,0,30,0,0,0,23),ncol=3,byrow=TRUE)
callhaplotype(dd2)
callhaplotype(dd2)/(2*57)

### The second example
print("#####")
print("The second example: \n")
dd=matrix(c(1212, 2, 0, 679, 0,0,75,0,0), byrow=TRUE, nrow=3)
colnames(dd)=c("CC", "CT", "TT")
rownames(dd)=c("CC", "CT", "TT")
callhaplotype(dd)
print("#####")
```

---

cubic

*Function that can resolve the cubic equation numerical stable and any lower dimensional case except unsolvable cases.*

---

**Description**

$A*x^3+B*x^2+C*x+D=0$ . All coefficients had to be numeric or integers. This function calculates from 4 coefficient all possible and senfully solutions.  $D=0$  returns no values at all. This would be a impossibel case. It returns upto 3 potential complex solutions. Less solutions are copied to get the tripple solution format.

**Usage**

cubic(A, B, C, D)

**Arguments**

A	The coefficient of $x^3$ .
B	The coefficient of $x^2$ .
C	The coefficient of $X$ .
D	The constant.

**Details**

This function is called by haplotypeit. The results are returned as vector of the three possible solutions: output[1],output[2],output[3]. Further data for checks of the roots. p,q and the discriminat. 10 and 11 are only usable for symmetry checks.

**Value**

Returns cubic(A,B,C,D)[c(1:3)] roots of the at most cubic equation.

**Note**

Using cardenian formular, a well known method.

**Author(s)**

Jan Wolfertz

**References**

Cardans formular as in e.g. The Mathematical Gazette (1993); 77 (Nov, No 480), 354-359 (jstor)  
<http://www.nickalls.org/dick/papers/math/cubic1993.pdf> or any other book for algebraic solutions.  
See also : [http://de.wikipedia.org/wiki/Cardanische\\_Formeln](http://de.wikipedia.org/wiki/Cardanische_Formeln) and [http://en.wikipedia.org/wiki/Cubic\\_equation](http://en.wikipedia.org/wiki/Cubic_equation)

**See Also**

haplotypeit,callhaplotype

**Examples**

```
cubic(1,0,0,-1)[c(1:3)]  
cubic(1,1,0,0)[c(1:3)]
```

---

 findoptimal

*Chose the most likely solution of the three potential onces.*


---

### Description

Starting with a 3x3 matrix, three potential haplotypes 2x2 matrices will be calculated and evaluated. The most likely one is chosen. A discrete solution values are not enforced. This reduces increases the right prediction on real data.

### Usage

```
findoptimal(A, B, C, D, mmorg, exact = 1e-05)
```

### Arguments

A	First entry of the 2x2 matrix.
B	Second numeric entry of the 2x2 matrix.
C	Third numeric entry of the 2x2 matrix.
D	Last numeric entry of the 2x2 matrix.
mmorg	3x3 matrix of the pair of original snip pairs.
exact	exact is a parameter when data are assumed to be equal. Actually 1e-5 is taken. Should not be larger than the inverse of four times the number of people.

### Details

It chose the 2x2 model of haplotypes with the smallest prediction error.

### Value

AA	Coefficient of $x^3$
BB	Coefficient of $x^2$
CC	Coefficient of $x$
DD	Coefficient of the intercept

### Author(s)

Jan wolfertz

### Examples

```
dd2=matrix(c(4,0,0,0,30,0,0,0,23),ncol=3,byrow=TRUE)
A=c(38+0i,2+12.1655i,2-12.1655i)
B=c(0+0i,36-12.1655i,36+12.1655i)
C=c(0+0i,36-12.1655i,36+12.1655i)
D=c(76+0i,40+12.1655i,40-12.1655i)
```

---

`haploypeit`*Haplotype a 3 x 3 counting matrix.*

---

**Description**

This functions recalculates the potential 2x2 haplotype matrices. It gets a 3x3 matrix and returns a list A,B,C,D of vectors. A[1],B[1],C[1],D[1] is the first solution of the matrix. There are always three solutions.

**Usage**

```
haploypeit(a, b, c, d, e, f, g, h, i)
```

**Arguments**

a	Number of counts of matching snip pairs.
b	Number of counts of matching snip pairs..
c	Number of counts of matching snip pairs.
d	Number of counts of matching snip pairs.
e	Number of counts of matching snip pairs.
f	Number of counts of matching snip pairs.
g	Number of counts of matching snip pairs.
h	Number of counts of matching snip pairs.
i	Number of counts of matching snip pairs.

**Details**

The software automatically resolves the cases  $e=0$  by circumventing the cubic equation. If the degree of the equation is lower additional copies of some solution will be made to produce the outputformat. The output format is a list of four vectors of coefficients. Each vector contains three complex numbers.

**Value**

output\$A is a vector of length 3. output\$B, output\$C, output\$D is a vector of length 3. One potential solution is A[1],B[1],C[1],D[1].

**Author(s)**

Jan wolfertz

**References**

This methods refers to an article: David Clayton. An r package for analysis of whole- genome association studies. Human Heredity, 64(1):45 - 51, 2007. doi: doi:10.1001/archgenpsychiatry.2010.25. URL <http://archpsyc.jamanetwork.com/article.aspx?articleid=210679>.

**See Also**

callhaplotype

**Examples**

```
haplotypeit(4,0,0,0,30,0,0,0,23)
print("#####")
print("This tests the cubic routine")
haplotypeit(4,0,0,0,30,0,0,0,23)
### Formated of 4 digits
print("Formated of 4 digits")
round(as.numeric(Re(haplotypeit(4,0,0,0,30,0,0,0,23)$A)),digit=4)
round(as.numeric(Re(haplotypeit(4,0,0,0,30,0,0,0,23)$B)),digit=4)
round(as.numeric(Re(haplotypeit(4,0,0,0,30,0,0,0,23)$C)),digit=4)
round(as.numeric(Re(haplotypeit(4,0,0,0,30,0,0,0,23)$D)),digit=4)
###
```

---

optimalfrequency      *Evaluate potential haplotypes.*

---

**Description**

Calculate the difference of a known haplotype and the resulting unordered pair of snip pairs.

**Usage**

```
optimalfrequency(mm, mmorg)
```

**Arguments**

mm	This is a contingency table of two haplotype snips. 2x2 matrix or ndata.frame
mmorg	This is a contingency table of two diplotype snips pairs.

**Details**

The average squared distance to the expected result 3x3 table is used as a T statistic. The p value not to be zero is calculated. The higher the p value the more exact is the haplotype.

**Value**

A list of values is returned.

result\$*LK*      Linkage disequilibrium

result\$*Testvalue*

The squared sitance multiplied by the number of entries in 3x3 matrix mmorg

result\$*prSimilarByChange*

The probability not o be equal to zero by change.

*optimalfrequency*

9

**Author(s)**

Jan wolfertz.

**References**

Stahel: Statistik fuer Naturwissenschaftler und Mediziner, pp. 107-120.

**See Also**

findoptimal

# Index

- \*Topic **'model-fit'**
    - optimalfrequency, 8
  - \*Topic **cubic**
    - callhaplotype, 3
    - cubic, 4
  - \*Topic **functions**
    - cubic, 4
  - \*Topic **haplotypeit**
    - callhaplotype, 3
  - \*Topic **haplotype**
    - optimalfrequency, 8
  - \*Topic **haplotyping**
    - callhaplotype, 3
    - findoptimal, 6
    - haplotypeit, 7
  - \*Topic **package**
    - AlgebraicHaploPackage-package, 2
- AlgebraicHaploPackage  
(AlgebraicHaploPackage-package),  
2
- AlgebraicHaploPackage-package, 2
- callhaplotype, 3  
cubic, 4
- findoptimal, 6
- haplotypeit, 7
- optimalfrequency, 8