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BCRA-package

A Package for Breast Cancer Risk Assessment

Description

This package is to project absolute risk of invasive breast cancer according to NCI's Breast Cancer Risk Assessment Tool (BCRAT) algorithm for specified race/ethnic groups and age intervals. The updated version 2.0 includes the new Hispanic model.

Details

This package can be used to estimate the risk of developing breast cancer over a predetermined time interval with risk factors. As the same as Breast Cancer Risk Assessment SAS Macro, the users can specify the time interval as appropriate, not only limited to the 5 years risk prediction available with BCRAT.

The main function in this package is absolute.risk, which is defined based on a statistical model known as the "Gail model". Parameters and constants needed in this function include initial and projection age, recoded covariates using function recode.check, relative risks of BrCa at age "<50" and ">=50" obtained from function relative.risk as well as other known constants listed from function list.constants like BrCa composite incidences, competing hazards, 1-attributable risk using in NCI BrCa Risk Assessment Tool (NCI BCRAT). With risk factors and projection interval ages for a group of women, the function absolute.risk will return the corresponding absolute risk projections. If the function returns any missing values, the function error.table or error.table.all is used to find where the errors occured. The function check.summary can give a quick check for errors of input file and missing values of risks.

For further analysis, a data frame is created from the function risk.summary, which includes age, duration of the projection time interval, covariates and the projected risk.

The version 2.0 includes absolute risk projections for Hispanic women (US born and Foreign born) based on race specific RR risk models developed on the San Francisco Bay Area Breast Cancer Study (SFBCS). Race specific attributable risks, breast cancer composite incidences and competing hazards are added to the updated package.

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Description

A function to estimate absolute risks of developing breast cancer

Usage

absolute.risk(data, Raw_Ind=1, Avg_White=0)

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.
Avg_White	Calculation indicator. Avg_White=0, calculate absolute risks; Avg_White=1, calculate average absolute risks based on the rates for average non-hispanic white women and average other (native american) women. The default value is 0.

Details

For the projection of absolute risks, this function is defined based on Gail Model. Parameters and constants needed in this function include initial and projection age, recoded covariates from function recode.check, relative risks of BrCa at age "<50" and ">=50" from function relative.risk as well as other known constants like BrCa composite incidences, competing hazards, 1-attributable risk using in NCI BrCa Risk Assessment Tool (NCI BCRAT).

Value

A vector which returns absolute risk values when Avg_White=0 or average absolute risk values when Avg_White=1.

See Also

recode.check,relative.risk

Examples

```
data(exampledata)
# calculate absolute risk
absolute.risk(exampledata)
# calculate average absolute risk
Avg_White <- 1
absolute.risk(exampledata, Raw_Ind=1, Avg_White)</pre>
```

BrCa_1_AR

Breast cancer 1-Attributable Risk

Description

1-Attributable Risk

Usage

data("BrCa_1_AR")

Format

A data frame with 2 observations on the following 5 variables.

Wh.Gail White

- AA.CARE African-American
- HU.Gail Hispanic-American (US born)
- NA.Gail Other (Native American and unknown race)
- HF.Gail Hispanic-American (Foreign born)
- Asian.AABCS Asian-American

BrCa_beta

Description

The logistic regression coefficients derived from the Gail model.

Usage

data("BrCa_beta")

Format

A data frame with 6 observations on the following 5 variables.

Wh.Gail White, Gail model

AA.CARE African-American, Care model

HU.Gail Hispanic-American (US born), Gail model

NA.Gail Other (Native American and unknown race), Gail model

HF.Gail Hispanic-American (Foreign born), Gail model

Asian.AABCS Asian-American, AABCS model

BrCa_lambda1 Breast cancer composite incidences

Description

Breast cancer composite incidences for different races and age groups from 20 to 90 by 5 years.

Usage

data("BrCa_lambda1")

Format

A data frame with 14 age groups on the following 12 variables.

Wh.1983_87 White SEER 1983:1987

AA.1994_98 African-American SEER 1994:1998

HU.1995_04 Hispanic-American (US born) 1995:2004

NA.1983_87 Native American and unknown race 1983:1987

HF.1995_04 Hispanic-American (Foreign born) 1995:2004

Ch.1998_02 Chinese-American SEER 18 1998:2002

Ja.1998_02 Japanese-American SEER 18 1998:2002 Fi.1998_02 Filipino-American SEER 18 1998:2002 Hw.1998_02 Hawaiian SEER 18 1998:2002 oP.1998_02 Other Pacific Islander SEER 18 1998:2002 oA.1998_02 Other Asian SEER 1998:2002 Wh_Avg.1992_96 Average White SEER 1992:1996

BrCa_lambda2 Breast cancer competing mortality

Description

Breast cancer competing mortality for different races and age groups from 20 to 90 by 5 years.

Usage

data("BrCa_lambda2")

Format

A data frame with 14 age groups on the following 12 variables.

Wh.1983_87 White SEER 1983:1987

- AA.1994_98 African-American SEER 1994:1998
- HU.1995_04 Hispanic-American (US born) 1995:2004
- NA.1983_87 Native American and unknown race 1983:1987
- HF.1995_04 Hispanic-American (Foreign born) 1995:2004
- Ch.1998_02 Chinese-American SEER 18 1998:2002
- Ja. 1998_02 Japanese-American SEER 18 1998:2002
- Fi.1998_02 Filipino-American SEER 18 1998:2002
- Hw.1998_02 Hawaiian SEER 18 1998:2002
- oP.1998_02 Other Pacific Islander SEER 18 1998:2002
- oA.1998_02 Other Asian SEER 1998:2002
- Wh_Avg.1992_96 Average White SEER 1992:1996

check.summary

Description

A function to show descriptive statistics by applying function mean and sd to the quantities Error_Ind, AbsRisk, RR_Star1 and RR_Star2.

Usage

check.summary(data, Raw_Ind=1, Avg_White=0)

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.
Avg_White	Calculation indicator. Avg_White=0, calculate absolute risks; Avg_White=1, calculate average absolute risks based on the rates for average non-hispanic white women and average other (native american) women. The default value is 0.

Details

When the mean and standard deviation for the variable Error_Ind is 0, implies that no errors have not been found. Otherwise when the mean and std for Error_Ind is not 0, implies that errors have been found. When errors are found, the number of records with errors is the count associated with AbsRisk listed under NMiss (number of missing).

Value

A summary table for error indicators, relative risks and absolute risks

See Also

recode.check, relative.risk, absolute.risk

```
error.table
```

Description

A function to list the records and errors for IDs with missing absolute risks. For each of the records with error, the record is listed followed by a line which gives some indication as to where the error occured. Relative risks and risk pattern numbers are also included.

Usage

```
error.table(data, Raw_Ind=1)
```

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.

Value

A data frame listing the raw records, errors, relative risks and pattern numbers for IDs with missing absolute risks. If there is nothing wrong with the input data, the function will return "NO ERROR!".

See Also

```
recode.check,error.table.all
```

error.table.all List all records and errors

Description

A function to list all records with both raw values and recoded values (or indications for errors). For each of the records, the record is listed followed by a line which gives some indication as to where the error occured.

Usage

```
error.table.all(data, Raw_Ind=1)
```

exampledata

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.

Value

A data frame listing all records and errors. If there is nothing wrong with the input data, the function will return "NO ERROR!".

See Also

recode.check,error.table

exampledata

Example data set

Description

A data set containing all the required input data needed to perform risk projections, such as initial age, projection age, BrCa relative risk covariates and race.

Usage

data("exampledata")

Format

A data frame with 26 observations on the following 9 variables.

- ID Woman's ID, positive integer 1, 2, 3,...
- T1 Initial age, all real numbers T1 in [20, 90).
- T2 BrCa projection age, all real numbers T2 in (20,90] such that T1<T2.

N_Biop The number of biopsies, 0, 1, 2,..., 99=unk (99 recoded to 0).

HypPlas Did biopsy display atypical hyperplasia? 0=no, 1=yes, 99=unk or not applicable.

AgeMen Age at menarchy, less than or equal to initial age, 99=unk.

Age1st Age at first live birth, greater or equal to age at menarchy and less than or equal to initial age, 98=nulliparous, 99=unk.

N_Rels The number of 1st degree relatives with BrCa, 0, 1, 2,... 99=unk.

Race Race, positive integer 1, 2, 3,...,11. See details.

Details

1=Wh	White 1983-87 SEER rates (rates used in NCI BCRAT)
2=AA	African-American
3=HU	Hispanic-American (US born) 1995-04
4=NA	Other (Native American and unknown race)
5=HF	Hispanic-American (Foreign born) 1995-04
6=Ch	Chinese-American
7=Ja	Japanese-American
8=Fi	Filipino-American
9=Hw	Hawaiian-American
10=oP	Other Pacific Islander
11=0A	Other Asian

list.constants

List all constants required for BrCa absolute risk projections

Description

A function to create a text file under user's working directory which contains all constants required for BrCa absolute risk projections.

Usage

list.constants(BrCa_lambda1, BrCa_lambda2, BrCa_beta, BrCa_1_AR)

Arguments

BrCa_lambda1	Breast Cancer Composite Incidences
BrCa_lambda2	Breast Cancer Competing Mortality
BrCa_beta	The logistic regression coefficients (beta) derived from the Gail model
BrCa_1_AR	1-Attributable Risk

Details

See "BrCa_lambda1.rda", "BrCa_lambda2.rda", "BrCa_beta.rda", "BrCa_1_AR.rda" in package data folder.

Value

A text file "list_all_constants.txt" exported under user's working directory for reading convenience.

recode.check

Description

A function to recode the relative risk covariates and check errors.

Usage

```
recode.check(data, Raw_Ind=1)
```

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.

Details

This function is to recode the following relative risk covariates. Recoded RR covariates are named as NB_Cat, AM_Cat, AF_Cat and NR_Cat for N_Biop, AgeMen, Age1st and N_Rels, respectively.

N_Biop:	The number of biopsies.
AgeMen:	Age at menarchy.
Age1st:	Age at first live birth.
N_Rels:	The number of first degree relatives with BrCa.

See the following table for recoding details.

Covariate	Raw Value	Recoded to
N_Biop	0 or 99 (unk or not applicable)	0
- 1	1	1
	2,3,4 and not 99	2
AgeMen	14.15.16 or 99 (unk)	0
0	12,13	1
	11 and younger	2
Age1st	19 and younger or 99 (unk)	0
0	20,21,22,23,24	1
	25,26,27,28,29 or 98 (nulliparous)	2
	30,31,32 and not 98 and not 99	3

recode.check

N_Rels	0 or 99 (unk)	0
	1	1
	2,3,4 and not 99	2

This function is also used to check consistency and errors of input data. Let $set_T1_missing$ and $set_T2_missing$ be the checking variables for T1 and T2. The constraint on T1 and T2 is 20 <= T1 < T2 <= 90. If it is violated, $set_T1_missing$ and $set_T2_missing$ and the absolute risk will be set to the missing value NA.

Let RacCat be the checking variable for Race. If the Race value is not included in the 11 races defined, the absolute risk will be set to the missing value NA and RacCat will be set to "U" (undefined). The corresponding character of Race CharRace will be set to "??".

Let set_HyperP_missing and set_R_Hyp_missing be the checking variables for HypPlas and R_Hyp. Consistency patterns for the number of Biopsies and Hyperplasia are:

Requirment (A) N_Biops=0 or 99, then HypPlas MUST = 99 (not applicable). Requirment (B) N_Biops>0 and <99, then HypPlas = 0, 1 or 99.

If ANY of the above 2 REQUIREMENTS is violated, NB_Cat, set_HyperP_missing and set_R_Hyp_missing will be set to the corresponding character "A" or "B" and the absolute risk will be set to the missing value NA. The consequences to the relative risk (RR) for the above two requirements are:

(A) N_Biops=0 or 99, HypPlas=99 (not applicable) inflates RR by 1.00.

(B) N_Biops>0 and <99, HypPlas=0 (no) inflates RR by 0.93;

N_Biops>0 and <99, HypPlas=1 (yes) inflates RR by 1.82;

N_Biops>0 and <99, HypPlas=99 (unk) inflates RR by 1.00.

For remaining relative risk covariates, AgeMen, Age1st and N_Rels:

AgeMen	Age at menarchy must be postive integer less than or equal to initial age T1. NOTE: (1) For African-American women AgeMen<=11 are grouped with AgeMen=12 or 13. (2) For US Born Hispanic women AgeMen is not included in the RR model
	and all values for this variable are recoded to 0.
Age1st	Age at 1st live birth must be postive integer greater than equal to AgeMen and less than or equal to initial age T1.
	NOTE: (1) For African-American women, Age1st is not included in the RR model and all values for this variable are recoded to 0. (2) For US Born and Foreign Born Hispanic women, the recoding for this variable follows:

Age1st	19 and younger or 99 (unk)	0
	20 - 29	1
	30+ or 98 (nulliparous) and not 99	2

relative.risk

N_Rels The number of 1st degree relatives with BrCa must be 0,1,2.... NOTE: For Asian-Americans Race=6-11 and Hispanic-Americans (US and foreign born), the number of 1st degree relative coded value of 2 gets grouped with 1.

Value

A data frame containing the error indictors, recoded covariates as well as other checking variables defined for checking the consistency of the input data.

See Also

error.table.all, error.table

Examples

data(exampledata)
recode.check(exampledata)

relative.risk Estimate relative risks

Description

A function to estimate relative risks for risk factor combinations

Usage

```
relative.risk(data, Raw_Ind=1)
```

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.

Details

The age is dichotomized as "age less than 50 years" and "age 50 years or more". The relative risks can be obtained from Gail Model, an unconditional logistic regression that included main effects NB_Cat, AM_Cat, AF_Cat, NR_Cat as well as interactions between AF_Cat and NR_Cat and between the age category and NR_Cat.

Value

RR_Star1	Relative risk for woman of interest at $ages < 50$.
RR_Star2	Relative risk for woman of interest at ages ≥ 50 .
PatternNumber	The sequence number of risk patterns. There are 3 levels for NB_Cat, 3 for AM_Cat, 4 for AF_Cat, 3 for NR_Cat, 3*3*4*3 = 108 patterns in total. Pattern Number=NB_Cat*3*3*4+AM_Cat*3*4+AF_Cat*3+NR_Cat*1+1.

See Also

recode.check

Examples

```
data(exampledata)
relative.risk(exampledata)
```

```
risk.summary
```

List the records with relative risks and absolute risks

Description

A function to list all the records with relative risks and absolute risks.

Usage

risk.summary(data, Raw_Ind=1)

Arguments

data	A data set containing all the required input data needed to perform risk projec- tions, such as initial age, projection age, BrCa relative risk covariates and race. See exampledata for details.
Raw_Ind	The raw file indicator with default value 1. Raw_Ind=1 means RR covariates are in raw/original format. Raw_Ind=0 means RR covariates have already been re-coded to 0, 1, 2 or 3.

Value

A data frame that includes age, duration of the projection time interval, covariates and the projected risk. A CSV file is created to save the data frame under user's working directory for reading convenience.

See Also

relative.risk, absolute.risk

risk.summary

Examples

data(exampledata)
risk.summary(exampledata)

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