

Package ‘iCARE’

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Title A Tool for Individualized Coherent Absolute Risk Estimation
(iCARE)

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Description An R package to compute Individualized Coherent Absolute Risk Estimators.

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Depends R (>= 3.3.0)

Suggests RUnit, BiocGenerics

License GPL-3 + file LICENSE

LazyData true

biocViews Software, StatisticalMethod, GenomeWideAssociation

NeedsCompilation yes

R topics documented:

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bc_data	<i>Data for examples</i>
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Description

Data for [computeAbsoluteRisk](#) and [computeAbsoluteRiskSplitInterval](#)

Details

"mort_inc" contains age-specific incidence rates of all-cause mortality from reference (1) below

"bc_15_snps" contains published SNP information from reference (2)

"bc_inc" contains age-specific incidence rates of breast cancer from reference (3)

"ref_cov_dat" contains a subsample of data imputed using reference (4) and (5)

References

- (1) Centers for Disease Control and Prevention (CDC), National Center for Health Statistics (NCHS). Underlying Cause of Death 1999-2011 on CDC WONDER Online Database, released 2014. Data are from the Multiple Cause of Death Files, 1999-2011, as compiled from data provided by the 57 vital statistics jurisdictions through the Vital Statistics Cooperative Program. Accessed at <http://wonder.cdc.gov/ucd-icd10.html> on Aug 26, 2014.
- (2) Michailidou K, Beesley J, Lindstrom S, et al. Genome-wide association analysis of more than 120,000 individuals identifies 15 new susceptibility loci for breast cancer. *Nature genetics* 2015;47:373-80.
- (3) Surveillance, Epidemiology, and End Results (SEER) Program SEER*Stat Database: Incidence - SEER 18 Regs Research Data, Nov 2011 Sub, Vintage 2009 Pops (2000-2009) <Katrina/Rita Population Adjustment> - Linked To County Attributes - Total U.S., 1969-2010 Counties. In: National Cancer Institute D, Surveillance Research Program, Surveillance Systems Branch, ed. SEER18 ed.
- (4) 2010 National Health Interview Survey (NHIS) Public Use Data Release, NHIS Survey Description. 2011. Accessed at ftp://ftp.cdc.gov/pub/Health_Statistics/NCHS/Dataset_Documentation/NHIS/2010/srvydesc.pdf.
- (5) Centers for Disease Control and Prevention (CDC). National Center for Health Statistics (NCHS). National Health and Nutrition Examination Survey Questionnaire. Hyattsville, MD: U.S. Department of Health and Human Services, Centers for Disease Control and Prevention; 2010.

Examples

```
temp <- data(bc_data, package="iCARE")

# Display the object names
temp
```

computeAbsoluteRisk *Building and Applying an Absolute Risk Model*

Description

This function is used to build absolute risk models and apply them to estimate absolute risks.

Usage

```
computeAbsoluteRisk(model.formula = NULL, model.cov.info = NULL,
  model.snp.info = NULL, model.log.RR = NULL, model.ref.dataset = NULL,
  model.ref.dataset.weights = NULL, model.disease.incidence.rates,
  model.competing.incidence.rates = NULL, model.bin.fh.name = NA,
  n.imp = 5, apply.age.start, apply.age.interval.length,
  apply.cov.profile = NULL, apply.snp.profile = NULL, use.c.code = 1,
  return.lp = FALSE, return.refs.risk = FALSE)
```

Arguments

- `model.formula` an object of class formula: a symbolic description of the model to be fitted, e.g. `Y~Parity+FamilyHistory`.
- `model.cov.info` contains information about the risk factors in the model ; a main list containing a list for each covariate, which must have the fields:
- "name" : a string with the covariate name, matching name in `model.formula`
 - "type" : a string that is either "continuous" or "factor".
- If factor variable, then:
- "levels" : vector with strings of level names
 - "ref" : optional field, string with name of referent level
- `model.snp.info` dataframe with three columns, named: ["snp.name", "snp.odds.ratio", "snp.freq"]
- `model.log.RR` vector with log odds ratios corresponding to the model params; no intercept; names must match design matrix arising from `model.formula` and `model.cov.info`; check names using function `check_design_matrix()`.
- `model.ref.dataset` dataframe of risk factors for a sample of subjects representative of underlying population, no missing values. Variables must be in same order with same names as in `model.formula`.
- `model.ref.dataset.weights` optional vector of sampling weights for `model.ref.dataset`.
- `model.disease.incidence.rates` two column matrix [integer ages, incidence rates] or three column matrix [start age, end age, rate] with incidence rate of disease. Must fully cover age interval for estimation.
- `model.competing.incidence.rates` two column matrix [integer ages, incidence rates] or three column matrix [start age, end age, rate] with incidence rate of competing events. Must fully cover age interval for estimation.
- `model.bin.fh.name` string name of family history variable, if in model. This must refer to a variable that only takes values 0,1, NA.
- `n.imp` integer value for number of imputations for handling missing SNPs.
- `apply.age.start` single integer or vector of integer ages for the start of the interval over which to compute absolute risk.
- `apply.age.interval.length` single integer or vector of integer years over which absolute risk should be computed.
- `apply.cov.profile` dataframe containing the covariate profiles for which absolute risk will be computed. Covariates must be in same order with same names as in `model.formula`.
- `apply.snp.profile` data frame with observed SNP data (coded 0,1, 2, or NA). May have missing values.


```

summary(results$risk)
plot(density(results$risk, na.rm=TRUE))
boxplot(results$risk ~ new_cov_prof$famhist, na.rm=TRUE)

apply.snp.profile = new_snp_prof,
return.refs.risk = TRUE)

```

```
computeAbsoluteRiskSplitInterval
```

Building and Applying an Absolute Risk Model: Compute Risk over Interval Split in Two Parts

Description

This function is used to build an absolute risk model that incorporates different input parameters before and after a given time point. The model is then applied to estimate absolute risks.

Usage

```

computeAbsoluteRiskSplitInterval(apply.age.start, apply.age.interval.length,
  apply.cov.profile, model.formula, model.disease.incidence.rates, model.log.RR,
  model.ref.dataset, model.ref.dataset.weights=NULL, model.cov.info, use.c.code=1,
  model.competing.incidence.rates=NULL, return.lp=FALSE, apply.snp.profile=NULL,
  model.snp.info=NULL, model.bin.fh.name=NULL, cut.time=NULL,
  apply.cov.profile.2=NULL, model.formula.2=NULL, model.log.RR.2=NULL,
  model.ref.dataset.2=NULL, model.ref.dataset.weights.2=NULL, model.cov.info.2=NULL,
  model.bin.fh.name.2=NULL, n.imp=5, return.refs.risk=FALSE)

```

Arguments

`apply.age.start`

single integer or vector of integer ages for the start of the interval over which to compute absolute risk.

`apply.age.interval.length`

single integer or vector of integer years over which absolute risk should be computed.

`apply.cov.profile`

dataframe containing the covariate profiles for which absolute risk will be computed. Covariates must be in same order with same names as in `model.formula`.

`model.formula` an object of class `formula`: a symbolic description of the model to be fitted, e.g. `Y~Parity+FamilyHistory`.

`model.disease.incidence.rates`

two column matrix [integer ages, incidence rates] or three column matrix [start age, end age, rate] with incidence rate of disease. Must fully cover age interval for estimation.

`model.log.RR`

vector with log odds ratios corresponding to the model params; no intercept; names must match design matrix arising from `model.formula` and `model.cov.info`; check names using function `check_design_matrix()`.

`model.ref.dataset`

dataframe of risk factors for a sample of subjects representative of underlying population, no missing values. Variables must be in same order with same names as in `model.formula`.

`model.ref.dataset.weights`
 optional vector of sampling weights for `model.ref.dataset`.

`model.cov.info` contains information about the risk factors in the model ; a main list containing a list for each covariate, which must have the fields:

- "name" : a string with the covariate name, matching name in `model.formula`
- "type" : a string that is either "continuous" or "factor".

If factor variable, then:

- "levels" : vector with strings of level names
- "ref" : optional field, string with name of referent level

`use.c.code` binary indicator of whether to run the c program for fast computation.

`model.competing.incidence.rates`
 two column matrix [integer ages, incidence rates] or three column matrix [start age, end age, rate] with incidence rate of competing events. Must fully cover age interval for estimation.

`return.lp` binary indicator of whether to return the linear predictor for each subject in `apply.cov.profile`.

`apply.snp.profile`
 data frame with observed SNP data (coded 0,1, 2, or NA). May have missing values.

`model.snp.info` dataframe with three columns [rs number, odds ratio, allele frequency]

`model.bin.fh.name`
 string name of family history variable, if in model. This must refer to a variable that only takes values 0,1, NA.

`cut.time` integer age for which to split computation into before and after

`apply.cov.profile.2`
 see `apply.cov.profile`, to be used for estimation in ages after the cutpoint

`model.formula.2`
 see `model.formula`, to be used for estimation in ages after the cutpoint

`model.log.RR.2` see `model.log.RR`, to be used for estimation in ages after the cutpoint

`model.ref.dataset.2`
 see `model.ref.dataset`, to be used for estimation in ages after the cutpoint

`model.ref.dataset.weights.2`
 see `model.ref.dataset.weights`, to be used for estimation in ages after the cutpoint

`model.cov.info.2`
 see `model.cov.info`, to be used for estimation in ages after the cutpoint

`model.bin.fh.name.2`
 see `model.bin.fh.name`, to be used for estimation in ages after the cutpoint

`n.imp` integer value for number of imputations for handling missing SNPs.

`return.refs.risk`
 binary indicator of whether to return the absolute risk prediction for each subject in `model.ref.dataset`.


```
summary(results$risk)
plot(density(results$risk, na.rm=TRUE))
boxplot(results$risk ~ new_cov_prof$famhist, na.rm=TRUE)

apply.cov.profile = new_cov_prof,
apply.snp.profile = new_snp_prof,
return.refs.risk = TRUE)
```

iCARE

A Tool for Individualized Coherent Absolute Risk Estimation (iCARE)

Description

Individualized Coherent Absolute Risk Estimators (iCARE) is a tool that allows researchers to quickly build models for absolute risk and apply them to estimate individuals' risk based on a set of user defined input parameters. The software gives users the flexibility to change or update models rapidly based on new risk factors or tailor models to different populations based on the specification of simply three input arguments: (1) a model for relative risk assumed to be externally derived (2) an age-specific disease incidence rate and (3) the distribution of risk factors for the population of interest. The tool can handle missing information on risk factors for risk estimation using an approach where all estimates are derived from a single model through appropriate model averaging.

Details

The main functions for building and applying an absolute risk model are [computeAbsoluteRisk](#) and [computeAbsoluteRiskSplitInterval](#). The first of these computes absolute risks over the specified time interval using a single set of parameters. The second provides more advanced functionality and computes absolute risk over the interval in two parts. [computeAbsoluteRiskSplitInterval](#) allows the user compute absolute risk over the interval in two parts, incorporating two different sets of parameters before and after a specified cutpoint. This function allows a different cutpoint for each covariate profile if desired.

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