Package ‘InterVA5’

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InterVA5-package

Perform InterVA5 algorithm and provide graphical summarization of COD distribution.

Description

Computes individual cause of death and population cause-specific mortality fractions using the InterVA5 algorithm. Provides a simple graphical representation of the result.

Details

To get the most up-to-date version of the package, as well as the past versions, please check the github repository at: https://github.com/jarathomas/InterVA5-R-Replicate/

Package: InterVA5
Type: Package
Version: 1.0
Date: 2018-02-01
License: GPL-3

Author(s)

Jason Thomas, Zehang Li, Tyler McCormick, Sam Clark
Maintainer: Jason Thomas <jarathomas@gmail.com>

References

http://www.interva.net/
Description

This is the translation of COD abbreviation codes into their corresponding full names.

Format

A data frame with the translation of codes to their names for 3 pregnancy statuses, 61 CODs (both the version of COD only and COD with group code), and 6 circumstances of mortality (COMCAT).

Examples

data(causetextV5)

COMCAT.interVA5  Summarize population level mortality fraction by Circumstance of Mortality Category

Description

The function takes input of a list of va object and calculates the mortality fraction by Circumstance of Mortality Category.

Usage

COMCAT.interVA5(va)

Arguments

va  The list of va object to summarize.

Value

dist.cod  The cause-specific mortality fraction (including undetermined category).

Author(s)

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

See Also

CSMF5
Examples

```r
## Not run:
data(RandomVA5)
# only fit first 20 observations for a quick illustration
RandomVA5 <- RandomVA5[1:20, ]

sample.output <- InterVA5(RandomVA5, HIV = "h", Malaria = "v",
                          write = TRUE, directory = tempdir(),
                          filename = "VA5_result", output = "extended", append = FALSE)
## Get CSMF without plots
comcat <- COMCAT.interVA5(sample.output$VA5)
## End(Not run)
```

---

**CSMF.interVA5**

Summarize population level cause-specific mortality fraction as InterVA5 suggested.

**Description**

The function takes input of a list of va object and calculates the cause-specific mortality fraction. It only calculates CSMF5 as aggregation of up to the third largest causes.

**Usage**

```r
CSMF.interVA5(va)
```

**Arguments**

- `va` The list of va object to summarize.

**Value**

- `dist.cod` The cause-specific mortality fraction (including undetermined category).

**Author(s)**

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

**See Also**

CSMF5
Examples

```r
## Not run:
data(RandomVA5)  
# only fit first 20 observations for a quick illustration
RandomVA5 <- RandomVA5[1:20, ]

sample.output <- InterVA5(RandomVA5, HIV = "h", Malaria = "v", write=TRUE,
                          directory = tempdir(), filename = "VA5_result", output = "extended", append = FALSE)
## Get CSMF without plots
csmf <- CSMF.interVA5(sample.output$VA5)
## End(Not run)
```

CSMF5

Summarize and plot a population level distribution of va probabilities.

Description

The function takes input of a list of va object and produces a summary plot for the population distribution.

Usage

```r
CSMF5(va, top.aggregate = NULL, InterVA.rule = FALSE, noplot = FALSE,
       type = "bar", top.plot = 10, min.prob = 0, ...)
```

Arguments

- **va**: The list of va object to summarize.
- **top.aggregate**: Integer indicating how many causes from the top need to go into summary. The rest of the probabilities goes into an extra category "Undetermined". When set to NULL, default is all causes to be considered. This is only used when InterVA.rule set to "FALSE".
- **InterVA.rule**: If it is set to "TRUE", only the top 3 causes reported by InterVA5 is calculated into CSMF as in InterVA5. The rest of probabilities goes into an extra category "Undetermined". Default set to "FALSE".
- **noplot**: A logical value indicating whether the plot will be shown. If it is set to "TRUE", only the CSMF will be returned.
- **type**: An indicator of the type of chart to plot. "pie" for pie chart; "bar" for bar chart.
- **top.plot**: the maximum number of causes to plot in bar plot
- **min.prob**: The minimum probability that is to be plotted in bar chart, or to be labeled in pie chart.
- **...**: Arguments to be passed to/from graphic function `barplot`, `pie`, and more graphical parameters (see `par`). They will affect the main title, size and font of labels, and the radius of the pie chart.
Value

dist.cod The population probability of CODs.

Author(s)

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

See Also

CSMF.interVA5

Examples

## Not run:
data(RandomVA5)
# only fit first 20 observations for a quick illustration
RandomVA5 <- RandomVA5[1:20, ]

sample.output <- InterVA5(RandomVA5, HIV = "h", Malaria = "v", write = FALSE,
directory = tempdir(), filename = "VA5_result", output = "extended",
append = FALSE)

## Get CSMF by considering only top 3 causes reported by InterVA5.
## This is equivalent to using CSMF.interVA5() command Note that
## it's different from using all top 3 causes, since they may not
## all be reported
CSMF.summary <- CSMF5(sample.output, InterVA.rule = TRUE,
noplot = TRUE)

## Population level summary using pie chart
CSMF.summary2 <- CSMF5(sample.output, type = "pie",
min.prob = 0.01, main = "population COD distribution using pie chart",
clockwise = FALSE, radius = 0.7, cex = 0.7, cex.main = 0.8)

## Population level summary using bar chart
CSMF.summary3 <- CSMF5(sample.output, type = "bar",
min.prob = 0.01, main = "population COD distribution using bar chart",
cex.main = 1)
CSMF.summary4 <- CSMF5(sample.output, type = "bar",
top.plot = 5, main = "Top 5 population COD distribution",
cex.main = 1)

## End(Not run)
DataCheck5

Data cleaning for InterVA-5 algorithm

Description

This function implements the data cleaning steps in the InterVA5 software.

Usage

DataCheck5(Input, id, probbaseV5, InSilico_check = FALSE, write)

Arguments

Input        original data vector for one observation coded by 0 (absence), 1 (presence), and NA (missing).
id           id for this observation
probbaseV5   matrix of probbaseV5
InSilico_check logical indicator for if the check uses InSilicoVA rule. InSilicoVA rule sets all symptoms that should not be asked to missing. In contrast, the default InterVA5 rule sets these symptoms to missing only when they take the substantive value.
write        logical indicator of writing to file

Value

Output       new data vector
firstPass    message for the first pass check
secondPass   message for the second pass check

Author(s)

Jason Thomas, Zehang Li, Tyler McCormick, Sam Clark

References

http://www.interva.net/

See Also

InterVA5.plot
## Examples

```r
data(RandomVA5)
data(probbaseV5)
probbaseV5 <- as.matrix(probbaseV5)
RandomVA5 <- as.matrix(RandomVA5)
input <- as.character(RandomVA5[1, ])
input[which(toupper(input) == "N")]<- "0"
input[which(toupper(input) == "Y")]<- "1"
input[which(input != "1" & input != "0")]<- NA
input <- as.numeric(input)
output <- DataCheck5(Input=input, id="d1", probbaseV5=probbaseV5, write=TRUE)
```

---

### Description

The function downloads the most recent version of InterVA5, extracts the probbase.xls file (i.e., the Symptom-Cause-Information [SCI] source), and returns SCI as a matrix.

### Usage

```r
download.SCI()
```

### Value

- **newProbbase**: The Symptom-Cause-Information ("Probbase") used to assign causes of death.

### Author(s)

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

### See Also

- `probbaseV5`

### Examples

```r
## Not run:
data(RandomVA5)
RandomVA5 <- RandomVA5[1:2, ]
newProbbase <- download.SCI()
out <- InterVA5(RandomVA5, sci = newProbbase, HIV = "h", Malaria = "l", write=FALSE,
directory = tempdir(), filename = "VA5_result", output = "extended", append = FALSE)
```
getTopSymptoms

## End(Not run)

---

getTopSymptoms

Get the symptoms with the largest conditional probability (symptom \ cause) for causes assigned by InterVA-5.

Description

The function takes an interVA5 object and the data used to assign the causes, and returns the the symptoms that contribute to the cause assignment (ranked in order of the conditional probabilities of observing a symptom, given the death is due to that particular cause).

Usage

```r
getTopSymptoms(object, data, IDs = NULL, pretty = TRUE, includeAll = FALSE)
```

Arguments

- **object**: An interVA5 object (i.e., the results returned from the InterVA5() function).
- **data**: The input data that InterVA5 used to assign the causes of death.
- **IDs**: A vector that contains the IDs for each death (note that all of IDs are contained in data$ID and object$ID).
- **pretty**: A logical indicating if you want the results in an easy-to-read format (default is `TRUE`).
- **includeAll**: A logical indicating if you want all of the symptoms included in the output (even those which are absent or have a value of missing/no) (default is `FALSE` which only includes symptoms that are present).

Value

- **dist.cod**: A list of results for each death (organized by ID). For each death, a list is returned that includes the death’s ID, the cause, and a vector of strings listing a symptom, it if contributes to the cause assignment (if includeAll = TRUE), and the conditional probability of observing the symptom given that the death is due to this cause.

Author(s)

Jason Thomas, Zehang Li, Tyler McCormick, Sam Clark

See Also

`InterVA5`, `getTopSymptoms`
Examples

```r
# Not run:
data(RandomVA5)
sample.output <- InterVA5(RandomVA5, HIV = "h", Malaria = "v", write=FALSE)
topSymptoms <- getTopSymptoms(object = sample.output,
data = RandomVA5,
IDs = sample.output$ID[1],
pretty = TRUE,
includeAll = FALSE)
```

# End(Not run)

---

**InterVA5**

*Provide InterVA5 analysis on the data input.*

### Description

This function implements the algorithm in the InterVA5 software. It produces individual cause of death (COD) and population cause-specific mortality fractions. The output is saved in a .csv file specified by user. The calculation is based on the conditional and prior distribution of 61 CODs. The function can also save the full probability distribution of each individual to file. All information about each individual is saved to a va class object.

### Usage

```r
InterVA5(Input, HIV, Malaria, write = TRUE, directory = NULL,
filename = "VA5_result", output = "classic", append = FALSE,
groupcode = FALSE, sci = NULL, returnCheckedData = FALSE, ...)
```

### Arguments

- **Input**: A matrix input, or data read from csv files in the same format as required by InterVA5. Sample input is included as data(RandomVA5).
- **HIV**: An indicator of the level of prevalence of HIV. The input should be one of the following: "h"(high), "l"(low), or "v"(very low).
- **Malaria**: An indicator of the level of prevalence of Malaria. The input should be one of the following: "h"(high), "l"(low), or "v"(very low).
- **write**: A logical value indicating whether or not the output (including errors and warnings) will be saved to file. If the value is set to TRUE, the user must also provide a value for the parameter "directory".
- **directory**: The directory to store the output from InterVA5. It should either be an existing valid directory, or a new folder to be created. If no path is given and the parameter for "write" is true, then the function stops and an error message is produced.
filename The filename the user wish to save the output. No extension needed. The output is in .csv format by default.

output "classic": The same deliminated output format as InterVA5; or "extended": delimited output followed by full distribution of cause of death probability.

append A logical value indicating whether or not the new output should be appended to the existing file.

groupcode A logical value indicating whether or not the group code will be included in the output causes.

sci A data frame that contains the symptom-cause-information (aka Probase) that InterVA uses to assign a cause of death.

returnCheckedData A logical indicating if the checked data (i.e., the data that have been modified by the consistency checks) should be returned.

... not used

Details

Be careful if the input file does not match InterVA5 input format strictly. The function will run normally as long as the number of symptoms are correct. Any inconsistent symptom names will be printed in console as warning. If there is a wrong match of symptom from warning, please change the input to the correct order.

Value

<table>
<thead>
<tr>
<th>ID</th>
<th>identifier from batch (input) file</th>
</tr>
</thead>
<tbody>
<tr>
<td>MALPREV</td>
<td>selected malaria prevalence</td>
</tr>
<tr>
<td>HIVPREV</td>
<td>selected HIV prevalence</td>
</tr>
<tr>
<td>PREGSTAT</td>
<td>most likely pregnancy status</td>
</tr>
<tr>
<td>PREGLIK</td>
<td>likelihood of PREGSTAT</td>
</tr>
<tr>
<td>PRMAT</td>
<td>likelihood of maternal death</td>
</tr>
<tr>
<td>INDET</td>
<td>indeterminate outcome</td>
</tr>
<tr>
<td>CAUSE1</td>
<td>most likely cause</td>
</tr>
<tr>
<td>LIK1</td>
<td>likelihood of 1st cause</td>
</tr>
<tr>
<td>CAUSE2</td>
<td>second likely cause</td>
</tr>
<tr>
<td>LIK2</td>
<td>likelihood of 2nd cause</td>
</tr>
<tr>
<td>CAUSE3</td>
<td>third likely cause</td>
</tr>
<tr>
<td>LIK3</td>
<td>likelihood of 3rd cause</td>
</tr>
<tr>
<td>COMCAT</td>
<td>most likely circumstance of mortality</td>
</tr>
<tr>
<td>COMNUM</td>
<td>likelihood of COMCAT</td>
</tr>
<tr>
<td>wholeprob</td>
<td>full distribution of causes of death</td>
</tr>
</tbody>
</table>

Author(s)

Jason Thomas, Zehang Li, Tyler McCormick, Sam Clark
References

http://www.interva.net/

See Also

InterVA5.plot

Examples

data(RandomVA5)
# only fit first 5 observations for a quick illustration
RandomVA5 <- RandomVA5[1:5, ]

## to get easy-to-read version of causes of death make sure the column
## orders match interVA5 standard input this can be monitored by checking
## the warnings of column names

sample.output1 <- InterVA5(RandomVA5, HIV = "h", Malaria = "l", write = FALSE,
directory = tempdir(), filename = "VA5_result", output = "extended", append = FALSE)

## Not run:
## to get causes of death with group code for further usage
sample.output2 <- InterVA5(RandomVA5, HIV = "h", Malaria = "l",
write = FALSE, directory = "VA test", filename = "VA5_result_wt_code", output = "classic",
append = FALSE, groupcode = TRUE)

## End(Not run)

InterVA5.plot

Plot an individual-level distribution of va probabilities.

Description

The function takes an input of a single va object and produces a summary plot for it.

Usage

InterVA5.plot(va, type = "bar", min.prob = 0.01, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>va</td>
<td>A va object</td>
</tr>
<tr>
<td>type</td>
<td>An indicator of the type of chart to plot. &quot;pie&quot; for pie chart; &quot;bar&quot; for bar chart.</td>
</tr>
<tr>
<td>min.prob</td>
<td>The minimum probability that is to be plotted in bar chart, or to be labeled in pie chart.</td>
</tr>
</tbody>
</table>
Arguments to be passed to/from graphic function `barplot, pie`, and more graphical parameters (see `par`). They will affect the main title, size and font of labels, and the radius of the pie chart.

See Also

CSMF5

Examples

```r
## Not run:
data(RandomVA5)
# only fit first 20 observations for a quick illustration
RandomVA5 <- RandomVA5[1:20, ]
#' sample.output <- InterVA5(RandomVA5, HIV = "h", Malaria = "v", write = FALSE,
#   directory = tempdir(), filename = "VA5_result", output = "extended", append = FALSE)

## Individual level summary using pie chart
InterVA5.plot(sample.output$VA5[[3]], type = "pie", min.prob = 0.01,
   main = "1st sample VA analysis using pie chart", clockwise = FALSE,
   radius = 0.6, cex = 0.6, cex.main = 0.8)

## Individual level summary using bar chart
InterVA5.plot(sample.output$VA5[[3]], type = "bar", min.prob = 0.01,
   main = "2nd sample VA analysis using bar chart", cex.main = 0.8)

## End(Not run)
```

print.interVA5_summary

*Print method for summary of the results obtained from InterVA5 algorithm*

Description

This function prints the summary message of the fitted results.

Usage

```r
## S3 method for class 'interVA5_summary'
print(x, ...)
```  

Arguments

- `x`: summary of InterVA5 results
- `...`: not used
Conditional probability of InterVA5 (version 17 – Sept. 9th, 2018)

Description

This is the table of conditional probabilities of symptoms given CODs, along with prior probabilities in the first row. The values are from InterVA-5

Format

A data frame with 354 observations on 87 variables. The first row contains observations corresponding to prior probabilities; while the subsequent observations (rows 2 - 354) are the conditional probabilities.

Examples

data(probbaseV5)

Version 14 of the conditional probability of InterVA5

Description

This is version 14 (February 15th, 2018) of the table of conditional probabilities of symptoms given CODs, along with prior probabilities in the first row. The values are from InterVA-5

Format

A data frame with 354 observations on 87 variables. The first row contains observations corresponding to prior probabilities; while the subsequent observations (rows 2 - 354) are the conditional probabilities.

Examples

data(probbaseV5_14)
**Description**

This is version 17 (Sept. 9th, 2018) of the table of conditional probabilities of symptoms given CODs, along with prior probabilities in the first row. The values are from InterVA-5.

**Format**

A data frame with 354 observations on 87 variables. The first row contains observations corresponding to prior probabilities; while the subsequent observations (rows 2 - 354) are the conditional probabilities.

**Examples**

```r
data(probbaseV5_17)
```

---

**RandomVA5**

200 records of Sample Input

**Description**

This is a dataset consisting of 200 arbitrary sample input deaths in the acceptable format of InterVA5. Any dataset that needs to be analyzed by this package should be in the same format. The order of the input fields must not be changed.

**Format**

200 arbitrary input records.

**Examples**

```r
data(RandomVA5)
```
summary.interVA5  

Summary of the results obtained from InterVA5 algorithm

Description
This function prints the summary message of the fitted results.

Usage

```r
## S3 method for class 'interVA5'
summary(object, top = 5, id = NULL,
        InterVA.rule = TRUE, ...)
```

Arguments

- `object`: fitted object from `InterVA5()`
- `top`: number of top CSMF to show
- `id`: the ID of a specific death to show
- `InterVA.rule`: If it is set to "TRUE", only the top 3 causes reported by InterVA5 is calculated into CSMF as in InterVA5. The rest of probabilities goes into an extra category "Undetermined". Default set to "TRUE".
- `...`: not used

References

http://www.interva.net/

Examples

```r
## Not run:
data(RandomVA5)
# only fit first 20 observations for a quick illustration
RandomVA5 <- RandomVA5[1:20, ]

## to get easy-to-read version of causes of death make sure the column
## orders match interVA5 standard input this can be monitored by checking
## the warnings of column names

sample.output1 <- InterVA5(RandomVA5, HIV = "h", Malaria = "l",
                           write = FALSE, directory = tempdir(), filename = "VA5_result",
                           output = "extended", append = FALSE)

summary(sample.output1)
summary(sample.output1, top = 10)
summary(sample.output1, id = "sample3")

## End(Not run)
```
Print the version of the Symptom-Cause-Information source (aka probbaseV5).

Description

The function takes

Usage

version.SCI(sci)

Arguments

sci a symptom-cause-information matrix

Value

Message stating the Probbase version.

Author(s)

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

See Also
download.SCI

Examples

## Not run:
data(probbaseV5)
version.SCI(sci = probbaseV5)

## End(Not run)
whyNotCOD

Get the symptoms with the largest conditional probability (symptom | cause) using VA data.

Description

The function takes verbal autopsy data (which can be passed to InterVA5() to assign causes of death), and returns the symptoms that contribute to the assignment of a particular cause of death. This function differs from getTopSymptom() in that the user specified the cause for which they would like the results. This is an interactive function in the sense that if a cause is not provided as an argument, then the function will print out a numbered list of possible causes and the user can enter in the number to identify the cause of interest.

Usage

whyNotCOD(data, IDs = NULL, cause = NULL, pretty = TRUE,
includeAll = FALSE)

Arguments

data 
The input data that InterVA5 used to assign the causes of death.
IDs 
A vector that contains the IDs for each death (note that all of IDs are contained in data$ID and object$ID).
cause 
A string giving the name of the cause for which the conditional probabilities will be returned.
pretty 
A logical indicating if you want the results in an easy-to-read format (default is ‘TRUE’).
includeAll 
A logical indicating if you want all of the symptoms included in the output (even those which are absent or have a value of missing/no) (default is ‘FALSE’ which only includes symptoms that are present).

Value

dist.cod 
A list of results for each death (organized by ID). For each death, a list is returned that includes the death’s ID, the cause, and a vector of strings listing a symptom, if it contributes to the cause assignment (if includeAll = TRUE), and the conditional probability of observing the symptom given that the death is due to this cause.

Author(s)

Jason Thomas, Zehang LI, Tyler McCormick, Sam Clark

See Also

InterVA5, whyNotCOD
## Examples

```r
## Not run:
data(RandomVA5)
whyNotCOD(data = RandomVA5,
    IDs = RandomVA5$ID[1],
    pretty = TRUE,
    includeAll = FALSE)

data(causetextV5)
causetextV5[22, 2]
whyNotCOD(data = RandomVA5,
    IDs = RandomVA5$ID[1],
    cause = causetextV5[22, 2],
    pretty = TRUE,
    includeAll = FALSE)

## End(Not run)
```
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