# **Package**

# October 21, 2020

Title PRIME -- Papillomavirus Rapid Interface for Modelling and Economics

**Version** 2.0.12

Author Modelling and Economic Evaluation of Vaccines group, LSHTM (https://meev.lshtm.ac.uk)
<b>Description</b> PRIME is a static model of HPV vaccination that uses proportional impact to estimate the health impact and cost-effectiveness of HPV vaccination in low- and middle-income countries.
<b>Depends</b> R (>= $3.4.3$ )
Imports data.table, foreach
License GPL-3
Encoding UTF-8
LazyData true
RoxygenNote 6.1.1
Suggests knitr, rmarkdown
VignetteBuilder knitr
R topics documented:
ageCoverage
analyseCosts
checkSize
CreatePsaData
data.cecx_1y_prevalence
data.cecx_3y_prevalence
data.cecx_5y_prevalence
data.costcecx
data.countryname
data.disability_weights
data.global
data.hpv_distribution
data.incidence
data.incidence_ui
data.mortall

 data.mortall.unwpp.nqx
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 ...
 .

2 ageCoverage

data.mortcecx_ui	16
data.pop	16
data.popproj	17
data.quality	18
data.sexual_debut	18
data.valid	19
dtAggregate	20
dtColMatch	21
EmulateVaccineImpactVimcStochastic	
Estimate Vaccine Impact Vimc Central	
getISO3	
lifeTable	24
monetary_to_number	25
OutputVimc	
prime	
propSexDebut	26
RegisterBatchData	27
RegisterBatchDataVimc	
RunCohort	29
RunCountry	30
writelog	31
Index	33

ageCoverage

Get age-specific coverage-rates

# Description

Get age-specific coverage-rates

## Usage

```
ageCoverage(ages, routine_coverage, vaccine_efficacy_nosexdebut,
  vaccine_efficacy_sexdebut, campaigns, lifetab, cohort, agevac,
  country_iso3 = NULL)
```

# Arguments

ages Numeric vector (required): ages in model

routine\_coverage

Number (required): proportion of population that receives routine vaccination

campaigns List or number (required): if a list, applies MAC vaccination (needs to change)

lifetab Data.table (required): lifetable generated with lifeTable()

cohort Number (required): cohort-size (only used in MAC campaigns)

agevac Number (required): target age for vaccination

vaccine\_efficacy

Number (required): proportion indicating vaccine-efficacy

analyseCosts 3

#### Value

Data.table with coverage and effective coverage by age. Used in RunCohort()

## **Examples**

```
ages <- c(0:100)
routine_coverage <- 0.75
vaccine_efficacy <- 0.8
lifetab <- lifeTable(unlist(data.mortall[iso3=="AFG", as.character(0:100),
    with=F], use.names=F), 9)
agevac <- 9
ageCoverage (ages, routine_coverage, vaccine_efficacy, -1,
    lifetab, cohort, agevac)</pre>
```

analyseCosts

Returns cost-effectiveness for a single birthcohort in a single country

## **Description**

Usually called using RunCountry(..., analyseCosts=TRUE)

# Usage

```
analyseCosts(results, vaccine_cost, gdp_per_capita)
```

# Arguments

```
results Data.table (required): results from RunCohort()
vaccine_cost Number (required): cost of a single vaccine
gdp_per_capita Number (required): GDP per capita
```

## Value

Data.table with cost-analysis

# **Examples**

```
analyseCosts(RunCountry("AFG"), 100, 561)
```

4 BatchRun

BatchRun

Run multiple cohorts in a batch

#### **Description**

Runs multiple cohorts in one batch, based on the data in .data.batch

#### Usage

```
BatchRun(countries = -1, coverage = -1, agevac = -1,
   agecohort = -1, canc.inc = "2018", sens = -1,
   unwpp_mortality = TRUE, year_born = -1, year_vac = -1, runs = 1,
   vaccine_efficacy_beforesexdebut = 1,
   vaccine_efficacy_aftersexdebut = 0, log = -1,
   by_calendaryear = FALSE, use_proportions = TRUE,
   analyseCosts = FALSE, canc.cost = "unadj", discounting = FALSE,
   disc.cost = 0.03, disc.ben = 0.03, psa = 0,
   psa_vals = ".data.batch.psa", disability.weights = "gbd_2017",
   wb.indicator = "NY.GDP.PCAP.PP.CD", wb.year = 2017,
   vaccine = "4vHPV")
```

#### **Arguments**

countries ignore, read from .data.batch ignore, read from .data.batch coverage agevac ignore, read from .data.batch agecohort ignore, read from .data.batch canc.inc year from where incidence data is read (2018; old data: 2012) - with updated 2018 Globocan data, DALY weights from GBD, and DALY estimation based on prevalence instead of age of incidence ignore, does not do anything anymore sens unwpp\_mortality logical, whether to create lifetables based on UNWPP mortality estimates or WHO data ignore year\_born year\_vac ignore ignore runs vaccine\_efficacy\_beforesexdebut vaccine efficacy before sexual debut vaccine\_efficacy\_aftersexdebut vaccine efficacy after sexual debut log name of log file by\_calendaryear logical, output values by calendar year or by year of birth cohort use\_proportions logical, output data as rates per capita or in totals logical, directly run cost-effectiveness analysis on output or not analyseCosts

checkSize 5

canc.cost	Character (optional): Is cost of cancer adjusted ("adj" for International \$) or not ("unadj" for US\$)	
discounting	Logical (optional): If TRUE, run cost-effectiveness analysis undiscounted and discounted. If FALSE, only uses undiscounted	
disc.cost	Number (optional): Discounting for health costs (only if discounting=TRUE)	
disc.ben	Number (optional): Discounting for health outcomes (only if discounting=TRUE)	
psa	integer, number of runs for probabilistic sensitivity analysis (PSA)	
psa_vals	data table with values to use in probabilistic sensitivity analysis, usually .data.batch.psa, generated by RegisterBatchData* functions (currently only RegisterBatchDataVimc)	
disability.weights		
	character, disability weights for cervical cancer from GBD 2017 or GBD 2001	
wb.indicator	character, World Bank indicator for GDP/GNI per capita in I\$/US\$ and current/constant data	
wb.year	numeric, year of the World Bank indicator value	
vaccine	character, bivalent/quadrivalent (4vHPV) or nonavalent (9vHPV) vaccine	

# Value

Returns combined results

# **Examples**

#

checkSize	Checks whether the size of a variable is larger than 0	
-----------	--	--

# Description

Used to determine that all required variables are passed to a function Checks whether a vector has length > 0 or a data.table/data.frame has nrow > 0

## Usage

checkSize(v)

# Arguments

v Variable (required)

## Value

Logical: TRUE if size is not 0, false if size is 0

6 CreatePsaData

#### **Examples**

```
x <- c()
checkSize(x)

x <- c(2,5)
checkSize(x)

A <- c()
B <- c(1,2,3)
sapply(c("A","B"),function(x){checkSize(get(x))})</pre>
```

CreatePsaData

Generate Latin hyper cube sample of parameters for sensitivity analysis

#### **Description**

Generate Latin hyper cube sample of input parameters based on their distributions for probabilistic sensitivity analysis.

#### Usage

```
CreatePsaData(country_codes, vaccine = "4vHPV", psa_runs = 0,
  seed_state = 1, psadat_file = "psadat.csv",
  psadat_vimc_file = "psadat_vimc.csv")
```

## **Arguments**

country\_codes ISO3 country codes of countries

vaccine bivalent/quadrivalent or nonavalent HPV vaccine

psa\_runs integer, simulation runs for sensitivity analysis

seed\_state integer, seed value for random number generator

psadat\_file character string, file to save Latin hyper cube sample of input parameters

psadat\_vimc\_file

character string, file to save Latin hyper cube sample of input parameters (VIMC format)

#### Value

Null return value; disease burden estimates are saved to corresponding files

## **Examples**

```
CreatePsaData (
  country_codes = c("AFG", "ALB"),
  vaccine = "4vHPV",
  psa_runs = 200,
  seed_state = 1,
  psadat_file = "psadat_csv",
  psadat_vimc_file = "psadat_vimc.csv")
```

 ${\tt data.cecx\_1y\_prevalence}$ 

1-year prevalence of cervical cancer

## Description

A dataset containing the 1-year prevalence (proportion) of cervical cancer in 185 countries, as reported by IARC's Globocan 2018 database.

## Usage

```
data.cecx_1y_prevalence
```

## **Format**

A data table containing 185 observations of 103 variables.

**Country** Country name **iso3** ISO3 country code **Source** Data source

Year Data source (year)

**0..100** Age 0-100

#### **Details**

As per IARC definition – The (1-year) prevalence of a given cancer is the number of individuals within a defined population who have been diagnosed with that cancer (within 1 year) and who are still alive at a given point in time (i.e. the survivors).

## Source

```
https://gco.iarc.fr/today/online-analysis-table
```

data.cecx\_3y\_prevalence

3-year prevalence of cervical cancer

# Description

A dataset containing the 3-year prevalence (proportion) of cervical cancer in 185 countries, as reported by IARC's Globocan 2018 database.

## Usage

```
data.cecx_3y_prevalence
```

#### **Format**

A data table containing 185 observations of 103 variables.

Country Country name iso3 ISO3 country code Source Data source Year Data source (year) 0..100 Age 0-100

#### Details

As per IARC definition – The (3-year) prevalence of a given cancer is the number of individuals within a defined population who have been diagnosed with that cancer (within 3 years) and who are still alive at a given point in time (i.e. the survivors).

#### **Source**

```
https://gco.iarc.fr/today/online-analysis-table
```

data.cecx\_5y\_prevalence

5-year prevalence of cervical cancer

#### **Description**

A dataset containing the 5-year prevalence (proportion) of cervical cancer in 185 countries, as reported by IARC's Globocan 2018 database.

#### Usage

```
data.cecx_5y_prevalence
```

#### **Format**

A data table with 185 observations of 103 variables.

Country Country name iso3 ISO3 country code Source Data source Year Data source (year) 0..100 Age 0-100

#### **Details**

As per IARC definition – The (5-year) prevalence of a given cancer is the number of individuals within a defined population who have been diagnosed with that cancer (within 5 years) and who are still alive at a given point in time (i.e. the survivors).

#### **Source**

```
https://gco.iarc.fr/today/online-analysis-table
```

data.costcecx 9

data.costcecx

Cost of cervical cancer treatment

#### **Description**

A dataset containing the cost of cervical cancer treatment.

## Usage

data.costcecx

#### **Format**

A data table with 194 observations of 7 variables.

country Country name

cancer\_cost cost per cancer episode, in 2017 US\$

cancer\_cost\_adj cost per cancer episode - adjusted, in 2017 US\$ - "adjusted" cancer costs are based on a GDP/capita based adjustment within the region

iso3 ISO3 country code

cancer\_cost\_2011 cost per cancer episode, in 2011 US\$

cancer\_cost\_adj\_2011 cost per cancer episode – adjusted, in 2011 US\$ – "adjusted" cancer costs
are based on a GDP/capita based adjustment within the region

inflation\_factor Inflation factor from 2011 to 2017 estimated from Inflation, GDP deflator (annual %) – https://data.worldbank.org/indicator/NY.GDP.DEFL.KD.ZG

data.countryname

Country names and codes

## **Description**

A dataset containing the country names and codes (ISO/WHO/WB/UN).

## Usage

 ${\tt data.countryname}$ 

#### **Format**

A data table with 251 observations of 13 variables.

name1, name2, name3, name4 Country names

iso2, iso3 ISO2 and ISO3 country codes

isonum ISO number

WHOcode WHO country code

who\_region WHO region

who\_mort WHO mortality stratum - A/B/C/D/E

WBincome WB income classification of countries

UNgroup UN group classification of countries

GDPpc2011id GDP per capita (2011)

10 data.global

data.disability\_weights

Disability weights and duration of cervical cancer stages

## **Description**

A dataset containing disability weights and duration of different phases of cervical cancer.

#### Usage

```
data.disability_weights
```

#### **Format**

A data table with 13 observations of 8 variables.

**Source** Source of disability weights - IHME / WHO

**Sequela** Sequelae / stage / phase of cervical cancer **Duration** Duration of cervical cancer phase

WHO\_MortalityStratum WHO moratlity stratum – applicable only for long term sequelae from

WHO source

Mid Disability weight (mid)

Low Disability weight (low)

High Disability weight (high)

**Description** Description of cervical cancer phase

data.global

Global data table

## Description

A dataset containing a global range of variables.

# Usage

data.global

#### **Format**

A data table with 194 observations of 35 variables.

Country Country name

iso2 ISO2 country code

WHO Region WHO regions

WHO Mortality Stratum WHO mortality stratum

World Bank Income Group (2011) World Bank income group levels

GAVI Eligibility GAVI eligibility status

PAHO Revolving Fund PAHO revolving fund status

Cohort size (2010) [1 ] Cohort size

Coverage (3 doses at year 10) [2 ] Vaccination coverage at age 10

Vaccine efficacy vs vaccine type infection [2 ] Vaccine efficacy

Duration of protection [2] Duration of vaccine protection

Age group [3] Age group

Vaccine price USD [4] Vaccine price for 2 doses US\$

Vaccine delivery/ operational/ admin costs (USD) [5] Vaccine delivery, operational and administration costs US\$

Cancer treatment costs - primary level hospital i\$ (per episode, over lifetime) [6] Cancer treatment costs - primary level hospital i\$ (per episode, over lifetime)

Cancer treatment costs - primary level hospital US\$ (per episode, over lifetime) [6] Cancer treatment costs - primary level hospital US\$ (per episode, over lifetime)

Cancer treatment costs - secondary level hospital i\$ (per episode, over lifetime) [6] Cancer treatment costs - secondary level hospital i\$ (per episode, over lifetime)

Cancer treatment costs - secondary level hospital US\$ (per episode, over lifetime) [6] Cancer treatment costs - secondary level hospital US\$ (per episode, over lifetime)

Cancer treatment costs - teaching hospital i\$ (per episode, over lifetime) [6] Cancer treatment costs - teaching hospital i\$ (per episode, over lifetime)

Cancer treatment costs - teaching hospital US\$ (per episode, over lifetime) [6] Cancer treatment costs - teaching hospital US\$ (per episode, over lifetime)

Discount rate [2] Discount rate

Perspective [2 ] Perspective

Costs [2]

Time horizon [2] Time horizon

"Percent" CeCx due to 16/18 Percentage of cervical cancer due to HPV strains 16 and 18

Vaccine programme Vaccine programme

Vaccine programme Oct2013 Vaccine programme Oct2013

Econ evaluation Economic evaluation

**GDP per capita (2011 US\$)** [7 ] GDP per capita (2011 US\$)

**GDP** per capita (2011 i\$) [7] GDP per capita (2011 i\$)

**GNI per capita (2011 i\$)** [7] GNI per capita (2011 i\$)

**GNI per capita (2011 US\$)** [7] GNI per capita (2011 US\$)

V33 International\$

V34 US\$

iso3 ISO3 country code

12 data.hpv\_distribution

data.hpv\_distribution Relative contribution of HPV 16/18/31/33/45/52/58 in ICC HPV-positive cases

#### **Description**

A dataset containing relative contribution of HPV 16/18/31/33/45/52/58 in cases of ICC HPV-positive, by region and country

## Usage

data.hpv\_distribution

#### **Format**

A data table with 249 observations of 12 variables.

Global World

Region UN region

Subregion UN subregion

Intermediate\_region UN intermediate region

Country Country name

iso3 ISO3 country code

**hpv\_4v** Relative contribution (%) of HPV 16/18 in ICC HPV-positive cases (mean)

**hpv\_4v\_low** Relative contribution (%) of HPV 16/18 in ICC HPV-positive cases (lower bound of 95% uncertainty interval)

**hpv\_4v\_high** Relative contribution (%) of HPV 16/18 in ICC HPV-positive cases (upper bound of 95% uncertainty interval)

**hpv\_9v** Relative contribution (%) of HPV 16/18/31/33/45/52/58 in ICC HPV-positive cases (mean)

**hpv\_9v\_low** Relative contribution (%) of HPV 16/18/31/33/45/52/58 in ICC HPV-positive cases (lower bound of 95% uncertainty interval)

**hpv\_9v\_high** Relative contribution (%) of HPV 16/18/31/33/45/52/58 in ICC HPV-positive cases (upper bound of 95% uncertainty interval)

#### Source

Serrano B, Alemany L, Tous S, Bruni L, Clifford GM, Weiss T, et al. Potential impact of a nine-valent vaccine in human papillomavirus related cervical disease. Infect Agents Cancer. 2012;7: 38. https://doi.org/10.1186/1750-9378-7-38

data.incidence 13

data.incidence Incidence of cervical cancer, by age and country
---

## **Description**

A dataset containing the incidence of cervical cancer in 185 countries, as reported by IARC's Globocan 2018 database. Crude rate, cervix uteri, females, by age.

## Usage

data.incidence

#### **Format**

A data table with 185 observations of 103 variables.

Country Country name

iso3 ISO3 country code

Source Data source

Year Data source (year)

**0..100** Age 0-100 – Crude rate, cervix uteri, females, by age; annual rate per individual

# Source

https://gco.iarc.fr/today/online-analysis-table

data.incidence_ui	Incidence of cervical cancer with uncertainty intervals, all ages and by country

## Description

A dataset containing the incidence of cervical cancer with uncertainty intervals in 185 countries, as reported by IARC's Globocan 2018 database. Estimated number of new cases in 2018, cervix uteri, females, all ages.

# Usage

data.incidence\_ui

## **Format**

A data table with 185 observations of 5 variables.

Country Country name

iso3 ISO3 country code

Mid Estimated number of new cases in 2018, cervix uteri, females, all ages (mean)

**Low** Estimated number of new cases in 2018, cervix uteri, females, all ages (lower bound of 95% uncertainty interval)

**High** Estimated number of new cases in 2018, cervix uteri, females, all ages (upper bound of 95% uncertainty interval)

**Source** Data source **Year** Data source (year)

#### **Source**

```
https://gco.iarc.fr/today/online-analysis-table
```

data.mortall

WHO life table

## Description

A dataset containing the WHO life table.

## Usage

data.mortall

#### **Format**

A data table with 196 observations of 107 variables.

Country | Age [12 ] Country name

**0..100** Age 0-100

V103..V106 na

iso3 ISO3 country code

## Source

https://www.who.int/gho/mortality\_burden\_disease/life\_tables/life\_tables/en/

```
data.mortall.unwpp.nqx
```

UNWPP life table

# Description

A dataset containing the UNWPP life table (World Population Prospects 2019).

## Usage

```
data.mortall.unwpp.nqx
```

data.mortcecx 15

#### **Format**

```
A data table with 122850 observations of 8 variables.
```

```
country_code_numeric Country code numeric
country_code ISO3 country code
country Country name
age_from age from (start-age)
age_to age to (end-age)
year Year
gender Gender
value nqx - probability of dying between ages x and x+n
```

#### **Source**

https://population.un.org/wpp/

data.mortcecx

Mortality from cervical cancer, by age and country

# Description

A dataset containing the mortality from cervical cancer in 185 countries, as reported by IARC's Globocan 2018 database.

## Usage

data.mortcecx

## **Format**

A data table with 185 observations of 103 variables.

Country Country name iso3 ISO3 country code Source Data source
Year Data source (year)
0..100 Age 0-100

#### **Source**

https://gco.iarc.fr/today/online-analysis-table

16 data.pop

 ${\tt data.mortcecx\_ui} \qquad {\tt \textit{Mortality from cervical cancer with uncertainty intervals, all ages and} \\ {\tt \textit{by country}}$ 

## Description

A dataset containing the number of deaths from cervical cancer with uncertainty intervals in 185 countries, as reported by IARC's Globocan 2018 database. Estimated number of deaths in 2018, cervix uteri, females, all ages.

## Usage

data.mortcecx\_ui

#### **Format**

A data table with 185 observations of 5 variables.

Country Country name

iso3 ISO3 country code

Mid Estimated number of deaths in 2018, cervix uteri, females, all ages (mean)

**Low** Estimated number of deaths in 2018, cervix uteri, females, all ages (lower bound of 95% uncertainty interval)

**High** Estimated number of deaths in 2018, cervix uteri, females, all ages (upper bound of 95% uncertainty interval)

Source Data source

Year Data source (year)

#### Source

https://gco.iarc.fr/today/online-analysis-table

data.pop

UNWPP population estimates

## **Description**

A dataset containing the UNWPP population estimates – World Population Prospects 2019.

## Usage

data.pop

data.popproj 17

## **Format**

```
A data table with 3392220 observations of 8 variables.
```

```
country_code_numeric Country code numeric
country_code ISO3 country code
country Country name
age_from age from (start-age)
age_to age to (end-age)
year Year
gender Gender
value Population size
```

## Source

data.popproj

Population projections of 5-year old girls

## Description

A dataset containing population projections of 5-year old girls. (not used – to be removed)

## Usage

```
data.popproj
```

#### **Format**

A data table with 98 observations of 91 variables.

```
iso3 ISO3 country code
2011..2100 Year – 2011..2100
```

18 data.sexual\_debut

data.quality

Data quality of incidence and mortality

#### **Description**

A dataset indicating data quality of cervical cancer incidence and mortality.

## Usage

```
data.quality
```

#### **Format**

A data table with 186 observations of 4 variables.

Country Country name

Incidence Quality of cervical cancer incidence data

Mortality Quality of cervical cancer mortality data

iso3 ISO3 country code

data.sexual\_debut

Sexual debut data

#### **Description**

A dataset containing sexual debut data and (2) parameters for the sexual debut curve (logistic model).

## Usage

```
data.sexual_debut
```

#### **Format**

A data table with 94 observations of 14 variables.

V1 Row number

iso2 ISO2 country code

country Country name

iso3 ISO3 country code

who WHO region

X15 Proportion of people who have sexually debuted at age 15

X18 Proportion of people who have sexually debuted at age 18

X20 Proportion of people who have sexually debuted at age 20

X22 Proportion of people who have sexually debuted at age 22

X25 Proportion of people who have sexually debuted at age 25

data.valid

**Never** Proportion of people who had not sexually debuted **cluster.id** Clustering countries with similar characteristics

a Parameter for sexual debut curve (logistic model)

**b** Parameter for sexual debut curve (logistic model)

data.valid

Model validation

#### **Description**

A dataset containing data for validation.

#### Usage

data.valid

#### **Format**

A data table with 26 observations of 49 variables.

Country Country name

iso2 ISO2 country code

WHO Region WHO region

World Bank Income Group (2011) World Bank income group classification (2011)

Author Author

Year Year of publication

Title Title of publication

**Currency** Currency

Currency year Currency year

Conversion to I\$2011 International dollar (I\$2011)

**ICER vs no prevention** Incremental cost-effectiveness ratio of vaccination versus no prevention (CHECK)

ICER vs screen Incremental cost-effectiveness ratio of vaccination versus no prevention (CHECK)

**Denominator** Denominator for health impact

Vaccine total costs Vaccine total costs

Vaccine coverage Vaccine coverage

Vaccine efficacy vs vaccine type infection Vaccine efficacy versus vaccine type infection

**Duration of protection** Duration of protective immunity from vaccination

Cohort size Cohort size

Age at vaccination Age at vaccination, years

Cancer treatment cost per episode Cancer treatment cost per episode, dollars

Discount rate: costs Discount rate for costs

**Discount rate: benefits** Discount rate for benefits **Perspective** Perspective of economic evaluation

20 dtAggregate

Comparator is no screening Comparator refers to no scereening, logical (Y/N)

Time horizon Time horizon of analysis

GDP per capita GDP per capita

Cervical cancer due to 16/18 Proportion of cervical cancer due to HPV types 16 and 18

CeCx cost low original Cervical cancer cost / low / original

CeCx cost high original Cervical cancer cost / high / original

CeCx cost low Cervical cancer cost / low

CeCx cost high Cervical cancer cost / high

CeCx data available Cervical cancer data available, logical (Y/N)

**0-4** 0-4 years

**5-9** 5-9 years

**9-14** 9-14 years

**15-19** 15-19 years

**20-24** 20-24 years

**25-29** 25-29 years

**30-34** 30-34 years

**35-39** 35-39 years

**40-44** 40-44 years

**45-49** 45-49 years

**50-54** 50-54 years

**55-59** 55-59 years

**60-64** 60-64 years

**65-69** 65-69 years

**70-74** 70-74 years

**75-79** 75-79 years

**80+** 80+ years

dtAggregate

Collapse data-tables

## **Description**

Collapse data-tables

## Usage

```
dtAggregate(DT, aggr_on, measure.vars = c(), id.vars = c(),
func = "sum", na.rm = TRUE)
```

dtColMatch 21

#### **Arguments**

DT Data-table (required) Character string (required): column-name that will be used to collapse on (i.e. aggr\_on combine all age-strata) Character string (optional): column-names that will be collapsed (function will measure.vars be applied to all these columns) id.vars Character string (optional): column-names that will remain stratified N.b. if measure.vars is not provided, all columns that are not in id.vars and aggr\_on will be assumed to be assumed func Character string (optional): function that will be applied to data (if optional, values will be summed) Logical (optional): if TRUE, removes NA from measure.vars columns before na.rm

#### Value

Returns collapsed data.table

dtColMatch Match two data-tables on multiple columns

applying function (or passes na.rm=TRUE to function)

#### **Description**

Returns vector with column-of-interest where columns match

## Usage

```
dtColMatch(input, input_match_on, reference, reference_match_on,
  reference_return)
```

## **Arguments**

input Data.table (required): input-table to match

input\_match\_on Character vector (required): column-names in input-table to match

Data.table (required): reference-table to match reference

reference\_match\_on

Character vector (required): column-names in reference-table to match

reference\_return

Character string (required): column-name in reference-table that is returned

(where values match)

#### **Details**

If at least one value in any of the input\_match\_on columns matches with a value in any of the reference\_match\_on columns, the two rows will match

#### Value

Character vector with values from reference\_return column in reference\_match\_on data.table where values match

#### **Examples**

```
dtColMatch (data.global, c("Country"), data.countryname,
   c("name1", "name2", "name3", "name4"), "iso3")
```

EmulateVaccineImpactVimcStochastic

Emulate vaccine impact estimates (VIMC stochastic runs)

## **Description**

Emulate vaccine impact estimates for VIMC stochastic runs/sensitivity analysis. The inputs are central disease burden estimates, input parameter distributions (latin hyper sampling), runs for sensitivity analysis, and filename for stochastic burden estimates. The outputs are stochastic disease burden estimates (full results file plus a file per country).

## Usage

```
EmulateVaccineImpactVimcStochastic(disease_burden_template_file,
  centralBurdenResultsFile, psaData, diseaseBurdenStochasticFolder,
  diseaseBurdenStochasticFile, psa_runs, countryCodes = -1,
  vaccination_scenario)
```

#### **Arguments**

disease\_burden\_template\_file

csv file (required), central disease burden template (vimc format); add column with run\_id to get stochastic disease burden template

central Burden Results File

csv file (required), central disease burden estimates pre and post vaccination

psaData data table (required), latin hyper cube sample of input parameters

 ${\tt diseaseBurdenStochasticFolder}$ 

character string (required), stochastic disease burden estimates folder (output)

diseaseBurdenStochasticFile

character string (required), stochastic disease burden estimates file(s) (output)

psa\_runs

integer (required), simulation runs for sensitivity analysis

countryCodes

list (optional), If country codes are provided, stochastic burden estimates are generated for these countries. If set to -1, then stochastic burden estimates are generated for the countries included in the central burden estimates.

vaccination\_scenario

logical (required), generate stochastic burden estimates for (vaccination) or (no vaccination) scenario

#### **Details**

Stochastic disease burden estimates are generated. (i) full results files 1 full results file for prevaccination (optional) 1 full results file for post-vaccination (ii) 1 file per country for all runs 1 file per country for all runs – pre-vaccination (optional) 1 file per country for all runs – post-vaccination

#### Value

Null return value; disease burden estimates are saved to corresponding files

EstimateVaccineImpactVimcCentral

Generate vaccine impact estimates (VIMC central run)

#### **Description**

Generate vaccine impact estimates for VIMC central runs. The inputs are vaccine coverage and disease burden template files and outputs are disease burden estimates (pre-vaccination and post-vaccination).

#### Usage

```
EstimateVaccineImpactVimcCentral(vaccine_coverage_file, disease_burden_template_file, disease_burden_no_vaccination_file, disease_burden_vaccination_file, disease_burden_results_file, campaign_vaccination, routine_vaccination, vaccine = "4vHPV")
```

#### **Arguments**

#### **Details**

Three disease burden estimates are generated. (i) disease burden estimates for no vaccination (vimc format) (ii) disease burden estimates for vaccination (vimc format) (iii) disease burden estimates for vaccination (pre- and post-vaccination) and includes YLDs and YLLs

## Value

Null return value; disease burden estimates are saved to corresponding files

24 lifeTable

#### **Examples**

```
EstimateVaccineImpactVimcCentral (
  vaccine_coverage_file = "coverage_hpv-routine-default.csv",
  disease_burden_template_file = "central-burden-template.csv",
  disease_burden_no_vaccination_file = "central_burden_no_vaccination.csv",
  disease_burden_vaccination_file = "central_burden_vaccination.csv",
  disease_burden_results_file = "central_burden_results.csv",
  routine_vaccination = TRUE,
  campaign_vaccination = TRUE)
```

getISO3

Retrieve ISO3-code of country

## **Description**

Retrieve ISO3-code of country

#### Usage

```
getISO3(countryname, name = FALSE)
```

#### **Arguments**

countryname Character string (required): Full name of the country

name Logical (optional): If TRUE, returns full name and alternative names of returned

country (may be useful to double-check that it is the correct country)

#### Value

Character string with ISO3 code. Will also return full name if name=TRUE.

#### **Examples**

```
getISO3("Afghanistan")
getISO3("Congo",name=TRUE)
```

lifeTable

Construct lifetable based on qx-column

#### **Description**

```
qx = age-specific probability of dying
```

## Usage

```
lifeTable(qx = NULL, mx = NULL, agecohort = 0)
```

## **Arguments**

qx Numeric vector (required): Age-specific probabilities of dying

agecohort Number (optional): Age at which cohort is started

monetary\_to\_number 25

#### Value

Data.table with lifetable

#### **Examples**

```
qx <- unlist(data.mortall[iso3=="AFG", as.character(0:100), with=F], use.names=F) lifeTable(qx, 9)
```

monetary\_to\_number

Convert monetary character-strings to numeric values

## Description

Convert monetary character-strings to numeric values

#### Usage

```
monetary_to_number(x)
```

## **Arguments**

Х

Character string to convert

#### Value

Returns number with value, stripped from any currency symbols and thousand-seperators (i.e. "B#2,010.50" becomes 2010.5)

## **Examples**

```
monetary_to_number ("$2,200.20")

# Note that values using German or Dutch notation (i.e. using a comma to separate decimals and a dot to seperate thousands) are converted as well. monetary_to_number ("$2.200,20")
```

OutputVimc

Formatting output for VIMC Montagu

# Description

OutputVimc takes result of BatchRun and outputs it in format to be uploaded to VIMC Montagu.

## Usage

```
OutputVimc(DT, age_stratified = TRUE, calendar_year = FALSE,
  vimc_template = -1)
```

26 propSexDebut

## **Arguments**

DT data table with results

age\_stratified logical, whether output should be stratified by age

calendar\_year logical, whether output should be given by calendar year of event OR by year of

birth of cohort

vimc\_template data table with template file downloaded from montagu

#### Value

#

## **Examples**

#

prime prime: Papillomavirus Rapid Interface for Modelling and Economics

(PRIME).

#### **Description**

PRIME stands for "Papillomavirus Rapid Interface for Modelling and Economics". The R package is based of the spreadsheet-based tool (see http://primetool.org).

## PRIME provides estimates of

The magnitude of the burden of cervical cancer.

The impact of introducing HPV vaccination for girls prior to sexual debut.

Healthcare costs incurred as a result of cervical cancer treatment.

Costs associated with vaccination.

Long-term savings which may result from a vaccination program.

propSexDebut Proportion of girls sexually debuted

# Description

propSexDebut returns proportion of girls sexually debuted in country\_iso3 at age age.

# Usage

```
propSexDebut(age, country_iso3)
```

## **Arguments**

age age of girls

country\_iso3 ISO3 country code

RegisterBatchData 27

#### Value

Returns proportion of girls in a given country that has sexually debuted at a given age.

## **Examples**

```
propSexDebut (20, "IND")
propSexDebut (30, "ETH")
```

RegisterBatchData

Creates .data.batch for running multiple birth cohorts

#### **Description**

Creates .data.batch which is used when running/looping over multiple birth cohorts ( runCohort() ) at once.

## Usage

```
RegisterBatchData(coverage_data, reporting_years = -1, force = FALSE)
```

## **Arguments**

 $coverage\_data \quad Data\ table\ with\ columns\ country\_code,\ year\ (of\ vaccination),\ age\_first,\ age\_last,$ 

coverage.

reporting\_years

Numeric\_vector, years that should be reported (parameter: not required)

force Logical, whether .data.batch should be overwritten if it already exists (parame-

ter: not required)

#### **Details**

.data.batch is based on the data.table (DT) coverage\_data, which is a DT with columns country\_code (ISO3), year (of vaccination), age\_first (age at vaccination), age\_last (age at vaccination), coverage (in proportion, for all the agegroups specified).

If you only want to run 1 age in this country/coverage combination, age\_first==age\_last

#### Value

batch data of cohorts with vaccination coverage

## **Examples**

#

RegisterBatchDataVimc Creates .data.batch for running multiple birth cohorts (VIMC runs)

## Description

Creates .data.batch which is used when running/looping over multiple birth cohorts ( runCohort() ) at once. Similar to RegisterBatchData, but for when we make runs for VIMC.

# Usage

```
RegisterBatchDataVimc(vimc_coverage, vimc_template, use_campaigns,
  use_routine, restrict_to_coverage_data = FALSE, force = FALSE,
  psa = 0)
```

## **Arguments**

vimc_coverage	data table with coverage estimates as downloaded from VIMC montagu	
vimc_template	data table with reporting template as downloaded from VIMC montagu	
use_campaigns	logical, whether campaigns as stated in coverage files should be modelled	
use_routine	logical, whether routine vaccination as stated in coverage file should be modelled	
restrict_to_coverage_data		
	logical, whether the first birth-cohort should be the first cohort that is mentioned in the coverage data. If TRUE, restrict to coverage data. If FALSE, restrict to cohorts provided in vimc_template.	
force	logical, whether .data.batch should be overwritten if it already exists	
psa	integer, indicating how many runs for probabilistic sensitivity analysis (PSA). 0 to run no PSA.	

## **Details**

.data.batch is based on the data.table (DT) coverage\_data, which is a DT with columns country\_code (ISO3), year (of vaccination), age\_first (age at vaccination), age\_last (age at vaccination), coverage (in proportion, for all the age groups specified).

## Value

batch data of cohorts with vaccination coverage

# Examples

#

RunCohort 29

RunCohort	Run PRIME for a single birth-cohort	
-----------	-------------------------------------	--

## Description

Runs PRIME for one birth-cohort. Usually called by another function such as RunCountry().

## Usage

```
RunCohort(lifetab, cohort, incidence, mortality_cecx, prevalence, agevac,
  coverage, campaigns, vaccine_efficacy_nosexdebut,
  vaccine_efficacy_sexdebut, cost_cancer, discounting = FALSE,
  disc.cost = 0.03, disc.ben = 0.03, country_iso3 = NULL,
  run_country = FALSE, disability.weights = "gbd_2017")
```

## Arguments

lifetab	Data.table: The life-table for this cohort. Can be created using the lifeTable() function.	
cohort	Number: The cohort-size of this birth-cohort at the time where the lifetable starts.	
incidence	Numeric vector: Age-specific CeCx(16/18) incidence-rates.	
mortality_cecx	Numeric vector: Age-specific CeCx(16/18) mortality-rates.	
prevalence	Numeric vector: Age-specific CeCx(16/18) prevalence rates (5-year prevalence) – referring to people who are alive within 5 years of diagnosis.	
agevac	Number: Age at which the cohort is vaccinated.	
coverage	Number: Proportion of the cohort that will receive a vaccination.	
campaigns	List or number: MAC cohort-vaccinations (needs to be changed).	
vaccine_efficacy_nosexdebut  Number: proportion indicating vaccine-efficacy before sexual debut.		
vaccine_efficacy_sexdebut  Number: proportion indicating vaccine-efficacy after sexual debut.		
cost_cancer	Number: total per capita cost of cancer.	
discounting	Logical (optional): If TRUE, run cost-effectiveness analysis undiscounted and discounted. If FALSE, only uses undiscounted	
disc.cost	Number (optional): Discounting for health costs (only if discounting=TRUE)	
disc.ben	Number (optional): Discounting for health outcomes (only if discounting=TRUE)	

## Value

Returns a data.table with size of the birth-cohort and age-specific incidence-rates, mortality-rates, years-of-life-lost, years-of-healthy-life-lost, and cancer-costs before and after vaccination. Also displays whether discounting has been used ("type" column).

30 RunCountry

#### **Examples**

```
lifetab <- lifeTable(unlist(data.mortall[iso3=="AFG",</pre>
  as.character(0:100), with=F], use.names=F), 9)
incidence <- unlist(data.incidence[iso3=="AFG", as.character(0:100), with=F],</pre>
 use.names=F)
mortality_cecx <- unlist(data.mortall[iso3=="AFG", as.character(0:100), with=F],</pre>
 use.names=F)
prevalence <- unlist(data.cecx_5y_prevalence[iso3=="AFG",</pre>
 as.character(0:100), with=F], use.names=F)
agevac <- 9
coverage <- 0.8
campaigns <- -1
vaccine_efficacy_nosexdebut <- 0.95</pre>
vaccine_efficacy_sexdebut <- 0</pre>
cost_cancer <- 100
RunCohort(lifetab, cohort, incidence, mortality_cecx, prevalence, agevac,
  coverage, campaigns, vaccine_efficacy_nosexdebut, vaccine_efficacy_sexdebut,
  cost_cancer, disc.cost=0.03, disc.ben=0.03,
  discounting=FALSE, country_iso3="AFG", run_country=FALSE)
```

RunCountry

Run PRIME for a specific country

## **Description**

Runs RunCohort() using country-specific estimates. If year\_born and year\_vac are not provided, assumes vaccination occurs in the current year.

## Usage

```
RunCountry(country_iso3, vaceff_beforesexdebut = 1,
  vaceff_aftersexdebut = 0, cov = 1, agevac = 10, agecohort = 10,
  cohort = -1, canc.inc = "2018", sens = -1,
  unwpp_mortality = TRUE, year_born = -1, year_vac = -1,
  campaigns = -1, analyseCosts = FALSE, canc.cost = "unadj",
  discounting = FALSE, disc.cost = 0.03, disc.ben = 0.03,
  run_batch = FALSE, psadat = -1, disability.weights = "gbd_2017",
  wb.indicator = "NY.GDP.PCAP.PP.CD", wb.year = 2017,
  vaccine = "4vHPV")
```

#### **Arguments**

country_iso3	Character string (required): ISO3 code of the country	
cov	Number (optional): Proportion with routine coverage	
agevac	Integer (optional): Target age for HPV vaccination	
agecohort	Integer (optional): Reference age for cohort-size (only used when 'cohort' is not provided)	
cohort	Integer (optional): Cohort-size1 if unknown	

writelog 31

canc.inc	Integer (optional): Reference year for cancer incidence rates (Globocan: 2018 or 2012)		
sens	Numeric-vector (optional): Specific values to be used in a PSA1 if PSA's are not used		
unwpp_mortality	<b>y</b>		
	Logical (optional): If TRUE, uses year-specific UNWPP mortality estimates to construct life-tables. If FALSE, use WHO based mortality estimates.		
year_born	Integer (optional): Year in which cohort is born		
year_vac	Integer (optional): Year in which cohort is vaccinated		
campaigns	List (optional): Multi-Age-Cohort campaigns (needs to be changed)		
analyseCosts	Logical (optional): If FALSE, returns result from RunCohort() function. If TRUE, runs analyseCosts() with country-specific results.		
canc.cost	Character (optional): Is cost of cancer adjusted ("adj" for International \$) or not ("unadj" for US\$)		
discounting	Logical (optional): If TRUE, run cost-effectiveness analysis undiscounted and discounted. If FALSE, only uses undiscounted		
disc.cost	Number (optional): Discounting for health costs (only if discounting=TRUE)		
disc.ben	Number (optional): Discounting for health outcomes (only if discounting=TRUE)		
disability.weights			
	character, disability weights for cervical cancer from GBD 2017 or GBD 2001		
wb.indicator	character, World Bank indicator for GDP/GNI per capita in I\$/US\$ and current/constant data		
wb.year	numeric, year of the World Bank indicator value		
vaccine	character, bivalent/quadrivalent (4vHPV) or nonavalent (9vHPV) vaccine		
vaceff	Number (optional): Proportion indicating vaccine-efficacy		

## Value

data.table with country-specific results of HPV vaccination. Returns cost-analysis if analyseCosts=TRUE

# **Examples**

```
RunCountry("AFG")
RunCountry("AFG", year_vac=2020, agevac=10, cov=0.75, vaceff=0.88)
RunCountry("AFG", year_vac=2020, agevac=10, cov=0.75, vaceff=0.88, analyseCosts=TRUE)
```

writelog Simulation log reporting

## Description

Appends message of simulation run (x) to log file (logname).

## Usage

```
writelog(logname, x)
```

32 writelog

# Arguments

logname log filename

x message of simulation run

Value

None

# **Examples**

#

# Index

*Topic datasets	data.popproj,17
data.cecx_1y_prevalence,7	data.quality, 18
<pre>data.cecx_3y_prevalence, 7</pre>	data.sexual_debut, 18
data.cecx_5y_prevalence, 8	data.valid, 19
data.costcecx, 9	dtAggregate, 20
data.countryname, 9	dtColMatch, 21
data.disability_weights, 10	
data.global, 10	EmulateVaccineImpactVimcStochastic, 22
data.hpv_distribution, 12	EstimateVaccineImpactVimcCentral, 23
data.incidence, 13	
data.incidence_ui, 13	getIS03, 24
data.mortall, 14	lifoToble 24
data.mortall.unwpp.nqx, 14	lifeTable, 24
data.mortcecx, 15	monetary_to_number, 25
data.mortcecx_ui, 16	monetar y_to_number, 25
data.pop, 16	OutputVimc, 25
data.popproj, 17	odepatrime, 25
data.quality, 18	prime, 26
data.sexual_debut, 18	prime-package (prime), 26
data.valid, 19	propSexDebut, 26
data. varia, 17	p. opcons as a c, 20
ageCoverage, 2	RegisterBatchData, 27
analyseCosts, 3	RegisterBatchDataVimc, 28
	RunCohort, 29
BatchRun, 4	RunCountry, 30
checkSize, 5	writelog, 31
CreatePsaData, 6	
data.cecx_1y_prevalence,7	
data.cecx_3y_prevalence,7	
data.cecx_5y_prevalence, 8	
data.costcecx, 9	
data.countryname, 9	
data.disability_weights, 10	
data.global, 10	
data.hpv_distribution, 12	
data.incidence, 13	
data.incidence_ui, 13	
data.mortall, 14	
data.mortall.unwpp.nqx, 14	
data.mortcecx, 15	
data.mortcecx_ui, 16	
data.pop, 16	
uu uu , pop, 10	