

Package ‘cemtool’

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Type Package

Title Step by step guide to build a Markov model

Version 0.3

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Description Markov models are increasingly used to determine the cost-effectiveness of medical innovations. However, medical researchers still use closed source software to build such models. The ceom-package was build to guide researchers to build their own Markov model in R. The current version supports Markov models that consist of 3 to 6 health-states. More options will be added in the future. Do you want to join development? Let us know by contacting the package author.

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Depends shiny, rhandsontable, diagram, shinydashboard, utils

Encoding UTF-8

LazyData false

RoxygenNote 6.1.1

URL <https://github.com/StanWijn/cemtool>

Suggests knitr,
rmarkdown

VignetteBuilder knitr

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cemprob

Second step of the cemtool: Define the transition probabilities, cost and utility values

Description

The `cemtool()` package provides a step-by-step tool to guide users in building a default Markov model. The tool guides the user through the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

Usage

`cemprob()`

Details

There are multiple software systems that can be used to build Markov models for cost-effectiveness analyses like TreeAge, Excel or R. Although there are numerous advantages to use R over the others, the biggest downside is the steep learning curve from R. The `cemtool()` package aims to close this gap by introducing a step-by-step tool to guide users in building a default Markov models. The tool guides the user through the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

– **Required variables in the Global Environment before the `cemprob()` function will work:**

- Number of healthstates (HS)
- Names of the healthstates (HS1, HS2 ... HS_n, dead)
- Names of both the intervention (intervention) and usual care strategy (control)
- Number of cycles (n.t)
- Vector of the strategy names (Strategies)
- Vector of the healthstate names (v.n)

These can be generated with the `cemtool()` function.

Value

The following variables will be created in the Global Environment:

- Empty Markov trace matrices for both strategies (`m.M` and `m.M_treatment`)
- A dataframe with the modelinput (`modelinput`)
- Discount rate for costs (`d.rc`) and effects (`d.re`)

The function will automatically run `cemtpm()` after finishing

Author(s)

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Examples

```
cemtool() # Start from scratch, please clear the objects from the Global Enviroment
cemprob() # Start from the second phase (definding the parameters)
cemtpm() # Start from the third phase (modify the transition probability matrix)
cemrun() # Run the model with the current m.M markov trace and m.P transition probability matrix
```

cemrun	<i>Run the Markov model with input generated from cemtool(), cemprob(),cemtpm()</i>
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Description

The cemtool() package provides a step-by-step tool to guide users in building a default Markov model. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

Usage

```
cemrun()
```

Details

There are multiple software systems that can be used to build Markov models for cost-effectiveness analyses like TreeAge, Excel or R. Although there are numerous advantages to use R over the others, the biggest downside is the steep learning curve from R. The cemtool() package aims to close this gap by introducing a step-by-step tool to guide users in building a default Markov models. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

– **Required variables in the Global Enviroment before the cemprob() function will work:**

- Number of healhstates (HS)
- Names of the healthstates (HS1, HS2 ... HS_n, dead)
- Names of both the intervention (intervention) and usual care strategy (control)
- Number of cycles (n.t)
- A vector of the strategy names (Strategies)
- A vector of the healhstate names (v.n)

These can be generated with the cemtool() function.

Value

The following variables will be created in the Global Enviroment:

- Full Markov trace (m.M and m.M_treatment)
- Calculate the costs and effects for both strategies.
- Results are saved (table_output) and shown in the console. The model structure and markov trace are both plotted and saved (plot1 and plot2)#'

Author(s)

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Examples

```
cemtool() # Start from scratch, please clear the objects from the Global Enviroment
cemprob() # Start from the second phase (definding the parameters)
cemtpm() # Start from the third phase (modify the transition probability matrix)
cemrun() # Run the model with the current m.M markov trace and m.P transition probability matrix
```

cemtool

Default function to start the cemtool package

Description

The cemtool() package provides a step-by-step tool to guide users in building a default Markov model. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

Usage

```
cemtool()
```

Details

There are multiple software systems that can be used to build Markov models for cost-effectiveness analyses like TreeAge, Excel or R. Although there are numerous advantages to use R over the others, the biggest downside is the steep learning curve from R. The cemtool() package aims to close this gap by introducing a step-by-step tool to guide users in building a default Markov models. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

– Required knowledge / information:*Model structure*

The structure of the Markov model: how many healthstates, the names of the healthstates etc.

Transition probabilities

The probabilities to move from one healthstate to another

Costs

The costs for a patient to be on the healhstates for one cycle / year

Effects

The effect size associated with the healthstate. For QALY a value between 1 and 0, from perfect health to death.

Value

The following variables will be created in the Global Enviroment:

- Number of healhstates (HS)
- Names of the healthstates (HS1, HS2 ... HS_n, dead)
- Names of both the intervention (intervention) and usual care strategy (control)
- Number of cycles (n.t)
- Vector of the strategy names (Strategies) and a vector of the healhstate names (v.n).

The function will automatically run cemprob() after finishing

Author(s)

S.R.W. Wijn MSc <stan.wijn@radboudumc.nl>

Examples

```
cemtool() # Start from stratch, please clear the objects from the Global Enviroment
cemprob() # Start from the second phase (definding the parameters)
cemtpm() # Start from the third phase (modify the transition probability matrix)
cemrun() # Run the model with the current m.M markov trace and m.P transition probability matrix
```

cemtpm

OPTIONAL: Third step of the cemtool: Alter the transition probability matrix

Description

The cemtool() package provides a step-by-step tool to guide users in building a default Markov model. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

Usage

```
cemtpm()
```

Details

There are multiple software systems that can be used to build Markov models for cost-effectiveness analyses like TreeAge, Excel or R. Although there are numerous advantages to use R over the others, the biggest downside is the steep learning curve from R. The cemtool() package aims to close this gap by introducing a step-by-step tool to guide users in building a default Markov models. The tool guides the user though the steps of the development of a Markov model and will present the final result using graphs and tables. The only prerequisite is that the user knows the structure of the model and the transition probabilities, no calculations or coding is required.

– **Required variables in the Global Enviroment before the cemprob() function will work:**

- Number of healthstates (HS)
- Names of the healthstates (HS1, HS2 ... HS_n, dead)
- Names of both the intervention (intervention) and usual care strategy (control)
- Number of cycles (n.t)
- Vector of the strategy names (Strategies)
- Vector of the healthstate names (v.n)

These can be generated with the `cemtool()` function.

Value

The following variables will be created in the Global Environment:

— Modified transition probability matrix for both strategies (M.P and m.P_treatment).

The function will automatically run `cemrun()` after finishing

Author(s)

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Examples

```
cemtool() # Start from scratch, please clear the objects from the Global Environment
cemprob() # Start from the second phase (defining the parameters)
cemtpm()  # Start from the third phase (modify the transition probability matrix)
cemrun()  # Run the model with the current m.M markov trace and m.P transition probability matrix
```

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