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# mimR – A Package for Graphical Modelling in R

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## Abstract

The `mimR` package (version 1.1.1) for graphical modelling in R is introduced. We present some facilities of `mimR`, namely those relating specifying models, editing models, fitting models and doing model search. We also discuss the entities needed for flexible graphical modelling in terms of an object structure. An example about a latent variable model is presented.

## 1 Introduction and background

The `mimR` package provides facilities for graphical modelling in the statistical program R (<http://www.R-project.org/>), both are available from the Comprehensive R Archive Network CRAN at <http://CRAN.R-project.org/>. The `mimR` package has also its own homepage at <http://www.jbs.agrsci.dk/~sorenh/mimR/> and is furthermore a part of the gR-project (<http://www.R-project.org/gR/>) which is a project to make graphical models available in R.

The statistical foundation for `mimR` is Mixed Interaction Models, a very general class of statistical models for mixed discrete and continuous variables. Statistical inference in mixed interaction models can be made by the stand-alone program MIM (available from <http://www.hypergraph.dk/>), and the core of `mimR` is an interface from R to the MIM program. The reader is assumed familiar with mixed interaction models, and to have a working knowledge of the MIM program. Edwards (2000) described both in a very clear way. For a comprehensive account of graphical models we refer to Lauritzen (1996). Other important references are Edwards (1990) and Lauritzen and Wermuth (1989).

There are two aims of this paper. First, to present the `mimR` package as a simple, yet operational package for graphical modelling in R. Second, to present some tentative ideas regarding an object structure for graphical modelling in R.

## 2 Preliminaries

### 2.1 Mixed interaction models *à vol d’oiseau*

Mixed interaction models include as special cases log–linear models for contingency tables and covariance selection models for the multivariate normal distribution. More importantly, however, is that the models allow a simultaneous modelling of discrete *and* continuous variables. Focus in mixed interaction models is often (although not exclusively) on conditional independence restrictions.

Within mixed interaction models, one can treat problems where all variables are treated on equal footing (i.e., there are no distinction between variables as being explanatory or responses). Such models are below referred to as *undirected models*. It is however also possible to work with problems where some variables are purely explanatory, other are purely responses and others play both roles. Such models are denoted *block recursive models*. Block recursive models contain undirected models as special cases, and – perhaps more importantly – can be established by a careful combination of undirected models through conditioning.

### 2.2 MIM as inference engine

From the users perspective, the MIM stand alone program can be regarded as an “inference engine” with which the user (at least in principle) needs not be concerned with. However, in practice it is worth keeping in mind that MIM is a stand alone program: In MIM there can be only 1) one specification of a data set, 2) one data set and 3) one model at any time. This means that in general when fitting a new model, a lot of information may have to be conveyed to MIM, and this may take some time (if the data set is large). Likewise, receiving fitted values from MIM to R may take some time too.

### 2.3 Getting help

In addition to the documentation in the `mimR` package, the MIM program itself contains a comprehensive help function which the user of `mimR` is encouraged to make use of.<sup>1</sup>

### 2.4 Ways of accessing MIM from R

There are two levels of accessing MIM from R.

- The “high level approach” is by creating model objects much like one does when working with linear, generalized linear and other models in R. This approach is the core contents of this paper.
- The “low level approach” is by sending commands directly to MIM using the `mim.cmd` and the `mcm` functions, see Appendices [C.1](#) and [C.2](#).

## 3 The rats dataset

Some features of `mimR` will be illustrated in the present paper on the basis of the `rats` dataset in the `mimR` package. The `rats` dataset is from a hypothetical drug

<sup>1</sup>To access the help function in MIM go to the MIM program and press F1.

trial, where the weight losses of male and female rats under three different drug treatments have been measured after one and two weeks. See [Edwards \(2000\)](#) for more details. The first rows of the data are:

```

      Sex Drug W1 W2
1     M   D1  5  6
2     M   D1  7  6
3     M   D1  9  9
4     M   D1  5  4
5     M   D2  9 12
6     M   D2  7  7
7     M   D2  7  6
8     M   D2  6  8
      .....

```

## 4 Objects in `mimR`

The core of `mimR` are the `gmData` and `mim` objects.

**gmData objects:** A `gmData` object contains information about variables, their labels, their levels (for the discrete variables) etc. A `gmData` object may also contain data, but need not do so. The name “`gmData`” can be taken to be short for “graphical model data” or “graphical meta data” (where we prefer the latter). The idea behind separating the specification of the variables from data is that some properties of a model can be investigated without any reference to data, for example decomposability and collapsibility.

**mim objects:** Links a model formula to a `gmData` object. Since a `gmData` object need not contain any data, fitting a `mim` model is separate process. (This is an important difference between model in `mimR` and e.g. linear models in R.) When the model has been fitted (provided that there are data in the `gmData` object), the `mim` object also contains the fitted values, parameter estimates etc.

## 5 gmData objects – graphical meta data

A `gmData` object contains information about variables, their labels, their levels (for the discrete variables) etc. A `gmData` object may also contain data, but need not do so.

### 5.1 Creating a `gmData` object manually

A `gmData` object (without data) can be created by

```

gmD.rats.nodata <- gmData(c("Sex", "Drug", "W1", "W2"),
  factor = c(2, 3, FALSE, FALSE),
  vallabels = list(c("M", "F"), c("D1", "D2", "D3")))

```

The `gmData` object looks like

```

  name letter factor levels
1 Sex      a   TRUE      2
2 Drug     b   TRUE      3
3 W1      c  FALSE     NA
4 W2      d  FALSE     NA

```

Data origin: no.data

To see the values of the factors use the 'vallabels' function

To see the data use the 'observations' function

To each variable, there is associated a letter. It is possible use the letters in specifying models, see the examples below.

## 5.2 Making a gmData object from a data frame or a table

Typically one will create a gmData object (with data) from a data frame (or a table) as follows:

```

data(rats)
gmd.rats <- as.gmData(rats)
data(HairEyeColor)
gmd.hec <- as.gmData(HairEyeColor)

```

## 6 Models in mimR

Currently, only undirected models are available in mimR. That is, models in which all variables are treated on equal footing as response variables. (Thus models where a possible response structure has not been accounted for can currently not be dealt with in mimR).

An undirected model is created using the mim function (which returns a mim object). The following two specifications are equivalent:

```

m1 <- mim("Sex:Drug/Sex:Drug:W1+Sex:Drug:W2/W1:W2", data = gmd.rats)
m2 <- mim("ab/abc+abd/cd", data = gmd.rats, letter = TRUE)

```

It is possible to specify 1) the main effects, 2) the saturated and 3) the homogeneous saturated models (possibly for only a subset of the variables) in short form:

```

m.main <- mim(".", data = gmd.rats, marginal = c("Sex", "Drug", "W1"))
m.sat <- mim("*", data = gmd.rats, marginal = c("Sex", "Drug", "W1"))
m.hsats <- mim("*h", data = gmd.rats, marginal = c("Sex", "Drug", "W1"))

```

## 7 Model fitting

Model fitting is separated from model specification, so the models created above are not fitted to data. For model fitting two functions are available: fit and emfit (emfit will be discussed later).

```

m1f <- fit(m1)
m1f
Model: Sex:Drug/Sex:Drug:W1+Sex:Drug:W2/W1:W2
Model(letter): ab/abc,abd/cd
likelihood: 273.705 DF: 15

```

## 8 Model summary

A summary (including certain model properties) of a `mim` can be achieved using the `summary()` function:

```
summary(m1f)
Formula: Sex:Drug/Sex:Drug:W1+Sex:Drug:W2/W1:W2
Formula(letter): ab/abc,abd/cd
likelihood: 273.705 DF: 15
Model properties:
  Variables in model:      Sex Drug W1 W2
  Is graphical:           TRUE
  Is decomposable:        TRUE
  Is mean linear:         TRUE
  Is homogeneous:         TRUE
  Is delta-collapsible:   TRUE
  Degrees of freedom:     15
  Cliques:
[1] "Sex:Drug:W1:W2"
```

## 9 Model selection and model editing

### 9.1 Editing models directly

Models can be edited directly, using the `editMIM` function by which one can 1) delete, 2) add and 3) homogeneously add interactions:

```
m.main <- mim(".", data = gmd.rats)
m2 <- editmim(m.main, add = c("Sex:Drug", "Sex:W2"))
m3 <- editmim(m.main, add = c("Sex:Drug", "Sex:W2"),hadd = "Drug:W1:W2")
```

### 9.2 Stepwise model selection

To a `mimModel` object the function `stepwise` applies which takes as additional arguments all arguments that the `STEPWISE` command in `MIM` does. The `stepwise` function returns a new `mimModel` object.

```
data(car carcass)
gmd.carc <- as.gmData(car carcass)
m.main <- mim(".", data = gmd.carc)
m.sat <- mim("*", data = gmd.carc)
m.m <- stepwise(m.main, "f") # forward
m.s <- stepwise(m.sat, "s") # backward, exact tests
```

## 10 Fitted values (parameter estimates)

The fitted values (parameters estimates) can be obtained using the `fitted` function:

```
mf2 <- fit(m2)
parms <- fitted(mf2)
parms
```

Drug	Sex	Counts	W1	W2	W1:W1	W1:W2	W2:W1	W2:W2	
1	1	1	4	9.75	8.500	17.104	0	0	5.583
2	2	1	4	9.75	8.500	17.104	0	0	5.583
3	3	1	4	9.75	8.500	17.104	0	0	5.583
4	1	2	4	9.75	8.833	17.104	0	0	9.639
5	2	2	4	9.75	8.833	17.104	0	0	9.639
6	3	2	4	9.75	8.833	17.104	0	0	9.639

## 11 Simulating data from a fitted model

Simulating data from a fitted model can be done by the `simulate` function:

```
samp <- simulate(mf2, size = 10) # 'sample' is already used in R
```

## 12 Obtaining the linear predictor

The `linpredict` function can be used to get the linear predictor for a set  $y$  given another set  $x$  (possibly empty) of variables, for example

```
d1 <- linpredict(mf2, y = "W2", x = "W1:Sex")
d2 <- linpredict(mf2, y = "Sex", x = "W1:W2")
```

```
d1
Distribution of W2 given W1:Sex
Sex=1
  int W1
W2 8.5  0
      W2
W2 5.58333
Sex=2
  int W1
W2 8.83333 0
      W2
W2 9.63922
```

```
d2
Distribution of Sex given W1:W2
  Sex Constant      W1      W2
1   1  0.00000  0.00000  0.00000
2   2 -0.37203 -0.01595  0.06087
```

## 13 Missing values and/or latent variables

To fit a model with to incomplete data or to fit a latent variable model, use the `emfit` function. See e.g. the Example in Section 14.

## 14 Example – Mathematics marks

This dataset (taken from [Mardia, Kent, and Bibby \(1979\)](#)) contains the examination marks for 88 students in 5 different subjects. Data is contained the data set `mathmark` in the `mimR` package. [Edwards \(2000\)](#) also investigates these data.

We start out by specifying the saturated model and do a backward elimination:

```
data(mathmark)
gmd.math <- as.gmData(mathmark)
math1 <- mim("*", data=gmd.math)
math2 <- stepwise(math1)
math2
Formula: //mechanics:vectors:algebra+algebra:analysis:statistics
Formula(letter): //abc,cde
likelihood: 3391.021 DF: 4
```

The model `math2` is shown in Figure 1.

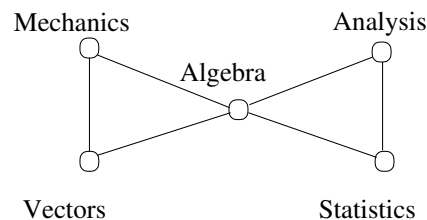


Figure 1: The “butterfly” model selected for the mathmarks data.

Next we consider a latent variable model: We suppose that there is a latent binary variable `L` such that the manifest variables are all conditionally independent given `L`. We fit such a model by:

```
math <- mathmark
math$L <- factor(NA, levels = 1:2) # L is binary and consists of NAs
gmd.math <- as.gmData(math)
latent(gmd.math) <- "L"
```

With the specification above, `L` is a binary variable consisting of `NA` values. The command `latent(gmd.math) <- "L"` makes it explicit in the `gmData` object, that `L` is indeed a latent variable. One consequence is, that the model can not be fitted using the `fit` function. Instead, the `emfit` function which uses the EM algorithm, [Dempster, Laird, and Rubin \(1977\)](#), must be used:

```
m1 <- mim("*", data = gmd.math)
m2 <- editmim(m1, del = paste(names(math)[1:5], ":", collapse = ''))
m2f <- emfit(m2, plot = TRUE)
d.imp <- retrieveData(gmd.math, impute = TRUE)
```

The argument `plot=TRUE` in `emfit()` creates the plot in Figure 2.

We plot the predicted value of `L` against the observation number:

```
plot(d.imp$L)
```

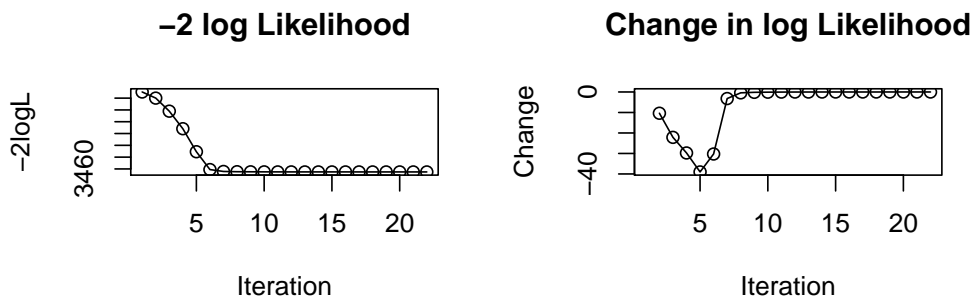


Figure 2: Convergence of the EM algorithm.

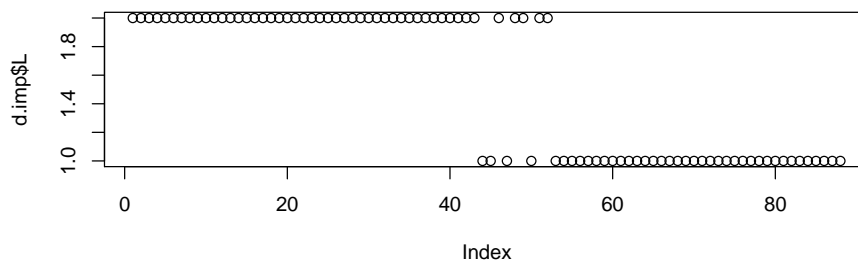


Figure 3: An index plot of the discrete latent variables.

The plot is shown in Figure 3. The grouping of the values of  $L$  suggests that data have been processed somehow prior to presentation. Edwards (2000), p. 181, conclude: “Certainly they (the data) have been mistreated in some way, doubtless by a statistician.”

The EM algorithm needs a set of initial values for the unobserved values to start from when calculating the first parameter estimates. It is well known, that the final estimate of the EM algorithm may depend on the initial values and that (especially in the case of latent variables) the likelihood may have multiple maxima. Default in the call of `emfit` is that MIM substitutes random values for the missing values. It is, however, possible to control the starting values of the EM algorithm as follows: The user can specify the values of  $L$  and subsequently declare  $L$  to be latent. In the call of `emfit`, the argument `S` causes the EM algorithm to take the specified values as starting point for the EM algorithm:

```
math[, "L"] <- factor(1:2, levels = 1:2) # L consists of 1,2,1,2...
gmd.math <- as.gmData(math)
latent(gmd.math) <- "L"
m1 <- mim("*", data = gmd.math)
m2 <- editmim(m1, del = paste(names(math)[1:5], ":", collapse = ''))
m2f <- emfit(m2, arg = "S", plot = TRUE)
d.imp <- retrieveData(gmd.math, impute = TRUE)
```

For this specific case, it turns out that the result is very insensitive to the initial

values.

## 15 Discussion

In this paper we have 1) illustrated some aspects of `mimR` package for graphical modelling in R, and 2) presented preliminary ideas regarding an object structure for graphical modelling in R. It is the hope that `mimR` will be obsolete in a not too distant future – not because of lack of relevance of being able to work with graphical models in R. Rather, it is the hope that a more proper package with this functionality will be implemented as an integrated part of R. That is one of the aims of the `gR`-project. Until that happens we will continue to develop `mimR`. `mimR` is currently at level of development where it is likely that significant changes (e.g. of names of functions and/or object classes) will occur.

## Acknowledgements

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## References

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## A Miscellaneous

**mimR mailing list:** To receive news about mimR send a mail to [mimr@genetics.agrsci.dk](mailto:mimr@genetics.agrsci.dk) with the text 'subscribe mimr' in the body.

**mimR Availability:** mimR is available only on Windows platforms because MIM only runs on Windows platforms.

**mimR and S-PLUS:** The current version of mimR does not run under S-PLUS. If sufficient interest appears, it may be considered to remedy this situation.

## B Using the appropriate version of MIM and R

To use mimR, the MIM program must be installed on your computer (Windows only). MIM is available from <http://www.hypergraph.dk/>. Upgrades of MIM are frequently released.

It is IMPORTANT to make sure that your version of MIM and R is in accordance with what mimR expects. When loading the mimR package using `library(mimR)` a message similar to the one below appears in R. From this one sees the earliest version of MIM and R with which the current version of mimR works.

```
-----
mimR: A package for graphical modelling in R
mimR, version 1.1.1 is now loaded
Copyright (C) 2002, Søren Højsgaard
Maintained by Søren Højsgaard <sorenh@agrsci.dk>
Webpage: http://www.jbs.agrsci.dk/~sorenh/mimR
```

```
Built: R 1.7.1; ; 2003-07-08 00:21:43
```

```
NOTICE:
```

- o mimR is available on Windows platforms only
  - o To use mimR the MIM program must be running
  - o The current version of mimR requires MIM version 3.1.2.20 or later
  - o MIM is available from <http://www.hypergraph.dk>.
  - o The current version of mimR requires R version 1.7.1 or later
- ```
-----
```

## C Low level access to MIM from R

### C.1 Primitive use of MIM from R – the `mim.cmd()` function

The core of mimR is the `mim.cmd` function. The arguments to `mim.cmd` are simply MIM commands (given as strings). For example:

```
>mim.cmd("fact a2 b2; statread ab; 25 2 17 8 !")
>mim.cmd("mod a,b; fit; print; print f")
```

The `mim.cmd` function returns the result of the commands submitted to MIM. The result of the last call of `mim.cmd` above is:

```

Deviance:          5.3111 DF: 1
The current model is: a,b.
Fitted counts, means and covariances.
  a b   Count
  1 1   21.808
  1 2    5.192
  2 1   20.192
  2 2    4.808

```

## C.2 Using MIM directly from `mimR`– the `mcm()` function

The `mcm` function (short for “MIM command mode”) provides a direct interface to MIM, i.e. the possibility to write MIM commands directly. The `mcm` function returns no value to R, and is intended only as an easy way to submit MIM commands without the overhead of wrapping them into the `mim.cmd` function (or submitting the commands directly to MIM). Hence, using `mcm`, the session above would be:

```

> mcm()
Enter MIM commands here. Type quit to return to R
MIM->fact a2 b2; statread ab
MIM->25 2 17 8 !
Reading completed.
MIM->mod a,b; fit
Deviance:          5.3111 DF: 1
MIM->print; print f
The current model is: a,b.
Fitted counts, means and covariances.
  a b   Count
  1 1   21.808
  1 2    5.192
  2 1   20.192
  2 2    4.808
MIM->quit
>

```

To return to R from the `mcm` function type `'quit'`, `'exit'`, `'end'`, `'q'` or `'e'` (i.e. the commands one would use to terminate MIM). These commands, however, do not terminate MIM – they only return control to R.