Linear Latent Variable Models: The lava-package

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Abstract
An R package for specifying and estimating linear latent variable models is presented. The philosophy of the implementation is to separate the model specification from the actual data, which leads to a dynamic and easy way of modeling complex hierarchical structures. Several advanced features are implemented including robust standard errors for clustered correlated data, multigroup analyses, non-linear parameter constraints, inference with incomplete data, maximum likelihood estimation with censored and binary observations, and instrumental variable estimators. In addition an extensive simulation interface covering a broad range of non-linear generalized structural equation models is described. The model and software are demonstrated in data of measurements of the serotonin transporter in the human brain.

Keywords: latent variable model, maximum likelihood, multigroup analysis, structural equation model, R, serotonin, SERT

1. Introduction

Multivariate data are often modelled using random effects in order to account for correlation between measurements in the statistical analysis. The dominating model is the linear mixed effects model (Laird and Ware, 1982), which is available in most standard statistical software packages, e.g. SAS PROC MIXED and in R in the packages nlme (Pinheiro and Bates, 2000) and lme4 (Bates and Maechler, 2009) with the latter one also offering some support for generalized linear mixed models.

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Another type of random effect model is the structural equation model (SEM) (Bollen, 1989), where the terminology latent variable often is used instead of random effects. While the mixed effect model and structural equation model have many aspects in common, the aim of a SEM analysis is typically to analyze the association between the latent variable, representing some process that is only partially observed, and some other variables (observed or latent). Thus, in SEMs focus is mainly on the latent variable, where one normally ascribes less interpretation to the random effects in a mixed effects model, which primarily serves as a way of capturing covariance between measurements. Because observed variables can be viewed as representations of underlying true variables SEMs offer a natural framework for handling measurement errors in study variables, and it often provides an efficient analysis of high dimensional data (Budtz-Jørgensen et al, 2003). The framework was pioneered by Jöreskog (Jöreskog, 1970) and since then it has been an active area of research with focus on relaxing linearity and distributional assumptions (Rabe-Hesketh et al, 2004).

SEMs have proven to be useful in many different fields of research. However, applications have been dominated by covariance structure analyses, and as residuals on the individual level are not available in this setup, model assessment has been based on more or less heuristic omnibus tests. The lack of profound model diagnostics have undoubtedly lead to several poor applications of SEMs (Steiger, 2001), and thus likely, however unjustified, giving the framework a somewhat bad reputation among groups of statisticians.

Several proprietary software solutions are available for analyzing SEMs with some of the most popular being LISREL, SAS PROC CALIS, AMOS, EQS, Stata 12 sem, Stata gllamm (Rabe-Hesketh et al, 2004) and Mplus (Muthén and Muthén, 2007), where the last two programs stand-out because of their general modelling framework. Common for all these solutions is that they are difficult to extend and therefore possibilities for examination of new methodological ideas are limited. In part because details of many features of proprietary software often remains hidden from the user. Implementations in an open-source environment such as R (R Development Core Team, 2010) directly address this problem. Currently two such solutions are available: the sem package (Fox, 2006, 2009) and the OpenMx package (Boker et al, 2011). While the former package is limited to standard covariance structure analysis, OpenMx offers sophisticated methods such as multiple group analysis, models for ordinal data and mixture models. The predecessor package Mx has been a popular for analyzing family data in epidemiological genetics and
OpenMx will undoubtedly become an important tool in that field of research.

This paper presents the lava-package for statistical analysis in a very general modelling framework known as the Linear Latent Variable Model, which includes structural equation models and mixed models as important special cases. This model class also allows for non-linear effects of covariates and non-linear parameter constraints. The lava-package offers a superior user interface for specifying, altering and visualizing the model design. Models are specified independently of data using commands that are similar to standard regression modeling in R. In addition path diagrams can be generated to help give the user an overview of the assumptions specified. Further, the package gives access to an extensive simulation procedure which covers, but is not limited to, linear latent variable models. This tool will be extremely useful e.g. for understanding the biases caused by different types of model misspecification. The lava-package also includes sophisticated inferential methods such as multigroup analyses, robust standard errors for clustered correlated data, maximum likelihood based inference with data missing at random and inference for indirect and total effects. In addition advanced model diagnostic techniques for structural equation models (fitted in lava) are available via the gof-package \cite{Holst2012}, and extensions to models for censored and binary outcomes are available via the package lava.tobit \cite{Holst2011} covered briefly in this article.

Modular programming has been a key concept during the software development thus making the process of extending the program (e.g. implementing new estimators, changing optimization routines etc.) easy, as exemplified by the above mentioned add-on packages.

Our hope is that the package will serve as a platform for testing, developing and sharing new ideas in the field of latent variable models.

2. Linear Latent Variable Models

We will define the Linear Latent Variable Model as the model defined by a Measurement part describing the responses $Y_i = (Y_{i1}, \ldots, Y_{ip})'$:

$$Y_{ij} = \nu_j + \sum_{k=1}^{l} \lambda_{jk} \eta_{ik} + \sum_{r=1}^{q} \kappa_{jr} X_{ir} + \sum_{k=1}^{l} \delta_{jk} V_{ijk} \eta_{ik} + \epsilon_{ij},$$

and a structural part describing the latent variables $\eta_i = (\eta_{i1}, \ldots, \eta_{il})'$:

$$\eta_{is} = \alpha_s + \sum_{k=1}^{l} \beta_{sk} \eta_{ik} + \sum_{r=1}^{q} \gamma_{sr} X_{ir} + \sum_{k=1}^{l} \tau_{sk} W_{isk} \eta_{ik} + \zeta_{is},$$
where \( i = 1, \ldots, n \) is the index of the sampling unit (e.g. individuals), \( j = 1, \ldots, p \) is the index of the observed variables (measurements or within cluster observations) and \( s = 1, \ldots, l \) is the index of the \( l \) distinct latent variables. In a more compact matrix notation the model can be written as

\[
Y_i = \nu + \Lambda \eta_i + K X_i + (\Delta \odot V_i) \eta_i + \epsilon_i, \quad (3)
\]

\[
\eta_i = \alpha + B \eta_i + \Gamma X_i + (T \odot W_i) \eta_i + \zeta_i, \quad (4)
\]

where \( \nu \in \mathbb{R}^p \) and \( \alpha \in \mathbb{R}^l \) are intercepts, and \( \Lambda, \Delta \in \mathbb{R}^{p \times l}, K \in \mathbb{R}^{p \times q}, \Gamma \in \mathbb{R}^{l \times q}, B, T \in \mathbb{R}^{l \times l} \) are regression coefficient matrices (defining both fixed effects and random slopes), and \( X_i \in \mathbb{R}^q, V_i \in \mathbb{R}^{p \times l}, W_i \in \mathbb{R}^{l \times l} \) are covariates. The \( \odot \) denotes the Schur product (element-wise multiplication).

The residual terms follow multivariate normal distributions, \( \epsilon_i \sim \mathcal{N}_p(0, \Sigma_\epsilon) \) and \( \zeta_i \sim \mathcal{N}(0, \Psi) \), which typically are assumed to be independent.

Note that the terms including the covariates \( V_i \) and \( W_i \) define random slope components as in the Laird-Ware mixed model formulation and differentiates the LLVM from the usual SEM formulation (Sanchez et al, 2005). Many cases can however be modeled without such terms, resulting in a more computational efficient model formulation with constant variance between individuals. In the following we will therefore initially assume that the model is parameterized by some \( \theta \) defining the matrices

\[
(\nu, \alpha, \Lambda, K, \Gamma, \Sigma_\epsilon, \Psi), \quad (5)
\]

with some restrictions on the parameter space to guarantee identification (obviously zeroes in the diagonal of \( B \), and possibly non-linear constraints between the different parameters. In Section 2.3 we will return to the general case, and demonstrate how to write up the model in atoms adapted to the case without any interaction terms. Note that we will allow non-linear constraints on the parameters between any of the elements in (5).

In the setup with \( \Delta = 0 \) and \( T = 0 \) it follows that the mean and variance of \( Y_i \) given the covariates are

\[
\mu_i(\theta) = \mathbb{E}(Y_i \mid X_i) = \nu + \Lambda (1 - B)^{-1} \alpha + [\Lambda (1 - B)^{-1} \Gamma + K] X_i, \quad (6)
\]

\[
\Sigma_{\theta} = \mathbb{Var}(Y_i \mid X_i) = \Lambda (1 - B)^{-1} \Psi (1 - B)^{-1} \Lambda' + \Sigma_\epsilon, \quad (7)
\]
where the fundamental property of the normal distribution that the marginals also are normal is exploited. Inference about $\theta$ can then be obtained by maximizing the corresponding likelihood \cite{Bollen1989}:

$$L(\theta; Y, X) \propto \prod_{i=1}^{n} \exp \left\{ -\frac{1}{2} (Y_i - \mu_i(\theta))' \Sigma^{-1}_\theta (Y_i - \mu_i(\theta)) \right\} |\Sigma_\theta|^{-\frac{1}{2}}.$$ 

2.1. Implementation

From a mathematical and implementation-wise point of view it is convenient to supplement the model formulation with the equivalent Reticular Action Model (RAM) formulation \cite{McArdle1984, Fox2009}, which have a direct connection with the underlying path diagram and also explicitly covers path analysis models. Let $U$ be the stochastic vector including the latent variables, $\eta$:

$$U = (Z_1, \ldots, Z_{p+q}, \eta_1, \ldots, \eta_l)' = (Z', \eta')',$$

where $Z = (Z_1, \ldots, Z_{p+q})'$ is the stochastic vector containing all observed variables $Z = (Y_1, \ldots, Y_p, X_1, \ldots, X_q)'$. The RAM formulation states that

$$U = v_\theta + A_\theta U + \epsilon,$$

where $v_\theta$ describes the intercepts, and $\epsilon$ is a residual term assumed to follow a zero-mean normal distribution with

$$\text{Var}(\epsilon) = P_\theta.$$ 

Hence the model is completely specified by the matrices $v_\theta$, $P_\theta$ and $A_\theta$ where the matrices generally are sparse, and the latter have zeros in the diagonal. In graph-terms the matrix $A_\theta$ represents the asymmetric paths whereas $P_\theta$ represent the symmetric paths.

Let $k$ be the total number of variables in the model (i.e. $A_\theta \in \mathbb{R}^{k \times k}$). We let $J$ be the matrix that picks out the observed variables from $U$ (see Section \ref{AppendixA}), and define

$$G_\theta = J(1 - A_\theta)^{-1}.$$ 

Now it follows that

$$\text{Var}(Z) = \Omega_\theta = G_\theta P_\theta G_\theta'.$$
and similarly the mean of the observed variables is then specified by the model structure as:

$$E(Z) = \xi_\theta = G_\theta v_\theta.$$  

(13)

The parameter \( \theta \) can then be estimated by maximizing the log-likelihood, \( \ell \), for the full data vector \( Z \)

$$\hat{\theta}_{ML} = \arg \max_\theta \ell(\theta \mid Z = z).$$  

(14)

At a first glance this formulation seems restrictive in the sense that the covariates also have to be normally distributed. However, if we split the parameter vector into \( \theta = (\theta_1, \theta_2) \), where \( \theta_1 \) parameterizes the conditional distribution of \( Y \) given \( X \) and \( \theta_2 \) are the mean and variance parameters of the covariates, then by Bayes formula the probability density for the joint distribution can be written as the product of the conditional density multiplied by the marginal density:

$$f_{\theta_1, \theta_2}(y, x) = f_{\theta_1}(y \mid x)f_{\theta_2}(x).$$  

(15)

It follows that maximum likelihood inference about the parameters \( \theta_1 \) is independent of the model for the covariates, and hence finding the MLE of the conditional likelihood is equivalent to finding the MLE of the joint likelihood. If we fix \( \theta_2 \) to the corresponding MLE, then the expected information agrees in the two models:

$$-E\left( \frac{\partial^2 \log f_{\theta_1}(y \mid x)}{\partial \theta_1 \partial \theta_1'} \right) = -E\left( \frac{\partial^2 \log f_{\theta_1, \hat{\theta}_2}(y, x)}{\partial \theta_1 \partial \theta_1'} \right).$$  

(16)

An important advantage of the RAM formulation is that the empirical variance and mean are sufficient statistics for the parameters, thus giving the joint formulation a computational advantage over the conditional likelihood which requires explicit calculation of the likelihood contribution of each individual observation. In particular given the sufficient statistics, which are easily computed, the computational complexity in the RAM parameterization is independent of \( n \).

2.2. Inference - standard SEM

In the following we will describe inference under a priori non-linear constraints \( \Omega = \Omega_\theta \) (twice differentiable w.r.t. \( \theta \)). Letting \( \hat{\mu} \) denote the empirical mean of the observed variables, we define

$$W_\theta = [\hat{\mu} - \xi_\theta][\hat{\mu} - \xi_\theta]'$$  

(17)
and
\[ T_\theta = \hat{\Sigma} + W_\theta, \]  
(18)
where \( \hat{\Sigma} \) is the ML covariance matrix estimate (non-central estimate). We will exploit that
\[
\sum_{i=1}^{n} (z_i - \xi_\theta)(z_i - \xi_\theta)' = \sum_{i=1}^{n} (z_i - \hat{\mu} + \hat{\mu} - \xi_\theta)(z_i - \hat{\mu} + \hat{\mu} - \xi_\theta)'
\]
\[ = n(\hat{\Sigma} + W_\theta) + \]
\[ n[(\hat{\mu} - \hat{\mu})(\hat{\mu} - \xi_\theta)' + (\hat{\mu} - \xi_\theta)(\hat{\mu} - \hat{\mu})'] = nT_\theta. \]  
(19)
The complete log-likelihood then is given by
\[
\ell(\theta | z_1, \ldots, z_n) = \sum_{i=1}^{n} \left\{ -\frac{k}{2} \log(2\pi) - \frac{1}{2} \log |\Omega_\theta| - \frac{1}{2}(z_i - \xi_\theta)'\Omega_\theta^{-1}(z_i - \xi_\theta) \right\}
\]
\[ = -\frac{nk}{2} \log(2\pi) - \frac{n}{2} \log |\Omega_\theta| - \frac{n}{2} \text{tr}\{T_\theta\Omega_\theta^{-1}\}, \]  
(20)
with score
\[
\frac{\partial \ell(\theta)}{\partial \theta} = \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1}T_\theta\Omega_\theta^{-1} \right]
\]
\[ - \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1} \right] \]
\[ - \frac{n}{2} \left( \frac{\partial \text{vec} T_\theta}{\partial \theta'} \right)' \text{vec} \left( \Omega_\theta^{-1} \right), \]  
(21)
and the MLE is obtained by solving the corresponding score equation by Fisher scoring or a similar iterative procedure. See Appendix B for details w.r.t. expressions for the relevant matrix derivatives and information matrix.

In certain situations we may need to calculate conditional moments, for instance to calculate conditional residuals (model diagnostics) or the conditional likelihood given the covariates (likelihood ratio testing, model selection, etc.). Here we need to apply the selection matrix, \( J_Y \), that picks
out the endogenous variables, $Y$, of $U$, and the cancellation matrix, $p_X$, that sets all exogenous variables, $X$, of $U$ to zero (see Section Appendix A). Notice we do not put any distributional assumptions on the exogenous variables.

Now

$$\mu_i(\theta) = \mathbb{E}(Y \mid X = x) = J_Y(1 - A_\theta)^{-1}(p_Xv_\theta + v_x), \quad (22)$$

where $v_x$ is the $p$-vector which is zero everywhere but on the index of the exogenous variables where it is set to $x$, and

$$\Sigma_\theta = \text{Var}(Y \mid X) = J_Y(1 - A_\theta)^{-1}(p_XP_\theta p_X')(1 - A_\theta)^{-1'}J_Y'. \quad (23)$$

2.3. Interactions with latent variables

Including interactions between covariates and random effects in the model, i.e. with non-zero $\Delta$ and $T$ in (3) and (4), we clearly lose the property of constant variance between individuals and the empirical mean and variance are therefore not sufficient. With $\Delta$ and $T$ consisting of ones in all entries, the marginal variance of $Y_i$, $\text{Var}(Y_i \mid X_i, V_i, W_i)$, will however take the same form as (7), exchanging $B$ with

$$B_i = B + W_i, \quad (24)$$

and

$$\Lambda_i = \Lambda + Z_i, \quad (25)$$

hence the expression for the likelihood contribution and its derivatives of a single individual will take the same form as derived in Appendix B. With free parameters in $\Delta$ or $T$ we can still adapt the model by adding one or more degenerate random effects. For instance, the term

$$\xi_{ij} = \delta_{jk}V_{ijk}\eta_{ik} + \epsilon_{ij} \quad (26)$$

where $\eta_{ik}$ and $\xi_{ijk}$ follow normal distributions, can trivially be parameterized as the simultaneous equation

$$\xi_{ij} = \delta_{jk}\tilde{\eta}_{ik} + \epsilon_{ij}, \quad (27)$$

$$\tilde{\eta}_{ik} = V_{ijk}\eta_{ik} + 0, \quad (28)$$

hence the model is expanded to include a random effect with residual term with variance 0 with fixed slope parameter $Z_{ijk}$, and the variance of the
observed variables in such a model therefore takes the form as described above \(^{(24)}\), i.e. \(\Delta = 1\).

The possibility of including interactions with the random effects adds important flexibility to the model class for instance when modeling longitudinal data or to account for certain types of variance heterogeneity.

2.4. Non-linear effects

Allowing non-linear parameter constraints or non-linear effects of some covariates opens up for several interesting applications for instance in dose-response modeling. An important extension of the model framework is therefore to allow parametric non-linear functions \(\phi^{(j)}, j = 1, \ldots, p\) and \(\psi^{(s)}, s = 1, \ldots, l\) of the covariates to enter the model:

\[
Y_{ij} = \nu_j + \sum_{k=1}^{l} \lambda_{jk} \eta_{ik} + \sum_{r=1}^{q} \kappa_{jr} X_{ir} + \sum_{k=1}^{l} \delta_{jk} V_{ijk} \eta_{ik} + \phi^{(j)}(X_{i1}, \ldots, X_{iq}) + \epsilon_{ij},
\]

and

\[
\eta_{is} = \alpha_s + \sum_{k=1}^{l} \beta_{sk} \eta_{ik} + \sum_{r=1}^{q} \gamma_{sr} X_{ir} + \sum_{k=1}^{l} \tau_{sk} W_{isk} \eta_{ik} + \psi^{(s)}(X_{i1}, \ldots, X_{iq}) + \zeta_{is}.
\]

Though we introduce non-linear effects in the model, we will still denote this model a Linear Latent Variable Model, as there are only linear effects of latent variables and endogenous variables in the model. Hence the observed data likelihood and its derivatives still have a closed form solution.

2.5. Multigroup analysis

A useful generalization of the model framework is the multigroup model, where we have several groups of data and specify a LLVM for each group. This naturally leads to the log-likelihood

\[
\log L(\theta \mid Y, X, V, W) = \sum_{g \in G} \log L_g(\theta_g \mid Y_g, X_g, V_g, W_g)
\]

where the intersection of the parameters \((\theta_g)_{g \in G}\) is not empty. Unbalanced designs and data with values missing at random (Little and Rubin, 2002) is naturally handled by this extension by forming groups from the different missing data patterns. Note that the additive structure of the log-likelihood makes the calculation of score functions and information matrices for the multigroup model explicitly available.
3. Model specification

The lava package aims to deliver a dynamic model specification experience in the sense that adding or removing model elements should be as easy as possible, and the model specification should be familiar to users accustomed to specifying models in for example glm in R. In order to achieve this we have designed a formal system for interactively specifying the complex hierarchical structure of a latent variable model. We believe this to be an important novel contribution, since the difficult specification of models in other languages often proves to be a significant obstacle.

The implementation relies on R (R Development Core Team, 2010) and the following packages all available from the Comprehensive R Archive Network (CRAN) : mvtnorm (Genz et al, 2009), graph (Gentleman et al, 2009), survival (Therneau and original R port by Thomas Lumley, 2009), numDeriv (Gilbert, 2009) and gof (Holst, 2012). The graphical system builds on graphviz (Gansner and North, 1999) and the R-package Rgraphviz (Gentry et al, 2009) (available from Bioconductor Gentleman et al (2004)).

The specification of models in the lava-language is primarily achieved via the constructor function lvm and the two methods regression and covariance (see Table II). A new model object is initialized with the constructor lvm

```r
> m1 <- lvm()
```

which creates an empty lvm-object. Variables (or a multivariate regression formula as described below) can be fed to the lvm-function as arguments in order to control the order of entry in the graph layout. However, this is optional as variables automatically will be added during the process of defining the linear structure. A list of formulas is also valid as a shortcut for successive calls to regression (see below).

3.1. Specifying Linear Relationships

Linear associations between variables are specified via the member function regression taking the character vector arguments to and from. For convenience a replacement function, regression<-, is also available which in addition supports specification via the usual formula statements in R. As a simple example we will specify the following structural equation model
Function | Task
--- | ---
Primary functions
lvm | Constructor of new model
regression | Add regression association to model
covariance | Add correlation between residuals terms
intercept | Add intercept parameter
constrain | Add non-linear covariate effects or parameter constraints
Secondary functions
latent | Define latent variables in model
addvar | Add variable to model
parfix | Define equality constraints index of parameters
parameter | Add a parameter name (for use with constrain)
cancel | Remove previously defined associations
kill | Remove variables from model

Table 1: Model building blocks

with two measurement models

\[ Y_j = \mu_j + \lambda_{1j}U_1 + \epsilon_{1j}, \]
\[ Z_j = \nu_j + \lambda_{2j}U_2 + \epsilon_{2j}, \quad j = 1, \ldots, 3, \]

and structural model defined by

\[ U_1 = \delta_1X_1 + \delta_2X_2 + \zeta_1, \]
\[ U_2 = \beta_1X_1 + \beta_2X_2 + \zeta_2, \]

and \( \text{Cov}(\zeta_1, \zeta_2) \neq 0 \). The model is illustrated in the path diagram of Figure 1.

The following commands specifies a multivariate linear regression model with two covariates \( x_1 \) and \( x_2 \) and two outcomes \( u_1 \) and \( u_2 \)

\[
\text{> } m1 \leftarrow \text{regression}(m1, "u1", c("x1", "x2"))
\]
\[
\text{> } \text{regression}(m1) \leftarrow u2 \sim x1 + x2
\]
In the following we will focus on the replacement functions and the formula specification as defined by the second line, but generally for all the available methods a standard function is available and arguments can be given as character vectors as above. A more compact call would simply be

```r
> m1 <- lvm(c(u1, u2) ~ x1 + x2)
```

Next, we define `u1` and the `u2` as a latent/unobserved variables using the `latent`-function:

```r
> latent(m1) <- ~u1 + u2
```

Again arguments can generally be given as character vectors instead of a formulas. To remove the latent status from a variable we simply use the `cancel=TRUE` argument (`latent(m1,cancel=TRUE) <- ...`).

Next we define the measurement part of the model:

```r
> regression(m1) <- c(y1, y2, y3) ~ u1
> regression(m1) <- c(z1, z2, z3) ~ u2
```

Covariance between residual terms can be specified using the replacement function `covariance<-` (or the function `covariance`) where the argument is a vector (or formula) of variables that are assumed to be pairwise correlated. In the current model specification, the residuals of the two latent variables are assumed to be conditionally independent given the covariates, and in order to define correlation between the residual terms of `u1` and `u2` we write

```r
> covariance(m1) <- u1 ~ u2
```

which specifies $\text{Cov}(\zeta_1, \zeta_2) \neq 0$, thus completing the specification of the model defined by the path diagram in Figure [1]. Note that the model is specified independently of any data. The model is linked to data when parameters are estimated using the `estimate` function (see Section [6]). Here it is important that the manifest variable names used in the model specification corresponds to the variable names in the data-frame.

Removal of associations or variables can be achieved with the `cancel` (and `cancel<-`) function which takes a character vector (or formula) as argument, removing any associations between all the variables in the vector. To remove the previously specified correlation and instead add a regression association between `u1` and `u2`, we write

```r
> cancel(m1) <- ~u1 + u2
> regression(m1) <- u2 ~ u1
```
Notice that the last regression call defining the association between $u_1$ and $u_2$ does not cancel the earlier defined predictors of $u_2$. Hence the current definition says (see Figure 2) that

$$E(U_2 \mid U_1, X_1, X_2) = \mu + \beta_0 U_1 + \beta_1 X_1 + \beta_2 X_2$$

for some parameters $(\mu, \beta_0, \beta_1, \beta_2)$.

To completely remove one or more variables from the model we can use the `kill<-` function.

### 3.2. Constraining Parameters

Defining restrictions on some parameters is usually needed in order to obtain an identifiable model, and by default lava will automatically set reasonable restrictions when model parameters are estimated (see Section 6). Also, in situations where we need to test associations or use a priori knowledge in the model building, constraints on some parameters are needed. The lava package allows specification of completely general non-linear constraints on all parameters.

The most common type of constraints are identity constraints where one or more parameters are fixed to either a specific numerical value or
to be equal to a common free parameter (equality constraints). For covariance and regression parameters (slope-parameters) the `regression` and `covariance` function can be used, and for the intercepts $\alpha$ and $\nu$ in (34), the `intercept<-` and `intercept` functions are used.

As an continuing example we will specify a new multivariate regression model with three outcomes, $(Y_1,Y_2,Y_3)$, and two predictors, $(X_1,X_2)$:

$$Y_i = \beta_i X + \gamma_i Z + \epsilon_i, i = 1, \ldots, 3$$

```r
> mregr <- lvm(c(y1, y2, y3) ~ x + z)
```

### 3.2.1. Constraining regression parameters

The restrictions of slope parameters can be accomplished with the `regression` function. For example fixing the slopes of $Y_1$, $Y_2$ and $Y_3$ on $X$ to be identical, $b_1$, and defining $Z$ as an offset (i.e. slope 1), can be achieved with the calls

```r
> regression(mregr, c(y1, y2, y3) ~ x) <- "b1"
> regression(mregr, c(y1, y2, y3) ~ z) <- 1
```

To simultaneously define several different constraints a list can be given as the right-hand side argument

```r
> regression(mregr, c(y1, y2) ~ x + z) <- list(1, "a", 2, "b")
```

All parameters for the first response on the covariates are given first, then for the second response, and so on. Hence in the above example we have

$$Y_1 = X + aZ + \cdots \quad \text{and} \quad Y_2 = 2X + bZ + \cdots \quad (33)$$
When defining constraints (intercepts, covariance or regression constraints) any missing associations will automatically be added to the model object. Hence the following call will add an extra level to the model, with a top-level response \( W \) with identical effects of \( Y_1, Y_2 \) and \( Y_3 \) (see Figure 3):

\[
> \text{regression(mregr, } w \sim y1 + y2 + y3) \leftarrow \text{"beta"}
\]

To remove the constraints again (but not removing the associations) we simply fix to the logical constant \( \text{NA} \):

\[
> \text{regression(mregr, } w \sim y1 + y2 + y3) \leftarrow \text{NA}
\]

### 3.2.2. Constraining covariance parameters

Constraints on the covariance between residual terms are set with the `covariance` function, with a similar syntax. For example, to fix the covariance of the residuals terms of \( Y_1 \) and \( Y_2 \) to 0.5, and the variance of the residual term of \( Y_1 \) to a parameter \( v1 \):

\[
> \text{covariance(mregr, } y1 \sim y1 + y2) \leftarrow \text{list("v1", 0.5)}
\]

If we only need to constrain the variance parameters (i.e. not covariances) then the following syntax can be used

\[
> \text{covariance(mregr, } ^\sim y1 + y2) \leftarrow \text{"v"}
\]

here setting the residual variance of \( Y_1 \) and \( Y_2 \) to be identical.

To fix the variance of the variables to different values, we simply give a list of the correct length as argument, e.g.

\[
> \text{covariance(mregr, } ^\sim y1 + y2) \leftarrow \text{list("v", 0.3)}
\]

If we are interested in fixing only the covariance parameters (and not the diagonal) we can add the `pairwise=TRUE` argument. For instance to specify that the covariances between \( Y_1, Y_2, \) and \( Y_3 \) are the same, we can call

\[
> \text{covariance(mregr, } ^\sim y1 + y2 + y3, \text{ pairwise } = \text{ TRUE}) \leftarrow \text{"r1"}
\]

thus specifying a compound-symmetry structure.

Finally a syntax like the one used by the `regression`-function can be used, such that

\[
> \text{covariance(mregr, } c(y1, y2) \sim y2 + y3) \leftarrow \text{list}(0.5, \text{"r", "r0", + 0.3)}
\]
defines the following covariance structure between residual terms $\varepsilon_1, \varepsilon_2, \varepsilon_3$ corresponding to $Y_1, Y_2, Y_3$ (with the $\cdot$ denoting elements that are not affected by the call):

$$\text{Var}((\varepsilon_1, \varepsilon_2, \varepsilon_3)^\prime) = \begin{pmatrix} \cdot & 0.5 & r \\ 0.5 & r_0 & 0.3 \\ r & 0.3 & \cdot \end{pmatrix}$$ (34)

As with intercept and slope parameters, we can remove covariance constraints by fixing parameters to the value NA. We end up with a model as defined by the path-diagram in Figure 3.

Figure 3: plot(mregr, labels=TRUE, diag=TRUE)

3.2.3. Constraining intercepts

To fix the intercepts of the three outcomes to be identical, we can write

> intercept(mregr, ~y1 + y2 + y3) <- "mu"

Instead of a parameter name, mu, we could have chosen a numerical, say 0.

Notice that a character vector could also have been given instead of the formula. The value on the right can be given as a list, hence to fix the intercept of $Y_1$ and $Y_2$ to be identical and $Y_3$ to zero, we could call

> intercept(mregr, ~y1 + y2 + y3) <- list("mu", "mu", 0)

3.3. Simultaneously specifying constraints on intercepts, slopes and variances

Using the formula syntax with the regression method it is possible to simultaneously specify constraints on intercept, regression and covariance
parameters. A special function, $f$, can be used within the formula to specify the slope parameters, and further a pair of square brackets can be appended to each variable in the formula. Inside the square bracket the intercept of the variable can be defined, or alternatively both the intercept and residual variance separated by a colon. As an example we can specify

$$
Y_1 = a + bU + \epsilon_1, \quad \epsilon_1 \sim \mathcal{N}(0, 1),
Y_2 = a + bU + \epsilon_2, \quad \epsilon_2 \sim \mathcal{N}(0, v),
U = b_2X + \zeta,
$$

using the square-bracket syntax:

```r
m2 <- lvm()
regression(m2) <- c(y1[a:1], y2[a:v]) ~ f(u[0], b)
regression(m2) <- u ~ f(x, b2)
```

An equivalently even more compact model specification can be obtained using a list of formulas with the model initializer `lvm`, hence an equivalent way of specifying `m2` would be

```r
m2 <- lvm(list(c(y1[a:1], y2[a:v]) ~ f(u[0], b), u ~ f(x, b2)))
```

If the index of the parameter is known (see the `coef` method below) the `parfix` method can also be used to simultaneous constrain parameters. For example to fix the parameters of a model `m`, at positions 1, 4 and 5 to the values `a`, `a` and 1, we can call

```r
parfix(m, c(1, 4, 5)) <- list("a", "a", 1)
```

### 3.4. Random slopes

Random slope effects (i.e. the matrices $V_i$ and $W_i$) can be defined by constraining the slope parameter of a latent variable to the name of a covariate. The covariate does not necessarily need to be added to the model explicitly, as the slope parameters are matched to the column names of the `data.frame` that is used during estimation (see Section 6).

In a standard structural equation model covariates enter the model in order to describe differences in the mean structure. Another question might be whether covariates can influence the variation in an outcome. In a situation where the variance depends on a covariate, the usual assumptions of variance homogeneity of the latent variables are not met and the model is not a SEM. In the LLVM this situation can be handled by defining a regression on the variance component using the random slope specification.
Another important example is to use the random slopes to describe variation in longitudinal studies, and in combination with measurement models this allows us to formulate random regression models taking measurement error into account (see Figure 4).

Figure 4: Longitudinal analysis with measurement models. Here illustrated with three time points where $\eta_1$, $\eta_2$, $\eta_3$ are modeled by a random intercept, $\iota$, and a random slope defined by the covariate $z_i$, $i = 1, 2, 3$ and the random effect $\kappa_i$: $\eta_i = \iota + \kappa_i z_i + \zeta_i$. As the $\eta$'s are only indirectly observed a measurement model is employed at each time point.

3.5. Non-linear constraints and effects

Non-linear parameter constraints are defined using the `constrain<-function`. The syntax is

```r
> constrain(m, formula) <- function(x) ...
```

where `m` is the `lvm`-object, and the left-hand side in the `formula` specifies the parameter that is a (non-linear) function of the parameters or covariates defined by the right-hand side, and the `function` defines this association. The result of the function can optionally be given the attributes `grad`, defining the analytic gradient, and `inv`, defining a monotonic transformation (typically the inverse function) used in conjunction with confidence limit calculations (see Section 6 on Statistical Inference).
As an example we will define the model

\[ Y_{ij} = \mu + b_j X_i + \epsilon_{ij}, \quad j = 1, 2 \]  

(35)

with \( \text{Cov}(\epsilon_{i1}, \epsilon_{i2}) = 0, \text{Var}(\epsilon_{i}) = \theta \):

```r
> mconstr <- lvm()
> regression(mconstr, c(y1, y2) ~ x) <- list("b1", "b2")
> intercept(mconstr, "y1 + y2") <- "mu"
> covariance(mconstr, "y1 + y2") <- "v"
```

To restrict the variance-parameter to live on the positive real axis, we add the constraint

\[ v = \exp(\alpha) \]  

(36)

with

```r
> constrain(mconstr, v ~ alpha) <- exp
```

using a log-link, and the parameter \( \alpha \) (log-variance) is added to list of model parameters. In this example we do not set any attributes and numerical derivatives of the constraint will therefore be used (based on the package `numDeriv`) during estimation. Notice that constraints on the variance parameters can be set automatically by the `estimate`-function (see Section 6). For general domain constraints the function `range.lvm` can be used as the right-hand side argument, e.g. `range.lvm(a=1,b=Inf)` will bound the parameter to the interval \((1, \infty)\).

Continuing the example, we could define the intercept, \( \mu \), to be the product of the two slope-parameters,

```r
> constrain(mconstr, mu ~ b1 + b2) <- prod
```

Constraints can be removed by letting the RHS be `NULL` (or `NA`)

```r
> constrain(mconstr, "mu") <- NULL
```

If we instead wish to add a non-linear effect of \( x \) on \( y_i \) and \( y_2 \):

\[ Y_{ij} = \alpha + \Phi(\beta X_{ij}) + b_j X_{ij} + \epsilon_{ij}, \quad i = 1, 2 \]  

(37)

with \( \Phi \) denoting the standard normal cumulative distribution function, we can make the call

```r
> constrain(mconstr, mu ~ alpha + beta + x) <- function(x) x[1] + pnorm(x[2] * x[3])
```
3.6. Complex models with feedback or co-existence of regression associations and covariance between residuals

The linear latent variable model framework in principle allows pathways going in both directions between two variables, i.e., feedback, and also simultaneous presence of both a regression association and covariance between residual terms of the same two variables. Note that both cases are different from a simple correlation between the residuals.

A simple example of an identified model is illustrated in Figure 5, where the aim is to estimate the effect of $Y$ on $X$ while taking the unmeasured confounder $C$ into account. This can be achieved using an instrumental variable, $I$, which by assumption must be (strongly) correlated with $X$, independent with the confounder, and conditionally independent with the $Y$ given $X$. We can model the inflation in covariance between $Y$ and $X$ using a correlation between their residual terms, which can be implemented as

```r
> m <- lvm()
> regression(m) <- y ~ x
> regression(m) <- x ~ i
> covariance(m) <- x ~ y
```

Figure 5: Path diagram showing the association between the variables $Y$ and $X$ with unmeasured confounder $C$ and instrumental variable $I$.

4. Inspecting the model assumptions

The philosophy of the `lava` package is to separate the model specification from the actual data, since examination of the model structure before actual
| Function   | Task                                          |
|------------|----------------------------------------------|
| plot       | Plot the graph of the model                   |
| regression | Display parameter restrictions                |
| intercept  | Display parameter restrictions                |
| covariance | Display parameter restrictions                |
| exogenous  | Extract (or define) exogenous variables (predictors) |
| endogenous | Extract set of endogenous variables (responses) |
| latent     | Extract set of latent variables               |
| manifest   | Extract set of manifest (observed) variables  |
| vars       | Extract all variables                         |
| children   | Extract children of a node                    |
| parents    | Extract parents of a node                     |
| coef       | Get list of parameters                        |
| constrain  | Display non-linear constraints                |
| path       | Extract direct and indirect unidirectional pathways between nodes |
| subset     | Extract sub-model                             |
| merge, %+% | Merge models                                  |

Table 2: Model inspection functions.

estimation is often an important aspect of modelling within this class of statistical models. The lava package includes several functions as an aid to obtain an overview of the model assumptions (see Table 2). In general, the below described methods also applies for a lvmfit-object (see Section 6).

The plot-method (see Figure 1 and 3) visualizes the model using a path-diagram, i.e. a graph structure where linear (causal) associations are shown with directed edges and covariance between residuals are shown as bidirectional edges. Manifest variables are shown as rectangles and latent variables as ellipsoids. The parameter constraints can be added as labels on the edges with the argument labels=TRUE and variance parameters can be added with the argument diag=TRUE (see Figure 3). The plot-function will be explained in more details in Section 10.

Via the summary function, a complete overview of the model and (identity) parameter constraints can be obtained. Returning to the model mregr defined in Figure 3:

> summary(mregr)

Latent Variable Model
with: 6 variables.
Npar=11+2

Regression parameters:
\[
\begin{align*}
y_1 & \quad \ast \\
y_2 & \quad \ast \\
y_3 & \quad \ast \\
x & \quad 1 \ 2 \ b_1 \\
z & \quad a \ b \ 1 \\
w & 
\end{align*}
\]

Covariance parameters:
\[
\begin{align*}
y_1 & \quad v \ 0.5 \ r \\
y_2 & \quad 0.5 \ r_0 \ 0.3 \\
y_3 & \quad r \ 0.3 \ \ast \\
w & \quad \ast 
\end{align*}
\]

Intercept parameters:
\[
\begin{align*}
y_1 & \quad y_2 \ y_3 \ w \\
\mu & \quad \mu \ 0 \ \ast 
\end{align*}
\]

Here the adjacency-matrix for the graph of all the unidirectional edges of the path-diagram can be read off under the title Regression parameters (i.e. slopes). Similarly the covariance structure of the residual terms and the intercept structure are shown. An empty element indicates that there is no direct association. A star indicates a free parameter and all other entries are either fixed numerical values or parameter names as defined by identity constraints during model specification. These three matrices can also be extracted via calls to regression, covariance or intercept.

The exogenous variables (covariates) of a lvm object can be identified with the exogenous function. Similarly a list of the endogenous (manifest) variables can be obtained with the function endogenous and the subset of these variables that do not predict other variables (top-level outcomes) can be shown by including the argument top=TRUE. In a similar way the observed and latent variables can be shown with manifest and latent. All variables of the model are listed with the vars function

> exogenous(mregr)

[1] "x" "z"

> endogenous(mregr, top = TRUE)
The `children` and `parents` functions extracts the children respectively the parents of one or several nodes in the unidirectional graph of the model, e.g.

```r
> children(mregr, ~x + y1)
[1] "y1" "y2" "y3" "w"
> parents(mregr, ~w)
[1] "y1" "y2" "y3"
```

The pathways from one variable to another can be viewed with the `path` function which returns a list of character vectors indicating the (causal) path.

```r
> path(mregr, w ~ x)
[[1]]
[1] "x" "y1" "w"
[[2]]
[1] "x" "y2" "w"
[[3]]
[1] "x" "y3" "w"
```

The function `subset` can be used to extract subsets of a model. To extract the upper level of the path analysis we call

```r
> subset(mregr, ~y1 + y2 + y3 + w)
```

which keeps all parameter restrictions of the original model. Conversely, `lvm` models can be merged with the `merge` method (or using the operator syntax: `m%++%m2`).

To examine the parameters (and in particular their order) one can call the `coef`-function.

```r
> coef(mregr)

         m1     m2   p1   p2   p3   p4   p5   p6
"y1" "w" "y1<-z" "y2<-z" "y3<-x" "w<-y1" "w<-y2" "w<-y3"
    p7   p8  p9  p10  p11
"y1<->y1" "y2<->y2" "y3<->y3" "w<->w" "y1<->y3"
```

23
where "<−" represents slope parameters (e.g. \( z \) on \( y_1 \)) and "<−>" represents covariance (See also the describecoef function). With the argument labels=TRUE we can get the same vector but with all parameter labels substituted by their constraints.

```r
> coef(mregr, labels = TRUE)
```

```
m1  m2  p1  p2  p3  p4  p5  p6
"mu" "w" "a" "b" "b1" "w<−y1" "w<−y2" "w<−y3"
P7  P8  P9  P10  P11
"v" "r0" "y3<−>y3" "w<−>w" "r"
```

The non-linear parameter constraints or non-linear regression specifications can be shown with

```r
> constrain(mconstr)
```

```
$v
function (x) .Primitive("exp")
attr("args")
[1] "alpha"

$mu
function (x)
x[1] + pnorm(x[2] * x[3])
attr("args")
[1] "alpha" "beta" "x"
```

5. Simulation

Simulation is a major component of modern statistics, which allows us to experimentally study the properties of a statistical method under various alternatives and to verify (or reject) preliminary ideas. The lava package includes the sim method offering a convenient tool for performing simulation studies from very general models.

As an initial example we will create a `data.frame` with 100 observations from the structural equation model `m1` defined in Section 3.1.

```r
> mydata <- sim(m1, 100)
```

The default parameter values are that all intercepts are 0, slope and residual variance parameters are 1, and covariance parameters are 0.5. To change the simulation parameters one can either fix the relevant parameters of the
Function | Task
---|---
sim | Simulation method for lvm-objects
`functional,constrain` | Introduce non-linearities in simulation
distribution | Change distribution and link of variables
heavetail | Define heavy tailed distribution of a variable
normal.lvm | Normal distribution
poisson.lvm | Poisson distribution
binomial.lvm | Binomial distribution
uniform.lvm | Uniform distribution
weibull.lvm | Weibull accelerated failure times

Table 3: Simulation methods.

model to the desired numerical values as described in the previous section, or give the parameters as the argument `p` directly to the `sim` method. For instance the following two calls will both simulate 1000 observations from the model `mregr` with the default parameter values, except that the residual variance of `w` is set to 2, the intercepts of `Y_1` and `Y_2` (`µ`) are set to 1, and `β = 1.5` (the slope of `Y_1` on `W`)

```r
> d.mregr <- sim(mregr, 1000, p = c(mu = 2, beta = 1.5, `w<-w` = 2))
> d.mregr2 <- sim(mregr, 1000, p = c(y1 = 2, `w<-y1` = 1.5, `w<-w` = 2))
```

To simulate data with heavier tails than the normal distribution the `heavetail` method can be used. The following defines `y1` and `y2` to be realizations from an unstructured bivariate normal distribution $N(µ, Σ)$:

```r
> mhtail <- lvm()
> covariance(mhtail) <- y1 ~ y2
```

We let $Y$ be a stochastic variable with this distribution, then the following call

```r
> heavetail(mhtail, df = 3) <- `y1 + y2`
```

will allow us to draw simulations of $y1$ and $y2$ from the distribution of $Y_j(3/Q)^{0.5}$ where $Q \sim \chi_3^2$, i.e., leading to a multivariate $t$-distribution with covariance matrix $Σ$, mean $µ$, and $ν = 3$ degrees of freedom, described by the density

$$f(x \mid µ, Σ, ν) = \frac{Γ((k + ν)/2)}{ν^{k/2}Γ(ν/2)Γ(1/2)^k} \frac{|Σ|^{-1/2}}{(1 + \frac{1}{ν}(x - µ)Σ^{-1}(x - µ))^{(ν+k)/2}}$$

(38)
where $k = 2$ is the dimension. The same realization of $Q$ will be used on both $y_1$ and $y_2$ in the above code. To make simulations where a different realization of the $\chi^2$-distribution is used for each outcome (leading to a star-shaped distribution), one simply has to make separate `heavytail` calls as in

```r
> heavytail(mhtail, df = 3) <- ~y1
> heavytail(mhtail, df = 3) <- ~y2
```

The method can be used with different degrees of freedom for different variables in the model, and thus gives access to an easy way of simulating models with various degrees of outlier contamination.

To allow simulations from quite general models, two additional replacement functions are available, `functional` and `distribution`. The `functional` replacement function is used for defining (nonlinear) functional relationships between variables and has the syntax

```r
functional(x, to, from) <- value
```

where `x` is a `lvm`-object, `from` and `to` are predictor and outcome, respectively, and `value` is a real function describing the functional form. In the model `mregr` (Figure 3) we have

$$E(Y_3|X, Z) = b_1X + Z$$

(39)

With the call

```r
> functional(mregr, y3 ~ x) <- function(x) x^2
```

we can simulate from the model

$$E(Y_3|X, Z) = b_1X^2 + Z$$

(40)

with the coefficient $b_1$ defined by the earlier identity constraint. To define a more complex polynomial effect of $X$ we can make a copy of the predictor with the `copy` function and apply `functional` on the copy

```r
> copy(mregr) <- x ~ x^2
> regression(mregr, y3 ~ x^2) <- "b2"
> functional(mregr, y3 ~ x^2) <- function(x) x^3
```

leading to the mean-structure

$$E(Y_3|X, Z) = b_1X^2 + b_2X^3 + Z$$

(41)

An alternative approach is to use the `constrain` function, e.g.
would define the model
\[ \mathbb{E}(Y_3|X, Z) = b_0 + b_1 X + b_2 X^2 + b_3 X^3 + Z \] (42)

The major difference between the two methods is that `functional` only has an impact on the `sim` method and not on the inferential methods, whereas `constrain` alters the model fitted by `estimate` (see Section 6).

The `distribution` replacement function is used for defining the link/distribution of variables, with syntax

\[
distribution(x, \text{variable}) \leftarrow \text{value}
\]

where \( x \) is a `lvm`-object, \( \text{variable} \) is the variable to define the distribution of, and \( \text{value} \) is a function defining the random generator for \( \text{variable} \) taking the form

\[
\text{function}(n, \mu, \var)
\]

where \( n \) defines the number of samples, \( \mu \) is the mean, and \( \var \) is the variance as defined by the latent variable model. Some of the most common distributions have been predefined in the functions `uniform.lvm`, `normal.lvm`, `binomial.lvm`, `poisson.lvm`, `weibull.lvm`.

As an example we will define a simple hierarchical model structure (path analysis)

\[
> \text{msim} \leftarrow \text{lvm}(t \sim y + u + z) \\
> \text{regression(msim)} \leftarrow y \sim u + x + z \\
> \text{regression(msim)} \leftarrow c(z, u) \sim x
\]

To change the distribution of \( Y \) to a Bernoulli distribution we simply call

\[
> \text{distribution(msim, \sim y) \leftarrow \text{binomial.lvm()}}
\]

The default link is logit

\[
\mathbb{P}(Y = 1 | U, X, Z) = \frac{\exp(\beta_0 + \beta_1 X + \beta_2 U + \beta_3 Z)}{1 + \exp(\beta_0 + \beta_1 X + \beta_2 U + \beta_3 Z)},
\] (43)

but a complementary log-log (cloglog) or probit link can be chosen via the link argument, e.g.
Similarly we can define the conditional distribution of $Z$ as Poisson

$$Z \sim \text{pois}(\mathbb{E}(Z \mid X)),$$

$$\log(\mathbb{E}(Z \mid X)) = \gamma_0 + \gamma_1 X,$$

and the conditional distribution of $U$ as uniform

$$U = \lambda_0 + \lambda_1 X + \sqrt{\sigma_U^2} U_0,$$

$$U_0 \sim \text{unif}(-1, 1),$$

and let $T$ follow a proportional hazards model with Weibull baseline with scale parameter 1.25 and shape parameter 2 (and no censoring)

$$\lambda(t) = \lambda_0 \text{Weibull}(1.25; 2) \exp(-\alpha_0 + \alpha_1 U + \alpha_2 Y + \alpha_3 Z).$$

The default simulation parameter values leads to intercepts $\beta_0 = \lambda_0 = \gamma_0 = \alpha_0 = 0$ and all the remaining parameters are 1 (including the residual variance $\sigma_U^2$).

The default distribution of exogenous variables is the standard normal distribution, $X \sim \mathcal{N}(0, 1)$ (and independence between the exogenous variables). It is also possible to let a variable be deterministic by simply assigning a list encapsulating the data:

$$\text{distribution}(\text{msim}, \sim x) <- \text{list} (\text{seq}(0, 1, \text{length.out}=100))$$
Obviously an error will occur if this variable definition differs in length from the number of samples to be drawn in the simulation.

Parameters can be altered in the same manner as endogenous variables (e.g. `sim(msim, p=c(x=3,'x<->x'=2))`), but they can also be fixed directly via the function `distribution`, e.g.

```r
> distribution(msim, ~x) <- binomial.lvm(p = 0.4)
```

defines $X$ to be simulated from a Bernoulli-process ($P(X = 1) = 0.4$), with similar options for the other methods (e.g. `poisson.lvm(lambda=2)`).

For a `lvmfit`-object (see Section 6) the `sim` method by default simulates observations with the parameter vector $p$ set equal to the estimated parameter vector and with the same number of observations as in the original data set. In this case the defaults causes the simulations to be drawn from the conditional distribution given the exogenous variables, hence the exogenous variables will be fixed at their original values (can be changed with the argument `xfix=FALSE`, in which case the covariates are simulated with mean and variance parameters set to the empirical mean and covariance).

### 6. Inference

| Function    | Task                              |
|-------------|-----------------------------------|
| `estimate`  | Estimates parameters of `lvm`-objects |
| `effects`   | Calculates direct and indirect effects |
| `compare`   | Likelihood ratio, Wald and Score tests |
| `gof`       | Goodness-of-fit measures |
| `logLik`    | Extracts individual Likelihood values |
| `information` | Extracts information matrix |
| `score`     | Extracts individual contribution to score |
| `bootstrap` | Non-parametric bootstrap |
| `confint`   | Calculates confidence limits (Wald and profile) |
| `constraints` | Extracts non-linear parameter constraints |
| `modelsearch` | Model searching via score tests |
| `equivalence` | Finds empirically equivalent models |

Table 4: Inferential tools.

Parameter estimation is achieved with the `estimate`-function which returns an object of class `lvmfit`. The syntax of the estimation procedure is:
estimate(x, data, estimator, control=list(),
  missing, weight, cluster, fix, ...)
is obtained with `lava.options(param="relative")` and loading parameters and intercepts are then interpreted as differences compared to the reference indicator. Setting `param="absolute"` will result in a parameterization where the variance of the latent variables are fixed to 1 and the intercept to 0 if not already fixed at some other values. `lava.options(param="hybrid")` alters the model such that the intercept of latent variables are set to 0, and one factor loading in each measurement model is set to 1.

Calling `lava.option(param="none")` has the same effect as `fix=FALSE` (in which case the user has to manually define parameter constraints that guarantee model identification).

The optional `control` argument must be a list of parameters for the optimization procedure. The element `method` should be a string pointing to a generic optimizer conforming to the syntax of `stats::nlminb` (the default optimizer), i.e. accepting the objective function (e.g. the log likelihood), the gradient, the Hessian and control parameters. Setting `method="nlminb0"` will only use the objective function during optimization, where `method="nlminb1"` also uses the gradient, and `method="nlminb2"` the Hessian as well. Defining `method="NR"` will use an alternative Newton-Raphson algorithm. Variance parameters are as default modeled using a log-link. This can be disabled by setting `constrain=FALSE`. Additional options like the number of iterations (`iter.max`), turning trace information on (`trace=1`), starter function (`starterfun` (a string pointing to a function generating starting values for the optimization), convergence tolerance (`tol`), reduction of step-size of the "NR" optimizer (`gamma`), etc. can also be defined (see the R help page of `nlminb` and `estimate`).

The control parameters can be set globally via the function `lava.options`. For instance to turn on the trace information of the optimizer as default in the current session, we would submit `lava.options(trace=1).

We demonstrate the procedure in the ongoing example (see Figure 2) with the data obtained from the simulation in Section 5 using the hybrid parameterization:

```
> lava.options(param = "hybrid")
> e <- estimate(m1, mydata)
> summary(e)
```
Latent variables: u1 u2
Number of rows in data=100

| Measurements: | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|---------------|----------|------------|---------|----------|--------|
| y1<-u1        | 1.00000  | 0.85423    |         |          |        |
| y2<-u1        | 0.85609  | 0.07774    | 11.01174| <1e-12   | 0.85902|
| y3<-u1        | 0.88990  | 0.08488    | 10.48415| <1e-12   | 0.83429|
| z1<-u2        | 1.00000  | 0.93812    |         |          |        |
| z2<-u2        | 1.09038  | 0.05153    | 21.16099| <1e-12   | 0.96411|
| z3<-u2        | 1.02444  | 0.04998    | 20.49764| <1e-12   | 0.95745|

| Regressions:  | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|---------------|----------|------------|---------|----------|--------|
| u1<-x1        | 1.11118  | 0.13232    | 8.39775 | <1e-12   | 0.61181|
| u1<-x2        | 0.96907  | 0.13392    | 7.23601 | <1e-12   | 0.51159|
| u2<-u1        | 0.94651  | 0.17037    | 5.55569 | 2.765e-08| 0.56706|
| u2<-x1        | 0.72832  | 0.21753    | 3.34815 | 0.0008135| 0.24025|
| u2<-x2        | 1.05500  | 0.20428    | 5.16434 | 2.413e-07| 0.33367|

| Intercepts:   | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|---------------|----------|------------|---------|----------|--------|
| u1            | 0.00000  | 0.00000    |         |          |        |
| u2            | 0.00000  | 0.00000    |         |          |        |
| y1            | 0.06805  | 0.15264    | 0.44582 | 0.6557   | 0.03110|
| y2            | 0.07461  | 0.12917    | 0.57759 | 0.5635   | 0.04005|
| y3            | -0.08326 | 0.14244    | -0.58453| 0.5589   | -0.04176|
| z1            | 0.07929  | 0.17709    | 0.44776 | 0.6543   | 0.02384|
| z2            | 0.07514  | 0.17400    | 0.43186 | 0.6658   | 0.02130|
| z3            | 0.07933  | 0.16811    | 0.47185 | 0.637    | 0.02376|

| Residual Variances: | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|---------------------|----------|------------|---------|----------|--------|
| u1                  | 1.02921  | 0.23949    | 4.29753 | 0.29462  |
| u2                  | 0.87832  | 0.23741    | 3.69960 | 0.09024  |
| y1                  | 1.29397  | 0.23560    | 5.49234 | 0.27029  |
| y2                  | 0.90931  | 0.16777    | 5.41987 | 0.26209  |
| y3                  | 1.20812  | 0.21027    | 5.74543 | 0.30396  |
| z1                  | 1.32627  | 0.23349    | 5.68022 | 0.11993  |
| z2                  | 0.87762  | 0.19360    | 4.53323 | 0.07050  |
| z3                  | 0.92797  | 0.18768    | 4.94448 | 0.08328  |

Estimator: gaussian

Number of observations = 100
Log-Likelihood = -1010.407
BIC = 2167.944
The parameter estimates in the output are divided into slope parameters belonging to the measurements part of the model (Measurements) (i.e. factor loadings), the structural part (Regressions) and the intercepts and residual (co)variances of the model. The summary method outputs all parameters of the model including the parameters that were fixed in contrast to the print method that only outputs the canonical parameters of the model. Notice that a regression parameter in each of the measurement models has been fixed to one in order to identify the model, and the slope parameter of $u_2$ on $u_1$ is therefore interpreted on the scale of $Y_{11}$ and $Y_{21}$. The standardized coefficients in the last column are interpreted as the change in standard deviation of the outcome when increasing the predictor one standard deviation. These parameters can be used to compare effects of predictor variables measured on different scales. If non-linear constraints were defined then the relevant estimates and approximate standard errors will be shown in the last part of the summary output. These effects can also be extracted with the constraints function.

The p-values for the variance parameters are deliberately omitted from the output as the asymptotic distribution under the null is non-standard (derived in special-cases such as the random intercept model as an equal mixture of a $\chi^2$-distribution and the Dirac measure in 0. Hence the naive p-values based on a $\chi^2$-distribution will tend to be conservative). Different versions of the information matrix can be used, via the argument type \{"E","hessian","outer","robust"\} (expected information (default), minus the second derivative of the log-likelihood, outer product of the score, and the robust/sandwich variant [White, 1982]), to calculate the asymptotic standard errors of the parameters (see also the information function).

The last part of the output includes some fit criteria (Akaike and Bayesian) and the omnibus goodness-of-fit $\chi^2$-test which is a likelihood ratio test of the current model against the saturated model structure. In the conditional model formulation, the least restrictive model allows covariance between residuals of all endogenous variables and the mean-vector $\mu$ to be a general linear combination of all the covariates. In the unconditional model formulation (15), this corresponds to a completely free mean structure and
a covariance $\Sigma$ that can be any symmetric positive definite matrix. Clearly
the maximum likelihood is attained at the sample mean and non-central
empirical covariance matrix, $S$. Hence the log-likelihood of the saturated
linear model is

$$-\frac{n}{2} \left( k \log(2\pi) + \log(|S|) + \frac{n-1}{n}(p + q) \right).$$

(49)

This part of the output can also be obtained with the call

> gof(e, chisq = TRUE)

Number of observations = 100
Log-Likelihood = -1010.407
BIC = 2167.944
AIC = 2066.814
log-Likelihood of model = -1010.407
log-Likelihood of saturated model = -1003.438
Chi-squared statistic: q = 13.93726 , df = 16 , P(Q>q) = 0.6033881
RMSEA (90% CI): 0 (0;0.08127)
rank(Information) = 23 (p=23)
condition(Information) = 0.007602741
||score||^2 = 1.306582e-08

Here the omnibus test gives a p-value of 0.60, thus indicating a reasonable
agreement between the model implied and empirical covariance structure.

Byproducts of the maximum likelihood estimation such as the score,
information and log-likelihood can be obtained with the functions score,
information (see also vcov), and logLik. The individual contribution to
score and log-likelihood are calculated with the argument indiv=TRUE. The
methods also allow altering the parameter, data, weights and type of esti-
mator (arguments p, data, weight and estimator). Predictions can be ob-
tained via the predict method (and conditional residuals via residuals),
where the latent variables are predicted by the conditional expectation given
all manifest variables (the prediction can be based on conditioning on a sub-
set of the manifest variables defined by the second argument of predict).
For instance to estimate $E(U_1 \mid Y_1, Y_2, Y_3, X_1, X_2)$ we can call

> u1hat <- predict(e, ~y1 + y2 + y3)[, "u1"]

Prediction of the residual terms can be obtained by setting the argument
residual=TRUE.

> zeta1hat <- predict(e, ~y1 + y2 + y3, residual = TRUE)[, "u1"]
Assessment of linearity and distributional assumptions can be based on examination of different residuals and their association with for example covariates in the model. We refer to the \texttt{gof} package (Holst, 2012) for residual based goodness-of-fit methods for structural equation models fitted with \texttt{lava}.

6.1. Direct and indirect effects

One of the strengths of the structural equation model framework is the possibility of decomposing the effects of a predictor into direct and indirect effects. In the model $m1$ (Figure 2) we have the following relations

\begin{align}
Z_3 &= \lambda_{23} U_2 + \epsilon_{23}, \\
U_2 &= \beta_1 X_1 + \beta_2 X_2 + \gamma U_1 + \xi_2, \\
U_1 &= \delta_1 X_1 + \delta_2 X_2 + \xi_1,
\end{align}

and we wish to quantify the effect of $X_1$ on $Z_3$. The direct effect is zero as $X_1$ is not present in (50). By substituting (51) and (52) into (50), it follows that

\begin{align}
Z_3 &= \beta_1 \lambda_{23} X_1 + \beta_2 \lambda_{23} X_2 + \beta_2 \lambda_{23} X_1 + \delta_1 \gamma \lambda_{23} X_1 \\
&\quad + \delta_2 \gamma \lambda_{23} X_2 + \gamma \lambda_{23} \xi_1 + \lambda_{23} \xi_2 + \epsilon_{23}.
\end{align}

Hence the total effect of $X_1$ is the sum of the two indirect effects $\beta_1 \lambda_{23}$ and $\delta_1 \gamma \lambda_{23}$. The estimation uncertainty of this effect can be approximated by the delta method. In general the distribution of a product of estimators can be approximated in the following way. Let

\begin{align}
f(\hat{\beta}) = f(\hat{\beta}_1, \ldots, \hat{\beta}_k) = \prod_{i=1}^k \hat{\beta}_i,
\end{align}

where the estimated parameters $\hat{\beta}$ are asymptotically normally distributed with covariance matrix $\Sigma_\beta$. Now

\begin{align}
\nabla f(\beta) = \begin{pmatrix}
\Pi_{i \neq 1} \beta_i \\
\vdots \\
\Pi_{i \neq k} \beta_i
\end{pmatrix}
\end{align}

and

\begin{align}
\sqrt{n} \left( f(\hat{\beta}) - f(\beta_0) \right) \xrightarrow{D} \mathcal{N}\left( 0, \nabla f(\hat{\beta})' \Sigma_\beta \nabla f(\hat{\beta}) \right)
\end{align}
The approximate distribution of linear combinations of products is obtained by straightforward calculations (i.e. $\nabla f_1 + \nabla f_2$).

The `effects` function can be used to estimate the (direct and indirect) effect of one variable on another together with approximate standard errors, e.g.

```r
> (f <- effects(e, z3 ~ x1))
```

**Total effect of 'x1' on 'z3':**

1.823569 (Approx. Std.Err = 0.1524713)

**Direct effect of 'x1' on 'z3':**

0 (Approx. Std.Err = NA)

**Indirect effects:**

Effect of 'x1' via x1->u1->u2->z3:

1.077448 (Approx. Std.Err = 0.2170743)

Effect of 'x1' via x1->u2->z3:

0.7461208 (Approx. Std.Err = 0.222262)

```r
> coef(f)
```

|        | Estimate | Std.Err | z value | Pr(>|z|) |
|--------|----------|---------|---------|----------|
| Total  | 1.823569 | 0.1524713 | 11.960083 | 0.000000e+00 |
| Direct | 0.000000 | NA      | NA      | NA       |
| z3<-u2<-u1<-x1 | 1.0774481 | 0.2170743 | 4.963458 | 6.923458e-07 |
| z3<-u2<-x1 | 0.7461208 | 0.2222620 | 3.356943 | 7.880941e-04 |

6.2. Hypothesis testing

Next we estimate the parameters of two nested models where we have restricted all factor loadings to 1 and in addition removed the effect from $U_1$ to $U_2$

```r
> m1a <- m1

> regression(m1a, c(z1, z2, z3) ~ u2) <- 1

> regression(m1a, c(y1, y2, y3) ~ u1) <- 1

> m1b <- m1a

> cancel(m1b) <- u2 ~ u1

> ea <- estimate(m1a, mydata)

> eb <- estimate(m1b, mydata)
```
6.2.1. Likelihood Ratio Test

For nested models, $M_1 \subseteq M_2$, the natural test is the \textit{likelihood ratio test} (LRT)

$$-2 \left[ \log L_1(\hat{\theta}_1) - \log L_2(\hat{\theta}_2) \right] \approx_{\text{approx.}} \chi^2_{\Delta d f}. \quad (58)$$

For non-nested models, one choice is the Akaike’s Information Criterion (AIC) favoring models with low values of

$$AIC = -2 \log(L) + 2n_{\text{par}}, \quad (59)$$

where $n_{\text{par}}$ is the number of parameters in the model (implemented in the \texttt{AIC} function), or the Bayesian Information Criterion

$$BIC = -2 \log(L) + n_{\text{par}} \log(N), \quad (60)$$

where $N$ denotes the total number of observations \cite{Raftery1993}, i.e. the number of endogenous variables times the number of individuals.

Successive LRT between nested models can be calculated with

\begin{verbatim}
> (LRT1 <- compare(e, ea, eb))

[[1]]

Likelihood ratio test
data: chisq = 6.6542, df = 4, p-value = 0.1553sample estimates:
log likelihood (model 1) log likelihood (model 2)
-1010.407 -1013.734

[[2]]

Likelihood ratio test
data: chisq = 33.9868, df = 1, p-value = 5.549e-09sample estimates:
log likelihood (model 1) log likelihood (model 2)
-1013.734 -1030.728
\end{verbatim}
Hence we accept the hypothesis that all factor loadings are equal (the model \texttt{m1a}) but reject the hypothesis that the two latent variables $U_1$ and $U_2$ are conditional independent given the covariates.

6.2.2. Wald Test

The \texttt{compare} method can also deal with hypothesis testing via Wald or Score tests. The hypothesis

\[ H_0: \beta = \beta_0 \]  

for a subset, $\beta$, of all the parameters, can be tested with a Wald test using the \texttt{par} and \texttt{null} arguments (the latter defaults to 0), for instance to test if all loading parameters are 1 (equivalent to the LRT of \texttt{m1} against \texttt{m1a}), we can write

\begin{verbatim}
> (W1 <- compare(e, par = c("y2<-u1", "y3<-u1", "z2<-u2", "z3<-u2"),
+    null = rep(1, 4)))

Wald test

data:  chisq = 7.2687, df = 4, p-value = 0.1223

For a general estimable contrast $C$, we can also test the hypothesis

\[ H_0: C\beta = \tilde{\beta}_0, \]  

where $C$ is matrix (or vector) with the same number of columns as the number of parameters, or alternatively a sub-matrix with column names given by parameter names (implicitly assuming that omitted columns are zero), leading to the test statistic

\[ (C\hat{\beta} - \tilde{\beta}_0)'(C\Sigma_\hat{\beta}C')^{-1}(C\hat{\beta} - \tilde{\beta}_0) \sim \chi^2_{\text{rank}(C)} \]  

The covariance matrix $\Sigma_\hat{\beta}$ will by default be the variance matrix as defined by the chosen estimator (for the linear gaussian models, \texttt{estimator="gaussian"}, this is the inverse of the expected information), but it can optionally be given as the argument \texttt{Sigma}, e.g. to use a robust variance estimate (see \texttt{information} method).

To test whether all the intercepts of the outcomes sum to zero, we can write

\begin{verbatim}

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\end{verbatim}
```r
> C <- rep(1, 6)
> names(C) <- endogenous(m1)
> (W2 <- compare(e, contrast = C))

Wald test

data:  
chisq = 0.1769, df = 1, p-value = 0.6741

hence we accept the hypothesis of equal intercepts.

6.2.3. Score Test

With the `scoretest` argument we can conduct Score tests. Letting \( \tilde{\theta} \) be the parameter belonging to the less restrictive model \( M_2 \), which is equal to \( \hat{\theta}_1 \), the MLE of the restrictive model \( M_1 \), for all the parameters shared by \( M_2 \) and zero elsewhere. The test statistic is then given by

\[
S = S_2(\tilde{\theta})'I_2^{-1}(\tilde{\theta})S_2(\tilde{\theta})
\]

with approximate \( \chi^2_{\Delta d f} \)-distribution under the null, where \( S_2 \) and \( I_2 \) are the score and information matrix of model \( M_2 \).

We will test whether adding correlation between the residuals terms of \( Z_3 \) and \( Z_2 \) significantly improves the model fit:

```r
> (S1 <- compare(e, scoretest = z3 ~ z2))

Score test

data: z3 ~ z2  
chisq = 0.1186, df = 1, p-value = 0.7306

which does not indicate evidence against the conditional independence assumption.

Similarly we can test the statistical significance of simultaneously adding two extra correlation parameters:

```r
> (S2 <- compare(e, scoretest = c(z3 ~ z2, z1 ~ z2)))

Score test

data: z3 ~ z2 z1 ~ z2  
chisq = 2.03, df = 2, p-value = 0.3624
```

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6.2.4. Model searching with the Score test

An advantage of the Score test over the LRT is that the MLE is only needed in the more restrictive model making it an ideal instrument for model searching, in order to check that important aspects of the covariance structure has not been neglected in the model specification. The \texttt{modelsearch} function can be used to examine all one-parameter extensions of the model. The following call give the 5 most significant one-parameter extensions

\begin{verbatim}
> print(ms <- modelsearch(e), tail = 5)
\end{verbatim}

\begin{verbatim}
Score: S  P(S>s)  Index  holm  BH
2.024  0.1549  u2<->y1  1  0.9031
2.124  0.145  y1<->z1  1  0.9031
2.283  0.1308  y1<->y2  1  0.9031
2.892  0.08904  y2<->z2  1  0.9031
4.207  0.04025  y2<->z1  1  0.9031
\end{verbatim}

As expected we do not see any significant improvements of the model among the 5 most significant Score tests (with the first and second column being the test statistic and corresponding p-value, and the last two columns being the p-values adjusted by the Bonferroni-Holm procedure to control the overall Type I error (Holm, 1979), and q-values of the Benjamini-Hochberg procedure controlling the FDR). Similarly, the most important $k$-parameter extensions to the model, can be examined with the argument \texttt{k}, but the number of models to search through will increase dramatically with \texttt{k}.

6.3. Model equivalence

A challenge in multivariate modeling is the problem of equivalent models, where two different parameterizations leads to identical model fit (likelihood) for all data sets. Hence without strong a priori knowledge of the model structure, e.g. based on other scientific evidence, the interpretation of model parameters must be made cautiously. Formal proofs of model equivalence can be difficult (Bollen, 1989). To identify candidates of equivalent models we suggest using the Score test. The idea is to study all one-parameter extensions of a given model using the score test. Two models are said to be empirically equivalent if the score tests agree. This can be achieved with the \texttt{equivalence} function. Two variables of the model are chosen, which not necessarily are defined as being directly related in the model structure. The score function for the model including covariance between the residuals of the two selected variables is then compared with score functions of models omitting this association, but with the same number of parameters.
As a simple example we will investigate the structural equation model in the path diagram of the left panel of Figure 7.

```r
> mEq <- lvm(list(c(y1, y2, y3) ~ u, u ~ x))
> latent(mEq) <- u
> covariance(mEq) <- y1 ~ y2
> dEq <- sim(mEq, 100)
> est.mEq <- estimate(mEq, dEq)
```

Below we are examining whether the inclusion of a residual correlation between \( Y_1 \) and \( Y_2 \) has any equivalent formulations.

```r
> (Eq <- equivalence(est.mEq, y1 ~ y2))
```

In fact, an equivalent model is defined by instead adding a direct effect of \( X \) on \( Y_3 \) (see Figure 7).

| Empirical equivalent models: |
|-----------------------------|
| 0) \( y_1 <\sim y_2 \) (10.31) |

Candidates for model improvement:

| none |

![Figure 7: Example of two equivalent models (mEq).](image)

6.4. Confidence limits

Wald confidence limits can be created using the method `confint`. However, for some parameters better coverage can be achieved with alternative methods. One method is the non-parametric bootstrap which can be calculated with the function `bootstrap`. The bootstrap is a computational intensive method, and parallel computation can be done by registering a `foreach` [REvolution Computing, 2009] parallel adaptor. In this example we will compute the bootstrap in parallel using the `parallel` and `doParallel` packages distributing the bootstrap computations across the available CPU cores.
> library(doParallel)
> registerDoParallel()
> (B <- bootstrap(e, 500))

Non-parametric bootstrap statistics (R=500):

|        | Estimate | Bias  | Std.Err | 2.5 %  | 97.5 % |
|--------|----------|-------|---------|--------|--------|
| y1     | 0.08467677 | 1.662669e-02 | 0.15922250 | -0.2417240 | 0.3787349 |
| y2     | 0.07691911 | 2.312410e-03 | 0.12146566 | -0.1457868 | 0.3101638 |
| y3     | -0.07196788 | 1.129167e-02 | 0.13941055 | -0.3536971 | 0.2112750 |
| z1     | 0.08407469 | 4.780799e-03 | 0.18185690 | -0.2799668 | 0.4421753 |
| z2     | 0.08527725 | 1.013330e-02 | 0.17283489 | -0.2966441 | 0.3995207 |
| z3     | 0.09493044 | 1.560519e-02 | 0.2081544 | 0.4407723 |
| u1<-x1 | 1.11860351 | 7.421090e-03 | 0.15172036 | 0.8141602 | 1.4084714 |
| u1<-x2 | 0.96908130 | 1.303164e-04 | 0.15096572 | 0.6741721 | 1.2717508 |
| u2<-u1 | 0.96947448 | 2.296893e-02 | 0.16712619 | 0.6754198 | 1.3478912 |
| u2<-x1 | 0.69630433 | -3.201448e-02 | 0.1494670 | 1.0941781 |
| u2<-x2 | 1.02390307 | 3.109418e-02 | 0.20826885 | 0.5773626 | 1.4170756 |
| y2<-u1 | 0.85971870 | 3.632407e-03 | 0.10370758 | 0.6779335 | 1.0642511 |
| y3<-u1 | 0.89440961 | 4.508961e-04 | 0.08414864 | 0.7388379 | 1.0566335 |
| z2<-u2 | 1.09661468 | 6.232894e-03 | 0.05341720 | 0.9971030 | 1.2131320 |
| z3<-u2 | 1.02856111 | 4.118107e-03 | 0.04733460 | 0.9419200 | 1.1232677 |
| u1<-u1 | 0.99249288 | -3.671648e-02 | 0.27738180 | 0.5390826 | 1.5590308 |
| u2<-u2 | 0.81896688 | -5.935355e-02 | 0.26481969 | 0.3523745 | 1.3589482 |
| y1<-y1 | 1.28947917 | -4.494201e-02 | 0.26570998 | 0.8302451 | 1.8369823 |
| y2<-y2 | 0.89753510 | -1.177998e-02 | 0.17311848 | 0.5861767 | 1.2511752 |
| y3<-y3 | 1.18530385 | -2.281194e-02 | 0.18883990 | 0.8092845 | 1.5525326 |
| z1<-z1 | 1.30673170 | -1.953404e-02 | 0.21030950 | 0.9245536 | 1.7315556 |
| z2<-z2 | 0.84766364 | -2.995799e-02 | 0.17431201 | 0.4971759 | 1.1894273 |
| z3<-z3 | 0.92206325 | -5.906032e-02 | 0.18632891 | 0.5572348 | 1.3095068 |

To bootstrap other statistics a user-defined function can be supplied as the argument `fun`. A parametric bootstrap can be computed setting the argument `parametric=TRUE` and setting the argument `p` to parameter values of the null model from which to simulate from. The parallel computation functionality can be disabled via the call `lava.options(parallel=FALSE)`.

As an alternative to the resample-based approach, we can also calculate the confidence limits based on the profile likelihood:

```r
> (ci <- confint(e, profile = TRUE, parm = "u2<-u1", level = 0.95))

2.5 % 97.5 %
u2<-u1 0.6392854 1.332559
```
where the parameter of interest can be given as the index or label name.

A third option is to use a variance stabilizing transform of the parameter. As an example we will calculate the confidence limits of the partial correlation (or conditional correlation) between two outcomes $Y_1$ and $Y_2$ given covariates $X$. Hence we assume that

$$Y_i = \beta'_i X + \varepsilon_i, \quad i = 1, 2$$

and aim to estimate the correlation, $\rho$, between $\varepsilon_1$ and $\varepsilon_2$.

We define this model with a single covariate and simulate some observations and find the corresponding MLE:

```r
> m <- lvm(c(y1, y2) ~ x)
> covariance(m, y1 ~ y2) <- "C"
> covariance(m, ~y1 + y2) <- list("v1", "v2")
> d <- sim(m, 100)
> e.pcor <- estimate(m, d)
```

Note with the default parameter values the correlation between $\varepsilon_1$ and $\varepsilon_2$ is 0.5. Next we define the correlation parameter using a non-linear parameter constraint and obtain the estimate with confidence limits based on the delta method.

```r
> constrain(e.pcor, rho ~ C + v1 + v2) <- function(x) x[1]/(x[2] * x[3])^0.5
> constraints(e.pcor)
```

```
Estimate Std. Error  Z value Pr(>|z|)     2.5%    97.5%
  rho 0.4449733 0.08019988  5.548303 2.884553e-08 0.2877844 0.6021621
```

Near the boundary of the parameter space these limits will tend to perform poorly and a better approach is to apply the variance stabilizing arctanh transform (Fisher's z-transform):

$$Z: \rho \rightarrow \frac{1}{2} \log \left( \frac{1 + \rho}{1 - \rho} \right)$$

Here we also supply the analytical gradient (optional) calculated with the chain-rule and in addition we set the attribute inv defining the inverse transformation, thus giving us the confidence limits on original correlation scale:
> constrain(e.pcor, z ~ C + v1 + v2) <- function(x) {
+   f <- function(p) p[1]/sqrt(p[2] * p[3])
+   res <- atanh(f(x))
+   df <- function(p) c(1/sqrt(p[2] * p[3]), -f(p)/(2 * p[2]),
+                     -f(p)/(2 * p[3]))
+   datanh <- function(r) 1/(1 - r^2)
+   attributes(res)$grad <- function(p) datanh(f(p)) * df(p)
+   attributes(res)$inv <- tanh
+   return(res)
+ }
> constraints(e.pcor)

| Estimate | Std. Error | Z value | Pr(>|z|) | 2.5% | 97.5% |
|----------|------------|---------|----------|------|-------|
| rho      | 0.4449733  | 0.08019988| 5.548303 | 2.884553e-08 | 0.2877844 | 0.6021621 |
| z        | 0.4784149  | 0.10000000| 4.784149 | 1.717133e-06 | 0.2824185 | 0.6744113 |
| inv(z)   | 0.4449733  | NA      | NA       | 0.2751420 | 0.5878741 |

In fact \( \sqrt{n}Z(\hat{\rho}_n) \xrightarrow{D} \mathcal{N}(0,1) \) as \( n \to \infty \) (Lehmann and Romano, 2005). Note that the correlation method calculates the correlation coefficients of a lvmfit object in a more elegant way and with a slightly more precise variance estimate (Hotelling, 1953) (see also the partialcor function).

7. Multigroup models

Multigroup analysis (31) can be used to combine different models linked via some shared parameters. Among other things this extension can be useful in testing general hypotheses of linear interactions, and the lava package supports this generalization via the estimate-function taking a list of lvm-objects and a list of data.frame's as arguments and returning an object of class multigroupfit:

> estimate(list(m1, m2, m3, ...), list(d1, d2, d3, ...), ...)

The list of lvm objects can optionally be named, as in the example below, to enhance the output. Parameters that are shared across the models m1,m2,m3,... will be also be shared in the multigroup analysis, whereas all other parameters will be estimated independently between the groups. In many applications the first argument will therefore be repetitions of the same lvm-object. Note, that when the different datasets are defined from a single data.frame using a grouping variable, the function split can be applied to define the second argument. A typical multigroup analysis call will therefore resemble
where the data-frame d here is split into a list defined from the variable d$x (with, in this case, n distinct values).

As an example we will create two nearly identical lvm-objects describing simple factor models (see Figure 8):

```r
> mg1 <- lvm()
> regression(mg1, ~Y1 ~ H) <- 1
> intercept(mg1, ~Y1) <- 0
> regression(mg1) <- ~c(Y2, Y3) ~ H
> regression(mg1) <- ~H ~ E
> latent(mg1) <- ~H
> mg1 <- baptize(mg1)
> covariance(mg1, endogenous(mg1)) <- NA
> mg2 <- mg1
> intercept(mg2, ~Y2 + Y3) <- 0
```

The `baptize` function labels all free parameters of the model, giving the parameter the names as defined by the `coef` function ("Y1<-H", "H<-H" etc.). An optional argument `labels` can be given to define custom labels.

In the above example the restrictions of the variances of the residuals of the endogenous variables are removed, and hence the two models mg1 and mg2 share all parameters except for these variance parameters, and the intercepts which are identical in mg2.

Next we simulate two datasets from model 1 (thus in fact only a single group):

```r
> data1 <- sim(mg1, 200)[, manifest(mg1)]
> data2 <- sim(mg1, 200)[, manifest(mg1)]
```

To estimate parameters using MLE we simply type

```r
> (e.mg <- estimate(list("Arm 1" = mg1, "Arm 2" = mg2), list(data1, + data2)))
```

```
Group 1: Arm 1 (n=200)

Measurements:          Estimate Std. Error    Z value  Pr(>|z|)
                      Y2<-H   0.99827   0.06189  16.13044 <1e-12
                      Y3<-H   1.04265   0.06527  15.97469 <1e-12
```

45
Comparisons of multigroup structures can be conducted using a LRT. As an example we fit the single group LLVM and perform a LRT to test whether the residual variances are the same in both groups and the intercepts are zero.

```r
> e0 <- estimate(mg2, rbind(data1, data2))

> compare(e0, e.mg)
```

```
Likelihood ratio test

data:  chisq = 2.3744, df = 5, p-value = 0.7953
sample estimates:
log likelihood (model 1) log likelihood (model 2)
-2001.323 -2000.136
```
8. Data with missing values

Missing data are common in studies with multivariate outcomes and complete case analyses can in these settings become quite inefficient and are further only consistent when data are missing completely at random (MCAR).

Under the more general assumption that data are missing at random (MAR), i.e. that the missing data mechanism depends only on the observed variables (Little and Rubin, 2002), then inference can be based on the marginal likelihood, where the missing values has been integrated out

\[ f(y_{obs}; \theta) = \int f(y_{obs}, y_{mis}; \theta) \, dy_{mis}. \]  

(67)

Here \( f(y_{obs}, y_{mis}; \theta) \) is the full likelihood of both the observed data \( (y_{obs}) \) and the missing part \( (y_{mis}) \), parameterized by \( \theta \).

In lava, MLE under the MAR assumption can be obtained by adding the `missing=TRUE` argument to `estimate` (both for `lvm` and multigroup objects) using the multigroup framework on the different missing patterns of the data.

To demonstrate this, we will imitate a MCAR missing data mechanism on the first of the datasets simulated from `mg1` (see Section 7), with a massive 30% probability of missingness on each outcome

```r
> d0 <- makemissing(data1, p = 0.3, cols = endogenous(mg1))
```

and the full-information maximum likelihood estimates can then be achieved with the call:

```r
> e.mis <- estimate(mg1, d0, missing = TRUE)
> summary(e.mis, std = NULL, labels = FALSE)
```

Latent variables: H

Number of rows in data=199 (73 complete cases, 7 groups)

|                  | Estimate | Std. Error | Z value | Pr(>|z|) |
|------------------|----------|------------|---------|----------|
| Measurements:    |          |            |         |          |
| Y1<-H            | 1.00000  |            |         |          |
| Y2<-H            | 0.90265  | 0.10861    | 8.31067 | <1e-12   |
| Y3<-H            | 0.95524  | 0.11812    | 8.08673 | <1e-12   |
| Regressions:     |          |            |         |          |
| H<-E             | 0.83840  | 0.11018    | 7.60938 | <1e-12   |
|                  |          |            |         |          |
Intercepts:
  Y1  0.00000
  H  -0.14287  0.11042 -1.29388  0.1957
  Y2  0.05932  0.11132  0.53288  0.5941
  Y3  -0.01611  0.12958 -0.12433  0.9011
Residual Variances:
  Y1  0.68217  0.17881  3.81499
  H  1.20761  0.21942  5.50375
  Y2  1.02999  0.19284  5.34120
  Y3  1.37374  0.23948  5.73625

Estimator: gaussian

Log-Likelihood = -730.9461
BIC = 1525.811
AIC = 1481.892

with standard errors based on the observed information (Kenward and Molenberghs, 1998).

9. Beyond the standard linear Gaussian case

While linear Gaussian models cover many useful situations there are clearly cases where these models are no longer adequate. In this section we will briefly describe extensions of lava that covers some of these cases.

9.1. Clustered correlated data

Models with very complex hierarchical structures can be estimated in lava. However, the full specification of such a model can be challenging and perhaps more importantly, as the lowest level in such a model is often not of primary interest, it can be more natural to relax the model assumptions for this part of the model.

As a hypothetical example we can imagine that the aim of a study is to estimate the association between noise levels and health. In practice this is done by measuring the average noise level, $E_i$, in different neighborhoods. We assume that the health status is measured indirectly for each subject by three proxy measures, $Y_{1i}$, $Y_{2i}$, $Y_{3i}$ (e.g. blood pressure and cholesterol levels), and that the within-subject correlation between these measurements can be described by a single latent variable, $H_i$ (see Figure 8). The effect of
noise on health is quantified as the linear association between $H_i$ and $E$. The study is complicated by the fact that measurements within neighborhoods are correlated beyond what $H_i$ is capturing (air pollution, crime levels, traffic and other factors that could affect stress levels) and disregarding this within-cluster correlation will generally lead to too optimistic standard errors of the noise effect.

Inference can instead be based on the i.i.d. decomposition of the score leading to a sandwich estimator (GEE-type) of the variance (Williams, 2000)

$$
\left( \frac{\partial S(\theta)}{\partial \theta} \right)^{-1} \left( \frac{K}{K-1} \sum_{c=1}^{K} S_{(c)}(\theta)^{\otimes 2} \right) \left( \frac{\partial S(\theta)}{\partial \theta} \right)^{-1}, \tag{68}
$$

where $S$ is the total score and $S_{(c)}$ is the sum of the scores within cluster $c$, and $K$ denotes the total number of clusters. Simulation studies (Yan and Fine, 2004; Paik, 1988) indicate that the sandwich estimator works well with $K > 50$.

With 5 individuals sampled from each cluster/neighborhood, the above model could specified with

```r
> K <- 5
> mclust1 <- lvm()
> for (i in 1:K) {
+   xy <- c("Y1", "Y2", "Y3") # i
+   h <- "H" # i
+   regression(mclust1, to = c(xyz), from = h) <- list(1, "l1",
+                                                   "l2")
+   regression(mclust1, to = h, from = c("U", "E")) <- list(1,
+                                                        "b")
+   intercept(mclust1, c(xyz)) <- list("mx", "my", "mz")
+   covariance(mclust1, c(xyz, h)) <- list("vx", "vy", "vz",
+                                          "v")
+ }
> latent(mclust1) <- c("H" # all 1:K, "U")
> intercept(mclust1, latent(mclust1)) <- 0
```

We simulate data from 250 clusters and obtain the MLE

```r
> dclust <- sim(mclust1, 250, p = c(b = 0.3))[, manifest(mclust1)]
> eclust <- estimate(mclust1, dclust)
```

and we wish to compare this with the marginal model (see Figure 8):
The marginal estimates with robust standard errors are obtained easily by giving the name of the column in the data.frame that specifies the clusters (or an actual vector) as argument to the estimate function.

```r
> eclust1 <- estimate(mclust, data = dclustWide, cluster = "id")
```

In this example we see a substantial under-estimation of the standard errors of the pollution effect estimate in the naïve approach (with covariates varying within clusters the bias could go in the opposite direction as well). In contrast, the results of the marginal approach is close to those the full model.

```r
> res <- rbind(coef(eclust, 2)["b", ], coef(eclust1, 2)["H<-E", + ], coef(eclust0, 2)["H<-E", ])
> rownames(res) <- c("full MLE", "Marg.robust", "Marg.naive")
> res

|             | Estimate  | Std. Error | Z value | Pr(>|z|) |
|-------------|-----------|------------|---------|----------|
| full MLE    | 0.3683574 | 0.07816049 | 4.712835| 2.442944e-06 |
| Marg.robust | 0.3702012 | 0.07658741 | 4.833708| 1.340131e-06 |
| Marg.naive  | 0.3702012 | 0.04843684 | 7.642967| 2.131628e-14 |
```
Typically the loss of power in this marginal approach is modest, and is countered by circumventing the need for explicit (mis)specification of the distribution of the cluster random effect.

9.2. Mixture models

The normal distribution is a central assumption in (3) and (4), and one way to relax this assumption while still avoiding the need for computational intense numerical approximations of the likelihood function of the observed data, is to allow mixtures of normal distributions in the model. Applications include pattern recognition and cluster analysis (machine learning), outlier analysis and modeling of heterogeneity, e.g. adjusting for unknown subpopulations in a sample.

In general we will allow models to be described by the convex combination

\[
f_\theta(y \mid z) = \sum_{j=1}^{K} \pi_j f_{\theta_j}^{(j)}(y \mid z), \quad \sum_{j=1}^{K} \pi_j = 1, \quad \pi_j \in ]0, 1]\]

where \( f_{j, \theta_j} \) is a probability density of a LLVM with responses \( y \) and covariates \( z = (x', v', w')' \), and \( \theta \) is the parameter-vector \( (\cup_j \theta_j, \pi_1, \ldots, \pi_{K-1}) \), noting that the \( \theta_j \)'s need not to be disjoint. We denote the number classes \( K \).

The likelihood for the mixture model is therefore

\[
L(\theta \mid y, z) = \prod_{i=1}^{n} \sum_{j=1}^{K} \pi_j f_{\theta_j}^{(j)}(y_i \mid z_i)
\]

To solve the corresponding score equation, the EM algorithm is typically applied (Dempster et al., 1977). We introduce latent indicator variables \( \xi_{ij} \), \( i = 1, \ldots, n \), describing the class membership of the observation \( (y_i, z_i) \).

\[
\xi_{ij} = 1 \{ y_{ij} \text{ belongs to class } j \}
\]

and hence \( \mathbb{E}\xi_{ij} = \mathbb{P}(y_{ij} \text{ belongs to class } j) = \pi_j \). We can then treat the mixture analysis as a missing data problem, \( v = (y, z, \xi) \), and complete-data log-likelihood:

\[
\log L_C(\theta \mid v) = \sum_{i=1}^{n} \sum_{j=1}^{K} \xi_{ij} \log \left( f_{\theta_j}^{(j)}(y_i \mid z_i) \right)
\]
While the EM-algorithm is generally slower than Newton-Raphson (sub-linear vs. quadratic convergence), this disadvantage is compensated by the fact that the EM-algorithm tends to be less sensitive to choice of starting values as the algorithm guarantees a non-decreasing likelihood in each iteration. In contrast, NR can behave erratically for poor choices of starting values. To address possible problems with convergence to a local maximum, it is still advisable to start the algorithm at several different starting points in the parameter space. The EM algorithm also implicitly constraints the probability vector to the correct parameter space, as the E-step at iteration \( l \) leads to a simple expression of the posterior class probabilities

\[
\hat{\pi}^{(l)}_{ji} = \frac{\hat{\pi}^{(l)}_{j} f_{j,\hat{\theta}^{(l)}}(y_i | z_i)}{\sum_{j=1}^{K} \hat{\pi}^{(l)}_{j} f_{j,\hat{\theta}^{(l)}}(y_i | z_i)}
\] (73)

In the M-step we obtain the new class probabilities

\[
\hat{\pi}^{(l+1)}_{j} = \frac{1}{n} \sum_{i=1}^{n} \hat{\pi}^{(l)}_{ji}
\] (74)

and \( \hat{\theta}^{(l+1)} \) is derived by solving

\[
\arg \max_{\theta} Q(\theta; \hat{\theta}^{(l)}) = \sum_{i=1}^{n} \sum_{j=1}^{K} \hat{\pi}^{(l)}_{ji} \log (f_{j,\theta}(y_i | z_i))
\] (75)

which for a general LLVM mixture is optimized iteratively (NR pr. default).

In principle a model relating class probabilities to covariates could also be included in this setup leading to a M-step where we instead of the simple expression (73) would have to maximize a weighted multinomial logit model.

To analyze a mixture model in lava the plugin package lava.mixture needs to be loaded. The function mixture fits the mixture model to a list of lvm objects implicitly defining the number of mixture components of the model. Via the control argument the parameters of the EM algorithm can be adjusted, such as the starting values (start), number of random starting points (nstart), convergence tolerance (tol, change in log-likelihood), etc.

```r
> library(lava.mixture)
> mixture(list(m1,m2,...),data=mydata,control,...)
```

Instead of a list, a single lvm object can be given as argument with the argument k specifying the number of mixture components.
As an example we will simulate data from a simple model (see Figure 9), where we have a single dichotomous unmeasured confounder $z$ ($P(z = 1) = 0.5$), which have a direct linear effect on the outcome of interest $Y$

$$Y = \mu_Y + \beta X + \gamma_Y Z + \epsilon_Y$$  \hspace{1cm} (76)

and on the predictor $X$, which we assume is conditionally normally distributed given $Z$

$$X = \mu_X + \gamma_X Z + \epsilon_X$$  \hspace{1cm} (77)

In our simulation we will let all intercepts be zero, residual variances 1, and slopes as defined by Figure 9:

```r
> mix0 <- lvm(list(Y ~ X + Z, X ~ Z))
> distribution(mix0, ~Z) <- binomial.lvm()
> d0 <- sim(mix0, p = c(Y>Z = 2), n = 500)
```

Next, we will define the mixture regression that takes into account that we have unobserved heterogeneity caused by $Z$. The base model is the simple linear regression model of $Y$ given $X$, however, with a covariance call we define $X$ to be endogenous and let all parameters except for the intercepts of $Y$ and $X$ be fixed between the classes

```r
> mix1 <- lvm(Y ~ X)
> covariance(mix1, ~X) <- "v"
> mix1 <- baptize(mix1)
> intercept(mix1, ~Y + X) <- NA
```

Figure 9: Model mix0 with unmeasured confounder $z$. 

Next, we will define the mixture regression that takes into account that we have unobserved heterogeneity caused by $Z$. The base model is the simple linear regression model of $Y$ given $X$, however, with a covariance call we define $X$ to be endogenous and let all parameters except for the intercepts of $Y$ and $X$ be fixed between the classes

```r
> mix1 <- lvm(Y ~ X)
> covariance(mix1, ~X) <- "v"
> mix1 <- baptize(mix1)
> intercept(mix1, ~Y + X) <- NA
```
defining the model

\[ Y = \mu_{y,c} + \beta X + \epsilon, \quad (78) \]

\( \left( \begin{array}{c} \epsilon \\ X \end{array} \right) \sim \mathcal{N} \left( \left( \begin{array}{c} 0 \\ \mu_{x,c} \end{array} \right), \left( \begin{array}{cc} \sigma_y^2 & 0 \\ 0 & \sigma_x^2 \end{array} \right) \right) \quad (79) \)

with the index \( c \) denoting the class. The model with two classes is fitted with the call

\[
M <- \text{mixture}(\text{mix1, d0, k = 2})
\]

\[
(s <- \text{summary}(M))
\]

Cluster 1 (n=231, Prior=0.4687):

| Estimate  | Std. Error | Z value | Pr(>|z|) |
|-----------|------------|---------|----------|
| Y<-X      | 0.95114    | 0.07881 | 12.06853 | <1e-12   |
| Intercepts: |           |         |          |          |
| Y         | 2.11558    | 0.15240 | 13.88137 | <1e-12   |
| X         | 1.11484    | 0.11280 | 9.88306  | <1e-12   |
| Residual Variances: |       |         |          |          |
| Y         | 1.14553    | 0.12566 | 9.11583  |          |
| X         | 1.05875    | 0.11837 | 8.94435  |          |

Cluster 2 (n=269, Prior=0.5313):

| Estimate  | Std. Error | Z value | Pr(>|z|) |
|-----------|------------|---------|----------|
| Y<-X      | 0.95114    | 0.07881 | 12.06853 | <1e-12   |
| Intercepts: |           |         |          |          |
| Y         | 0.15417    | 0.11353 | 1.35802  | 0.1745   |
| X         | -0.10255   | 0.10257 | -0.99982 | 0.3174   |
| Residual Variances: |       |         |          |          |
| Y         | 1.14553    | 0.12566 | 9.11583  |          |
| X         | 1.05875    | 0.11837 | 8.94435  |          |

AIC= 3325.163
||score||^2= 0.0001193449

We see that the mixture model gives a regression coefficient of 0.951 which is close to the true value. This should be compared to the biased OLS estimate of \( \beta \):
In fact for the given set of parameters the bias is

\[
\text{bias}(\hat{\beta}_{\text{OLS}}) = \beta - \frac{(\beta \gamma_X^2 + \gamma_Y \gamma_X) \text{Var}(Z) + \beta \text{Var}(\epsilon_X)}{\gamma_X^2 \text{Var}(Z) + \text{Var}(\epsilon_X)} = \frac{2}{5}
\]  

(80)

and replicating the above simulation 1000 times we obtain the following statistics for the \( \beta \) parameter

| Estimator  | Bias  | Std.Err | MSE  |
|------------|-------|---------|------|
| OLS        | 0.403 | 0.0510  | 0.165|
| Mixture    | 0.002 | 0.0822  | 0.068|

which clearly shows the advantage of the mixture regression model.

The example generalizes to several binary confounders (\( 2^k \) classes with \( k \) confounders, and the continuous case could likewise be approximated by a finite number of mixtures) and if we were suspecting an interaction effect between \( X \) and \( Z \) we could also have allowed the slope parameter (\( \beta \)) to vary freely in the two classes. The difficult task of choosing an optimal number of components in the mixture could be based on cross-validation, but currently this is not implemented in \texttt{lava.mixture} (nor is the problem of adjusting the standard errors for this model selection which could be based on a bootstrap resampling). Still we believe that a mixture analysis as in the previous example could serve as an important sensitivity analysis in many applications.

Two variants of the EM-algorithm are also implemented in the \texttt{mixture} function (via the argument \texttt{type}): CEM (Classification EM) where each observation in the E-step is assigned to the class with the highest maximum posterior probability (73), and StEM (Stochastic EM) where each observation is assigned randomly to a class based on a draw from a multinomial distribution with the posterior probabilities as parameter. In both cases, the M-step reduces to the maximization of a simple multigroup LLVM (see Section [7]). The latter approach leads to a time-homogeneous Markov chain of the parameters, which under certain regularity conditions is ergodic with a normal stationary distribution with a mean that is a consistent estimate of the mixture parameters (Celeux and Diebolt, 1986). In some
In models with an unstructured mean and covariance in each class, the function `mvnmix` can be used because it exploits the fact, that the M-step has a closed-form solution (in fact the likelihood is unbounded, however, this is of more technical than practical interest as we typically can disregard the obviously wrong solutions to the score equations). As an example we will fit a two-component Gaussian mixture model to the waiting times between eruptions and the durations of the eruptions for the Old Faithful geyser in Yellowstone National Park:

```r
> data(faithful)
> mixff <- mvnmix(faithful, k = 2)
> (s <- summary(mixff))
```

Cluster 1 (n=97, Prior=0.3559):

| Estimate  | Std. Error | Z value | Pr(>|z|) |
|-----------|------------|---------|----------|
| Intercepts: |            |        |          |
| eruptions  | 2.03639    | 0.03495 | 58.27283 | <1e-12   |
| waiting    | 54.47852   | 0.62846 | 86.68510 | <1e-12   |
| Residual Variances: |        |        |          |
| eruptions  | 0.06917    | 0.01074 | 6.43961  |
| eruptions,waiting | 0.43517 | 0.17984 | 2.41979  | 0.01553  |
| waiting    | 33.69728   | 5.77754 | 5.83246  |

Cluster 2 (n=175, Prior=0.6441):

| Estimate  | Std. Error | Z value | Pr(>|z|) |
|-----------|------------|---------|----------|
| Intercepts: |            |        |          |
| eruptions  | 4.28966    | 0.03349 | 128.06900| <1e-12   |
| waiting    | 79.96812   | 0.47131 | 169.67075| <1e-12   |
| Residual Variances: |        |        |          |
| eruptions  | 0.16997    | 0.02112 | 8.04628  |
| eruptions,waiting | 0.94061  | 0.18679 | 5.03562  | 4.763e-07|
| waiting    | 36.04621   | 4.09090 | 8.81132  |
9.3. Binary data

In epidemiology binary data are among the most common types of endpoints and often correlated binary data are collected. Several methods have been proposed to deal with this sort of data, e.g. marginal models [Liang and Zeger, 1986], conditional maximum likelihood estimation [Andersen, 1971] or numerical integration to obtain the marginal likelihood of the observed data [Pinheiro and Chao, 2006]. Via the package lava.tobit [Holst, 2011] we can estimate LLVMs where a subset of the endogenous variables are binary using a probit-link (only subtle differences with a logit-link resulting in a scaling of roughly 1.7 of the parameters).

As an example we will look at a simple factor analysis model with a covariate

$$P(Y_{ij} = 1 | \eta_i, X_i) = \Phi (\mu_j + \lambda_j \eta_i),$$

$$\eta_i = \gamma X_i + \zeta_i,$$

where $\zeta_i \sim N(0, \sigma^2)$, and $\Phi$ is the standard normal cumulative distribution function.

There are several ways to specify this model in lava, but here we will use the binary function:
> mprobit <- lvm(list(c(y1, y2, y3) ~ eta, eta ~ x))
> latent(mprobit) <- ~eta
> binary(mprobit) <- endogenous(mprobit)
> set.seed(1)
> dprobit <- sim(mprobit, 500)

With the binary call the endogenous variables, \(Y_{ij}\), are changed from being continuous to being dichotomous, where we assume that there exists a latent conditionally normally distributed variable, \(Y^*_{ij}\), such that

\[
Y_{ij} = \begin{cases} 
1, & Y^*_{ij} > 0 \\
0, & Y^*_{ij} \leq 0. 
\end{cases}
\]  

(83)

The MLE is obtained as usual

> lava.options(param = "hybrid", trace = 1, method = "NR")
> (eprobit <- estimate(mprobit, dprobit))

| Measurements:        | Estimate | Std. Error | Z value | Pr(>|z|) |
|----------------------|----------|------------|---------|----------|
| y2<~eta              | 0.78017  | 0.15378    | 5.07336 | 3.909e-07|
| y3<~eta              | 0.92687  | 0.18950    | 4.89114 | 1.003e-06|
| Regressions:         |          |            |         |          |
| eta<-x               | 1.09228  | 0.15999    | 6.82737 | 8.649e-12|
| Intercepts:          |          |            |         |          |
| y1                   | 0.12358  | 0.09374    | 1.31829 | 0.1874   |
| y2                   | -0.06024 | 0.08104    | -0.74330| 0.4573   |
| y3                   | 0.04414  | 0.08930    | 0.49432 | 0.6211   |
| Residual Variances:  |          |            |         |          |
| eta                  | 1.32052  | 0.40634    | 3.24979 |          |

Another interesting example is the logit-probit-normal model (Caffo and Griswold, 2006) which is a conditional model with marginal fixed effects on a logit scale. E.g. a random intercept model with a single covariate \(X_{ij}\) can be formulated as

\[
P(Y_{ij} = 1 \mid \eta_i, X_{ij}) = \Phi \left( \Phi^{-1} \left( \frac{1}{1 + \exp(-\mu - \beta X_{ij})} \right) \sqrt{1 + \sigma^2 + \eta_i} \right),
\]

(84)

where \(\sigma^2\) is the variance of the random effect \(\eta_i\). The nonlinear parameterization ensures the condition

\[
\logit P(Y_{ij} = 1 \mid X_{ij}) = \mu + \beta X_{ij}.
\]

(85)
Specification is straightforward using the `constrain` method, and standard likelihood theory can be applied on this model in contrast to the generalized estimating equation framework, e.g. likelihood ratio testing, profile likelihood confidence limits, and analysis with data missing at random as described in Section [7].

As an example we simulate observations from a simple random intercept probit model

```r
> margm <- lvm(c(y1, y2) ~ x)
> regression(margm, c(y1, y2) ~ u) <- "gamma"
> intercept(margm, endogenous(margm)) <- "mu"
> binary(margm) <- endogenous(margm)
> covariance(margm, ~u) <- 1
> latent(margm) <- ~u
> dmarg <- sim(margm, 100)
```

The logit-probit-normal model is then specified as

```r
> regression(margm, c(y1, y2) ~ x) <- 0
> constrain(margm, mu ~ x + alpha + beta + gamma) <- function(z) qnorm(1/(1 + exp(-z[2] - z[3] * z[1]))) * sqrt(1 + z[4]^2)
```

and estimates are obtained in the usual way

```r
> (emargm <- estimate(margm, dmarg))
```

| Estimate  | Std. Error | Z value | Pr(>|z|) |
|-----------|------------|---------|----------|
| **Measurements:** | | | |
| y1<-u | 2.08070 | 0.57583 | 3.61339 | 0.0003022 |
| **Intercepts:** | | | |
| alpha | 0.04207 | 0.20986 | 0.20049 | 0.8411 |
| beta | 1.35056 | 0.25772 | 5.24039 | 1.602e-07 |

9.4. Censored data

Censoring is a complication that is often encountered in cohort studies but can also be seen in experimental studies where thresholding of measurements may occur due to limiting precision of laboratory equipment.

We will assume that the data-generating mechanism is defined by [8] and [4] and that a subset of the endogenous variables $Y_{ij}^*$, $j \in J$ are censored such that for given censoring times $C_{ij}$, $j \in J$ we only observe

$$Y_{ij} = \begin{cases} Y_{ij}^*, & Y_{ij}^* < C_{ij} \\ C_{ij}, & Y_{ij}^* \geq C_{ij} \end{cases}$$

(86)
As an example we will simulate data from a regression model (see Figure 9), where we have a single dichotomous mediator $Z$ ($P(Z = 1) = 0.5$), which have a direct linear effect on the outcome of interest $Y$

$$Y = \mu_Y + \beta X + \gamma_Y Z + \epsilon_Y$$ (87)

and on the predictor $X$, which we assume is conditionally normally distributed given $Z$

$$X = \mu_X + \gamma_X Z + \epsilon_X$$ (88)

In our simulation we will let all intercepts be zero, residual variances 1, and slopes as defined by Figure 9:

```r
> med0 <- lvm(list(Y ~ X + Z, X ~ Z))
> distribution(med0, ~Z) <- binomial.lvm()
> d0 <- sim(med0, p = c(Y<Z, Y<2), n = 500)
```

We further assume we only observe the thresholded versions of $Y$ and $X$:

```r
> dtobit <- transform(d0, X = as.factor((X > 0) * 1), Y = Surv(pmin(Y, + 2), Y < 2))
```

As $X$ is coded as a factor and $Y$ is coded as a right-censored Surv-object (combinations of left and right censoring are allowed) the estimate method automatically applies a Probit and Tobit model respectively

```r
> (etobit <- estimate(med0, dtobit))
```

| Estimate | Std. Error | Z value | Pr(>|z|) |
|----------|------------|---------|----------|
| Y<-X     | 0.94429    | 0.07790 | 12.12215 | <1e-12   |
| Y<-Z     | 2.04975    | 0.13527 | 15.15247 | <1e-12   |
| X<-Z     | 1.00076    | 0.12154 | 8.23419  | <1e-12   |

| Intercept | Estimate | Std. Error | Z value | Pr(>|z|) |
|-----------|----------|------------|---------|----------|
| Y         | -0.02887 | 0.07733    | -0.37334 | 0.7089   |
| X         | -0.07362 | 0.07994    | -0.92096 | 0.3571   |

Residual Variances:

| Y | 0.97204 | 0.10225 | 9.50687 |

The Probit/Tobit model framework also has important applications in the causal modeling framework where it allows us to elegantly define direct and indirect effects for binary and censored data in complex path analyses.
The interpretation in this setup is linked to the assumption that the observations are generated by an unobserved continuous variable following a conditional normal distribution, and the indirect effects can therefore be quantified via the \texttt{effects} method:

\begin{verbatim}
> effects(etobit, Y ~ Z)
Total effect of 'Z' on 'Y':
  2.994759 (Approx. Std.Err = 0.1509939)
Direct effect of 'Z' on 'Y':
  2.04975 (Approx. Std.Err = 0.135275)
Indirect effects:
  Effect of 'Z' via Z->X->Y:
  0.9450087 (Approx. Std.Err = 0.1358391)
\end{verbatim}

9.4.1. Inverse probability weights

For models with complex sampling (survey data) and as an alternative method to deal with censored or missing data it is convenient to introduce Inverse Probability Weights in the estimation procedure (Horvitz and Thompson, 1952; Rotnitzky and Robins, 1995). The estimating equations in this situation becomes

\begin{equation}
\dot{U}_i^W(\theta; Y_i, Z_i) = -\frac{1}{2} \left\{ \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \left( \text{vec} \left[ (\Omega_\theta^{-1} \Omega_\theta^{-1}) W_i \right] - \Omega_\theta^{-1}(Y_i - \xi_{\theta,i})(Y_i - \xi_{\theta,i})' \Omega_\theta^{-1} \right) \right\} W_i = 0 \tag{89}
\end{equation}

where $W_i$ is the weight-matrix for the $i$th observation, and $\xi_{\theta,i}$ and $\Omega_\theta$ are the model-specific mean and covariance matrix of $Y_i$ given covariates $Z_i = (X_i', V_i', W_i')'$. In \texttt{lava}, equation (89) can be solved with a diagonal weight matrix using the estimator \texttt{weighted} and using the argument \texttt{weight} with the \texttt{estimate} method. The \texttt{weight} argument should be either a matrix with the weights of the endogenous variables of the model (or a named matrix with a subset of the variables) or alternatively a character vector with the names in the data.frame that corresponds to \texttt{weight1}1 (the weights are then

\footnote{1For the \texttt{lava.tobit} package the \texttt{weight} argument is already reserved and the \texttt{weight2} argument should be used instead and further the estimator should not be changed from the default.}
assigned to the variables in the models in the order they appear, see e.g the \texttt{vars} function. For multigroup models a list of matrices or character-vectors is expected.

9.5. Instrumental variables

In econometrics \textit{Instrumental Variable} (IV) estimators are popular tools for dealing with the problem of covariates that are correlated with the residual term of the response variable. In this situation ordinary linear regression analysis will yield biased estimates. The idea is to identify an IV, which is a variable fulfilling the conditions

1. Correlated with the problematic covariate, $X$, given all other covariates
2. Uncorrelated with the residual error, $\epsilon$, of the response, $Y$.

The estimator can be then be formulated as Two-Stage Least-Squares (2SLS) approach. In the first step, regress $X$ on the IV(s) and obtain the predicted covariate, $\hat{X}$. In the second stage, regress $Y$ on $\hat{X}$ (and other covariates). Consistency follows under very weak assumptions \cite{Greene2002, Angrist2001}.

The method can also be applied to estimate parameters in SEMs \cite{Bollen1996}. In the following we will assume that the model of interest is a SEM (no random slopes, single group) without linear or non-linear parameter constraints

\begin{equation}
Y = \nu + \Lambda \eta + KX + \epsilon, \tag{90}
\end{equation}
\begin{equation}
\eta = \alpha + B\eta + \Gamma X + \zeta, \tag{91}
\end{equation}

with a zero diagonal of $B$. We further assume that we have at least one indicator for each latent variable. This means that there exists a subset of endogenous variables $\tilde{Y}$ of $Y$, with no other predictors than the latent variable. From these we can identify the latent variables

$$\eta = \tilde{Y} - \tilde{\epsilon}. \tag{92}$$

Substituting \textit{(92)} into \textit{(90)} and \textit{(91)} we obtain the following equation for one of the endogenous variables, $Y_j$ (not part of $\tilde{Y}$):

\begin{equation}
Y_j = \nu_j + \Lambda_j \tilde{Y} + K_j X + \epsilon_j - \Lambda_j\tilde{\epsilon}, \tag{93}
\end{equation}
and similarly for the surrogate

\[ \tilde{Y}_r = \alpha_r + B_r \tilde{Y} + \Gamma_r X + \zeta_r + \tilde{\epsilon}_r - B_r \tilde{\epsilon}_r. \]  

Equations (93) and (94) only includes observed predictors but parameters cannot be estimated using OLS because of obvious correlations between predictors and residuals. Instruments have to be identified that are uncorrelated with the residual terms \((u_r \text{ or } u_j)\) in (93)-(94), while at the same time being correlated with the part of \(\tilde{Y}\) that are entering the equation as predictors. In \texttt{lava} these conditions are checked automatically via the model-implied covariance structure, selecting the largest possible set of instrument variables for each equation. To implicitly select the instruments for the different regression equations, covariance between specific residual terms should be added to the model structure via the \texttt{covariance} method (thereby indirectly disqualifying a variable as an instrument candidate). As all equations are solved simultaneously (Bollen, 2001) the covariance matrix of the estimates are available (i.e. \texttt{vcov}) and Wald tests via the \texttt{compare} method are possible. Consistent estimates of the variance parameters, are obtained via MLE in the model with all other parameters fixed at the IV estimates (can be disable with the control parameter \texttt{variance=FALSE}).

To illustrate the method we simulate data from a complex latent model with three measurement models (see Figure 11):

```r
> mIV <- lvm()
> regression(mIV) <- c(y1, y2, y3) ~ eta1
> regression(mIV) <- c(v1, v2, v3) ~ eta2
> regression(mIV) <- c(z1, z2, z3) ~ eta3
> latent(mIV) <- ~eta1 + eta2 + eta3
> regression(mIV) <- eta1 ~ eta2 + eta3 + x2
> regression(mIV) <- eta2 ~ eta3 + x1
> regression(mIV) <- eta3 ~ x1
> covariance(mIV) <- y1 ~ v1
> regression(mIV) <- y2 ~ x2
> regression(mIV, c(y1[0], v1[0], z1[0])) ~ eta1 + eta2 + eta3) <- c(1, 1, 1)
> dIV <- sim(mIV, 1000)[, manifest(mIV)]
```

In this model we explicitly defined \(y1, v1\) and \(z1\) as the indicators in (92) setting the factor loadings to one and intercepts to zero. To apply the instrumental variable estimator using the model-implied instruments, we
simply add the argument `estimator="IV"` to the `estimate` function, and as we have already chosen the indicator-set we set `fix=FALSE`:

```r
> summary(eIV <- estimate(mIV, dIV, estimator = "IV", fix = FALSE))
```

Latent variables: eta1 eta2 eta3
Number of rows in data=1000

| Measurements: | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|---------------|----------|------------|---------|---------|--------|
| y1<-eta1 | 1.00000 | 0.96794 | 1.00000 | 0.96794 |
| y2<-eta1 | 0.99358 | 0.01199 | 82.89643 | <1e-12 | 0.89359 |
| y3<-eta1 | 0.98599 | 0.01175 | 83.90153 | <1e-12 | 0.96603 |
| v1<-eta2 | 1.00000 | 0.91585 | 1.00000 | 0.91585 |
| v2<-eta2 | 0.99870 | 0.01972 | 50.64819 | <1e-12 | 0.92661 |
| v3<-eta2 | 1.02062 | 0.02023 | 50.45033 | <1e-12 | 0.92472 |
| z1<-eta3 | 1.00000 | 0.81644 | 1.00000 | 0.81644 |
| z2<-eta3 | 0.94467 | 0.03249 | 29.07141 | <1e-12 | 0.80369 |
| z3<-eta3 | 0.98653 | 0.03382 | 29.17123 | <1e-12 | 0.81647 |

| Regressions: | Estimate | Std. Error | Z value | Pr(>|z|) | std.xy |
|--------------|----------|------------|---------|---------|--------|
| y2<-x2 | 0.98771 | 0.04606 | 21.44170 | <1e-12 | 0.22739 |
| eta1<-eta2 | 1.05111 | 0.06144 | 17.10780 | <1e-12 | 0.64088 |
| eta1<-eta3 | 0.89998 | 0.10608 | 8.48410 | <1e-12 | 0.32826 |
| eta1<-x2 | 0.97850 | 0.05490 | 17.82364 | <1e-12 | 0.25048 |
| eta2<-eta3 | 0.90539 | 0.06139 | 14.74821 | <1e-12 | 0.54161 |
| eta2<-x1 | 1.05768 | 0.07972 | 13.26707 | <1e-12 | 0.44770 |
| eta3<-x1 | 0.97241 | 0.04543 | 21.40496 | <1e-12 | 0.68806 |
Intercepts:
\[
\begin{align*}
y_1 & = 0.00000 & 0.00000 \\
y_2 & = -0.03743 & 0.04464 & -0.83837 & 0.4018 & -0.00861 \\
y_3 & = -0.07527 & 0.04534 & 1.66018 & 0.09688 & -0.01886 \\
\eta_1 & = 0.01279 & 0.05512 & 0.23202 & 0.8165 & 0.00327 \\
v_1 & = 0.00000 & 0.00000 \\
v_2 & = -0.03401 & 0.04679 & -0.74560 & 0.4559 & -0.01323 \\
v_3 & = -0.02803 & 0.04534 & -0.59910 & 0.5491 & -0.01065 \\
\eta_2 & = 0.09849 & 0.05343 & 1.84351 & 0.06526 & 0.04131 \\
z_1 & = 0.00000 & 0.00000 \\
z_2 & = -0.00774 & 0.04259 & -0.18173 & 0.8558 & -0.00462 \\
z_3 & = 0.02848 & 0.04488 & 0.63450 & 0.5258 & 0.01652 \\
\eta_3 & = 0.05345 & 0.04585 & 1.16578 & 0.2437 & 0.03747
\end{align*}
\]

Residual Variances:
\[
\begin{align*}
y_1 & = 1.02973 & 0.06310 \\
y_{1,v1} & = 0.53963 & 0.05131 \\
y_2 & = 0.98617 & 0.05217 \\
y_3 & = 0.6375 & 0.06678 \\
\eta_1 & = 1.01587 & 0.06644 \\
v_1 & = 1.09261 & 0.16123 \\
v_2 & = 0.93365 & 0.14140 \\
v_3 & = 1.00333 & 0.14490 \\
\eta_2 & = 0.98078 & 0.17254 \\
z_1 & = 1.01747 & 0.33342 \\
z_2 & = 0.99507 & 0.35408 \\
z_3 & = 0.99004 & 0.33338 \\
\eta_3 & = 1.07110 & 0.52657
\end{align*}
\]

Estimator: IV

Latent variables: \( \eta_1, \eta_2, \eta_3 \)

Surrogate variables: \( y_1, v_1, z_1 \)

Response Instruments
\[
\begin{align*}
y_2 & = y_3, v_2, v_3, z_1, z_2, z_3, x_2, x_1 \\
y_3 & = y_2, v_2, v_3, z_1, z_2, z_3, x_2, x_1 \\
\eta_1 & = v_2, v_3, z_2, z_3, x_2, x_1 \\
v_2 & = y_2, y_3, v_3, z_1, z_2, z_3, x_2, x_1 \\
v_3 & = y_2, y_3, v_2, z_1, z_2, z_3, x_2, x_1 \\
\eta_2 & = z_2, z_3, x_2, x_1
\end{align*}
\]
The IV estimator has the advantage of being a non-iterative procedure and requires weaker assumptions than MLE. There are also indications that IV estimators are performing well in low sample-sizes and are generally more robust to structural model mis-specification than MLE (Bollen et al, 2007). Where applicable an IV analysis is therefore a good choice of method for analyzing the model fit of a structural equation model fitted with MLE (e.g. based on a Hausman type test statistics).

10. Graphics

| Function      | Task                                              |
|---------------|---------------------------------------------------|
| plot          | Plots path diagram of model                       |
| labels        | Defines labels for variables                      |
| edgelabels    | Defines labels and style for edges of the graph   |
| nodecolor     | Defines color and style of nodes/variables        |
| Graph         | Extracts graph (graphNEL object)                  |

Table 5: Graphics functions.

A plot method is available for both the lvm, multigroup and lvmfit classes. Plotting a lvmfit object shows the user whether lava has linked the model to the data as intended. A common mistake is that a variable name in the model specification does not appear in the data. In this case lava consider the corresponding variable to be latent which is easily identified from the plot.

Layout and rendering of the graphs are achieved via Rgraphviz (Gentry et al, 2009), which in combination with the tikzDevice (Sharpsteen and Bracken, 2010) makes it possible to produce publication quality path diagrams.

In the following we will use model m1 defined in Section 3.1 as the example. To enhance the graph we will add some labels to the nodes using the function labels.

---

for more information on mathematical annotation in R we refer to the plotmath help-page.
> labels(m1) <- c(u1 = expression(eta[1]), u2 = expression(eta[2]))

Similarly, subscripted versions of the observed variables could be defined but we will keep them as they are for now. The labels of an object can be examined with `labels(m)`.

Labels of edges can be defined with the `edgelabels`-function, e.g. to define new labels, colors and line width for the edges from \( x_1 \) to \( u_1 \) and from \( u_1 \) to \( y_{11} \) we call

```r
> m1 <- m0
> edgelabels(m1, u1~x1, lwd=3, col="blue") <- expression(beta[1])
> edgelabels(m1, y11~u1, lwd=3, col="red",
+        labcol="red") <- expression(beta[2])
```

In addition the argument `labels=TRUE` can be parsed to the `plot` method to add parameter names (named \( p_1, p_2, \ldots \) if no names were previously given using, e.g., `regression`) to the edges of the plot. This will override (but not delete) previously defined edge label attributes.

The color of the nodes is automatically added to the graph. To disable this functionality the argument `addcolor=FALSE` should be parsed to the `plot`-method. New coloring and style can be added via `nodecolor`:

```r
nodecolor(m1) <- "indianred"
nodecolor(m1, ~y1+y12+y13,
+        labcol="white", lwd=c(3,1,1)) <- "lightblue"
nodecolor(m1, ~x1+x2, labcol="red",
+        border=c("black","white")) <- "white"
```

### 10.1. Graph Attributes

Specific attributes of the graph such as font size, line width etc. can also be controlled via the `nodeRenderInfo` and `edgeRenderInfo` functions called on the `graph` object in a `lvm`-object accessible via the `Graph` function (also available for `lvmfit`-objects). E.g. to set the font size to 2 of all edge-labels (see Figure[12]) we would write

```r
> edgeRenderInfo(Graph(m1))$cex <- 1.5
```

Methods for visualizing `lvmfit` are also available. As an example we will extract the pathways from \( x_1 \) to \( z_3 \) with the `path` method and highlight and label the corresponding arrows in the path-diagram (see Figure[13]):

```r
Use the `path` function to extract pathways:```
Figure 12: plot(m1)

```r
Graph(e) <- Graph(e, add=TRUE)
labels(Graph(e)) <- c(u1=expression(eta[1]), u2=expression(eta[2]))
mypath <- path(e, z3~x1)
edgeRenderInfo(Graph(e))$lwd[unlist(mypath$edges)] <- 2
edgeRenderInfo(Graph(e))$label <- NA
edgelabels(e, edges=unique(unlist(mypath$edges)), cex=1.1) <-
+ c("beta[1]", "gamma[1]", "gamma[2]", "beta[2]")
```

10.2. Graph Layout

Several automatic graph layout algorithms are available through the `layoutType` argument (see Figure 14). Additionally, the graph (obtainable via the `Graph` method) can be saved with the `doDot` function and be processed in an external program supporting the `dot`-format (graphviz).
\[ \eta_1 = 0.9465 \]
\[ \eta_2 = 1.024 \]
\[ \beta_1 = 1.111 \]
\[ \beta_2 = 0.7283 \]

Figure 13: Selected estimates from model \texttt{m1 (plot(e,diag=FALSE))}

Figure 14: Different graph layout algorithms from top left to bottom right: dot, neato, twopi, fdp, circo.
11. Application: Brain serotonin transporter imaging data

We consider 54 observations from [Kalbitzer et al., 2010] with measurements of the serotonin transporter (SERT) in the human brain as measured by Positron Emission Tomography (PET) techniques. The outcome of interest, SERT, was quantified as binding potential of the specific tracer binding (BP_{ND}) in four regions of interest, which a priori were identified as high-binding and reliable measurements of SERT. The serotonergic system has been suggested to have a strong impact on mood and as a potential predictor of the development of seasonal affective disorder (SAD). The aim of the original study was to explore the association between levels of SERT and the interaction between seasonality and a repeat polymorphism in the promoter region of the serotonin transporter gene (5-HTTLPR). This was achieved by linear regression on each regional outcome. However, these data are characterized by high inter-regional correlation, and in the following we will supplement the original analysis with a multivariate analysis taking this aspect into account. It has been demonstrated that SERT levels respond to chronic changes in brain serotonin (5-HT) levels as suggested by studies of the effect of SSRI treatment and in animal studies. It has therefore been suggested that the common regulator of SERT is this underlying 5-HT level [Erritzoe et al., 2010] which, however, cannot be measured directly in vivo. This biological model can be captured by a structural equation model, with a simple measurement model describing the four regions of interest, and a structural model in which exogenous variables affects the SERT measurements indirectly through the intermediate latent variable as shown on Figure 15.

We chose to model the seasonal effect using a harmonic function with a period of one year, described by the amplitude, A, and the translation parameter, δ (time of peak):

\[
A \cos \left( \frac{2\pi(t - \delta)}{365} \right) = A \cos \left( \frac{2\pi\delta}{365} \right) \cos \left( \frac{2\pi t}{365} \right) + A \sin \left( \frac{2\pi\delta}{365} \right) \sin \left( \frac{2\pi t}{365} \right),
\]

(95)

where \( t \) is the date of the scan. This parameterization approximately embeds a simpler model, where the seasonal effect is described as a linear function of the amount of daylight minutes on the date of the PET scan. However, the harmonic curve adds the flexibility of letting the time of peak be a free parameter which allows a possible delayed seasonal effects on the serotonin.
transporter to be taken into account. In lava the seasonal effect could be modeled directly (via `constrain`) as the left-hand-side of (95), thus directly quantifying $A$ and $\delta$. Here we use the linear parameterization on the right-hand-side using the cosine and sine transformed time variables as predictors. As in the original study, we also adjust for possible main effects of age, gender and the 5-HTTLPR polymorphism (dichotomized as carriers of the (short) s-variation vs. non-carriers (long-long alleles)).

```r
ttmod <- lvm(c(cau,th,put,mid) ~ eta)
regression(httmod) <- eta ~ f(cos,b1)+f(sin,b2)+age+sex+gene
```

As described in Section [6](#), we assessed the model fit via residual plots, score tests and the global $\chi^2$-test, and concluded that the local independence between the midbrain and thalamus region was not plausible. Significantly different age and gender effects on caudate nucleus were also identified in this process and therefore added to the model.

```r
regression(httmod) <- cau~sex+age
covariance(httmod) <- th~mid
```

No additional evidence against the model was identified, thus leading to a final model as illustrated in Figure 16:

```r
tikz(file="sertsem1.tex",8,8,standAlone=TRUE)
m <- lvm(~cau+th+put+mid)
regression(m) <- c(cau,mid,th,put)~eta
latent(m) <- `eta; labels(m) <- c(eta="$\eta$")
regression(m) <- eta~sex+age+date
```

Figure 15: Brain regions of interests and initial SERT model.
> nodecolor(m,vars(m)) <- 0
> nodecolor(m,endogenous(m),labcol="white") <-
> + c("red","blue","green","orange")
> plot(m)
> dev.off()

Figure 16: Final SERT-seasonality model.

Parameter estimates are obtained by maximum likelihood using a Newton-Raphson algorithm:

> lava.options(method="NR",trace=1,tol=1e-12,param="relative")
> httmod.fit <- estimate(httmod,data=dasb)

One of the aims of the analysis is to quantify the seasonal effect, therefore we will also estimate the translation and amplitude (unsigned)

\[
\hat{\delta} = \arctan(\hat{\beta}_2/\hat{\beta}_1)365/(2\pi) \tag{96}
\]

\[
|\hat{A}| = \sqrt{\hat{\beta}_1^2 + \hat{\beta}_2^2} \tag{97}
\]

> constrain(httmod.fit,delta~b1+b2) <- function(x) atan(x[2]/x[1])*365/(2*pi)
> constrain(httmod.fit,A~b1+b2) <- function(x) sqrt(x[1]^2+x[2]^2)
> summary(httmod.fit, std=NULL)
Score: $1.171417e^{-11}$

Latent variables: $\eta$

Number of rows in data: 54

| Measurements: | Estimate | Std. Error | Z value | Pr(|z|) |
|---------------|----------|------------|---------|---------|
| $\text{cau} \leftarrow \eta$ | $1.0000e+00$ |  |  | |
| $\text{th} \leftarrow \eta$ | $9.6834e-01$ | $3.0972e-01$ | $3.1264e+00$ | $1.7693e-03$ |
| $\text{put} \leftarrow \eta$ | $1.2119e+00$ | $3.5519e-01$ | $3.4120e+00$ | $6.4499e-04$ |
| $\text{mid} \leftarrow \eta$ | $1.1317e+00$ | $7.5972e-01$ | $1.4896e+00$ | $1.3633e-01$ |

| Regressions: | Estimate | Std. Error | Z value | Pr(|z|) |
|--------------|----------|------------|---------|---------|
| $\text{cau} \leftarrow \text{age}$ | $6.7822e-03$ | $1.9574e-03$ | $3.4648e+00$ | $5.3054e-04$ |
| $\text{cau} \leftarrow \text{sex}$ | $-3.2907e-01$ | $7.4087e-02$ | $-4.4416e+00$ | $8.9284e-06$ |
| $b1: \eta \leftarrow \text{cos}$ | $1.0722e-01$ | $3.9546e-02$ | $2.7112e+00$ | $6.7043e-03$ |
| $b2: \eta \leftarrow \text{sin}$ | $-3.7658e-02$ | $4.1421e-02$ | $-9.0915e-01$ | $3.6327e-01$ |
| $\eta \leftarrow \text{age}$ | $-2.9471e-03$ | $1.4856e-03$ | $-1.9838e+00$ | $4.7282e-02$ |
| $\eta \leftarrow \text{sex}$ | $-7.8173e-03$ | $4.9608e-02$ | $-1.5758e+00$ | $8.7439e-01$ |
| $\eta \leftarrow \text{GnonLL}$ | $-4.5062e-02$ | $4.7688e-02$ | $-9.4494e-01$ | $3.4469e-01$ |

| Intercepts: | Estimate | Std. Error | Z value | Pr(|z|) |
|-------------|----------|------------|---------|---------|
| $\text{cau}$ | $0.0000e+00$ |  |  | |
| $\text{th}$ | $2.8698e-01$ | $5.9078e-01$ | $4.8577e-01$ | $6.2713e-01$ |
| $\text{put}$ | $-4.1726e-01$ | $6.8945e-01$ | $-6.0521e-01$ | $5.4504e-01$ |
| $\text{mid}$ | $2.0487e+00$ | $1.4175e+00$ | $1.4453e+00$ | $1.4838e+01$ |
| $\eta$ | $1.9647e+00$ | $8.7535e-02$ | $2.2445e+01$ | $<1e-16$ |

| Residual Variances: | Estimate | Std. Error | Z value | Pr(|z|) |
|---------------------|----------|------------|---------|---------|
| $\text{cau}$ | $5.4901e-02$ | $1.2519e-02$ | $4.3855e+00$ | |
| $\text{th}$ | $6.9573e-02$ | $1.4911e-02$ | $4.6658e+00$ | |
| $\text{th,mid}$ | $9.6725e-02$ | $3.4453e-02$ | $2.8074e+00$ | $4.9940e-03$ |
| $\text{put}$ | $1.0455e-02$ | $9.2801e-03$ | $1.1266e+00$ | |
| $\text{mid}$ | $6.9218e-01$ | $1.3472e-01$ | $5.1377e+00$ | |
| $\eta$ | $2.0335e-02$ | $9.8548e-03$ | $2.0635e+00$ | |

Non-linear constraints:

| Estimate | Std. Error | Z value | Pr(|z|) | 2.5% | 97.5% |
|----------|------------|---------|--------|------|------|
| $\delta$ | $-19.6215810$ | $19.5444140$ | $-1.0039483$ | $0.3154035$ | $-57.9279285$ | $18.6848$ |
| $A$ | $0.1136378$ | $0.0426363$ | $2.6652834$ | $0.0076923$ | $0.0300722$ | $0.1972$ |

Estimator: gaussian

Number of observations: 54

Log-Likelihood: $-57.88782$
As an alternative to the Wald test we can conduct a LRT to test the significance of the seasonal parameters:

```r
> httmod0 <- httmod; kill(httmod0) <- ~cos+sin
> httmod0.fit <- estimate(httmod0,dd,control=list(start=coef(httmod.fit)))
> compare(httmod0.fit,httmod.fit)
```

Likelihood ratio test

data:  
chisq = 12.3978, df = 2, p-value = 0.002032
sample estimates:
log likelihood (model 1) log likelihood (model 2)
-64.53133     -58.33241

Thus, we find a highly significant seasonal effect ($p = 0.002$). The estimated time of peak and 95% Wald confidence limits can be quantified as

```r
> format(as.Date("2010-1-1")+constraints(httmod.fit)["delta",c(1,5:6)],
+ "%d%b")
```

Estimate  2.5%  97.5%
"12Dec"  "04Nov"  "19Jan"

and with an estimate around middle of December this is in reasonable agreement with a suggested peak around winter solstice (about December 21). The parameter estimates of the initial model (Figure 15) yielded very similar seasonal parameter estimates and confidence limits.

Next, we will examine the interaction between season and the 5-HTTLPR polymorphism. This is done via a multigroup analysis and hence we divide the data into two groups defined by the genetic variable

```r
> d1 <- subset(dd, G="LL")
> d2 <- subset(dd, G="nonLL")
```
A standard multigroup analysis could be conducted where the parameters $\beta_1$ and $\beta_2$ would be allowed to vary freely in the two groups. However, a more biological plausible model is to fix the translation parameter $\delta$ in the two groups to be the same and let the amplitude $A$ be free. This can be implemented via the left-hand-side of (95) and the `constrain`-method:

```r
> m <- baptize(kill(httmod, "G*cos*sin"))
> intercept(m, "cau+eta") <- list(0, "mu1")
> regression(m, cau~eta) <- 1
> regression(m, eta~Day) <- 0
> m2 <- m
> intercept(m2, "eta") <- "mu2"
> mycos <- function(x) x[2] + x[3] * cos(2*pi*(x[1]-x[4])/365)
> constrain(m, mu~Day+nu1+A1+delta) <- mycos
> constrain(m2, mu2~Day+nu2+A2+delta) <- mycos
```

Here we explicitly chose caudate nucleus as our reference region by fixing the factor loading to 1 and intercept to 0. The intercept parameters $\nu_1$ and $\nu_2$ describes the main effect of the genotype. The significance of the interaction can then be examined with a LRT against the first model:

```r
> httmod.2 <- estimate(list(m, m2), list(d1, d2))
> compare(httmod.2, httmod.fit)
```

Likelihood ratio test

data:
chisq = 4.7267, df = 1, p-value = 0.0297

sample estimates:
log likelihood (model 1) log likelihood (model 2)
-55.52446 -57.88782

The estimated amplitude parameters with 95% confidence limits are

```r
> confint(httmod.2)[c("A1", "A2"), c(1, 3, 4)]
```

|         | Estimate | Pr(>|z|) | 2.5%     | 97.5%     |
|----------|----------|----------|----------|-----------|
| A1       | 0.01672275 | 0.691577803 | -0.06589577 | 0.10934077 |
| A2       | 0.14670814 | 0.002547141 | 0.05142227 | 0.24203387 |

and the difference in amplitude can be quantified as

```r
> constrain(httmod.2, dA~A1+A2) <- diff
> constraints(httmod.2)["dA",]
```
with peak time around January 1:

\[
\text{coef(httmod.2)[[1]]["delta",]}
\]

| Estimate  | Std. Error  | Z value | Pr(>|z|) | 2.5%      | 97.5%     |
|-----------|-------------|---------|---------|-----------|-----------|
| 0.129985393 | 0.061819428 | 2.102662508 | 0.035495282 | 0.008821541 | 0.251149245 |

Hence we see a statistical significant difference in amplitude between the two genotypes \( (p = 0.03, \text{ 95\% confidence limits [0.01; 0.25]}), \) with carriers of the short 5-HTTLPR allele having on average a higher seasonal variation in SERT binding.

To visualize the harmonic curves of the two genotypes we need to predict the parameters \( \mu_1 \) and \( \mu_2 \). This can also be achieved with `constraints` where we supply the argument `idx` indicating which non-linear parameter to extract. For a `multigroup` model we also need to specify which group to extract this parameter from via the argument `k`. For instance predicting the two means on each day of the year with 95% confidence limits we can write

\[
\begin{align*}
\text{mu1} &\leftarrow \text{constraints}(e, k=1, idx="\mu", data=data.frame(Day=1:365), level=0.95) \\
\text{mu2} &\leftarrow \text{constraints}(e, k=2, idx="\mu2", data=data.frame(Day=1:365), level=0.95)
\end{align*}
\]

See Figure [17] for a plot of these two curves.

This study shows that S-allele carriers (nonLL) on average have a much more varying SERT binding level which could suggest a decreased serotonin concentration in the winter in this group. This could be caused by the decreasing amount of daylight in this period in the study country (Denmark) and supplements previous studies showing that S-allele carriers have higher risk of developing SAD. A detailed discussion can be found in the original paper [Kalbitzer et al, 2010].

12. Conclusion

A package `lava` has been developed which covers the classical covariance structure analysis and which hopefully can serve as a platform for methodological development in the field of structural equation models and related models.

The key features of the package is
1. Easy interactive specification and visualization of complex models
2. Simulation routines (for a broad class of models beyond the LLVM)
3. Extensions to binary and censored data via \texttt{lava.tobit}
4. Multigroup analyses
5. MLE with data missing at random
6. Non-linear parameter constraints and covariate effects
7. Asymptotically correct standard errors for clustered correlated data

Further, the program is built up around a series of modules (optimizers, estimators, plot hooks, simulations hooks, pre and post estimation hooks) which should ensure that future extensions can be written quite easily. This has been one of the main aims during the development of the \texttt{lava} package.

Additional extensions of the package is currently under preparation including non-linear random effects and MLE for a broader class of the exponential family with estimation based on adaptive quadrature rules. In this process, we are preparing to export loop-intensive parts of the program to C++, which also should give a considerable computational and memory (via call-by-reference) improvement for some of the closed-form likelihood models (e.g. random slope models) and models with large number of variables.

If you use \texttt{lava} please cite this paper and the R software in publications.
13. Acknowledgments

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Appendix A. Some zero-one matrices

In this section we will define a few matrix-operators in order to define various conditional moments. Let $B \in \mathbb{R}^{n \times m}$ be a matrix, and define the indices $x = \{x_1, \ldots, x_k\} \in \{1, \ldots, n\}$, and $y = \{y_1, \ldots, y_l\} \in \{1, \ldots, m\}$. We define $J_{n,x} = J_x \in \mathbb{R}^{(n-k) \times n}$ as the $n \times n$ identity matrix with rows $x$ removed. E.g.

$$J_{6,(3,4)} = \begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 1
\end{pmatrix}. $$

To remove rows $x$ from $B$ we simply multiply from the left with $J_{n,x}$. If we in addition want to remove columns $y$ we multiply with the transpose of $J_{n,y}$ from the right:

$$J_{n,x}B J_{n,y}'. \quad (A.1)$$

We will use the notation $J$ to denote the matrix that removes all latent variables ($\eta$) from the vector of all variables, $U$ as defined in (8). We denote $J_Y$ the matrix that only keeps endogenous variables ($Y$).

We also need an operator that cancels out rows or columns of a matrix/vector. Define the square matrix $p_{n,x} \in \mathbb{R}^{n \times n}$ as the identity-matrix with diagonal elements at position $x$ canceled out:

$$p_{n,x}(i,j) = \begin{cases}
1, & i = j, i \notin x, \\
0, & \text{else}.
\end{cases} \quad (A.2)$$

E.g.

$$p_{6,(3,4)} = \begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0
\end{pmatrix}. $$

To cancel out rows $x$ and columns $y$ of the matrix $B \in \mathbb{R}^{n \times m}$ we calculate

$$p_{n,x}B p_{m,y}'. $$

We will use $p_X$ and $p_{\bar{X}}$ as the matrix that cancels out the rows corresponding to the index of the exogenous variables ($X$) respectively the matrix that cancels out all rows but the ones corresponding to $X$. 

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Appendix B. The Score Function and Information

In this section we will calculate the analytical derivatives of the log-likelihood. In order to obtain these results we will first introduce the notation of some common matrix operations. Let \( A \in \mathbb{R}^{m \times n} \), then we define the column-stacking operation:

\[
\text{vec}(A) = \begin{pmatrix} a_1 \\ \vdots \\ a_n \end{pmatrix},
\]

where \( a_i \) denotes the \( i \)th column of \( A \). The unique commutation matrix, \( \mathbb{R}^{mn \times mn} \) is defined by

\[
K^{(m,n)} \text{vec}(A) = \text{vec}(A').
\]

Letting \( H^{(i,j)} \) be the \( m \times n \)-matrix with one at position \((i, j)\) and zero elsewhere, then

\[
K^{(m,n)} = \sum_{i=1}^{m} \sum_{j=1}^{n} (H^{(i,j)} \otimes H^{(i,j)'})\),
\]

e.g.

\[
K^{(2,3)} = \begin{pmatrix} 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{pmatrix}.
\]

It should be noted that product with a commutation matrix can be implemented very efficiently instead of relying on a direct implementation of the above mathematical definition.

Let \( A \in \mathbb{R}^{m \times n} \) and \( B \in \mathbb{R}^{p \times q} \) then the Kronecker product is the \( mp \times nq \)-matrix:

\[
A \otimes B = \begin{pmatrix} a_{11}B & \cdots & a_{1n}B \\ \vdots & & \vdots \\ a_{m1}B & \cdots & a_{mn}B \end{pmatrix}
\]
We will calculate the derivatives of (20) by means of matrix differential calculus. The *Jacobian matrix* of a matrix-function $F: \mathbb{R}^n \to \mathbb{R}^{m \times p}$ is the $mp \times n$ matrix defined by

$$DF(\theta) = \frac{\partial \text{vec } F(\theta)}{\partial \theta'}.$$ 

Letting $d$ denote the differential operator (see Magnus and Neudecker (1988)), the first identification rule states that $d \text{vec } F(\theta) = A(\theta) d \theta \Rightarrow DF(\theta) = A(\theta)$.

**Appendix B.0.1. Score function**

Using the identities $d \log |X| = \text{tr}(X^{-1} d X)$ and $d X^{-1} = -X^{-1}(d X)X^{-1}$, and applying the product rule we get

$$d \ell(\theta) = -\frac{n}{2} \text{tr}(\Omega^{-1}_\theta d \Omega_\theta) - \frac{n}{2} \text{tr}(\hat{\Sigma} \Omega^{-1}_\theta)$$  

where

$$d \Omega_\theta = \{d G_\theta \} P_\theta G'_\theta + G_\theta \{d P_\theta G'_\theta \}$$  

and

$$d G_\theta = J(1_m - A_\theta)^{-1} \{d A_\theta \} (1_m - A_\theta)^{-1}.$$  

Taking vec's it follows that

$$d \text{vec } G_\theta = [((1_m - A_\theta)^{-1})' \otimes G_\theta] \text{ d vec } A_\theta \text{ d } \theta,$$

hence by the first identification rule

$$\frac{\partial \text{vec } G_\theta}{\partial \theta'} = [((1_m - A_\theta)^{-1})' \otimes G_\theta] \frac{\partial \text{vec } A_\theta}{\partial \theta'},$$

and similarly

$$\frac{\partial \text{vec } \Omega_\theta}{\partial \theta'} = (1_{k^2} + K^{(k,k)}) [G_\theta P_\theta \otimes 1_k] \frac{\partial \text{vec } G_\theta}{\partial \theta'} + [G_\theta \otimes G_\theta] \frac{\partial \text{vec } P_\theta}{\partial \theta'},$$

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and finally (exploiting the symmetry of $\Omega_\theta$ and commutative property under the trace operator) we obtain the gradient

$$\frac{\partial \ell(\theta)}{\partial \theta} = \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1} \hat{\Sigma} \Omega_\theta^{-1} \right] - \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1} \right]. \tag{B.11}$$

Next we examine the model including a mean structure \cite{20}. W.r.t. to the first differential we observe that

$$d \text{tr} \left\{ T_\theta \Omega_\theta^{-1} \right\} = - \text{tr} \left\{ T_\theta \Omega_\theta^{-1} (d \Omega_\theta) \Omega_\theta^{-1} \right\} + \text{tr} \left\{ (d T_\theta) \Omega_\theta^{-1} \right\}. \tag{B.12}$$

Hence

$$\frac{\partial \ell(\theta)}{\partial \theta} = \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1} T_\theta \Omega_\theta^{-1} \right] - \frac{n}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right)' \text{vec} \left[ \Omega_\theta^{-1} \right]$$

$$- \frac{n}{2} \left( \frac{\partial \text{vec} T_\theta}{\partial \theta'} \right)' \text{vec} \left( \Omega_\theta^{-1} \right). \tag{B.13}$$

Further by the chain-rule

$$\frac{\partial \text{vec} T_\theta}{\partial \theta'} = \frac{\partial (\hat{\mu} - v_\theta)}{\partial \theta'} \frac{\partial \text{vec} T_\theta}{\partial (\hat{\mu} - \xi_\theta)} = - [k_1 \otimes (\hat{\mu} - \xi_\theta)] \frac{\partial \xi_\theta}{\partial \theta'}, \tag{B.14}$$

and

$$d \xi_\theta = (d G_\theta) v_\theta + G_\theta (d v_\theta). \tag{B.15}$$

Taking vec ($G_\theta v_\theta = 1 G_\theta v_\theta$):

$$\frac{\partial \xi_\theta}{\partial \theta'} = (v'_\theta \otimes 1_k) \frac{\partial \text{vec} G_\theta}{\partial \theta'} + G_\theta \frac{\partial v_\theta}{\partial \theta'}. \tag{B.16}$$

We have calculated the full score but in some situations it will be useful to evaluate the score in a single point. The contribution of a single observation to the log-likelihood is

$$\ell(\theta \mid z_i) \propto \frac{1}{2} \log |\Omega_\theta| + \frac{1}{2} (z_i - \xi_\theta)' \Omega_\theta^{-1} (z_i - \xi_\theta), \tag{B.17}$$

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or as in (20) where we simply exchange $T_\theta$ with $T_{z_i, \theta} = (z_i - \xi_\theta)(z_i - \xi_\theta)'$, hence the score is as in (B.13) where (B.14) is calculated with $z_i$ instead of $\hat{\mu}$. Alternatively, letting $z_i - \xi_\theta = u_\theta = u_\theta(i)$:

$$d(u_\theta' \Omega_\theta^{-1} u_\theta) = u_\theta' [(2\Omega_\theta^{-1}) d u_\theta + (d \Omega_\theta^{-1}) u_\theta] = -u_\theta' [(2\Omega_\theta^{-1}) d \xi_\theta + \Omega_\theta^{-1}(d \Omega_\theta) \Omega_\theta^{-1} u_\theta],$$

(B.18)

where we used that for constant symmetric $A$ the differential of a quadratic form is

$$d(u' A u) = 2u' [A^2] d u.$$  

(B.19)

Hence the contribution to the score function of the $i$th observation is

$$S_i(\theta) = -\frac{1}{2} \left\{ \text{vec}(\Omega_\theta^{-1}) \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} - 2u_\theta' \Omega_\theta^{-1} \frac{\partial \text{vec} \xi_\theta}{\partial \theta'} - (u_\theta' \Omega_\theta^{-1} \otimes u_\theta' \Omega_\theta^{-1}) \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right\},$$

(B.20)

where the score-function evaluated in $\theta$ is $S(\theta) = \sum_{i=1}^n S_i(\theta)$.

Appendix B.0.2. The Information matrix

The second order partial derivative is given by

$$\frac{\partial \ell(\theta)}{\partial \theta_i \theta_j} = -\frac{1}{2} \frac{\partial}{\partial \theta_i} \left\{ \text{vec}(\Omega_\theta^{-1}) - \text{vec}(\Omega_\theta^{-1} u_\theta u_\theta') \right\} \frac{\partial \text{vec} \Omega_\theta}{\partial \theta_j} - 2u_\theta' \Omega_\theta^{-1} \frac{\partial \text{vec} \xi_\theta}{\partial \theta_j}. $$

(B.21)

Taking negative expectation with respect to the true parameter $\theta_0$ we obtain the expected information (Magnus and Neudecker, 1988), which get rid of all second order derivatives

$$\mathcal{I}(\theta_0) = \frac{1}{2} \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right|_{\theta=\theta_0} ' \left( \Omega_{\theta_0}^{-1} \otimes \Omega_{\theta_0}^{-1} \right) \left( \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'} \right|_{\theta=\theta_0} ' + \left( \frac{\partial \xi_\theta}{\partial \theta'} \right|_{\theta=\theta_0} ' \Omega_{\theta_0}^{-1} \left( \frac{\partial \xi_\theta}{\partial \theta'} \right|_{\theta=\theta_0} ').$$

(B.22)
We will further derive the observed information in the case where the second derivatives vanishes in the case of the matrix functions $A_\theta, P_\theta$ and $v_\theta$. Now
\[
d^2G_\theta = d \left[ J (1 - A_\theta)^{-1} (dA_\theta)(1 - A_\theta)^{-1} \right]. \tag{B.24}
\]
Hence
\[
\frac{\partial^2 G_\theta}{\partial \theta_i \partial \theta_j} = G_\theta \left[ \frac{\partial A_\theta}{\partial \theta_i} (1 - A_\theta)^{-1} \frac{\partial A_\theta}{\partial \theta_j} + \frac{\partial A_\theta}{\partial \theta_j} (1 - A_\theta)^{-1} \frac{\partial A_\theta}{\partial \theta_i} \right] (1 - A_\theta)^{-1}. \tag{B.25}
\]
Next we will find the derivative of (B.10). We let $m$ denote the number of variables, $p$ the number of parameters, and $k$ the number of observed variable (e.g. $G_\theta \in \mathbb{R}^{k \times m}$ and the number of columns in the derivatives are $p$). We have $G_\theta P_\theta \in \mathbb{R}^{k \times m}$ and using rules for evaluating the differential of Kronecker-product (see Magnus and Neudecker (1988) pp. 184) we obtain
\[
\frac{\partial \text{vec}}{\partial \theta'} (G_\theta P_\theta \otimes 1_k) = \left( 1_m \otimes K^{(k,k)} \otimes 1_k \right) (1_km \otimes \text{vec} 1_k) \frac{\partial \text{vec} G_\theta P_\theta}{\partial \theta'} \\
= \left( 1_m \otimes K^{(k,k)} \otimes 1_k \right) (1_km \otimes \text{vec} 1_k) \times \left[ (P_\theta \otimes 1_k) \frac{\partial \text{vec} G_\theta}{\partial \theta'} + (1_m \otimes G_\theta) \frac{\partial \text{vec} P_\theta}{\partial \theta'} \right]. \tag{B.26}
\]
And
\[
\frac{\partial \text{vec}}{\partial \theta'} \left[ (G_\theta \otimes G_\theta) \frac{\partial \text{vec} P_\theta}{\partial \theta'} \right] = \left[ \frac{\partial \text{vec} P_\theta'}{\partial \theta'} \otimes 1_k \right] \frac{\partial \text{vec} G_\theta \otimes G_\theta}{\partial \theta'} \\
= \left[ \frac{\partial \text{vec} P_\theta'}{\partial \theta'} \otimes 1_k \right] \left( 1_m \otimes K^{(m,k)} \otimes 1_k \right) \times \left( 1_km \otimes \text{vec} G_\theta + \text{vec} G_\theta \otimes 1_km \right) \frac{\partial \text{vec} G_\theta}{\partial \theta'}. \tag{B.27}
\]
Hence from (B.26) and (B.27) and using rules for applying the vec operator
on products of matrices we obtain

\[ \frac{\partial^2 \text{vec} \Omega_\theta}{\partial \theta \partial \theta'} = \left[ \left( \frac{\partial \text{vec} G_\theta}{\partial \theta'} \right)' \otimes (1_k \otimes K^{(k,k)}) \right] (1_m \otimes \Omega_\theta^{-1} \otimes 1_k) \times \]
\[ \left[ (P_\theta \otimes 1_k) \frac{\partial \text{vec} G_\theta}{\partial \theta'} + (1_m \otimes G_\theta') \frac{\partial \text{vec} P_\theta}{\partial \theta'} \right] + \]
\[ \left( 1_p \otimes (G_\theta P_\theta \otimes 1_k) \right) \frac{\partial^2 \text{vec} G_\theta}{\partial \theta \partial \theta'} + \]
\[ \left( \left( \frac{\partial \text{vec} G_\theta}{\partial \theta'} \right)' \otimes 1_k \right) (1_m \otimes \Omega_\theta^{-1} \otimes 1_k) \times \]
\[ (1_k \otimes \text{vec} G_\theta \otimes 1_k) \frac{\partial \text{vec} G_\theta}{\partial \theta'}. \]

(B.28)

with the expressions for the derivatives and second derivatives of \( G_\theta \) given in (B.9) and (B.25). Further

\[ \frac{\partial^2 \xi_\theta}{\partial \theta \partial \theta'} = \frac{\partial \text{vec} (v_\theta' \otimes 1_k)}{\partial \theta'} \frac{\partial \text{vec} G_\theta}{\partial \theta'} + \frac{\partial \text{vec} G_\theta}{\partial \theta'} \frac{\partial \text{vec} v_\theta}{\partial \theta'} = \]
\[ \left( \left( \frac{\partial \text{vec} G_\theta}{\partial \theta'} \right)' \otimes 1_k \right) \left( 1_m \otimes \text{vec} 1_k \right) \frac{\partial \text{vec} v_\theta}{\partial \theta'} \]
\[ + (1_p \otimes (v_\theta' \otimes 1_k)) \frac{\partial^2 \text{vec} G_\theta}{\partial \theta \partial \theta'} \]
\[ + \left( \frac{\partial \text{vec} G_\theta}{\partial \theta'} \right)' \frac{\partial \text{vec} v_\theta}{\partial \theta'}. \]

(B.29)

and

\[ \frac{\partial \text{vec} \Omega_\theta^{-1}}{\partial \theta'} = -(\Omega_\theta^{-1} \otimes \Omega_\theta^{-1}) \frac{\partial \text{vec} \Omega_\theta}{\partial \theta'}. \]

(B.30)

and

\[ \text{d} (\Omega_\theta^{-1} u_\theta u_\theta' \Omega_\theta^{-1}) = -\Omega_\theta^{-1} (\text{d} \Omega_\theta) \Omega_\theta^{-1} u_\theta u_\theta' \Omega_\theta + \Omega_\theta^{-1} (\text{d} u_\theta) \mu_\theta \Omega_\theta^{-1} \]
\[ + \Omega_\theta^{-1} u_\theta (\text{d} u_\theta') \Omega_\theta^{-1} - \Omega_\theta^{-1} u_\theta u_\theta' \Omega_\theta (\text{d} \Omega_\theta) \Omega_\theta^{-1}. \]

(B.31)
By using the identity $\text{vec}(ABC) = (C' \otimes A) \text{vec}(B)$ several times we obtain

$$
\frac{\partial \text{vec}}{\partial \theta'} \Omega_{\theta}^{-1} u_{\theta} u_{\theta}' \Omega_{\theta}^{-1} = - \left( [\Omega_{\theta}^{-1} u_{\theta} u_{\theta}' \Omega_{\theta}^{-1}]' \otimes \Omega_{\theta}^{-1} \right) \frac{\partial \text{vec} \Omega_{\theta}}{\partial \theta'} \quad (B.33)
$$

$$
- \left( [u_{\theta}' \Omega_{\theta}^{-1}]' \otimes \Omega_{\theta}^{-1} \right) \frac{\partial \text{vec} \xi_{\theta}}{\partial \theta'} \quad (B.34)
$$

$$
- (\Omega_{\theta}^{-1} \otimes [\Omega_{\theta}^{-1} u_{\theta}]) \frac{\partial \text{vec} \xi_{\theta}}{\partial \theta'} \quad (B.35)
$$

$$
- (\Omega_{\theta}^{-1} \otimes [\Omega_{\theta}^{-1} u_{\theta} u_{\theta}' \Omega_{\theta}^{-1}]) \frac{\partial \text{vec} \Omega_{\theta}}{\partial \theta'} \quad (B.36)
$$

and the second order derivative of the log-likelihood (B.21) now follows from applying the product rule with (B.28), (B.29), (B.30) and (B.33).
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