

# Package ‘lava’

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**Type** Package

**Title** Latent Variable Models

**Version** 1.9.0

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**Description** A general implementation of Structural Equation Models with latent variables (MLE, 2SLS, and composite likelihood estimators) with both continuous, censored, and ordinal outcomes (Holst and Budtz-Joergensen (2013) <doi:10.1007/s00180-012-0344-y>). Mixture latent variable models and non-linear latent variable models (Holst and Budtz-Joergensen (2020) <doi:10.1093/biostatistics/kxy082>). The package also provides methods for graph exploration (d-separation, back-door criterion), simulation of general non-linear latent variable models, and estimation of influence functions for a broad range of statistical models.

**URL** <https://kkholst.github.io/lava/>, <https://github.com/kkholst/lava>

**BugReports** <https://github.com/kkholst/lava/issues>

**License** Apache License (== 2.0)

**LazyLoad** yes

**Depends** R (>= 3.5.0)

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---

addvar	<i>Add variable to (model) object</i>
--------	---------------------------------------

---

**Description**

Generic method for adding variables to model object

**Usage**

```
addvar(x, ...)
```

**Arguments**

x	Model object
...	Additional arguments

**Author(s)**

Klaus K. Holst

---

backdoor	<i>Backdoor criterion</i>
----------	---------------------------

---

**Description**

Check backdoor criterion of a lvm object

**Usage**

```
backdoor(object, f, cond, ..., return.graph = FALSE)
```

**Arguments**

object	lvm object
f	formula. Conditioning, z, set can be given as $y \sim x   z$
cond	Vector of variables to condition on
...	Additional arguments to lower level functions
return.graph	Return moral ancestral graph with z and effects from x removed

**Examples**

```
m <- lvm(y~c2,c2~c1,x~c1,m1~x,y~m1, v1~c3, x~c3,v1~y,
         x~z1, z2~z1, z2~z3, y~z3+z2+g1+g2+g3)
ll <- backdoor(m, y~x)
backdoor(m, y~x|c1+z1+g1)
```

baptize *Label elements of object*

---

**Description**

Generic method for labeling elements of an object

**Usage**

```
baptize(x, ...)
```

**Arguments**

x	Object
...	Additional arguments

**Author(s)**

Klaus K. Holst

---

binomial.rd *Define constant risk difference or relative risk association for binary exposure*

---

**Description**

Set up model as defined in Richardson, Robins and Wang (2017).

**Usage**

```
binomial.rd(  
  x,  
  response,  
  exposure,  
  target.model,  
  nuisance.model,  
  exposure.model = binomial.lvm(),  
  ...  
)
```

**Arguments**

x	model
response	response variable (character or formula)
exposure	exposure variable (character or formula)
target.model	variable defining the linear predictor for the target model
nuisance.model	variable defining the linear predictor for the nuisance model
exposure.model	model for exposure (default binomial logit link)
...	additional arguments to lower level functions

---

blockdiag	<i>Combine matrices to block diagonal structure</i>
-----------	---

---

**Description**

Combine matrices to block diagonal structure

**Usage**

```
blockdiag(x, ..., pad = 0)
```

**Arguments**

x	matrix
...	additional matrices
pad	value outside block-diagonal

**Author(s)**

Klaus K. Holst

**Examples**

```
A <- diag(3)+1
blockdiag(A,A,A,pad=NA)
```

bmd

*Longitudinal Bone Mineral Density Data (Wide format)*

---

**Description**

Bone Mineral Density Data consisting of 112 girls randomized to receive calcium or placebo. Longitudinal measurements of bone mineral density ( $\text{g/cm}^2$ ) measured approximately every 6th month in 3 years.

**Format**

data.frame

**Source**

Vonesh & Chinchilli (1997), Table 5.4.1 on page 228.

**See Also**calcium

---

bmidata

*Data*

---

**Description**

Description

**Format**data.frame

---

bootstrap

*Generic bootstrap method*

---

**Description**

Generic method for calculating bootstrap statistics

**Usage**

```
bootstrap(x, ...)
```

**Arguments**

x                    Model object  
 ...                  Additional arguments

**Author(s)**

Klaus K. Holst

**See Also**

bootstrap.lvm bootstrap.lvmfit

---

bootstrap.lvm	<i>Calculate bootstrap estimates of a lvm object</i>
---------------	--

---

**Description**

Draws non-parametric bootstrap samples

**Usage**

```
## S3 method for class 'lvm'
bootstrap(x,R=100,data,fun=NULL,control=list(),
          p, parametric=FALSE, bollenstine=FALSE,
          constraints=TRUE,sd=FALSE, mc.cores,
          future.args=list(future.seed=TRUE),
          ...)

## S3 method for class 'lvmfit'
bootstrap(x,R=100,data=model.frame(x),
          control=list(start=coef(x)),
          p=coef(x), parametric=FALSE, bollenstine=FALSE,
          estimator=x$estimator,weights=Weights(x),...)
```

**Arguments**

x                    lvm-object.  
 R                    Number of bootstrap samples  
 data                The data to resample from  
 fun                 Optional function of the (bootstrapped) model-fit defining the statistic of interest  
 control             Options to the optimization routine  
 p                    Parameter vector of the null model for the parametric bootstrap  
 parametric         If TRUE a parametric bootstrap is calculated. If FALSE a non-parametric (row-sampling) bootstrap is computed.

bollenstine	Bollen-Stine transformation (non-parametric bootstrap) for bootstrap hypothesis testing.
constraints	Logical indicating whether non-linear parameter constraints should be included in the bootstrap procedure
sd	Logical indicating whether standard error estimates should be included in the bootstrap procedure
mc.cores	Optional number of cores for parallel computing. If omitted future.apply will be used (see future::plan)
future.args	arguments to future.apply::future_lapply
...	Additional arguments, e.g. choice of estimator.
estimator	String defining estimator, e.g. 'gaussian' (see estimator)
weights	Optional weights matrix used by estimator

**Value**

A bootstrap.lvm object.

**Author(s)**

Klaus K. Holst

**See Also**

[confint.lvmfit](#)

**Examples**

```
m <- lvm(y~x)
d <- sim(m,100)
e <- estimate(lvm(y~x), data=d)
## Reduce Ex.Timings
B <- bootstrap(e,R=50,mc.cores=1)
B
```

---

brisa

*Simulated data*

---

**Description**

Simulated data

**Format**

data.frame

**Source**

Simulated

---

By *Apply a Function to a Data Frame Split by Factors*

---

**Description**

Apply a Function to a Data Frame Split by Factors

**Usage**

```
By(x, INDICES, FUN, COLUMNS, array = FALSE, ...)
```

**Arguments**

x	Data frame
INDICES	Indices (vector or list of indices, vector of column names, or formula of column names)
FUN	A function to be applied to data frame subsets of 'data'.
COLUMNS	(Optional) subset of columns of x to work on
array	if TRUE an array/matrix is always returned
...	Additional arguments to lower-level functions

**Details**

Simple wrapper of the 'by' function

**Author(s)**

Klaus K. Holst

**Examples**

```
By(datasets::CO2, ~Treatment+Type, colMeans, ~conc)
By(datasets::CO2, ~Treatment+Type, colMeans, ~conc+uptake)
```

---

calcium *Longitudinal Bone Mineral Density Data*

---

**Description**

Bone Mineral Density Data consisting of 112 girls randomized to receive calcium or placebo. Longitudinal measurements of bone mineral density (g/cm<sup>2</sup>) measured approximately every 6th month in 3 years.

**Format**

A data.frame containing 560 (incomplete) observations. The 'person' column defines the individual girls of the study with measurements at visiting times 'visit', and age in years 'age' at the time of visit. The bone mineral density variable is 'bmd' (g/cm<sup>2</sup>).

**Source**

Vonesh & Chinchilli (1997), Table 5.4.1 on page 228.

---

cancel	<i>Generic cancel method</i>
--------	------------------------------

---

**Description**

Generic cancel method

**Usage**

cancel(x, ...)

**Arguments**

x	Object
...	Additional arguments

**Author(s)**

Klaus K. Holst

---

children	<i>Extract children or parent elements of object</i>
----------	--

---

**Description**

Generic method for memberships from object (e.g. a graph)

**Usage**

children(object, ...)

**Arguments**

object	Object
...	Additional arguments

**Author(s)**

Klaus K. Holst

---

click *Identify points on plot*

---

### Description

Extension of the identify function

### Usage

```
## Default S3 method:  
click(x, y=NULL, label=TRUE, n=length(x), pch=19, col="orange", cex=3, ...)  
idplot(x, y, ..., id=list(), return.data=FALSE)
```

### Arguments

x	X coordinates
...	Additional arguments parsed to plot function
y	Y coordinates
label	Should labels be added?
n	Max number of inputs to expect
pch	Symbol
col	Colour
cex	Size
id	List of arguments parsed to click function
return.data	Boolean indicating if selected points should be returned

### Details

For the usual 'X11' device the identification process is terminated by pressing any mouse button other than the first. For the 'quartz' device the process is terminated by pressing either the pop-up menu equivalent (usually second mouse button or 'Ctrl'-click) or the 'ESC' key.

### Author(s)

Klaus K. Holst

### See Also

[idplot](#), [identify](#)

**Examples**

```

if (interactive()) {
  n <- 10; x <- seq(n); y <- runif(n)
  plot(y ~ x); click(x,y)

  data(iris)
  l <- lm(Sepal.Length ~ Sepal.Width*Species,iris)
  res <- plotConf(l,var2="Species")## ylim=c(6,8), xlim=c(2.5,3.3))
  with(res, click(x,y))

  with(iris, idplot(Sepal.Length,Petal.Length))
}

```

---

closed_testing	<i>Closed testing procedure</i>
----------------	---------------------------------

---

**Description**

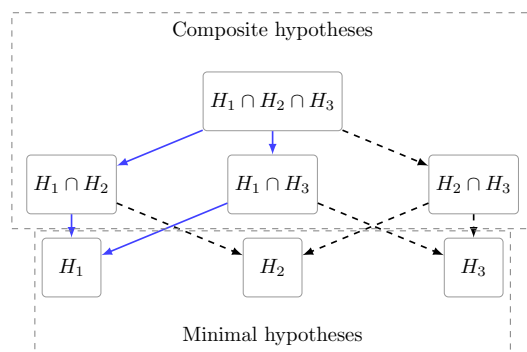
Given  $p$  hypotheses  $H_1, \dots, H_p$  all  $2^p - 1$  intersection hypotheses are calculated and adjusted p-values are obtained for  $H_j$  is calculated as the max p-value of all intersection hypotheses containing  $H_j$ . Example, for  $p=3$ , the adjusted p-value for  $H_1$  will be obtained from  $\{(H_1, H_2, H_3), (H_1, H_2), (H_1, H_3), (H_1)\}$ .

**Usage**

```
closed_testing(object, test = test_wald, ...)
```

**Arguments**

object	estimate object
test	function that conducts hypothesis test. See details below.
...	Additional arguments passed to test

**Details**

The function test should be a function function(object, index, ...) which as its first argument takes an estimate object and with an argument index which is a integer vector specifying

which subcomponents of object to test. The ellipsis argument can be any other arguments used in the test function. The function `test_wald()` is an example of valid test function (which has an additional argument `null` in reference to the above mentioned ellipsis arguments).

## References

Marcus, R; Peritz, E; Gabriel, KR (1976). "On closed testing procedures with special reference to ordered analysis of variance". *Biometrika*. 63 (3): 655–660.

## Examples

```
m <- lvm()
regression(m, c(y1,y2,y3,y4)~x) <- c(0, 0.25, 0, 0.25)
regression(m, to=endogenous(m), from="u") <- 1
variance(m,endogenous(m)) <- 1
set.seed(1)
d <- sim(m, 200)
l1 <- lm(y1~x,d)
l2 <- lm(y2~x,d)
l3 <- lm(y3~x,d)
l4 <- lm(y4~x,d)

(a <- merge(l1, l2, l3, l4, subset=2))
if (requireNamespace("mets",quietly=TRUE)) {
  alpha_zmax(a)
}
adj <- closed_testing(a)
adj
adj$p.value
summary(adj)
```

---

Col

*Generate a transparent RGB color*


---

## Description

This function transforms a standard color (e.g. "red") into an transparent RGB-color (i.e. `alpha-blend<1`).

## Usage

```
Col(col, alpha = 0.2, locate = 0)
```

## Arguments

<code>col</code>	color (numeric or character)
<code>alpha</code>	degree of transparency (0-1)
<code>locate</code>	Choose colour (with mouse)

**Details**

This only works for certain graphics devices (Cairo-X11 (x11 as of R>=2.7), quartz, pdf, ...).

**Value**

A character vector with elements of 7 or 9 characters, # followed by the red, blue, green and optionally alpha values in hexadecimal (after rescaling to '0 ... 255').

**Author(s)**

Klaus K. Holst

**Examples**

```
plot(runif(1000),cex=runif(1000,0,4),
     col=Col(c("darkblue", "orange"),0.5),pch=16)
```

---

colorbar

*Add color-bar to plot*

---

**Description**

Add color-bar to plot

**Usage**

```
colorbar(
  clut = Col(rev(rainbow(11, start = 0, end = 0.69))), alpha),
  x.range = c(-0.5, 0.5),
  y.range = c(-0.1, 0.1),
  values = seq(clut),
  digits = 2,
  label.offset,
  srt = 45,
  cex = 0.5,
  border = NA,
  alpha = 0.5,
  position = 1,
  direction = c("horizontal", "vertical"),
  ...
)
```

**Arguments**

clut	Color look-up table
x.range	x range
y.range	y range
values	label values
digits	number of digits
label.offset	label offset
srt	rotation of labels
cex	text size
border	border of color bar rectangles
alpha	Alpha (transparency) level 0-1
position	Label position left/bottom (1) or top/right (2) or no text (0)
direction	horizontal or vertical color bars
...	additional low level arguments (i.e. parsed to text)

**Examples**

```
## Not run:
plotNeuro(x, roi=R, mm=-18, range=5)
colorbar(clut=Col(rev(rainbow(11, start=0, end=0.69)), 0.5),
         x=c(-40, 40), y.range=c(84, 90), values=c(-5:5))

colorbar(clut=Col(rev(rainbow(11, start=0, end=0.69)), 0.5),
         x=c(-10, 10), y.range=c(-100, 50), values=c(-5:5),
         direction="vertical", border=1)

## End(Not run)
```

---

Combine

*Report estimates across different models*


---

**Description**

Report estimates across different models

**Usage**

```
Combine(x, ...)
```

**Arguments**

x	list of model objects
...	additional arguments to lower-level functions

**Author(s)**

Klaus K. Holst

**Examples**

```

data(serotonin)
m1 <- lm(cau ~ age*gene1 + age*gene2,data=serotonin)
m2 <- lm(cau ~ age + gene1,data=serotonin)
m3 <- lm(cau ~ age*gene2,data=serotonin)

Combine(list(A=m1,B=m2,C=m3),fun=function(x)
  c("_____"="","R2=" %+%format(summary(x)$r.squared,digits=2)))

```

---

 commutation

*Finds the unique commutation matrix*


---

**Description**

Finds the unique commutation matrix  $K$ :  $Kvec(A) = vec(A^t)$

**Usage**

```
commutation(m, n = m)
```

**Arguments**

m	rows
n	columns

**Author(s)**

Klaus K. Holst

---

 compare

*Statistical tests*


---

**Description**

Performs Likelihood-ratio, Wald and score tests

**Usage**

```
compare(object, ...)
```

**Arguments**

object            lvmfit-object  
 ...              Additional arguments to low-level functions

**Value**

Matrix of test-statistics and p-values

**Author(s)**

Klaus K. Holst

**See Also**

[modelsearch](#), [equivalence](#)

**Examples**

```
m <- lvm();
regression(m) <- c(y1,y2,y3) ~ eta; latent(m) <- ~eta
regression(m) <- eta ~ x
m2 <- regression(m, c(y3,eta) ~ x)
set.seed(1)
d <- sim(m,1000)
e <- estimate(m,d)
e2 <- estimate(m2,d)

compare(e)

compare(e,e2) ## LRT, H0: y3<-x=0
compare(e,scoretest=y3~x) ## Score-test, H0: y3~x=0
compare(e2,par=c("y3~x")) ## Wald-test, H0: y3~x=0

B <- diag(2); colnames(B) <- c("y2~eta","y3~eta")
compare(e2,contrast=B,null=c(1,1))

B <- rep(0,length(coef(e2))); B[1:3] <- 1
compare(e2,contrast=B)

compare(e,scoretest=list(y3~x,y2~x))
```

**Description**

Estimate parameters in a probit latent variable model via a composite likelihood decomposition.

**Usage**

```
complik(
  x,
  data,
  k = 2,
  type = c("all", "nearest"),
  pairlist,
  messages = 0,
  estimator = "normal",
  quick = FALSE,
  ...
)
```

**Arguments**

x	lvm-object
data	data.frame
k	Size of composite groups
type	Determines number of groups. With type="nearest" (default) only neighboring items will be grouped, e.g. for k=2 (y1,y2),(y2,y3),... With type="all" all combinations of size k are included
pairlist	A list of indices specifying the composite groups. Optional argument which overrides k and type but gives complete flexibility in the specification of the composite likelihood
messages	Control amount of messages printed
estimator	Model (pseudo-likelihood) to use for the pairs/groups
quick	If TRUE the parameter estimates are calculated but all additional information such as standard errors are skipped
...	Additional arguments parsed on to lower-level functions

**Value**

An object of class `estimate.complik` inheriting methods from `lvm`

**Author(s)**

Klaus K. Holst

**See Also**

`estimate`

**Examples**

```

m <- lvm(c(y1,y2,y3)~b*x+1*u[0],latent=~u)
ordinal(m,K=2) <- ~y1+y2+y3
d <- sim(m,50,seed=1)
if (requireNamespace("mets", quietly=TRUE)) {
  e1 <- complik(m,d,control=list(trace=1),type="all")
}

```

---

confband	<i>Add Confidence limits bar to plot</i>
----------	--

---

**Description**

Add Confidence limits bar to plot

**Usage**

```

confband(
  x,
  lower,
  upper,
  center = NULL,
  line = TRUE,
  delta = 0.07,
  centermark = 0.03,
  pch,
  blank = TRUE,
  vert = TRUE,
  polygon = FALSE,
  step = FALSE,
  ...
)

```

**Arguments**

x	Position (x-coordinate if vert=TRUE, y-coordinate otherwise)
lower	Lower limit (if NULL no limits is added, and only the center is drawn (if not NULL))
upper	Upper limit
center	Center point
line	If FALSE do not add line between upper and lower bound
delta	Length of limit bars
centermark	Length of center bar
pch	Center symbol (if missing a line is drawn)
blank	If TRUE a white ball is plotted before the center is added to the plot

vert	If TRUE a vertical bar is plotted. Otherwise a horizontal bar is used
polygon	If TRUE polygons are added between 'lower' and 'upper'.
step	Type of polygon (step-function or piecewise linear)
...	Additional low level arguments (e.g. col, lwd, lty,...)

**Author(s)**

Klaus K. Holst

**See Also**

confband

**Examples**

```

plot(0,0,type="n",xlab="",ylab="")
confband(0.5,-0.5,0.5,0,col="darkblue")
confband(0.8,-0.5,0.5,0,col="darkred",vert=FALSE,pch=1,cex=1.5)

set.seed(1)
K <- 20
est <- rnorm(K)
se <- runif(K,0.2,0.4)
x <- cbind(est,est-2*se,est+2*se,runif(K,0.5,2))
x[c(3:4,10:12),] <- NA
rownames(x) <- unlist(lapply(letters[seq(K)],function(x) paste(rep(x,4),collapse="")))
rownames(x)[which(is.na(est))] <- ""
signif <- sign(x[,2])==sign(x[,3])
forestplot(x,text.right=FALSE)
forestplot(x[, -4],sep=c(2,15),col=signif+1,box1=TRUE,delta=0.2,pch=16,cex=1.5)
forestplot(x,vert=TRUE,text=FALSE)
forestplot(x,vert=TRUE,text=FALSE,pch=NA)
##forestplot(x,vert=TRUE,text.vert=FALSE)
##forestplot(val,vert=TRUE,add=TRUE)

z <- seq(10)
zu <- c(z[-1],10)
plot(z,type="n")
confband(z,zu,rep(0,length(z)),col=Col("darkblue"),polygon=TRUE,step=TRUE)
confband(z,zu,zu-2,col=Col("darkred"),polygon=TRUE,step=TRUE)

z <- seq(0,1,length.out=100)
plot(z,z,type="n")
confband(z,z,z^2,polygon="TRUE",col=Col("darkblue"))

set.seed(1)
k <- 10
x <- seq(k)
est <- rnorm(k)
sd <- runif(k)
val <- cbind(x,est,est-sd,est+sd)

```

```

par(mfrow=c(1,2))
plot(0,type="n",xlim=c(0,k+1),ylim=range(val[,-1]),axes=FALSE,xlab="",ylab="")
axis(2)
confband(val[,1],val[,3],val[,4],val[,2],pch=16,cex=2)
plot(0,type="n",ylim=c(0,k+1),xlim=range(val[,-1]),axes=FALSE,xlab="",ylab="")
axis(1)
confband(val[,1],val[,3],val[,4],val[,2],pch=16,cex=2,vert=FALSE)

x <- seq(0, 3, length.out=20)
y <- cos(x)
yl <- y - 1
yu <- y + 1
plot_region(x, y, yl, yu)
plot_region(x, y, yl, yu, type='s', col="darkblue", add=TRUE)

```

---

confint.lvmfit

*Calculate confidence limits for parameters*


---

## Description

Calculate Wald og Likelihood based (profile likelihood) confidence intervals

## Usage

```

## S3 method for class 'lvmfit'
confint(
  object,
  parm = seq_len(length(coef(object))),
  level = 0.95,
  profile = FALSE,
  curve = FALSE,
  n = 20,
  interval = NULL,
  lower = TRUE,
  upper = TRUE,
  ...
)

```

## Arguments

object	lvm-object.
parm	Index of which parameters to calculate confidence limits for.
level	Confidence level
profile	Logical expression defining whether to calculate confidence limits via the profile log likelihood
curve	if FALSE and profile is TRUE, confidence limits are returned. Otherwise, the profile curve is returned.

n	Number of points to evaluate profile log-likelihood in over the interval defined by interval
interval	Interval over which the profiling is done
lower	If FALSE the lower limit will not be estimated (profile intervals only)
upper	If FALSE the upper limit will not be estimated (profile intervals only)
...	Additional arguments to be passed to the low level functions

### Details

Calculates either Wald confidence limits:

$$\hat{\theta} \pm z_{\alpha/2} * \hat{\sigma}_{\hat{\theta}}$$

or profile likelihood confidence limits, defined as the set of value  $\tau$ :

$$\log Lik(\hat{\theta}_{\tau}, \tau) - \log Lik(\hat{\theta}) < q_{\alpha}/2$$

where  $q_{\alpha}$  is the  $\alpha$  fractile of the  $\chi_1^2$  distribution, and  $\hat{\theta}_{\tau}$  are obtained by maximizing the log-likelihood with tau being fixed.

### Value

A 2xp matrix with columns of lower and upper confidence limits

### Author(s)

Klaus K. Holst

### See Also

[bootstrap{lvm}](#)

### Examples

```
m <- lvm(y~x)
d <- sim(m,100)
e <- estimate(lvm(y~x), d)
confint(e,3,profile=TRUE)
confint(e,3)
## Reduce Ex.timings
B <- bootstrap(e,R=50)
B
```

---

confpred	<i>Conformal prediction</i>
----------	-----------------------------

---

**Description**

Conformal predictions using locally weighted conformal inference with a split-conformal algorithm

**Usage**

```
confpred(object, data, newdata = data, alpha = 0.05, mad, ...)
```

**Arguments**

object	Model object (lm, glm or similar with predict method) or formula (lm)
data	data.frame
newdata	New data.frame to make predictions for
alpha	Level of prediction interval
mad	Conditional model (formula) for the MAD (locally-weighted CP)
...	Additional arguments to lower level functions

**Value**

data.frame with fitted (fit), lower (lwr) and upper (upr) predictions bands.

**Examples**

```
set.seed(123)
n <- 200
x <- seq(0,6,length.out=n)
delta <- 3
ss <- exp(-1+1.5*cos((x-delta)))
ee <- rnorm(n,sd=ss)
y <- (x-delta)+3*cos(x+4.5-delta)+ee
d <- data.frame(y=y,x=x)

newd <- data.frame(x=seq(0,6,length.out=50))
cc <- confpred(lm(y~splines::ns(x,knots=c(1,3,5)),data=d), data=d, newdata=newd)
if (interactive()) {
plot(y~x,pch=16,col=lava::Col("black"),ylim=c(-10,10),xlab="X",ylab="Y")
with(cc,
      lava::confband(newd$x,lwr,upr,fit,
                    lwd=3,polygon=TRUE,col=Col("blue"),border=FALSE))
}
```

---

constrain<-                    *Add non-linear constraints to latent variable model*

---

### Description

Add non-linear constraints to latent variable model

### Usage

```
## Default S3 replacement method:
constrain(x,par,args,endogenous=TRUE,...) <- value

## S3 replacement method for class 'multigroup'
constrain(x,par,k=1,...) <- value

constraints(object,data=model.frame(object),vcov=object$vcov,level=0.95,
            p=pars.default(object),k,idx,...)
```

### Arguments

x	lvm-object
...	Additional arguments to be passed to the low level functions
value	Real function taking args as a vector argument
par	Name of new parameter. Alternatively a formula with lhs specifying the new parameter and the rhs defining the names of the parameters or variable names defining the new parameter (overruling the args argument).
args	Vector of variables names or parameter names that are used in defining par
endogenous	TRUE if variable is endogenous (sink node)
k	For multigroup models this argument specifies which group to add/extract the constraint
object	lvm-object
data	Data-row from which possible non-linear constraints should be calculated
vcov	Variance matrix of parameter estimates
level	Level of confidence limits
p	Parameter vector
idx	Index indicating which constraints to extract

### Details

Add non-linear parameter constraints as well as non-linear associations between covariates and latent or observed variables in the model (non-linear regression).

As an example we will specify the follow multiple regression model:

$$E(Y|X_1, X_2) = \alpha + \beta_1 X_1 + \beta_2 X_2$$

$$V(Y|X_1, X_2) = v$$

which is defined (with the appropriate parameter labels) as

```
m <- lvm(y ~ f(x, beta1) + f(x, beta2))
```

```
intercept(m) <- y ~ f(alpha)
```

```
covariance(m) <- y ~ f(v)
```

The somewhat strained parameter constraint

$$v = \frac{(\beta_1 - \beta_2)^2}{\alpha}$$

can then specified as

```
constrain(m, v ~ beta1 + beta2 + alpha) <- function(x) (x[1]-x[2])^2/x[3]
```

A subset of the arguments `args` can be covariates in the model, allowing the specification of non-linear regression models. As an example the non-linear regression model

$$E(Y | X) = \nu + \Phi(\alpha + \beta X)$$

where  $\Phi$  denotes the standard normal cumulative distribution function, can be defined as

```
m <- lvm(y ~ f(x, 0)) # No linear effect of x
```

Next we add three new parameters using the parameter assignment function:

```
parameter(m) <- ~nu+alpha+beta
```

The intercept of  $Y$  is defined as  $\mu$

```
intercept(m) <- y ~ f(mu)
```

And finally the newly added intercept parameter  $\mu$  is defined as the appropriate non-linear function of  $\alpha$ ,  $\nu$  and  $\beta$ :

```
constrain(m, mu ~ x + alpha + nu) <- function(x) pnorm(x[1]*x[2])+x[3]
```

The `constrain` function can be used to show the estimated non-linear parameter constraints of an estimated model object (`lvmfit` or `multigroupfit`). Calling `constrain` with no additional arguments beyond `x` will return a list of the functions and parameter names defining the non-linear restrictions.

The gradient function can optionally be added as an attribute `grad` to the return value of the function defined by `value`. In this case the analytical derivatives will be calculated via the chain rule when evaluating the corresponding score function of the log-likelihood. If the gradient attribute is omitted the chain rule will be applied on a numeric approximation of the gradient.

## Value

A `lvm` object.

## Author(s)

Klaus K. Holst

**See Also**

[regression](#), [intercept](#), [covariance](#)

**Examples**

```
#####
### Non-linear parameter constraints 1
#####
m <- lvm(y ~ f(x1,gamma)+f(x2,beta))
covariance(m) <- y ~ f(v)
d <- sim(m,100)
m1 <- m; constrain(m1,beta ~ v) <- function(x) x^2
## Define slope of x2 to be the square of the residual variance of y
## Estimate both restricted and unrestricted model
e <- estimate(m,d,control=list(method="NR"))
e1 <- estimate(m1,d)
p1 <- coef(e1)
p1 <- c(p1[1:2],p1[3]^2,p1[3])
## Likelihood of unrestricted model evaluated in MLE of restricted model
logLik(e,p1)
## Likelihood of restricted model (MLE)
logLik(e1)

#####
### Non-linear regression
#####

## Simulate data
m <- lvm(c(y1,y2)~f(x,0)+f(eta,1))
latent(m) <- ~eta
covariance(m,~y1+y2) <- "v"
intercept(m,~y1+y2) <- "mu"
covariance(m,~eta) <- "zeta"
intercept(m,~eta) <- 0
set.seed(1)
d <- sim(m,100,p=c(v=0.01,zeta=0.01))[,manifest(m)]
d <- transform(d,
               y1=y1+2*pnorm(2*x),
               y2=y2+2*pnorm(2*x))

## Specify model and estimate parameters
constrain(m, mu ~ x + alpha + nu + gamma) <- function(x) x[4]*pnorm(x[3]+x[1]*x[2])
## Reduce Ex.Timings
e <- estimate(m,d,control=list(trace=1,constrain=TRUE))
constraints(e,data=d)
## Plot model-fit
plot(y1~x,d,pch=16); points(y2~x,d,pch=16,col="gray")
x0 <- seq(-4,4,length.out=100)
lines(x0,coef(e)["nu"] + coef(e)["gamma"]*pnorm(coef(e)["alpha"]*x0))

#####
```

```

### Multigroup model
#####
### Define two models
m1 <- lvm(y ~ f(x,beta)+f(z,beta2))
m2 <- lvm(y ~ f(x,psi) + z)
### And simulate data from them
d1 <- sim(m1,500)
d2 <- sim(m2,500)
### Add 'non'-linear parameter constraint
constrain(m2,psi ~ beta2) <- function(x) x
## Add parameter beta2 to model 2, now beta2 exists in both models
parameter(m2) <- ~ beta2
ee <- estimate(list(m1,m2),list(d1,d2),control=list(method="NR"))
summary(ee)

m3 <- lvm(y ~ f(x,beta)+f(z,beta2))
m4 <- lvm(y ~ f(x,beta2) + z)
e2 <- estimate(list(m3,m4),list(d1,d2),control=list(method="NR"))
e2

```

---

contr *Create contrast matrix*

---

## Description

Create contrast matrix typically for use with 'estimate' (Wald tests).

## Usage

```
contr(p, n, diff = TRUE, ...)
```

## Arguments

p	index of non-zero entries (see example)
n	Total number of parameters (if omitted the max number in p will be used)
diff	If FALSE all non-zero entries are +1, otherwise the second non-zero element in each row will be -1.
...	Additional arguments to lower level functions

## Examples

```

contr(2,n=5)
contr(as.list(2:4),n=5)
contr(list(1,2,4),n=5)
contr(c(2,3,4),n=5)
contr(list(c(1,3),c(2,4)),n=5)
contr(list(c(1,3),c(2,4),5))

parsedesign(c("aa","b","c"),"?","?",diff=c(FALSE,TRUE))

```

```
## All pairs comparisons:
pdiff <- function(n) lava::contr(lapply(seq(n-1), function(x) seq(x, n)))
pdiff(4)
```

---

correlation	<i>Generic method for extracting correlation coefficients of model object</i>
-------------	---

---

### Description

Generic correlation method

### Usage

```
correlation(x, ...)
```

### Arguments

x	Object
...	Additional arguments

### Author(s)

Klaus K. Holst

---

covariance	<i>Add covariance structure to Latent Variable Model</i>
------------	--

---

### Description

Define covariances between residual terms in a lvm-object.

### Usage

```
## S3 replacement method for class 'lvm'
covariance(object, var1=NULL, var2=NULL, constrain=FALSE, pairwise=FALSE,...) <- value
```

### Arguments

object	lvm-object
...	Additional arguments to be passed to the low level functions
var1	Vector of variables names (or formula)
var2	Vector of variables names (or formula) defining pairwise covariance between var1 and var2)
constrain	Define non-linear parameter constraints to ensure positive definite structure
pairwise	If TRUE and var2 is omitted then pairwise correlation is added between all variables in var1
value	List of parameter values or (if var1 is unspecified)

**Details**

The covariance function is used to specify correlation structure between residual terms of a latent variable model, using a formula syntax.

For instance, a multivariate model with three response variables,

$$Y_1 = \mu_1 + \epsilon_1$$

$$Y_2 = \mu_2 + \epsilon_2$$

$$Y_3 = \mu_3 + \epsilon_3$$

can be specified as

```
m <- lvm(~y1+y2+y3)
```

Pr. default the two variables are assumed to be independent. To add a covariance parameter  $r = cov(\epsilon_1, \epsilon_2)$ , we execute the following code

```
covariance(m) <- y1 ~ f(y2, r)
```

The special function  $f$  and its second argument could be omitted thus assigning an unique parameter the covariance between  $y_1$  and  $y_2$ .

Similarly the marginal variance of the two response variables can be fixed to be identical ( $var(Y_i) = v$ ) via

```
covariance(m) <- c(y1, y2, y3) ~ f(v)
```

To specify a completely unstructured covariance structure, we can call

```
covariance(m) <- ~y1+y2+y3
```

All the parameter values of the linear constraints can be given as the right handside expression of the assignment function `covariance<-` if the first (and possibly second) argument is defined as well.

E.g:

```
covariance(m, y1~y1+y2) <- list("a1", "b1")
```

```
covariance(m, ~y2+y3) <- list("a2", 2)
```

Defines

$$var(\epsilon_1) = a1$$

$$var(\epsilon_2) = a2$$

$$var(\epsilon_3) = 2$$

$$cov(\epsilon_1, \epsilon_2) = b1$$

Parameter constraints can be cleared by fixing the relevant parameters to NA (see also the regression method).

The function `covariance` (called without additional arguments) can be used to inspect the covariance constraints of a `lvm`-object.

**Value**

A lvm-object

**Author(s)**

Klaus K. Holst

**See Also**

[regression<-](#), [intercept<-](#), [constrain<-](#) [parameter<-](#), [latent<-](#), [cancel<-](#), [kill<-](#)

**Examples**

```
m <- lvm()
### Define covariance between residuals terms of y1 and y2
covariance(m) <- y1~y2
covariance(m) <- c(y1,y2)~f(v) ## Same marginal variance
covariance(m) ## Examine covariance structure
```

---

csplit

*Split data into folds*

---

**Description**

Split data into folds

**Usage**

```
csplit(x, p = NULL, replace = FALSE, return.index = FALSE, k = 2, ...)
```

**Arguments**

x	Data or integer (size)
p	Number of folds, or if a number between 0 and 1 is given two folds of size p and (1-p) will be returned
replace	With or with-out replacement
return.index	If TRUE index of folds are returned otherwise the actual data splits are returned (default)
k	(Optional, only used when p=NULL) number of folds without shuffling
...	additional arguments to lower-level functions

**Author(s)**

Klaus K. Holst

**Examples**

```
foldr(5,2,rep=2)
csplit(10,3)
csplit(iris[1:10,]) ## Split in two sets 1:(n/2) and (n/2+1):n
csplit(iris[1:10,],0.5)
```

---

curly

*Adds curly brackets to plot*


---

**Description**

Adds curly brackets to plot

**Usage**

```
curly(
  x,
  y,
  len = 1,
  theta = 0,
  wid,
  shape = 1,
  col = 1,
  lwd = 1,
  lty = 1,
  grid = FALSE,
  npoints = 50,
  text = NULL,
  offset = c(0.05, 0)
)
```

**Arguments**

x	center of the x axis of the curly brackets (or start end coordinates (x1,x2))
y	center of the y axis of the curly brackets (or start end coordinates (y1,y2))
len	Length of the curly brackets
theta	angle (in radians) of the curly brackets orientation
wid	Width of the curly brackets
shape	shape (curvature)
col	color (passed to lines/grid.lines)
lwd	line width (passed to lines/grid.lines)
lty	line type (passed to lines/grid.lines)
grid	If TRUE use grid graphics (compatibility with ggplot2)
npoints	Number of points used in curves
text	Label
offset	Label offset (x,y)

**Examples**

```

if (interactive()) {
plot(0,0,type="n",axes=FALSE,xlab="",ylab="")
curly(x=c(1,0),y=c(0,1),lwd=2,text="a")
curly(x=c(1,0),y=c(0,1),lwd=2,text="b",theta=pi)
curly(x=-0.5,y=0,shape=1,theta=pi,text="c")
curly(x=0,y=0,shape=1,theta=0,text="d")
curly(x=0.5,y=0,len=0.2,theta=pi/2,col="blue",lty=2)
curly(x=0.5,y=-0.5,len=0.2,theta=-pi/2,col="red",shape=1e3,text="e")
}

```

---

deprdiag

*50 patients from Monash Medical Centre, Melbourne*


---

**Description**

Diagnosis of depression (DSM-III-R MDD, Dysthymia, Adjustment Disorder with Depressed Mood, Depression NOS), Beck Depression Inventory (BDI) (Beck et al., 1961) General Health Questionnaire (GHQ) (Goldberg & Williams, 1988)

**Format**

data.frame

**Source**

Clarke, D. M., Smith, G. C., & Herrman, H. E. (1993). A comparative study of screening instruments for mental disorders in general hospital patients. *International Journal Psychiatry in Medicine*, 23, pp. 323-337.

McKenzie et al. (1996). Comparing correlated Kappas by resampling: Is one level of agreement significantly different from another? *J. Psychiat. Res.* 30 (6), pp. 483-492.

---

devcoords

*Returns device-coordinates and plot-region*


---

**Description**

Returns device-coordinates and plot-region

**Usage**

```
devcoords()
```

**Value**

A list with elements

- dev.x1: device left x-coordinate
- dev.x2: device right x-coordinate
- dev.y1: device bottom y-coordinate
- dev.y2: device top y-coordinate
- fig.x1: plot left x-coordinate
- fig.x2: plot right x-coordinate
- fig.y1: plot bottom y-coordinate
- fig.y2: plot top y-coordinate

**Author(s)**

Klaus K. Holst

---

diagtest	<i>Calculate diagnostic tests for 2x2 table</i>
----------	---

---

**Description**

Calculate prevalence, sensitivity, specificity, and positive and negative predictive values

**Usage**

```
diagtest(
  table,
  positive = 2,
  exact = FALSE,
  p0 = NA,
  confint = c("logit", "arcsin", "pseudoscore", "exact"),
  ...
)
```

**Arguments**

table	Table or (matrix/data.frame with two columns)
positive	Switch reference
exact	If TRUE exact binomial proportions CI/test will be used
p0	Optional null hypothesis (test prevalenc, sensitivity, ...)
confint	Type of confidence limits
...	Additional arguments to lower level functions

**Details**

Table should be in the format with outcome in columns and test in rows. Data.frame should be with test in the first column and outcome in the second column.

**Author(s)**

Klaus Holst

**Examples**

```
M <- as.table(matrix(c(42,12,
                      35,28),ncol=2,byrow=TRUE,
                      dimnames=list(rater=c("no", "yes"),gold=c("no", "yes"))))
diagtest(M,exact=TRUE)
```

---

dsep.lvm

*Check d-separation criterion*

---

**Description**

Check for conditional independence (d-separation)

**Usage**

```
## S3 method for class 'lvm'
dsep(object, x, cond = NULL, return.graph = FALSE, ...)
```

**Arguments**

object	lvm object
x	Variables for which to check for conditional independence
cond	Conditioning set
return.graph	If TRUE the moralized ancestral graph with the conditioning set removed is returned
...	Additional arguments to lower level functions

**Details**

The argument x can be given as a formula, e.g.  $x \sim y | z+v$  or  $\sim x+y | z+v$  With everything on the rhs of the bar defining the variables on which to condition on.

**Examples**

```

m <- lvm(x5 ~ x4+x3, x4~x3+x1, x3~x2, x2~x1)
if (interactive()) {
plot(m,layoutType='neato')
}
dsep(m,x5~x1|x2+x4)
dsep(m,x5~x1|x3+x4)
dsep(m,~x1+x2+x3|x4)

```

---

equivalence

*Identify candidates of equivalent models*


---

**Description**

Identifies candidates of equivalent models

**Usage**

```
equivalence(x, rel, tol = 0.001, k = 1, omitrel = TRUE, ...)
```

**Arguments**

x	lvmfit-object
rel	Formula or character-vector specifying two variables to omit from the model and subsequently search for possible equivalent models
tol	Define two models as empirical equivalent if the absolute difference in score test is less than tol
k	Number of parameters to test simultaneously. For equivalence the number of additional associations to be added instead of rel.
omitrel	if k greater than 1, this boolean defines whether to omit candidates containing rel from the output
...	Additional arguments to be passed to the lower-level functions

**Author(s)**

Klaus K. Holst

**See Also**

[compare](#), [modelsearch](#)

---

estimate.array	<i>Estimate parameters and influence function.</i>
----------------	--

---

**Description**

Estimate parameters for the sample mean, variance, and quantiles

**Usage**

```
## S3 method for class 'array'
estimate(x, type = "mean", probs = 0.5, ...)
```

**Arguments**

x	numeric matrix
type	target parameter ("mean", "variance", "quantile")
probs	numeric vector of probabilities (for type="quantile")
...	Additional arguments to lower level functions (i.e., stats::density.default when type="quantile")

---

estimate.default	<i>Estimation of functional of parameters</i>
------------------	---

---

**Description**

Estimation of functional of parameters. Wald tests, robust standard errors, cluster robust standard errors

**Usage**

```
## Default S3 method:
estimate(
  x = NULL,
  f = NULL,
  ...,
  data,
  id,
  iddata,
  stack = TRUE,
  average = FALSE,
  subset,
  score.deriv,
  level = 0.95,
  IC = robust,
```

```

    type = c("robust", "df", "mbn"),
    keep,
    use,
    regex = FALSE,
    ignore.case = FALSE,
    contrast,
    null,
    vcov,
    coef,
    robust = TRUE,
    df = NULL,
    print = NULL,
    labels,
    label.width,
    only.coef = FALSE,
    back.transform = NULL,
    folds = 0,
    cluster,
    R = 0,
    null.sim
)

```

### Arguments

x	model object (glm, lvmfit, ...)
f	transformation of model parameters and (optionally) data, or contrast matrix (or vector)
...	additional arguments to lower level functions
data	data.frame
id	(optional) id-variable corresponding to ic decomposition of model parameters.
iddata	(optional) id-variable for 'data'
stack	if TRUE (default) the i.i.d. decomposition is automatically stacked according to 'id'
average	if TRUE averages are calculated
subset	(optional) subset of data.frame on which to condition (logical expression or variable name)
score.deriv	(optional) derivative of mean score function
level	level of confidence limits
IC	if TRUE (default) the influence function decompositions are also returned (extract with IC method)
type	type of small-sample correction
keep	(optional) index of parameters to keep from final result
use	(optional) index of parameters to use in calculations
regex	If TRUE use regular expression (perl compatible) for keep, use arguments

ignore.case	Ignore case-sensitiveness in regular expression
contrast	(optional) Contrast matrix for final Wald test
null	(optional) null hypothesis to test
vcov	(optional) covariance matrix of parameter estimates (e.g. Wald-test)
coef	(optional) parameter coefficient
robust	if TRUE robust standard errors are calculated. If FALSE p-values for linear models are calculated from t-distribution
df	degrees of freedom (default obtained from 'df.residual')
print	(optional) print function
labels	(optional) names of coefficients
label.width	(optional) max width of labels
only.coef	if TRUE only the coefficient matrix is return
back.transform	(optional) transform of parameters and confidence intervals
folds	(optional) aggregate influence functions (divide and conquer)
cluster	(obsolete) alias for 'id'.
R	Number of simulations (simulated p-values)
null.sim	Mean under the null for simulations

### Details

influence function decomposition of estimator  $\hat{\theta}$  based on data  $Z_1, \dots, Z_n$ :

$$\sqrt{n}(\hat{\theta} - \theta) = \frac{1}{\sqrt{n}} \sum_{i=1}^n IC(Z_i; P) + o_p(1)$$

can be extracted with the IC method.

### See Also

estimate.array

### Examples

```
## Simulation from logistic regression model
m <- lvm(y~x+z);
distribution(m,y~x) <- binomial.lvm("logit")
d <- sim(m,1000)
g <- glm(y~z+x,data=d,family=binomial())
g0 <- glm(y~1,data=d,family=binomial())

## LRT
estimate(g, g0)

## Plain estimates (robust standard errors)
estimate(g)
```

```

## Testing contrasts
estimate(g, null=0)
estimate(g, rbind(c(1,1,0), c(1,0,2)))
estimate(g, rbind(c(1,1,0), c(1,0,2)), null=c(1,2))
estimate(g, 2:3) ## same as cbind(0,1,-1)
estimate(g, as.list(2:3)) ## same as rbind(c(0,1,0),c(0,0,1))
## Alternative syntax
estimate(g, "z", "z"-x, 2*z"-3*x")
estimate(g, "?") ## Wildcards
estimate(g, "*Int*", "z")
estimate(g, "1", "2"-3, null = c(0,1))
estimate(g, 2, 3)

## Usual (non-robust) confidence intervals
estimate(g, robust=FALSE)

## Transformations
estimate(g, function(p) p[1]+p[2])

## Multiple parameters
e <- estimate(g, function(p) c(p[1]+p[2], p[1]*p[2]))
e
vcov(e)

## Label new parameters
estimate(g, function(p) list("a1"=p[1]+p[2], "b1"=p[1]*p[2]))
##'
## Multiple group
m <- lvm(y~x)
m <- baptize(m)
d2 <- d1 <- sim(m,50,seed=1)
e <- estimate(list(m,m),list(d1,d2))
estimate(e) ## Wrong
ee <- estimate(e, id=rep(seq(nrow(d1)), 2)) ## Clustered
ee
estimate(lm(y~x,d1))

## Marginalize
f <- function(p,data)
  list(p0=lava:::expit(p["(Intercept)"] + p["z"]*data["z"]),
        p1=lava:::expit(p["(Intercept)"] + p["x"] + p["z"]*data["z"]))
e <- estimate(g, f, average=TRUE)
e
estimate(e,diff)
estimate(e,cbind(1,1))

## Clusters and subset (conditional marginal effects)
d$id <- rep(seq(nrow(d)/4),each=4)
estimate(g,function(p,data)
  list(p0=lava:::expit(p[1] + p["z"]*data["z"]),
        subset=d$id>0, id=d$id, average=TRUE))

## More examples with clusters:

```

```

m <- lvm(c(y1,y2,y3)~u+x)
d <- sim(m,10)
l1 <- glm(y1~x,data=d)
l2 <- glm(y2~x,data=d)
l3 <- glm(y3~x,data=d)

## Some random id-numbers
id1 <- c(1,1,4,1,3,1,2,3,4,5)
id2 <- c(1,2,3,4,5,6,7,8,1,1)
id3 <- seq(10)

## Un-stacked and stacked i.i.d. decomposition
IC(estimate(l1,id=id1,stack=FALSE))
IC(estimate(l1,id=id1))

## Combined i.i.d. decomposition
e1 <- estimate(l1,id=id1)
e2 <- estimate(l2,id=id2)
e3 <- estimate(l3,id=id3)
(a2 <- merge(e1,e2,e3))

## If all models were estimated on the same data we could use the
## syntax:
## Reduce(merge,estimate(list(l1,l2,l3)))

## Same:
IC(a1 <- merge(l1,l2,l3,id=list(id1,id2,id3)))

IC(merge(l1,l2,l3,id=TRUE)) # one-to-one (same clusters)
IC(merge(l1,l2,l3,id=FALSE)) # independence

## Monte Carlo approach, simple trend test example

m <- categorical(lvm(),~x,K=5)
regression(m,additive=TRUE) <- y~x
d <- simulate(m,100,seed=1,'y~x'=0.1)
l <- lm(y~-1+factor(x),data=d)

f <- function(x) coef(lm(x~seq_along(x)))[2]
null <- rep(mean(coef(l)),length(coef(l)))
## just need to make sure we simulate under H0: slope=0
estimate(l,f,R=1e2,null.sim=null)

estimate(l,f)

# ----- influence function calculus -----
a <- estimate(coef = c("a" = 0.5), IC = rnorm(10), id = 1:10)
b <- estimate(coef = c("b" = 0.8), IC = rnorm(10), id = 1:10)

e <- c(a, b) # merge
merge(a, b)
c(e1=a, b) # naming of par

```

```

labels(e, c("p1", "p2")) # renaming parameters
e["a"] # subset
subset(e, "a")

# pipes
# c(a, b) |>
# transform(function(x) x^2) |>
# subset("a") |>
# labels("sq")

# Parameter transformation with automatic calculation of derivatives
a * b
(3 * cos(a) / sqrt(b) + 1) / a
expit(c(a,b))
c(sum=sum(e), sum2=a+b,
  prod=prod(e), prod2=a*b)
e %*% e # inner prod.
c(1, 2) %*% e
c(pow = a^b)
a^c(0.5, 2)
c(b=e["a"] * e["b"] / a, also.b=e["b"])

B <- rbind(c(1,-1), c(1,0), c(0,1))
B %*% e
e == 1 # wald-test, null-hypothesis H0: b=1
e == c(1,2)
B %*% e == 1

```

---

estimate.lvm

*Estimation of parameters in a Latent Variable Model (lvm)*


---

## Description

Estimate parameters. MLE, IV or user-defined estimator.

## Usage

```

## S3 method for class 'lvm'
estimate(
  x,
  data = parent.frame(),
  estimator = NULL,
  control = list(),
  missing = FALSE,
  weights,
  weightsname,
  data2,
  id,
  fix,

```

```

    index = !quick,
    graph = FALSE,
    messages = lava.options()$messages,
    quick = FALSE,
    method,
    param,
    cluster,
    p,
    ...
  )

```

### Arguments

x	lvm-object
data	data.frame
estimator	String defining the estimator (see details below)
control	control/optimization parameters (see details below)
missing	Logical variable indicating how to treat missing data. Setting to FALSE leads to complete case analysis. In the other case likelihood based inference is obtained by integrating out the missing data under assumption the assumption that data is missing at random (MAR).
weights	Optional weights to used by the chosen estimator.
weightsname	Weights names (variable names of the model) in case weights was given as a vector of column names of data
data2	Optional additional dataset used by the chosen estimator.
id	Vector (or name of column in data) that identifies correlated groups of observations in the data leading to variance estimates based on a sandwich estimator
fix	Logical variable indicating whether parameter restriction automatically should be imposed (e.g. intercepts of latent variables set to 0 and at least one regression parameter of each measurement model fixed to ensure identifiability.)
index	For internal use only
graph	For internal use only
messages	Control how much information should be printed during estimation (0: none)
quick	If TRUE the parameter estimates are calculated but all additional information such as standard errors are skipped
method	Optimization method
param	set parametrization (see help(lava.options))
cluster	Obsolete. Alias for 'id'.
p	Evaluate model in parameter 'p' (no optimization)
...	Additional arguments to be passed to lower-level functions

## Details

A list of parameters controlling the estimation and optimization procedures is parsed via the `control` argument. By default Maximum Likelihood is used assuming multivariate normal distributed measurement errors. A list with one or more of the following elements is expected:

**start:** Starting value. The order of the parameters can be shown by calling `coef` (with `mean=TRUE`) on the `lvm`-object or with `plot(..., labels=TRUE)`. Note that this requires a check that it is actual the model being estimated, as `estimate` might add additional restriction to the model, e.g. through the `fix` and `exo.fix` arguments. The `lvm`-object of a fitted model can be extracted with the `Model`-function.

**starterfun:** Starter-function with syntax `function(lvm, S, mu)`. Three builtin functions are available: `startvalues`, `startvalues0`, `startvalues1`, ...

**estimator:** String defining which estimator to use (Defaults to “gaussian”)

**meanstructure** Logical variable indicating whether to fit model with meanstructure.

**method:** String pointing to alternative optimizer (e.g. `optim` to use simulated annealing).

**control:** Parameters passed to the optimizer (default `stats::nlminb`).

**tol:** Tolerance of optimization constraints on lower limit of variance parameters.

## Value

A `lvmfit`-object.

## Author(s)

Klaus K. Holst

## See Also

`estimate.default` `score`, `information`

## Examples

```
dd <- read.table(header=TRUE,
text="x1 x2 x3
  0.0 -0.5 -2.5
-0.5 -2.0  0.0
  1.0  1.5  1.0
  0.0  0.5  0.0
-2.5 -1.5 -1.0")
e <- estimate(lvm(c(x1,x2,x3)~u),dd)

## Simulation example
m <- lvm(list(y~v1+v2+v3+v4,c(v1,v2,v3,v4)~x))
covariance(m) <- v1~v2+v3+v4
dd <- sim(m,10000) ## Simulate 10000 observations from model
e <- estimate(m, dd) ## Estimate parameters
e

## Using just sufficient statistics
```

```

n <- nrow(dd)
e0 <- estimate(m,data=list(S=cov(dd)*(n-1)/n,mu=colMeans(dd),n=n))
rm(dd)

## Multiple group analysis
m <- lvm()
regression(m) <- c(y1,y2,y3)~u
regression(m) <- u~x
d1 <- sim(m,100,p=c("u,u"=1,"u~x"=1))
d2 <- sim(m,100,p=c("u,u"=2,"u~x"=-1))

mm <- baptize(m)
regression(mm,u~x) <- NA
covariance(mm,~u) <- NA
intercept(mm,~u) <- NA
ee <- estimate(list(mm,mm),list(d1,d2))

## Missing data
d0 <- makemissing(d1,cols=1:2)
e0 <- estimate(m,d0,missing=TRUE)
e0

```

---

eventTime

---

*Add an observed event time outcome to a latent variable model.*


---

## Description

For example, if the model 'm' includes latent event time variables are called 'T1' and 'T2' and 'C' is the end of follow-up (right censored), then one can specify

## Usage

```
eventTime(object, formula, eventName = "status", ...)
```

## Arguments

object	Model object
formula	Formula (see details)
eventName	Event names
...	Additional arguments to lower levels functions

## Details

```
eventTime(object=m, formula=ObsTime~min(T1=a,T2=b,C=0,"ObsEvent"))
```

when data are simulated from the model one gets 2 new columns:

- "ObsTime": the smallest of T1, T2 and C
- "ObsEvent": 'a' if T1 is smallest, 'b' if T2 is smallest and '0' if C is smallest

Note that "ObsEvent" and "ObsTime" are names specified by the user.

**Author(s)**

Thomas A. Gerds, Klaus K. Holst

**Examples**

```
# Right censored survival data without covariates
m0 <- lvm()
distribution(m0,"eventtime") <- coxWeibull.lvm(scale=1/100,shape=2)
distribution(m0,"censtime") <- coxExponential.lvm(rate=1/10)
m0 <- eventTime(m0,time~min(eventtime=1,censtime=0),"status")
sim(m0,10)

# Alternative specification of the right censored survival outcome
## eventTime(m,"Status") <- ~min(eventtime=1,censtime=0)

# Cox regression:
# lava implements two different parametrizations of the same
# Weibull regression model. The first specifies
# the effects of covariates as proportional hazard ratios
# and works as follows:
m <- lvm()
distribution(m,"eventtime") <- coxWeibull.lvm(scale=1/100,shape=2)
distribution(m,"censtime") <- coxWeibull.lvm(scale=1/100,shape=2)
m <- eventTime(m,time~min(eventtime=1,censtime=0),"status")
distribution(m,"sex") <- binomial.lvm(p=0.4)
distribution(m,"sbp") <- normal.lvm(mean=120,sd=20)
regression(m,from="sex",to="eventtime") <- 0.4
regression(m,from="sbp",to="eventtime") <- -0.01
sim(m,6)
# The parameters can be recovered using a Cox regression
# routine or a Weibull regression model. E.g.,
## Not run:
  set.seed(18)
  d <- sim(m,1000)
  library(survival)
  coxph(Surv(time,status)~sex+sbp,data=d)

  sr <- survreg(Surv(time,status)~sex+sbp,data=d)
  library(SurvRegCensCov)
  ConvertWeibull(sr)

## End(Not run)

# The second parametrization is an accelerated failure time
# regression model and uses the function weibull.lvm instead
# of coxWeibull.lvm to specify the event time distributions.
# Here is an example:

ma <- lvm()
distribution(ma,"eventtime") <- weibull.lvm(scale=3,shape=1/0.7)
distribution(ma,"censtime") <- weibull.lvm(scale=2,shape=1/0.7)
```

```

ma <- eventTime(ma,time~min(eventtime=1,censtime=0),"status")
distribution(ma,"sex") <- binomial.lvm(p=0.4)
distribution(ma,"sbp") <- normal.lvm(mean=120,sd=20)
regression(ma,from="sex",to="eventtime") <- 0.7
regression(ma,from="sbp",to="eventtime") <- -0.008
set.seed(17)
sim(ma,6)
# The regression coefficients of the AFT model
# can be transformed into log(hazard ratios):
# coef.coxWeibull = - coef.weibull / shape.weibull
## Not run:
  set.seed(17)
  da <- sim(ma,1000)
  library(survival)
  fa <- coxph(Surv(time,status)~sex+sbp,data=da)
  coef(fa)
  c(0.7,-0.008)/0.7

## End(Not run)

# The following are equivalent parametrizations
# which produce exactly the same random numbers:

model.aft <- lvm()
distribution(model.aft,"eventtime") <- weibull.lvm(intercept=-log(1/100)/2,sigma=1/2)
distribution(model.aft,"censtime") <- weibull.lvm(intercept=-log(1/100)/2,sigma=1/2)
sim(model.aft,6,seed=17)

model.aft <- lvm()
distribution(model.aft,"eventtime") <- weibull.lvm(scale=100^(1/2), shape=2)
distribution(model.aft,"censtime") <- weibull.lvm(scale=100^(1/2), shape=2)
sim(model.aft,6,seed=17)

model.cox <- lvm()
distribution(model.cox,"eventtime") <- coxWeibull.lvm(scale=1/100,shape=2)
distribution(model.cox,"censtime") <- coxWeibull.lvm(scale=1/100,shape=2)
sim(model.cox,6,seed=17)

# The minimum of multiple latent times one of them still
# being a censoring time, yield
# right censored competing risks data

mc <- lvm()
distribution(mc,~X2) <- binomial.lvm()
regression(mc) <- T1~f(X1,-.5)+f(X2,0.3)
regression(mc) <- T2~f(X2,0.6)
distribution(mc,~T1) <- coxWeibull.lvm(scale=1/100)
distribution(mc,~T2) <- coxWeibull.lvm(scale=1/100)
distribution(mc,~C) <- coxWeibull.lvm(scale=1/100)
mc <- eventTime(mc,time~min(T1=1,T2=2,C=0),"event")
sim(mc,6)

```

---

Expand

*Create a Data Frame from All Combinations of Factors*

---

### Description

Create a Data Frame from All Combinations of Factors

### Usage

```
Expand(`_data`, ...)
```

### Arguments

<code>_data</code>	Data.frame
<code>...</code>	vectors, factors or a list containing these

### Details

Simple wrapper of the 'expand.grid' function. If x is a table then a data frame is returned with one row pr individual observation.

### Author(s)

Klaus K. Holst

### Examples

```
dd <- Expand(iris, Sepal.Length=2:8, Species=c("virginica","setosa"))
summary(dd)
```

```
T <- with(warpbreaks, table(wool, tension))
Expand(T)
```

---

`fplot`*fplot*

---

**Description**

Faster plot via RGL

**Usage**

```
fplot(  
  x,  
  y,  
  z = NULL,  
  xlab,  
  ylab,  
  ...,  
  z.col = topo.colors(64),  
  data = parent.frame(),  
  add = FALSE,  
  aspect = c(1, 1),  
  zoom = 0.8  
)
```

**Arguments**

<code>x</code>	X variable
<code>y</code>	Y variable
<code>z</code>	Z variable (optional)
<code>xlab</code>	x-axis label
<code>ylab</code>	y-axis label
<code>...</code>	additional argument to lower-level plot functions
<code>z.col</code>	color (use argument <code>alpha</code> to set transparency)
<code>data</code>	<code>data.frame</code>
<code>add</code>	if TRUE use current active device
<code>aspect</code>	aspect ratio
<code>zoom</code>	zoom level

**Examples**

```
if (interactive()) {  
  data(iris)  
  fplot(Sepal.Length ~ Petal.Length+Species, data=iris, size=2, type="s")  
}
```

---

getSAS	<i>Read SAS output</i>
--------	------------------------

---

**Description**

Run SAS code like in the following:

**Usage**

```
getSAS(infile, entry = "Parameter Estimates", ...)
```

**Arguments**

infile	file (csv file generated by ODS)
entry	Name of entry to capture
...	additional arguments to lower level functions

**Details**

```
ODS CSVALL BODY="myest.csv"; proc nlmixed data=aj qpoints=2 dampstep=0.5; ... run; ODS  
CSVALL Close;
```

and read results into R with:

```
getsas("myest.csv", "Parameter Estimates")
```

**Author(s)**

Klaus K. Holst

---

gof	<i>Extract model summaries and GOF statistics for model object</i>
-----	--

---

**Description**

Calculates various GOF statistics for model object including global chi-squared test statistic and AIC. Extract model-specific mean and variance structure, residuals and various predictions.

**Usage**

```

gof(object, ...)

## S3 method for class 'lvmfit'
gof(object, chisq=FALSE, level=0.90, rmsea.threshold=0.05,all=FALSE,...)

moments(x,...)

## S3 method for class 'lvm'
moments(x, p, debug=FALSE, conditional=FALSE, data=NULL, latent=FALSE, ...)

## S3 method for class 'lvmfit'
logLik(object, p=coef(object),
        data=model.frame(object),
        model=object$estimator,
        weights=Weights(object),
        data2=object$data$data2,
        ...)

## S3 method for class 'lvmfit'
score(x, data=model.frame(x), p=pars(x), model=x$estimator,
      weights=Weights(x), data2=x$data$data2, ...)

## S3 method for class 'lvmfit'
information(x,p=pars(x),n=x$data$n,data=model.frame(x),
          model=x$estimator,weights=Weights(x), data2=x$data$data2, ...)

```

**Arguments**

object	Model object
...	Additional arguments to be passed to the low level functions
x	Model object
p	Parameter vector used to calculate statistics
data	Data.frame to use
latent	If TRUE predictions of latent variables are included in output
data2	Optional second data.frame (only for censored observations)
weights	Optional weight matrix
n	Number of observations
conditional	If TRUE the conditional moments given the covariates are calculated. Otherwise the joint moments are calculated
model	String defining estimator, e.g. "gaussian" (see estimate)
debug	Debugging only
chisq	Boolean indicating whether to calculate chi-squared goodness-of-fit (always TRUE for estimator='gaussian')

level            Level of confidence limits for RMSEA  
 rmsea.threshold    Which probability to calculate,  $\Pr(\text{RMSEA} < \text{rmsea.threshold})$   
 all              Calculate all (ad hoc) FIT indices: TLI, CFI, NFI, SRMR, ...

**Value**

A htest-object.

**Author(s)**

Klaus K. Holst

**Examples**

```
m <- lvm(list(y~v1+v2+v3+v4,c(v1,v2,v3,v4)~x))
set.seed(1)
dd <- sim(m,1000)
e <- estimate(m, dd)
gof(e,all=TRUE,rmsea.threshold=0.05,level=0.9)

set.seed(1)
m <- lvm(list(c(y1,y2,y3)~u,y1~x)); latent(m) <- ~u
regression(m,c(y2,y3)~u) <- "b"
d <- sim(m,1000)
e <- estimate(m,d)
rsq(e)
##'
rr <- rsq(e,TRUE)
rr
estimate(rr,contrast=rbind(c(1,-1,0),c(1,0,-1),c(0,1,-1)))
```

---

Graph

*Extract graph*

---

**Description**

Extract or replace graph object

**Usage**

Graph(x, ...)

Graph(x, ...) <- value

**Arguments**

x	Model object
...	Additional arguments to be passed to the low level functions
value	New graphNEL object

**Author(s)**

Klaus K. Holst

**See Also**

[Model](#)

**Examples**

```
m <- lvm(y~x)
Graph(m)
```

---

Grep

*Finds elements in vector or column-names in data.frame/matrix*

---

**Description**

Pattern matching in a vector or column names of a data.frame or matrix.

**Usage**

```
Grep(x, pattern, subset = TRUE, ignore.case = TRUE, ...)
```

**Arguments**

x	vector, matrix or data.frame.
pattern	regular expression to search for
subset	If TRUE returns subset of data.frame/matrix otherwise just the matching column names
ignore.case	Default ignore case
...	Additional arguments to 'grep'

**Value**

A data.frame with 2 columns with the indices in the first and the matching names in the second.

**Author(s)**

Klaus K. Holst

**See Also**

[grep](#), and [agrep](#) for approximate string matching.

**Examples**

```
data(iris)
head(Grep(iris,"(len)|(sp)"))
```

---

hubble	<i>Hubble data</i>
--------	--------------------

---

**Description**

Velocity (v) and distance (D) measures of 36 Type Ia super-novae from the Hubble Space Telescope

**Format**

data.frame

**Source**

Freedman, W. L., et al. 2001, *AstroPhysicalJournal*, 553, 47.

---

hubble2	<i>Hubble data</i>
---------	--------------------

---

**Description**

Hubble data

**Format**

data.frame

**See Also**

[hubble](#)

---

IC.default	<i>Extract influence function from model object</i>
------------	---

---

**Description**

Extract i.i.d. decomposition (influence function) from model object

**Usage**

```
## Default S3 method:
IC(x, bread, id = NULL, ...)
```

**Arguments**

x	model object
bread	(optional) Inverse of derivative of mean score function
id	(optional) id/cluster variable
...	additional arguments

**Examples**

```
m <- lvm(y~x+z)
distribution(m, ~y+z) <- binomial.lvm("logit")
d <- sim(m, 1e3)
g <- glm(y~x+z, data=d, family=binomial)
var_ic(IC(g))
```

---

iid	<i>Extract i.i.d. decomposition from model object</i>
-----	---

---

**Description**

This function extracts

**Usage**

```
iid(x, ...)
```

**Arguments**

x	Model object
...	Additional arguments (see the man-page of the IC method)

---

 images

---

*Organize several image calls (for visualizing categorical data)*


---

**Description**

Visualize categorical by group variable

**Usage**

```
images(
  x,
  group,
  ncol = 2,
  byrow = TRUE,
  colorbar = 1,
  colorbar.space = 0.1,
  label.offset = 0.02,
  order = TRUE,
  colorbar.border = 0,
  main,
  rowcol = FALSE,
  plotfun = NULL,
  axis1,
  axis2,
  mar,
  col = list(c("#EFF3FF", "#BDD7E7", "#6BAED6", "#2171B5"), c("#FEE5D9", "#FCAE91",
    "#FB6A4A", "#CB181D"), c("#EDF8E9", "#BAE4B3", "#74C476", "#238B45"), c("#FEEDDE",
    "#FDBE85", "#FD8D3C", "#D94701")),
  ...
)
```

**Arguments**

x	data.frame or matrix
group	group variable
ncol	number of columns in layout
byrow	organize by row if TRUE
colorbar	Add color bar
colorbar.space	Space around color bar
label.offset	label offset
order	order
colorbar.border	Add border around color bar
main	Main title

rowcol	switch rows and columns
plotfun	Alternative plot function (instead of 'image')
axis1	Axis 1
axis2	Axis 2
mar	Margins
col	Colours
...	Additional arguments to lower level graphics functions

**Author(s)**

Klaus Holst

**Examples**

```
X <- matrix(rbinom(400,3,0.5),20)
group <- rep(1:4,each=5)
images(X,colorbar=0,zlim=c(0,3))
images(X,group=group,zlim=c(0,3))
## Not run:
images(X,group=group,col=list(RColorBrewer::brewer.pal(4,"Purples"),
                             RColorBrewer::brewer.pal(4,"Greys"),
                             RColorBrewer::brewer.pal(4,"YlGn"),
                             RColorBrewer::brewer.pal(4,"PuBuGn")),colorbar=2,zlim=c(0,3))

## End(Not run)
images(list(X,X,X,X),group=group,zlim=c(0,3))
images(list(X,X,X,X),ncol=1,group=group,zlim=c(0,3))
images(list(X,X),group,axis2=c(FALSE,FALSE),axis1=c(FALSE,FALSE),
       mar=list(c(0,0,0,0),c(0,0,0,0)),yaxs="i",xaxs="i",zlim=c(0,3))
```

---

index*Generic method for extract index of an object*

---

**Description**

Generic method for extract index of an object

Generic method for setting the index of an object

**Usage**

index(x, ...)

index(x, ...) &lt;- value

**Arguments**

x	object on which to set the index
...	further arguments to be passed to or from other methods.
value	the new index

**See Also**

[index<-](#)  
[index](#)

---

index.lvm

---

*Extract the parameter indicies of a lvm object*


---

**Description**

Extracts the matrices with indices of model parameters from a latent variable model (lvm). Returns a list with

- A: Matrix with fixed parameters and ones where parameters are free
- J: Manifest variable selection matrix
- M0: Index of free regression parameters
- M1: Index of free and *unique* regression parameters
- P: Matrix with fixed variance parameters and ones where parameters are free
- P0: Index of free variance parameters
- P1: Index of free and *unique* regression parameters
- npar.var: Number of covariance parameters

**Usage**

```
## S3 method for class 'lvm'
index(x, ...)

## S3 replacement method for class 'lvm'
index(x, ...) <- value
```

**Arguments**

x	object on which to set the index
...	further arguments to be passed to or from other methods.
value	new index

**See Also**

[modelPar\(\)](#)

---

 indoorenv

*Data*


---

**Description**

Description

**Format**

data.frame

**Source**

Simulated

---

 intercept

*Fix mean parameters in 'lvm'-object*


---

**Description**

Define linear constraints on intercept parameters in a lvm-object.

**Usage**

```
## S3 replacement method for class 'lvm'
intercept(object, vars, ...) <- value
```

**Arguments**

object	lvm-object
...	Additional arguments
vars	character vector of variable names
value	Vector (or list) of parameter values or labels (numeric or character) or a formula defining the linear constraints (see also the regression or covariance methods).

**Details**

The intercept function is used to specify linear constraints on the intercept parameters of a latent variable model. As an example we look at the multivariate regression model

$$E(Y_1|X) = \alpha_1 + \beta_1 X$$

$$E(Y_2|X) = \alpha_2 + \beta_2 X$$

defined by the call

```
m <- lvm(c(y1,y2) ~ x)
```

To fix  $\alpha_1 = \alpha_2$  we call

```
intercept(m) <- c(y1,y2) ~ f(mu)
```

Fixed parameters can be reset by fixing them to NA. For instance to free the parameter restriction of  $Y_1$  and at the same time fixing  $\alpha_2 = 2$ , we call

```
intercept(m, ~y1+y2) <- list(NA,2)
```

Calling `intercept` with no additional arguments will return the current intercept restrictions of the lvm-object.

### Value

A lvm-object

### Note

Variables will be added to the model if not already present.

### Author(s)

Klaus K. Holst

### See Also

[covariance<-](#), [regression<-](#), [constrain<-](#), [parameter<-](#), [latent<-](#), [cancel<-](#), [kill<-](#)

### Examples

```
## A multivariate model
m <- lvm(c(y1,y2) ~ f(x1,beta)+x2)
regression(m) <- y3 ~ f(x1,beta)
intercept(m) <- y1 ~ f(mu)
intercept(m, ~y2+y3) <- list(2,"mu")
intercept(m) ## Examine intercepts of model (NA translates to free/unique paramete##r)
```

---

intervention.lvm

*Define intervention*

---

### Description

Define intervention in a lvm object

**Usage**

```
## S3 method for class 'lvm'
intervention(object, to, value, dist = none.lvm(), ...)
```

**Arguments**

object	lvm object
to	String defining variable or formula
value	function defining intervention
dist	Distribution
...	Additional arguments to lower level functions

**See Also**

regression lvm sim

**Examples**

```
m <- lvm(y ~ a + x, a ~ x)
distribution(m, ~a+y) <- binomial.lvm()
mm <- intervention(m, "a", value=3)
sim(mm, 10)
mm <- intervention(m, a~x, function(x) (x>0)*1)
sim(mm, 10)
```

---

Inverse

*Generalized matrix inverse*

---

**Description**

Generalized matrix inverse

**Usage**

```
Inverse(
  X,
  tol = lava.options()$itol,
  det = TRUE,
  names = !chol,
  chol = FALSE,
  symmetric = FALSE
)
```

**Arguments**

<code>X</code>	nxn matrix
<code>tol</code>	tolerance for pseudo inverse
<code>det</code>	logical, if true the determinant is returned
<code>names</code>	preserve dimnames
<code>chol</code>	use Cholesky decomposition for calculating inverse otherwise SVD
<code>symmetric</code>	set to true if matrix is symmetric

---

<code>ksmooth2</code>	<i>Plot/estimate surface</i>
-----------------------	------------------------------

---

**Description**

Plot/estimate surface

**Usage**

```
ksmooth2(
  x,
  data,
  h = NULL,
  xlab = NULL,
  ylab = NULL,
  zlab = "",
  gridsize = rep(51L, 2),
  ...
)
```

**Arguments**

<code>x</code>	formula or data
<code>data</code>	data.frame
<code>h</code>	bandwidth
<code>xlab</code>	X label
<code>ylab</code>	Y label
<code>zlab</code>	Z label
<code>gridsize</code>	grid size of kernel smoother
<code>...</code>	Additional arguments to graphics routine (persp3d or persp)

**Examples**

```

if (requireNamespace("KernSmooth")) {##'
ksmooth2(rmvn0(1e4,sigma=diag(2)*.5+.5),c(-3.5,3.5),h=1,
         rgl=FALSE,theta=30)
##'
if (interactive()) {
  ksmooth2(rmvn0(1e4,sigma=diag(2)*.5+.5),c(-3.5,3.5),h=1)
  ksmooth2(function(x,y) x^2+y^2, c(-20,20))
  ksmooth2(function(x,y) x^2+y^2, xlim=c(-5,5), ylim=c(0,10))

  f <- function(x,y) 1-sqrt(x^2+y^2)
  surface(f,xlim=c(-1,1),alpha=0.9,aspect=c(1,1,0.75))
  surface(f,xlim=c(-1,1),clut=heat.colors(128))
  ##play3d(spin3d(axis=c(0,0,1), rpm=8), duration=5)
}

if (interactive()) {
  surface(function(x) dmvn0(x,sigma=diag(2)),c(-3,3),lit=FALSE,smooth=FALSE,box=FALSE,alpha=0.8)
  surface(function(x) dmvn0(x,sigma=diag(2)),c(-3,3),box=FALSE,specular="black")##'
}

if (!inherits(try(find.package("fields"),silent=TRUE),"try-error")) {
  f <- function(x,y) 1-sqrt(x^2+y^2)
  ksmooth2(f,c(-1,1),rgl=FALSE,image=fields::image.plot)
}
}

```

---

labels<-

*Define labels of graph*


---

**Description**

Alters labels of nodes and edges in the graph of a latent variable model

**Usage**

```

## Default S3 replacement method:
labels(object, ...) <- value
## S3 replacement method for class 'lvm'
edgelabels(object, to, ...) <- value
## Default S3 replacement method:
nodecolor(object, var=vars(object),
border, labcol, shape, lwd, ...) <- value

```

**Arguments**

```

object      lvm-object.
...         Additional arguments (lwd, cex, col, labcol), border.

```

value	node label/edge label/color
to	Formula specifying outcomes and predictors defining relevant edges.
var	Formula or character vector specifying the nodes/variables to alter.
border	Colors of borders
labcol	Text label colors
shape	Shape of node
lwd	Line width of border

**Author(s)**

Klaus K. Holst

**Examples**

```

m <- lvm(c(y,v)~x+z)
regression(m) <- c(v,x)~z
labels(m) <- c(y=expression(psi), z=expression(zeta))
nodecolor(m,~y+z+x,border=c("white","white","black"),
          labcol="white", lwd=c(1,1,5),
          lty=c(1,2)) <- c("orange","indianred","lightgreen")
edgelabels(m,y~z+x, cex=c(2,1.5), col=c("orange","black"),labcol="darkblue",
           arrowhead=c("tee","dot"),
           lwd=c(3,1)) <- expression(phi,rho)
edgelabels(m,c(v,x)~z, labcol="red", cex=0.8,arrowhead="none") <- 2
if (interactive()) {
  plot(m,addstyle=FALSE)
}

m <- lvm(y~x)
labels(m) <- list(x="multiple\nlines")
if (interactive()) {
  op <- par(mfrow=c(1,2))
  plot(m,plain=TRUE)
  plot(m)
  par(op)

d <- sim(m,100)
e <- estimate(m,d)
plot(e,type="sd")
}

```

---

lava.options

*Set global options for lava*

---

**Description**

Extract and set global parameters of lava. In particular optimization parameters for the estimate function.

**Usage**

```
lava.options(...)
```

**Arguments**

```
... Arguments
```

**Details**

- param: 'relative' (factor loading and variance of one endogenous variables in each measurement model are fixed to one), 'absolute' (mean and variance of latent variables are set to 0 and 1, respectively), 'hybrid' (intercept of latent variables is fixed to 0, and factor loading of at least one endogenous variable in each measurement model is fixed to 1), 'none' (no constraints are added)
- layout: One of 'dot', 'fdp', 'circo', 'twopi', 'neato', 'osage'
- messages: Set to 0 to disable various output messages
- ...

see control parameter of the estimate function.

**Value**

list of parameters

**Author(s)**

Klaus K. Holst

**Examples**

```
## Not run:
lava.options(iter.max=100,messages=0)

## End(Not run)
```

---

 lvm

---

*Initialize new latent variable model*


---

**Description**

Function that constructs a new latent variable model object

**Usage**

```
lvm(x = NULL, ..., latent = NULL, messages = lava.options()$messages)
```

**Arguments**

x	Vector of variable names. Optional but gives control of the sequence of appearance of the variables. The argument can be given as a character vector or formula, e.g. $\sim y_1 + y_2$ is equivalent to $c("y_1", "y_2")$ . Alternatively the argument can be a formula specifying a linear model.
...	Additional arguments to be passed to the low level functions
latent	(optional) Latent variables
messages	Controls what messages are printed (0: none)

**Value**

Returns an object of class lvm.

**Author(s)**

Klaus K. Holst

**See Also**

[regression](#), [covariance](#), [intercept](#), ...

**Examples**

```
m <- lvm() # Empty model
m1 <- lvm(y~x) # Simple linear regression
m2 <- lvm(~y1+y2) # Model with two independent variables (argument)
m3 <- lvm(list(c(y1,y2,y3)~u,u~x+z)) # SEM with three items
```

---

makemissing

*Create random missing data*

---

**Description**

Generates missing entries in data.frame/matrix

**Usage**

```
makemissing(
  data,
  p = 0.2,
  cols = seq_len(ncol(data)),
  rowwise = FALSE,
  nafun = function(x) x,
  seed = NULL
)
```

**Arguments**

data	data.frame
p	Fraction of missing data in each column
cols	Which columns (name or index) to alter
rowwise	Should missing occur row-wise (either none or all selected columns are missing)
nafun	(Optional) function to be applied on data.frame before return (e.g. na.omit to return complete-cases only)
seed	Random seed

**Value**

data.frame

**Author(s)**

Klaus K. Holst

---

measurement.error      *Two-stage (non-linear) measurement error*

---

**Description**

Two-stage measurement error

**Usage**

```
measurement.error(
  model1,
  formula,
  data = parent.frame(),
  predictfun = function(mu, var, data, ...) mu[, 1]^2 + var[1],
  id1,
  id2,
  ...
)
```

**Arguments**

model1	Stage 1 model
formula	Formula specifying observed covariates in stage 2 model
data	data.frame
predictfun	Predictions to be used in stage 2
id1	Optional id-vector of stage 1
id2	Optional id-vector of stage 2
...	Additional arguments to lower level functions

**See Also**

stack.estimate

**Examples**

```
m <- lvm(c(y1,y2,y3)~u,c(y3,y4,y5)~v,u~~v,c(u,v)~x)
transform(m,u2~u) <- function(x) x^2
transform(m,uv~u+v) <- prod
regression(m) <- z~u2+u+v+uv+x
set.seed(1)
d <- sim(m,1000,p=c("u,u"=1))

## Stage 1
m1 <- lvm(c(y1[0:s],y2[0:s],y3[0:s])~1*u,c(y3[0:s],y4[0:s],y5[0:s])~1*v,u~b*x,u~~v)
latent(m1) <- ~u+v
e1 <- estimate(m1,d)

pp <- function(mu,var,data,...) {
  cbind(u=mu["u"],u2=mu["u"]^2+var["u","u"],
        v=mu["v"],uv=mu["u"]*mu["v"]+var["u","v"])
}
(e <- measurement.error(e1, z~1+x, data=d, predictfun=pp))

## uu <- seq(-1,1,length.out=100)
## pp <- estimate(e,function(p,...) p["(Intercept)"+p["u"]*uu+p["u2"]*uu^2]$coefmat
if (interactive()) {
  plot(e,intercept=TRUE,line=0)
  dev.off()
  f <- function(p) p[1]+p["u"]*u+p["u2"]*u^2
  u <- seq(-1,1,length.out=100)
  plot(e, f, data=data.frame(u), ylim=c(-.5,2.5))
}
```

---

Missing

*Missing value generator*

---

**Description**

Missing value generator

**Usage**

```
Missing(object, formula, Rformula, missing.name, suffix = "0", ...)
```

**Arguments**

object            lvm-object.

formula	The right hand side specifies the name of a latent variable which is not always observed. The left hand side specifies the name of a new variable which is equal to the latent variable but has missing values. If given as a string then this is used as the name of the latent (full-data) name, and the observed data name is 'missing.data'
Rformula	Missing data mechanism with left hand side specifying the name of the observed data indicator (may also just be given as a character instead of a formula)
missing.name	Name of observed data variable (only used if 'formula' was given as a character specifying the name of the full-data variable)
suffix	If missing.name is missing, then the name of the observed data variable will be the name of the full-data variable + the suffix
...	Passed to binomial.lvm.

### Details

This function adds a binary variable to a given lvm model and also a variable which is equal to the original variable where the binary variable is equal to zero

### Value

lvm object

### Author(s)

Thomas A. Gerds [tag@biostat.ku.dk](mailto:tag@biostat.ku.dk)

### Examples

```
library(lava)
set.seed(17)
m <- lvm(y0~x01+x02+x03)
m <- Missing(m, formula=x1~x01, Rformula=R1~0.3*x02+-0.7*x01, p=0.4)
sim(m, 10)

m <- lvm(y~1)
m <- Missing(m, "y", "r")
## same as
## m <- Missing(m, y~1, r~1)
sim(m, 10)

## same as
m <- lvm(y~1)
Missing(m, "y") <- r~x
sim(m, 10)

m <- lvm(y~1)
m <- Missing(m, "y", "r", suffix=".")
## same as
## m <- Missing(m, "y", "r", missing.name="y.")
```

```
## same as
## m <- Missing(m,y.~y,"r")
sim(m,10)
```

---

missingdata

*Missing data example*

---

### Description

Simulated data generated from model

$$E(Y_i | X) = X, \quad cov(Y_1, Y_2 | X) = 0.5$$

### Format

list of data.frames

### Details

The list contains four data sets

1. Complete data
2. MCAR
3. MAR
4. MNAR (missing mechanism depends on variable V correlated with Y1,Y2)

### Source

Simulated

### Examples

```
data(missingdata)
e0 <- estimate(lvm(c(y1,y2)~b*x,y1~~y2),missingdata[[1]]) ## No missing
e1 <- estimate(lvm(c(y1,y2)~b*x,y1~~y2),missingdata[[2]]) ## CC (MCAR)
e2 <- estimate(lvm(c(y1,y2)~b*x,y1~~y2),missingdata[[2]],missing=TRUE) ## MCAR
e3 <- estimate(lvm(c(y1,y2)~b*x,y1~~y2),missingdata[[3]]) ## CC (MAR)
e4 <- estimate(lvm(c(y1,y2)~b*x,y1~~y2),missingdata[[3]],missing=TRUE) ## MAR
```

---

mixture	<i>Estimate mixture latent variable model.</i>
---------	--

---

### Description

Estimate mixture latent variable model

### Usage

```
mixture(
  x,
  data,
  k = length(x),
  control = list(),
  vcov = "observed",
  names = FALSE,
  ...
)
```

### Arguments

x	List of lvm objects. If only a single lvm object is given, then a k-mixture of this model is fitted (free parameters varying between mixture components).
data	data.frame
k	Number of mixture components
control	Optimization parameters (see details) #type Type of EM algorithm (standard, classification, stochastic)
vcov	of asymptotic covariance matrix (NULL to omit)
names	If TRUE returns the names of the parameters (for defining starting values)
...	Additional arguments parsed to lower-level functions

### Details

Estimate parameters in a mixture of latent variable models via the EM algorithm.

The performance of the EM algorithm can be tuned via the `control` argument, a list where a subset of the following members can be altered:

**start** Optional starting values

**nstart** Evaluate `nstart` different starting values and run the EM-algorithm on the parameters with largest likelihood

**tol** Convergence tolerance of the EM-algorithm. The algorithm is stopped when the absolute change in likelihood and parameter (2-norm) between successive iterations is less than `tol`

**iter.max** Maximum number of iterations of the EM-algorithm

**gamma** Scale-down (i.e. number between 0 and 1) of the step-size of the Newton-Raphson algorithm in the M-step

**trace** Trace information on the EM-algorithm is printed on every `traceth` iteration

Note that the algorithm can be aborted any time (C-c) and still be saved (via `on.exit` call).

### Author(s)

Klaus K. Holst

### See Also

`mvnmix`

### Examples

```
m0 <- lvm(list(y~x+z,x~z))
distribution(m0,~z) <- binomial.lvm()
d <- sim(m0,2000,p=c("y~z"=2,"y~x"=1),seed=1)

## unmeasured confounder example
m <- baptize(lvm(y~x, x~1));
intercept(m,~x+y) <- NA

if (requireNamespace('mets', quietly=TRUE)) {
  set.seed(42)
  M <- mixture(m,k=2,data=d,control=list(trace=1,tol=1e-6))
  summary(M)
  lm(y~x,d)
  estimate(M,"y~x")
  ## True slope := 1
}
```

---

Model

*Extract model*

---

### Description

Extract or replace model object

### Usage

`Model(x, ...)`

`Model(x, ...) <- value`

**Arguments**

x	Fitted model
...	Additional arguments to be passed to the low level functions
value	New model object (e.g. lvm or multigroup)

**Value**

Returns a model object (e.g. lvm or multigroup)

**Author(s)**

Klaus K. Holst

**See Also**

[Graph](#)

**Examples**

```
m <- lvm(y~x)
e <- estimate(m, sim(m,100))
Model(e)
```

---

modelsearch

*Model searching*

---

**Description**

Performs Wald or score tests

**Usage**

```
modelsearch(x, k = 1, dir = "forward", type = "all", ...)
```

**Arguments**

x	lvmfit-object
k	Number of parameters to test simultaneously. For equivalence the number of additional associations to be added instead of rel.
dir	Direction to do model search. "forward" := add associations/arrows to model/graph (score tests), "backward" := remove associations/arrows from model/graph (wald test)
type	If equal to 'correlation' only consider score tests for covariance parameters. If equal to 'regression' go through direct effects only (default 'all' is to do both)
...	Additional arguments to be passed to the low level functions

**Value**

Matrix of test-statistics and p-values

**Author(s)**

Klaus K. Holst

**See Also**

[compare](#), [equivalence](#)

**Examples**

```
m <- lvm();
regression(m) <- c(y1,y2,y3) ~ eta; latent(m) <- ~eta
regression(m) <- eta ~ x
m0 <- m; regression(m0) <- y2 ~ x
dd <- sim(m0,100)[,manifest(m0)]
e <- estimate(m,dd);
modelsearch(e,messages=0)
modelsearch(e,messages=0,type="cor")
```

---

multinomial

*Estimate probabilities in contingency table*


---

**Description**

Estimate probabilities in contingency table

**Usage**

```
multinomial(
  x,
  data = parent.frame(),
  marginal = FALSE,
  transform,
  vcov = TRUE,
  IC = TRUE,
  ...
)
```

**Arguments**

x	Formula (or matrix or data.frame with observations, 1 or 2 columns)
data	Optional data.frame
marginal	If TRUE the marginals are estimated
transform	Optional transformation of parameters (e.g., logit)

```
vcov          Calculate asymptotic variance (default TRUE)
IC            Return ic decomposition (default TRUE)
...          Additional arguments to lower-level functions
```

**Author(s)**

Klaus K. Holst

**Examples**

```
set.seed(1)
breaks <- c(-Inf,-1,0,Inf)
m <- lvm(); covariance(m,pairwise=TRUE) <- ~y1+y2+y3+y4
d <- transform(sim(m,5e2),
               z1=cut(y1,breaks=breaks),
               z2=cut(y2,breaks=breaks),
               z3=cut(y3,breaks=breaks),
               z4=cut(y4,breaks=breaks))

multinomial(d[,5])
(a1 <- multinomial(d[,5:6]))
(K1 <- kappa(a1)) ## Cohen's kappa

K2 <- kappa(d[,7:8])
## Testing difference K1-K2:
estimate(merge(K1,K2,id=TRUE),diff)

estimate(merge(K1,K2,id=FALSE),diff) ## Wrong std.err ignoring dependence
sqrt(vcov(K1)+vcov(K2))

## Average of the two kappas:
estimate(merge(K1,K2,id=TRUE),function(x) mean(x))
estimate(merge(K1,K2,id=FALSE),function(x) mean(x)) ## Independence
##'
## Goodman-Kruskal's gamma
m2 <- lvm(); covariance(m2) <- y1~y2
breaks1 <- c(-Inf,-1,0,Inf)
breaks2 <- c(-Inf,0,Inf)
d2 <- transform(sim(m2,5e2),
               z1=cut(y1,breaks=breaks1),
               z2=cut(y2,breaks=breaks2))

(g1 <- gkgamma(d2[,3:4]))
## same as
## Not run:
gkgamma(table(d2[,3:4]))
gkgamma(multinomial(d2[,3:4]))

## End(Not run)

##partial gamma
d2$x <- rbinom(nrow(d2),2,0.5)
```

```
gkgamma(z1~z2|x,data=d2)
```

---

mvnmix

*Estimate mixture latent variable model*


---

## Description

Estimate mixture latent variable model

## Usage

```
mvnmix(
  data,
  k = 2,
  theta,
  steps = 500,
  tol = 1e-16,
  lambda = 0,
  mu = NULL,
  silent = TRUE,
  extra = FALSE,
  n.start = 1,
  init = "kmp",
  ...
)
```

## Arguments

data	data.frame
k	Number of mixture components
theta	Optional starting values
steps	Maximum number of iterations
tol	Convergence tolerance of EM algorithm
lambda	Regularisation parameter. Added to diagonal of covariance matrix (to avoid singularities)
mu	Initial centres (if unspecified random centres will be chosen)
silent	Turn on/off output messages
extra	Extra debug information
n.start	Number of restarts
init	Function to choose initial centres
...	Additional arguments parsed to lower-level functions

## Details

Estimate parameters in a mixture of latent variable models via the EM algorithm.

**Value**

A mixture object

**Author(s)**

Klaus K. Holst

**See Also**

mixture

**Examples**

```
data(faithful)
set.seed(1)
M1 <- mvnmix(faithful[, "waiting", drop=FALSE], k=2)
M2 <- mvnmix(faithful, k=2)
if (interactive()) {
  par(mfrow=c(2,1))
  plot(M1, col=c("orange", "blue"), ylim=c(0, 0.05))
  plot(M2, col=c("orange", "blue"))
}
```

---

na.pass0

*Handle Missing Values in Objects*

---

**Description**

Returns the object with missing data replaced by zeros. This is sometimes useful for example when working with inverse probability weighting of the complete-case data.

**Usage**

```
na.pass0(object, na.action = na.omit, ...)
```

**Arguments**

object	data.frame, vector
na.action	function used to identify missing values
...	additional arguments to lower level functions

**See Also**

[na.pass\(\)](#), [na.omit\(\)](#), [na.fail\(\)](#)

**Examples**

```
d <- data.frame(y=c(1,1,NA,2,NA,2), r=c(1,1,0,1,1,1))
na.pass0(d)
glm(y ~ 1, weights=d$r, data=d, na.action=na.pass0)
```

NA2x

*Convert to/from NA***Description**

Convert vector to/from NA

**Usage**

```
NA2x(s, x = 0)
```

**Arguments**

`s` The input vector (of arbitrary class)  
`x` The elements to transform into NA resp. what to transform NA into.

**Value**

A vector with same dimension and class as `s`.

**Author(s)**

Klaus K. Holst

**Examples**

```
##'
x2NA(1:10, 1:5)
NA2x(x2NA(c(1:10),5),5)##'
```

nldata

*Example data (nonlinear model)***Description**

Example data (nonlinear model)

**Format**

data.frame

**Source**

Simulated

---

NR	<i>Newton-Raphson method</i>
----	------------------------------

---

**Description**

Newton-Raphson method

**Usage**

```
NR(
  start,
  objective = NULL,
  gradient = NULL,
  hessian = NULL,
  control,
  args = NULL,
  ...
)
```

**Arguments**

start	Starting value
objective	Optional objective function (used for selecting step length)
gradient	gradient
hessian	hessian (if NULL a numerical derivative is used)
control	optimization arguments (see details)
args	Optional list of arguments parsed to objective, gradient and hessian
...	additional arguments parsed to lower level functions

**Details**

control should be a list with one or more of the following components:

- trace integer for which output is printed each 'trace'th iteration
- iter.max number of iterations
- stepsize: Step size (default 1)
- nstepsize: Increase stepsize every nstepsize iteration (from stepsize to 1)
- tol: Convergence criterion (gradient)
- epsilon: threshold used in pseudo-inverse
- backtrack: In each iteration reduce stepsize unless solution is improved according to criterion (gradient, armijo, curvature, wolfe)

**Examples**

```
# Objective function with gradient and hessian as attributes
f <- function(z) {
  x <- z[1]; y <- z[2]
  val <- x^2 + x*y^2 + x + y
  structure(val, gradient=c(2*x+y^2+1, 2*y*x+1),
            hessian=rbind(c(2,2*y),c(2*y,2*x)))
}
NR(c(0,0),f)

# Parsing arguments to the function and
g <- function(x,y) (x*y+1)^2
NR(0, gradient=g, args=list(y=2), control=list(trace=1,tol=1e-20))
```

---

nsem	<i>Example SEM data (nonlinear)</i>
------	-------------------------------------

---

**Description**

Simulated data

**Format**

data.frame

**Source**

Simulated

---

ordinal<-	<i>Define variables as ordinal</i>
-----------	------------------------------------

---

**Description**

Define variables as ordinal in latent variable model object

**Usage**

```
ordinal(x, ...) <- value
```

**Arguments**

x	Object
...	additional arguments to lower level functions
value	variable (formula or character vector)

**Examples**

```

if (requireNamespace("mets")) {
  m <- lvm(y + z ~ x + 1*u[0], latent=~u)
  ordinal(m, K=3) <- ~y+z
  d <- sim(m, 100, seed=1)
  e <- estimate(m, d)
}

```

---

ordreg

*Univariate cumulative link regression models*


---

**Description**

Ordinal regression models

**Usage**

```

ordreg(
  formula,
  data = parent.frame(),
  offset,
  family = stats::binomial("probit"),
  start,
  fast = FALSE,
  ...
)

```

**Arguments**

formula	formula
data	data.frame
offset	offset
family	family (default proportional odds)
start	optional starting values
fast	If TRUE standard errors etc. will not be calculated
...	Additional arguments to lower level functions

**Author(s)**

Klaus K. Holst

**Examples**

```

m <- lvm(y~x)
ordinal(m,K=3) <- ~y
d <- sim(m,100)
e <- ordreg(y~x,d)

```

---

parpos                      *Generic method for finding indeces of model parameters*

---

**Description**

Generic method for finding indeces of model parameters

**Usage**

```
parpos(x, ...)
```

**Arguments**

x	Model object
...	Additional arguments

**Author(s)**

Klaus K. Holst

---

partialcor                      *Calculate partial correlations*

---

**Description**

Calculate partial correlation coefficients and confidence limits via Fishers z-transform

**Usage**

```
partialcor(formula, data, level = 0.95, ...)
```

**Arguments**

formula	formula specifying the covariates and optionally the outcomes to calculate partial correlation for
data	data.frame
level	Level of confidence limits
...	Additional arguments to lower level functions

**Value**

A coefficient matrix

**Author(s)**

Klaus K. Holst

**Examples**

```
m <- lvm(c(y1,y2,y3)~x1+x2)
covariance(m) <- c(y1,y2,y3)~y1+y2+y3
d <- sim(m,500)
partialcor(~x1+x2,d)
```

---

path	<i>Extract pathways in model graph</i>
------	--

---

**Description**

Extract all possible paths from one variable to another connected component in a latent variable model. In an estimated model the effect size is decomposed into direct, indirect and total effects including approximate standard errors.

**Usage**

```
## S3 method for class 'lvm'
path(object, to = NULL, from, all=FALSE, ...)
## S3 method for class 'lvmfit'
effects(object, to, from, ...)
```

**Arguments**

object	Model object (lvm)
...	Additional arguments to be passed to the low level functions
to	Outcome variable (string). Alternatively a formula specifying response and predictor in which case the argument from is ignored.
from	Response variable (string), not necessarily directly affected by to.
all	If TRUE all simple paths (in undirected graph) is returned on/off.

**Value**

If object is of class `lvmfit` a list with the following elements is returned

idx	A list where each element defines a possible pathway via a integer vector indicating the index of the visited nodes.
V	A List of covariance matrices for each path.
coef	A list of parameters estimates for each path
path	A list where each element defines a possible pathway via a character vector naming the visited nodes in order.
edges	Description of 'comp2'

If object is of class `lvm` only the path element will be returned.

The effects method returns an object of class effects.

**Note**

For a `lvmfit`-object the parameters estimates and their corresponding covariance matrix are also returned. The `effects`-function additionally calculates the total and indirect effects with approximate standard errors

**Author(s)**

Klaus K. Holst

**See Also**

children, parents

**Examples**

```
m <- lvm(c(y1,y2,y3)~eta)
regression(m) <- y2~x1
latent(m) <- ~eta
regression(m) <- eta~x1+x2
d <- sim(m,500)
e <- estimate(m,d)

path(Model(e),y2~x1)
parents(Model(e), ~y2)
children(Model(e), ~x2)
children(Model(e), ~x2+eta)
effects(e,y2~x1)
## All simple paths (undirected)
path(m,y1~x1,all=TRUE)
```

---

pcor

*Polychoric correlation*

---

**Description**

Maximum likelihood estimates of polychoric correlations

**Usage**

```
pcor(x, y, X, start, ...)
```

**Arguments**

x	Variable 1
y	Variable 2
X	Optional covariates
start	Optional starting values
...	Additional arguments to lower level functions

---

 PD

*Dose response calculation for binomial regression models*


---

**Description**

Dose response calculation for binomial regression models

**Usage**

```
PD(
  model,
  intercept = 1,
  slope = 2,
  prob = NULL,
  x,
  level = 0.5,
  ci.level = 0.95,
  vcov,
  family,
  EB = NULL
)
```

**Arguments**

model	Model object or vector of parameter estimates
intercept	Index of intercept parameters
slope	Index of intercept parameters
prob	Index of mixture parameters (only relevant for zibreg models)
x	Optional weights $\text{length}(x)=\text{length}(\text{intercept})+\text{length}(\text{slope})+\text{length}(\text{prob})$
level	Probability at which level to calculate dose
ci.level	Level of confidence limits
vcov	Optional estimate of variance matrix of parameter estimates
family	Optional distributional family argument
EB	Optional ratio of treatment effect and adverse effects used to find optimal dose (regret-function argument)

**Author(s)**

Klaus K. Holst

---

pdfconvert                      *Convert pdf to raster format*

---

**Description**

Convert PDF file to print quality png (default 300 dpi)

**Usage**

```
pdfconvert(  
  files,  
  dpi = 300,  
  resolution = 1024,  
  gs,  
  gsopt,  
  resize,  
  format = "png",  
  ...  
)
```

**Arguments**

files	Vector of (pdf-)filenames to process
dpi	DPI
resolution	Resolution of raster image file
gs	Optional ghostscript command
gsopt	Optional ghostscript arguments
resize	Optional resize arguments (mogrify)
format	Raster format (e.g. png, jpg, tif, ...)
...	Additional arguments

**Details**

Access to ghostscript program 'gs' is needed

**Author(s)**

Klaus K. Holst

**See Also**

dev.copy2pdf, printdev

---

plot.estimate                      *Plot method for 'estimate' objects*

---

### Description

Plot method for 'estimate' objects

### Usage

```
## S3 method for class 'estimate'
plot(
  x,
  f,
  idx,
  intercept = FALSE,
  data,
  confint = TRUE,
  type = "l",
  xlab = "x",
  ylab = "f(x)",
  col = 1,
  add = FALSE,
  null = 0,
  ...
)
```

### Arguments

x	estimate object
f	function of parameter coefficients and data parsed on to 'estimate'. If omitted a forest-plot will be produced.
idx	Index of parameters (default all)
intercept	include intercept in forest-plot
data	data.frame
confint	Add confidence limits
type	plot type ('l')
xlab	x-axis label
ylab	y-axis label
col	color
add	add plot to current device
null	null value for forest-plot
...	additional arguments to lower-level functions

---

plot.lvm	<i>Plot path diagram</i>
----------	--------------------------

---

**Description**

Plot the path diagram of a SEM

**Usage**

```
## S3 method for class 'lvm'
plot(
  x,
  diag = FALSE,
  cor = TRUE,
  labels = FALSE,
  intercept = FALSE,
  addcolor = TRUE,
  plain = FALSE,
  cex,
  fontsize1 = 10,
  noplot = FALSE,
  graph = list(rankdir = "BT"),
  attrs = list(graph = graph),
  unexpr = FALSE,
  addstyle = TRUE,
  plot.engine = lava.options()$plot.engine,
  init = TRUE,
  layout = lava.options()$layout,
  edgecolor = lava.options()$edgecolor,
  graph.proc = lava.options()$graph.proc,
  ...
)
```

**Arguments**

<code>x</code>	Model object
<code>diag</code>	Logical argument indicating whether to visualize variance parameters (i.e. diagonal of variance matrix)
<code>cor</code>	Logical argument indicating whether to visualize correlation parameters
<code>labels</code>	Logical argument indicating whether to add labels to plot (Unnamed parameters will be labeled p1,p2,...)
<code>intercept</code>	Logical argument indicating whether to add intercept labels
<code>addcolor</code>	Logical argument indicating whether to add colors to plot (overrides nodecolor calls)
<code>plain</code>	if TRUE strip plot of colors and boxes

cex	Fontsize of node labels
fontsize1	Fontsize of edge labels
noplot	if TRUE then return graphNEL object only
graph	Graph attributes (Rgraphviz)
attrs	Attributes (Rgraphviz)
unexpr	if TRUE remove expressions from labels
addstyle	Logical argument indicating whether additional style should automatically be added to the plot (e.g. dashed lines to double-headed arrows)
plot.engine	default 'Rgraphviz' if available, otherwise visNetwork,igraph
init	Reinitialize graph (for internal use)
layout	Graph layout (see Rgraphviz or igraph manual)
edgecolor	if TRUE plot style with colored edges
graph.proc	Function that post-process the graph object (default: subscripts are automatically added to labels of the nodes)
...	Additional arguments to be passed to the low level functions

**Author(s)**

Klaus K. Holst

**Examples**

```

if (interactive()) {
m <- lvm(c(y1,y2) ~ eta)
regression(m) <- eta ~ z+x2
regression(m) <- c(eta,z) ~ x1
latent(m) <- ~eta
labels(m) <- c(y1=expression(y[scriptscriptstyle(1)]),
y2=expression(y[scriptscriptstyle(2)]),
x1=expression(x[scriptscriptstyle(1)]),
x2=expression(x[scriptscriptstyle(2)]),
eta=expression(eta))
edgelabels(m, eta ~ z+x1+x2, cex=2, lwd=3,
col=c("orange","lightblue","lightblue")) <- expression(rho,phi,psi)
nodecolor(m, vars(m), border="white", labcol="darkblue") <- NA
nodecolor(m, ~y1+y2+z, labcol=c("white","white","black")) <- NA
plot(m,cex=1.5)

d <- sim(m,100)
e <- estimate(m,d)
plot(e)

m <- lvm(c(y1,y2) ~ eta)
regression(m) <- eta ~ z+x2
regression(m) <- c(eta,z) ~ x1
latent(m) <- ~eta
plot(lava:::beautify(m,edgecol=FALSE))
}

```

---

`plot.sim`*Plot method for simulation 'sim' objects*

---

**Description**

Density and scatter plots

**Usage**

```
## S3 method for class 'sim'
plot(
  x,
  estimate,
  se = NULL,
  true = NULL,
  names = NULL,
  auto.layout = TRUE,
  byrow = FALSE,
  type = "p",
  ask = grDevices::dev.interactive(),
  col = c("gray60", "orange", "darkblue", "seagreen", "darkred"),
  pch = 16,
  cex = 0.5,
  lty = 1,
  lwd = 0.3,
  legend,
  legendpos = "topleft",
  cex.legend = 0.8,
  plot.type = c("multiple", "single"),
  polygon = TRUE,
  density = NULL,
  angle = -45,
  cex.axis = 0.8,
  alpha = 0.2,
  main,
  cex.main = 1,
  equal = FALSE,
  delta = 1.15,
  ylim = NULL,
  xlim = NULL,
  ylab = "",
  xlab = "",
  rug = FALSE,
  rug.alpha = 0.5,
  line.col = scatter.col,
  line.lwd = 1,
  line.lty = 1,
```

```

line.alpha = 1,
scatter.ylab = "Estimate",
scatter.ylim = NULL,
scatter.xlim = NULL,
scatter.alpha = 0.5,
scatter.col = col,
border = col,
true.lty = 2,
true.col = "gray70",
true.lwd = 1.2,
density.plot = TRUE,
scatter.plot = FALSE,
running.mean = scatter.plot,
...
)

```

### Arguments

x	sim object
estimate	columns with estimates
se	columns with standard error estimates
true	(optional) vector of true parameter values
names	(optional) names of estimates
auto.layout	Auto layout (default TRUE)
byrow	Add new plots to layout by row
type	plot type
ask	if TRUE user is asked for input, before a new figure is drawn
col	colour (for each estimate)
pch	plot symbol
cex	point size
lty	line type
lwd	line width
legend	legend
legendpos	legend position
cex.legend	size of legend text
plot.type	'single' or 'multiple' (default)
polygon	if TRUE fill the density estimates with colour
density	if non-zero add shading lines to polygon
angle	shading lines angle of polygon
cex.axis	Font size on axis
alpha	Semi-transparent level (1: non-transparent, 0: full)
main	Main title

cex.main	Size of title font
equal	Same x-axis and y-axis for all plots
delta	Controls the amount of space around axis limits
ylim	y-axis limits
xlim	x-axis limits
ylab	y axis label
xlab	x axis label
rug	if TRUE add rug representation of data to x-axis
rug.alpha	rug semi-transparency level
line.col	line colour (running mean, only for scatter plots)
line.lwd	line width (running mean, only for scatter plots)
line.lty	line type (running mean, only for scatter plots)
line.alpha	line transparency
scatter.ylab	y label for density plots
scatter.ylim	y-axis limits for density plots
scatter.xlim	x-axis limits for density plots
scatter.alpha	semi-transparency of scatter plot
scatter.col	scatter plot colour
border	border colour of density estimates
true.lty	true parameter estimate line type
true.col	true parameter colour
true.lwd	true parameter line width
density.plot	if TRUE add density plot
scatter.plot	if TRUE add scatter plot
running.mean	if TRUE add running average estimate to scatter plot
...	additional arguments to lower level functions

### Examples

```

n <- 1000
val <- cbind(est1=rnorm(n,sd=1),est2=rnorm(n,sd=0.2),est3=rnorm(n,1,sd=0.5),
            sd1=runif(n,0.8,1.2),sd2=runif(n,0.1,0.3),sd3=runif(n,0.25,0.75))

plot.sim(val,estimate=c(1,2),true=c(0,0),se=c(4,5),equal=TRUE,scatter.plot=TRUE)
plot.sim(val,estimate=c(1,3),true=c(0,1),se=c(4,6),xlim=c(-3,3),
scatter.ylim=c(-3,3),scatter.plot=TRUE)
plot.sim(val,estimate=c(1,2),true=c(0,0),se=c(4,5),equal=TRUE,
plot.type="single",scatter.plot=TRUE)
plot.sim(val,estimate=c(1),se=c(4,5,6),plot.type="single",scatter.plot=TRUE)
plot.sim(val,estimate=c(1,2,3),equal=TRUE,scatter.plot=TRUE)
plot.sim(val,estimate=c(1,2,3),equal=TRUE,byrow=TRUE,scatter.plot=TRUE)
plot.sim(val,estimate=c(1,2,3),plot.type="single",scatter.plot=TRUE)
plot.sim(val,estimate=1,se=c(3,4,5),plot.type="single",scatter.plot=TRUE)

density.sim(val,estimate=c(1,2,3),density=c(0,10,10),lwd=2,angle=c(0,45,-45),cex.legend=1.3)

```

---

plotConf	<i>Plot regression lines</i>
----------	------------------------------

---

**Description**

Plot regression line (with interactions) and partial residuals.

**Usage**

```
plotConf(  
  model,  
  var1 = NULL,  
  var2 = NULL,  
  data = NULL,  
  ci.lty = 0,  
  ci = TRUE,  
  level = 0.95,  
  pch = 16,  
  lty = 1,  
  lwd = 2,  
  npoints = 100,  
  xlim,  
  col = NULL,  
  colpt,  
  alpha = 0.5,  
  cex = 1,  
  delta = 0.07,  
  centermark = 0.03,  
  jitter = 0.2,  
  cidiff = FALSE,  
  mean = TRUE,  
  legend = ifelse(is.null(var1), FALSE, "topright"),  
  trans = function(x) {  
    x  
  },  
  partres = inherits(model, "lm"),  
  partse = FALSE,  
  labels,  
  vcov,  
  predictfun,  
  plot = TRUE,  
  new = TRUE,  
  ...  
)
```

**Arguments**

model	Model object (e.g. <code>lm()</code> )
-------	--

var1	predictor (Continuous or factor)
var2	Factor that interacts with var1
data	data.frame to use for prediction (model.frame is used as default)
ci.lty	Line type for confidence limits
ci	Boolean indicating whether to draw pointwise 95% confidence limits
level	Level of confidence limits (default 95%)
pch	Point type for partial residuals
lty	Line type for estimated regression lines
lwd	Line width for regression lines
npoints	Number of points used to plot curves
xlim	Range of x axis
col	Color (for each level in var2)
colpt	Color of partial residual points
alpha	Alpha level
cex	Point size
delta	For categorical var1
centermark	For categorical var1
jitter	For categorical var1
cidiff	For categorical var1
mean	For categorical var1
legend	Boolean (add legend)
trans	Transform estimates (e.g. exponential)
partres	Boolean indicating whether to plot partial residuals
partse	.
labels	Optional labels of var2
vcov	Optional variance estimates
predictfun	Optional predict-function used to calculate confidence limits and predictions
plot	If FALSE return only predictions and confidence bands
new	If FALSE add to current plot
...	additional arguments to lower level functions

### Value

list with following members:

- x Variable on the x-axis var1
- y Variable on the y-axis (partial residuals)
- predict Matrix with confidence limits and predicted startvalues.R

**Author(s)**

Klaus K. Holst

**See Also**[termplot\(\)](#)**Examples**

```

n <- 100
x0 <- rnorm(n)
x1 <- seq(-3,3, length.out=n)
x2 <- factor(rep(c(1,2),each=n/2), labels=c("A","B"))
y <- 5 + 2*x0 + 0.5*x1 + -1*(x2=="B")*x1 + 0.5*(x2=="B") + rnorm(n, sd=0.25)
dd <- data.frame(y=y, x1=x1, x2=x2)
lm0 <- lm(y ~ x0 + x1*x2, dd)
plotConf(lm0, var1="x1", var2="x2")
abline(a=5,b=0.5,col="red")
abline(a=5.5,b=-0.5,col="red")
### points(5+0.5*x1 -1*(x2=="B")*x1 + 0.5*(x2=="B") ~ x1, cex=2)

data(iris)
l <- lm(Sepal.Length ~ Sepal.Width*Species,iris)
plotConf(l,var2="Species")
plotConf(l,var1="Sepal.Width",var2="Species")

## Not run:
## lme4 model
dd$Id <- rbinom(n, size = 3, prob = 0.3)
lmer0 <- lme4::lmer(y ~ x0 + x1*x2 + (1|Id), dd)
plotConf(lmer0, var1="x1", var2="x2")

## End(Not run)

```

---

predict.lvm

*Prediction in structural equation models*


---

**Description**

Prediction in structural equation models

**Usage**

```

## S3 method for class 'lvm'
predict(
  object,
  x = NULL,
  y = NULL,
  residual = FALSE,

```

```

    p,
    data,
    path = FALSE,
    quick = is.null(x) & !(residual | path),
    ...
  )

```

### Arguments

object	Model object
x	optional list of (endogenous) variables to condition on
y	optional subset of variables to predict
residual	If true the residuals are predicted
p	Parameter vector
data	Data to use in prediction
path	Path prediction
quick	If TRUE the conditional mean and variance given covariates are returned (and all other calculations skipped)
...	Additional arguments to lower level function

### See Also

predictlvm

### Examples

```

m <- lvm(list(c(y1,y2,y3)~u,u~x)); latent(m) <- ~u
d <- sim(m,100)
e <- estimate(m,d)

## Conditional mean (and variance as attribute) given covariates
r <- predict(e)
## Best linear unbiased predictor (BLUP)
r <- predict(e,vars(e))
## Conditional mean of y3 giving covariates and y1,y2
r <- predict(e,y3~y1+y2)
## Conditional mean gives covariates and y1
r <- predict(e,~y1)
## Predicted residuals (conditional on all observed variables)
r <- predict(e,vars(e),residual=TRUE)

```

---

predictlvm                      *Predict function for latent variable models*

---

### Description

Predictions of conditional mean and variance and calculation of jacobian with respect to parameter vector.

### Usage

```
predictlvm(object, formula, p = coef(object), data = model.frame(object), ...)
```

### Arguments

object	Model object
formula	Formula specifying which variables to predict and which to condition on
p	Parameter vector
data	Data.frame
...	Additional arguments to lower level functions

### See Also

predict.lvm

### Examples

```
m <- lvm(c(x1,x2,x3)~u1,u1~z,
         c(y1,y2,y3)~u2,u2~u1+z)
latent(m) <- ~u1+u2
d <- simulate(m,10,"u2,u2"=2,"u1,u1"=0.5,seed=123)
e <- estimate(m,d)

## Conditional mean given covariates
predictlvm(e,c(x1,x2)~1)$mean
## Conditional variance of u1,y1 given x1,x2
predictlvm(e,c(u1,y1)~x1+x2)$var
```

---

Print	<i>Generic print method</i>
-------	-----------------------------

---

**Description**

Nicer print method for tabular data. Falls back to standard print method for all other data types.

**Usage**

```
Print(x, n = 5, digits = max(3, getOption("digits") - 3), ...)
```

**Arguments**

x	object to print
n	number of rows to show from top and bottom of tabular data
digits	precision
...	additional arguments to print method

---

Range.lvm	<i>Define range constraints of parameters</i>
-----------	---

---

**Description**

Define range constraints of parameters

**Usage**

```
Range.lvm(a = 0, b = 1)
```

**Arguments**

a	Lower bound
b	Upper bound

**Value**

function

**Author(s)**

Klaus K. Holst

---

rbind.Surv	<i>Appending Surv objects</i>
------------	-------------------------------

---

**Description**

rbind method for Surv objects

**Usage**

```
## S3 method for class 'Surv'
rbind(...)
```

**Arguments**

... Surv objects

**Value**

Surv object

**Author(s)**

Klaus K. Holst

**Examples**

```
y <- y1 <- yr <- rnorm(10)
y1[1:5] <- NA; yr[6:10] <- NA
S1 <- survival::Surv(y1,yr,type="interval2")
S2 <- survival::Surv(y,y>0,type="right")
S3 <- survival::Surv(y,y<0,type="left")

rbind(S1,S1)
rbind(S2,S2)
rbind(S3,S3)
```

---

regression<-	<i>Add regression association to latent variable model</i>
--------------	--

---

**Description**

Define regression association between variables in a lvm-object and define linear constraints between model equations.

**Usage**

```
## S3 method for class 'lvm'
regression(object = lvm(), to, from, fn = NA,
messages = lava.options()$messages, additive=TRUE, y, x, value, ...)
## S3 replacement method for class 'lvm'
regression(object, to=NULL, quick=FALSE, ...) <- value
```

**Arguments**

object	lvm-object.
...	Additional arguments to be passed to the low level functions
value	A formula specifying the linear constraints or if to=NULL a list of parameter values.
to	Character vector of outcome(s) or formula object.
from	Character vector of predictor(s).
fn	Real function defining the functional form of predictors (for simulation only).
messages	Controls which messages are turned on/off (0: all off)
additive	If FALSE and predictor is categorical a non-additive effect is assumed
y	Alias for 'to'
x	Alias for 'from'
quick	Faster implementation without parameter constraints

**Details**

The regression function is used to specify linear associations between variables of a latent variable model, and offers formula syntax resembling the model specification of e.g. `lm`.

For instance, to add the following linear regression model, to the lvm-object, `m`:

$$E(Y|X_1, X_2) = \beta_1 X_1 + \beta_2 X_2$$

We can write

```
regression(m) <- y ~ x1 + x2
```

Multivariate models can be specified by successive calls with `regression`, but multivariate formulas are also supported, e.g.

```
regression(m) <- c(y1, y2) ~ x1 + x2
```

defines

$$E(Y_i|X_1, X_2) = \beta_{1i} X_1 + \beta_{2i} X_2$$

The special function, `f`, can be used in the model specification to specify linear constraints. E.g. to fix  $\beta_1 = \beta_2$ , we could write

```
regression(m) <- y ~ f(x1, beta) + f(x2, beta)
```

The second argument of `f` can also be a number (e.g. defining an offset) or be set to `NA` in order to clear any previously defined linear constraints.

Alternatively, a more straight forward notation can be used:

```
regression(m) <- y ~ beta*x1 + beta*x2
```

All the parameter values of the linear constraints can be given as the right handside expression of the assignment function `regression<-` if the first (and possibly second) argument is defined as well. E.g:

```
regression(m,y1~x1+x2) <- list("a1","b1")
```

defines  $E(Y_1|X_1, X_2) = a_1X_1 + b_1X_2$ . The rhs argument can be a mixture of character and numeric values (and NA's to remove constraints).

The function `regression` (called without additional arguments) can be used to inspect the linear constraints of a `lvm`-object.

### Value

A `lvm`-object

### Note

Variables will be added to the model if not already present.

### Author(s)

Klaus K. Holst

### See Also

[intercept<-](#), [covariance<-](#), [constrain<-](#), [parameter<-](#), [latent<-](#), [cancel<-](#), [kill<-](#)

### Examples

```
m <- lvm() ## Initialize empty lvm-object
### E(y1|z,v) = beta1*z + beta2*v
regression(m) <- y1 ~ z + v
### E(y2|x,z,v) = beta*x + beta*z + 2*v + beta3*u
regression(m) <- y2 ~ f(x,beta) + f(z,beta) + f(v,2) + u
### Clear restriction on association between y and
### fix slope coefficient of u to beta
regression(m, y2 ~ v+u) <- list(NA,"beta")

regression(m) ## Examine current linear parameter constraints

## ## A multivariate model, E(yi|x1,x2) = beta[1i]*x1 + beta[2i]*x2:
m2 <- lvm(c(y1,y2) ~ x1+x2)
```

---

revdiag	<i>Create/extract 'reverse'-diagonal matrix or off-diagonal elements</i>
---------	--

---

**Description**

Create/extract 'reverse'-diagonal matrix or off-diagonal elements

**Usage**

```
revdiag(x,...)
offdiag(x,type=0,...)

revdiag(x,...) <- value
offdiag(x,type=0,...) <- value
```

**Arguments**

x	vector
...	additional arguments to lower level functions
value	For the assignment function the values to put in the diagonal
type	0: upper and lower triangular, 1: upper triangular, 2: lower triangular, 3: upper triangular + diagonal, 4: lower triangular + diagonal

**Author(s)**

Klaus K. Holst

---

rmvar	<i>Remove variables from (model) object.</i>
-------	--

---

**Description**

Generic method for removing elements of object

**Usage**

```
rmvar(x, ...) <- value
```

**Arguments**

x	Model object
...	additional arguments to lower level functions
value	Vector of variables or formula specifying which nodes to remove

**Author(s)**

Klaus K. Holst

**See Also**

cancel

**Examples**

```
m <- lvm()
addvar(m) <- ~y1+y2+x
covariance(m) <- y1~y2
regression(m) <- c(y1,y2) ~ x
### Cancel the covariance between the residuals of y1 and y2
cancel(m) <- y1~y2
### Remove y2 from the model
rmvar(m) <- ~y2
```

---

`rotate2`*Performs a rotation in the plane*

---

**Description**

Performs a rotation in the plane

**Usage**`rotate2(x, theta = pi)`**Arguments**

<code>x</code>	Matrix to be rotated (2 times n)
<code>theta</code>	Rotation in radians

**Value**Returns a matrix of the same dimension as `x`**Author(s)**

Klaus K. Holst

**Examples**

```
rotate2(cbind(c(1,2),c(2,1)))
```

---

 scheffe

*Calculate simultaneous confidence limits by Scheffe's method*


---

**Description**

Function to compute the Scheffe corrected confidence interval for the regression line

**Usage**

```
scheffe(model, newdata = model.frame(model), level = 0.95)
```

**Arguments**

model	Linear model
newdata	new data frame
level	confidence level (0.95)

**Examples**

```
x <- rnorm(100)
d <- data.frame(y=rnorm(length(x),x),x=x)
l <- lm(y~x,d)
plot(y~x,d)
abline(l)
d0 <- data.frame(x=seq(-5,5,length.out=100))
d1 <- cbind(d0,predict(l,newdata=d0,interval="confidence"))
d2 <- cbind(d0,scheffe(l,d0))
lines(lwr~x,d1,lty=2,col="red")
lines(upr~x,d1,lty=2,col="red")
lines(lwr~x,d2,lty=2,col="blue")
lines(upr~x,d2,lty=2,col="blue")
```

---

 semdata

*Example SEM data*


---

**Description**

Simulated data

**Format**

data.frame

**Source**

Simulated

serotonin

*Serotonin data***Description**

This simulated data mimics a PET imaging study where the 5-HT2A receptor and serotonin transporter (SERT) binding potential has been quantified into 8 different regions. The 5-HT2A cortical regions are considered high-binding regions measurements. These measurements can be regarded as proxy measures of the extra-cellular levels of serotonin in the brain

day	numeric	Scan day of the year
age	numeric	Age at baseline scan
mem	numeric	Memory performance score
depr	numeric	Depression (mild) status 500 days after baseline
gene1	numeric	Gene marker 1 (HTR2A)
gene2	numeric	Gene marker 2 (HTTTLPR)
cau	numeric	SERT binding, Caudate Nucleus
th	numeric	SERT binding, Thalamus
put	numeric	SERT binding, Putamen
mid	numeric	SERT binding, Midbrain
aci	numeric	5-HT2A binding, Anterior cingulate gyrus
pci	numeric	5-HT2A binding, Posterior cingulate gyrus
sfc	numeric	5-HT2A binding, Superior frontal cortex
par	numeric	5-HT2A binding, Parietal cortex

**Format**

data.frame

**Source**

Simulated

sim.default

*Monte Carlo simulation***Description**

Applies a function repeatedly for a specified number of replications or over a list/data.frame with plot and summary methods for summarizing the Monte Carlo experiment. Can be parallelized via the future package (use the `future::plan()` function).

**Usage**

```
## Default S3 method:
sim(
  x = NULL,
  R = 100,
  f = NULL,
  colnames = NULL,
  seed = NULL,
  args = list(),
  iter = FALSE,
  mc.cores,
  progressr.message = NULL,
  estimate.index = 1:2,
  ...
)
```

**Arguments**

x	function or 'sim' object
R	Number of replications or data.frame with parameters
f	Optional function (i.e., if x is a matrix)
colnames	Optional column names
seed	(optional) Seed (needed with cl=TRUE)
args	(optional) list of named arguments passed to (mc)mapply
iter	If TRUE the iteration number is passed as first argument to (mc)mapply
mc.cores	Optional number of cores. Will use parallel::mcmapply instead of future
progressr.message	Optional message for the progressr progress-bar
estimate.index	If return object inherits from estimate then only these column indices are extracted (estimate, se, lower, upper, p-val)
...	Additional arguments to <a href="#">future.apply::future_mapply()</a>

**Details**

To parallelize the calculation use the [future::plan\(\)](#) function (e.g., `future::plan(multisession())` to distribute the calculations over the R replications on all available cores). The output is controlled via the `progressr` package (e.g., `progressr::handlers(global=TRUE)` to enable progress information).

**See Also**

[summary.sim\(\)](#) [plot.sim\(\)](#) [sim.lvm\(\)](#)

**Examples**

```

m <- lvm(y~x+e)
distribution(m,~y) <- 0
distribution(m,~x) <- uniform.lvm(a=-1.1,b=1.1)
transform(m,e~x) <- function(x) (1*x^4)*rnorm(length(x),sd=1)

onerun <- function(iter=NULL,...,n=2e3,b0=1,idx=2) {
  d <- sim(m,n,p=c("y~x"=b0))
  l <- lm(y~x,d)
  res <- c(coef(summary(l))[idx,1:2],
          confint(l)[idx,],
          estimate(l,only.coef=TRUE)[idx,2:4])
  names(res) <- c("Estimate","Model.se","Model.lo","Model.hi",
                "Sandwich.se","Sandwich.lo","Sandwich.hi")
  res
}
val <- sim(onerun,R=10,b0=1)
val

val <- sim(val,R=40,b0=1) ## append results
summary(val,estimate=c(1,1),confint=c(3,4,6,7),true=c(1,1))

summary(val,estimate=c(1,1),se=c(2,5),names=c("Model","Sandwich"))
summary(val,estimate=c(1,1),se=c(2,5),true=c(1,1),
        names=c("Model","Sandwich"),confint=TRUE)

if (interactive()) {
  plot(val,estimate=1,c(2,5),true=1,
       names=c("Model","Sandwich"),polygon=FALSE)
  plot(val,estimate=c(1,1),se=c(2,5),main=NULL,
       true=c(1,1),names=c("Model","Sandwich"),
       line.lwd=1,col=c("gray20","gray60"),
       rug=FALSE)
  plot(val,estimate=c(1,1),se=c(2,5),true=c(1,1),
       names=c("Model","Sandwich"))
}

f <- function(a=1, b=1) {
  rep(a*b, 5)
}
R <- Expand(a=1:3, b=1:3)
sim(f, R)
sim(function(a,b) f(a,b), 3, args=c(a=5,b=5))
sim(function(iter=1,a=5,b=5) iter*f(a,b), iter=TRUE, R=5)

```

**Description**

Simulate data from a general SEM model including non-linear effects and general link and distribution of variables.

**Usage**

```
## S3 method for class 'lvm'
sim(x, n = NULL, p = NULL, normal = FALSE, cond = FALSE,
    sigma = 1, rho = 0.5, X = NULL, unlink=FALSE, latent=TRUE,
    use.labels = TRUE, seed=NULL, ...)
```

**Arguments**

x	Model object
n	Number of simulated values/individuals
p	Parameter value (optional)
normal	Logical indicating whether to simulate data from a multivariate normal distribution conditional on exogenous variables hence ignoring functional/distribution definition
cond	for internal use
sigma	Default residual variance (1)
rho	Default covariance parameter (0.5)
X	Optional matrix of fixed values of variables (manipulation)
unlink	Return Inverse link transformed data
latent	Include latent variables (default TRUE)
use.labels	convert categorical variables to factors before applying transformation
seed	Random seed
...	Additional arguments to be passed to the low level functions

**Author(s)**

Klaus K. Holst

**Examples**

```
#####
## Logistic regression
#####
m <- lvm(y~x+z)
regression(m) <- x~z
distribution(m,~y+z) <- binomial.lvm("logit")
d <- sim(m,1e3)
head(d)

e <- estimate(m,d,estimator="glm")
e
```



```

                                Binary.lvm(interval=list(c(0.3,0.5),c(0.8,1))))
sim(m,10)

#####
### Cox model
### piecewise constant hazard
#####
m <- lvm(t~x)
rates <- c(1,0.5); cuts <- c(0,5)
## Constant rate: 1 in [0,5), 0.5 in [5,Inf)
distribution(m,~t) <- coxExponential.lvm(rate=rates,timecut=cuts)

## Not run:
d <- sim(m,2e4,p=c("t~x"=0.1)); d$status <- TRUE
plot(timereg::aalen(survival::Surv(t,status)~x,data=d,
                    resample.iid=0,robust=0),spec=1)
L <- approxfun(c(cuts,max(d$t)),f=1,
               cumsum(c(0,rates*diff(c(cuts,max(d$t))))),
               method="linear")
curve(L,0,100,add=TRUE,col="blue")

## End(Not run)

#####
### Cox model
### piecewise constant hazard, gamma frailty
#####
m <- lvm(y~x+z)
rates <- c(0.3,0.5); cuts <- c(0,5)
distribution(m,~y+z) <- list(coxExponential.lvm(rate=rates,timecut=cuts),
                             loggamma.lvm(rate=1,shape=1))

## Not run:
d <- sim(m,2e4,p=c("y~x"=0,"y~z"=0)); d$status <- TRUE
plot(timereg::aalen(survival::Surv(y,status)~x,data=d,
                    resample.iid=0,robust=0),spec=1)
L <- approxfun(c(cuts,max(d$y)),f=1,
               cumsum(c(0,rates*diff(c(cuts,max(d$y))))),
               method="linear")
curve(L,0,100,add=TRUE,col="blue")

## End(Not run)
## Equivalent via transform (here with Aalens additive hazard model)
m <- lvm(y~x)
distribution(m,~y) <- aalenExponential.lvm(rate=rates,timecut=cuts)
distribution(m,~z) <- Gamma.lvm(rate=1,shape=1)
transform(m,t~y+z) <- prod
sim(m,10)
## Shared frailty
m <- lvm(c(t1,t2)~x+z)
rates <- c(1,0.5); cuts <- c(0,5)
distribution(m,~y) <- aalenExponential.lvm(rate=rates,timecut=cuts)
distribution(m,~z) <- loggamma.lvm(rate=1,shape=1)
## Not run:

```

```

mets::fast.reshape(sim(m,100),varying="t")

## End(Not run)

#####
### General multivariate distributions
#####
## Not run:
m <- lvm()
distribution(m,~y1+y2,oratio=4) <- VGAM::rbiplackcop
ksmooth2(sim(m,1e4),rgl=FALSE,theta=-20,phi=25)

m <- lvm()
distribution(m,~z1+z2,"or1") <- VGAM::rbiplackcop
distribution(m,~y1+y2,"or2") <- VGAM::rbiplackcop
sim(m,10,p=c(or1=0.1,or2=4))

## End(Not run)

m <- lvm()
distribution(m,~y1+y2+y3,TRUE) <- function(n,...) rmvN0(n,sigma=diag(3)+1)
var(sim(m,100))

## Syntax also useful for univariate generators, e.g.
m <- lvm(y~x+z)
distribution(m,~y,TRUE) <- function(n) rnorm(n,mean=1000)
sim(m,5)
distribution(m,~y,"m1",0) <- rnorm
sim(m,5)
sim(m,5,p=c(m1=100))

#####
### Regression design in other parameters
#####
## Variance heterogeneity
m <- lvm(y~x)
distribution(m,~y) <- function(n,mean,x) rnorm(n,mean,exp(x)^.5)
if (interactive()) plot(y~x,sim(m,1e3))
## Alternatively, calculate the standard error directly
addvar(m) <- ~sd ## If 'sd' should be part of the resulting data.frame
constrain(m,sd~x) <- function(x) exp(x)^.5
distribution(m,~y) <- function(n,mean,sd) rnorm(n,mean,sd)
if (interactive()) plot(y~x,sim(m,1e3))

## Regression on variance parameter
m <- lvm()
regression(m) <- y~x
regression(m) <- v~x
##distribution(m,~v) <- 0 # No stochastic term
## Alternative:
## regression(m) <- v[NA:0]~x
distribution(m,~y) <- function(n,mean,v) rnorm(n,mean,exp(v)^.5)
if (interactive()) plot(y~x,sim(m,1e3))

```

```

## Regression on shape parameter in Weibull model
m <- lvm()
regression(m) <- y ~ z+v
regression(m) <- s ~ exp(0.6*x-0.5*z)
distribution(m,~x+z) <- binomial.lvm()
distribution(m,~cens) <- coxWeibull.lvm(scale=1)
distribution(m,~y) <- coxWeibull.lvm(scale=0.1,shape=~s)
eventTime(m) <- time ~ min(y=1,cens=0)

if (interactive()) {
  d <- sim(m,1e3)
  require(survival)
  (cc <- coxph(Surv(time,status)~v+strata(x,z),data=d))
  plot(survfit(cc) ,col=1:4,mark.time=FALSE)
}

#####
### Categorical predictor
#####
m <- lvm()
## categorical(m,K=3) <- "v"
categorical(m,labels=c("A","B","C")) <- "v"

regression(m,additive=FALSE) <- y~v
## Not run:
plot(y~v,sim(m,1000,p=c("y~v:2"=3)))

## End(Not run)

m <- lvm()
categorical(m,labels=c("A","B","C"),p=c(0.5,0.3)) <- "v"
regression(m,additive=FALSE,beta=c(0,2,-1)) <- y~v
## equivalent to:
## regression(m,y~v,additive=FALSE) <- c(0,2,-1)
regression(m,additive=FALSE,beta=c(0,4,-1)) <- z~v
table(sim(m,1e4)$v)
glm(y~v, data=sim(m,1e4))
glm(y~v, data=sim(m,1e4,p=c("y~v:1"=3)))

transform(m,v2~v) <- function(x) x=='A'
sim(m,10)

#####
### Pre-calculate object
#####
m <- lvm(y~x)
m2 <- sim(m,'y~x'=2)
sim(m,10,'y~x'=2)
sim(m2,10) ## Faster

```

spaghetti

*Spaghetti plot***Description**

Spaghetti plot for longitudinal data

**Usage**

```
spaghetti(
  formula,
  data = NULL,
  id = "id",
  group = NULL,
  type = "o",
  lty = 1,
  pch = NA,
  col = 1:10,
  alpha = 0.3,
  lwd = 1,
  level = 0.95,
  trend.formula = formula,
  tau = NULL,
  trend.lty = 1,
  trend.join = TRUE,
  trend.delta = 0.2,
  trend = !is.null(tau),
  trend.col = col,
  trend.alpha = 0.2,
  trend.lwd = 3,
  trend.jitter = 0,
  legend = NULL,
  by = NULL,
  xlab = "Time",
  ylab = "",
  add = FALSE,
  ...
)
```

**Arguments**

formula	Formula (response ~ time)
data	data.frame
id	Id variable
group	group variable
type	Type (line 'l', stair 's', ...)

lty	Line type
pch	Colour
col	Colour
alpha	transparency (0-1)
lwd	Line width
level	Confidence level
trend.formula	Formula for trendline
tau	Quantile to estimate (trend)
trend.lty	Trend line type
trend.join	Trend polygon
trend.delta	Length of limit bars
trend	Add trend line
trend.col	Colour of trend line
trend.alpha	Transparency
trend.lwd	Trend line width
trend.jitter	Jitter amount
legend	Legend
by	make separate plot for each level in 'by' (formula, name of column, or vector)
xlab	Label of X-axis
ylab	Label of Y-axis
add	Add to existing device
...	Additional arguments to lower level arguments

**Author(s)**

Klaus K. Holst

**Examples**

```

if (interactive() & requireNamespace("mets")) {
  K <- 5
  y <- "y"%+%"seq(K)
  m <- lvm()
  regression(m,y=y,x=~u) <- 1
  regression(m,y=y,x=~s) <- seq(K)-1
  regression(m,y=y,x=~x) <- "b"
  N <- 50
  d <- sim(m,N); d$z <- rbinom(N,1,0.5)
  dd <- mets::fast.reshape(d); dd$num <- dd$num+3
  spaghetti(y~num,dd,id="id",lty=1,col=Col(1,.4),
            trend.formula=~factor(num),trend=TRUE,trend.col="darkblue")
  dd$num <- dd$num+rnorm(nrow(dd),sd=0.5) ## Unbalance
  spaghetti(y~num,dd,id="id",lty=1,col=Col(1,.4),
            trend=TRUE,trend.col="darkblue")
}

```

```
spaghetti(y~num,dd,id="id",lty=1,col=Col(1,.4),
          trend.formula=~num+I(num^2),trend=TRUE,trend.col="darkblue")
}
```

---

stack.estimate      *Stack estimating equations*

---

## Description

Stack estimating equations (two-stage estimator)

## Usage

```
## S3 method for class 'estimate'
stack(
  x,
  model2,
  D1u,
  inv.D2u,
  propensity,
  dpropensity,
  U,
  keep1 = FALSE,
  propensity.arg,
  estimate.arg,
  na.action = na.pass,
  ...
)
```

## Arguments

x	Model 1
model2	Model 2
D1u	Derivative of score of model 2 w.r.t. parameter vector of model 1
inv.D2u	Inverse of deri
propensity	propensity score (vector or function)
dpropensity	derivative of propensity score wrt parameters of model 1
U	Optional score function (model 2) as function of all parameters
keep1	If FALSE only parameters of model 2 is returned
propensity.arg	Arguments to propensity function
estimate.arg	Arguments to 'estimate'
na.action	Method for dealing with missing data in propensity score
...	Additional arguments to lower level functions

**Examples**

```

m <- lvm(z0~x)
Missing(m, z ~ z0) <- r~x
distribution(m,~x) <- binomial.lvm()
p <- c(r=-1, 'r~x'=0.5, 'z0~x'=2)
beta <- p[3]/2
d <- sim(m,500,p=p,seed=1)
m1 <- estimate(r~x,data=d,family=binomial)
d$w <- d$r/predict(m1,type="response")
m2 <- estimate(z~1, weights=w, data=d)
(e <- stack(m1,m2,propensity=TRUE))

```

subset.lvm

*Extract subset of latent variable model***Description**

Extract measurement models or user-specified subset of model

**Usage**

```

## S3 method for class 'lvm'
subset(x, vars, ...)

```

**Arguments**

x	lvm-object.
vars	Character vector or formula specifying variables to include in subset.
...	Additional arguments to be passed to the low level functions

**Value**

A lvm-object.

**Author(s)**

Klaus K. Holst

**Examples**

```

m <- lvm(c(y1,y2)~x1+x2)
subset(m,~y1+x1)

```

summary.sim

*Summary method for 'sim' objects***Description**

Summary method for 'sim' objects

**Usage**

```
## S3 method for class 'sim'
summary(
  object,
  estimate = NULL,
  se = NULL,
  confint,
  true = NULL,
  fun,
  names = NULL,
  unique.names = TRUE,
  minimal = FALSE,
  level = 0.95,
  quantiles = c(0, 0.025, 0.5, 0.975, 1),
  df = Inf,
  ...
)
```

**Arguments**

object	sim object
estimate	(optional) columns with estimates
se	(optional) columns with standard error estimates
confint	(optional) list of pairs of columns with confidence limits
true	(optional) vector of true parameter values
fun	(optional) summary function
names	(optional) names of estimates
unique.names	if TRUE, unique.names will be applied to column names
minimal	if TRUE, minimal summary will be returned
level	confidence level (0.95)
quantiles	quantiles (0,0.025,0.5,0.975,1)
df	degrees of freedom in t-distribution used for constructing CIs (default Gaussian approximation)
...	additional levels to lower-level functions

---

timedep	<i>Time-dependent parameters</i>
---------	----------------------------------

---

**Description**

Add time-varying covariate effects to model

**Usage**

```
timedep(object, formula, rate, timecut, type = "coxExponential.lvm", ...)
```

**Arguments**

object	Model
formula	Formula with rhs specifying time-varying covariates
rate	Optional rate parameters. If given as a vector this parameter is interpreted as the raw (baseline-)rates within each time interval defined by timecut. If given as a matrix the parameters are interpreted as log-rates (and log-rate-ratios for the time-varying covariates defined in the formula).
timecut	Time intervals
type	Type of model (default piecewise constant intensity)
...	Additional arguments to lower level functions

**Author(s)**

Klaus K. Holst

**Examples**

```
## Piecewise constant hazard
m <- lvm(y~1)
m <- timedep(m,y~1,timecut=c(0,5),rate=c(0.5,0.3))

## Not run:
d <- sim(m,1e4); d$status <- TRUE
dd <- mets::lifetable(Surv(y,status)~1,data=d,breaks=c(0,5,10));
exp(coef(glm(events ~ offset(log(atrisk)) + -1 + interval, dd, family=poisson)))

## End(Not run)

## Piecewise constant hazard and time-varying effect of z1
m <- lvm(y~1)
distribution(m,~z1) <- Binary.lvm(0.5)
R <- log(cbind(c(0.2,0.7,0.9),c(0.5,0.3,0.3)))
m <- timedep(m,y~z1,timecut=c(0,3,5),rate=R)
```

```

## Not run:
d <- sim(m,1e4); d$status <- TRUE
dd <- mets::lifetable(Surv(y,status)~z1,data=d,breaks=c(0,3,5,Inf));
exp(coef(glm(events ~ offset(log(atrisk)) + -1 + interval+z1:interval, dd, family=poisson)))

## End(Not run)

## Explicit simulation of time-varying effects
m <- lvm(y~1)
distribution(m,~z1) <- Binary.lvm(0.5)
distribution(m,~z2) <- binomial.lvm(p=0.5)
#variance(m,~m1+m2) <- 0
#regression(m,m1[m1:0] ~ z1) <- log(0.5)
#regression(m,m2[m2:0] ~ z1) <- log(0.3)
regression(m,m1 ~ z1,variance=0) <- log(0.5)
regression(m,m2 ~ z1,variance=0) <- log(0.3)
intercept(m,~m1+m2) <- c(-0.5,0)
m <- timedep(m,y~m1+m2,timecut=c(0,5))

## Not run:
d <- sim(m,1e5); d$status <- TRUE
dd <- mets::lifetable(Surv(y,status)~z1,data=d,breaks=c(0,5,Inf))
exp(coef(glm(events ~ offset(log(atrisk)) + -1 + interval + interval:z1, dd, family=poisson)))

## End(Not run)

```

---

toformula

*Converts strings to formula*


---

## Description

Converts a vector of predictors and a vector of responses (characters) into a formula expression.

## Usage

```
toformula(y = ".", x = ".")
```

## Arguments

y	vector of predictors
x	vector of responses

## Value

An object of class formula

**Author(s)**

Klaus K. Holst

**See Also**

[as.formula](#),

**Examples**

```
toformula(c("age", "gender"), "weight")
```

---

tr	<i>Trace operator</i>
----	-----------------------

---

**Description**

Calculates the trace of a square matrix.

**Usage**

```
tr(x, ...)
```

**Arguments**

x	Square numeric matrix
...	Additional arguments to lower level functions

**Value**

numeric

**Author(s)**

Klaus K. Holst

**See Also**

[crossprod](#), [tcrossprod](#)

**Examples**

```
tr(diag(1:5))
```

---

trim	<i>Trim string of (leading/trailing/all) white spaces</i>
------	---

---

**Description**

Trim string of (leading/trailing/all) white spaces

**Usage**

```
trim(x, all = FALSE, ...)
```

**Arguments**

x	String
all	Trim all whitespaces?
...	additional arguments to lower level functions

**Author(s)**

Klaus K. Holst

---

twindata	<i>Twin menarche data</i>
----------	---------------------------

---

**Description**

Simulated data

id	numeric	Twin-pair id
zyg	character	Zygosity (MZ or DZ)
twinnum	numeric	Twin number (1 or 2)
agemena	numeric	Age at menarche (or censoring)
status	logical	Censoring status (observed:=T,censored:=F)
bw	numeric	Birth weight
msmoke	numeric	Did mother smoke? (yes:=1,no:=0)

**Format**

data.frame

**Source**

Simulated

---

twostage	<i>Two-stage estimator</i>
----------	----------------------------

---

**Description**

Generic function.

**Usage**

```
twostage(object, ...)
```

**Arguments**

object	Model object
...	Additional arguments to lower level functions

**See Also**

twostage.lvm twostage.lvmfit twostage.lvm.mixture twostage.estimate

---

twostage.lvmfit	<i>Two-stage estimator (non-linear SEM)</i>
-----------------	---

---

**Description**

Two-stage estimator for non-linear structural equation models

**Usage**

```
## S3 method for class 'lvmfit'
twostage(
  object,
  model2,
  data = NULL,
  predict.fun = NULL,
  id1 = NULL,
  id2 = NULL,
  all = FALSE,
  formula = NULL,
  std.err = TRUE,
  ...
)
```

**Arguments**

object	Stage 1 measurement model
model2	Stage 2 SEM
data	data.frame
predict.fun	Prediction of latent variable
id1	Optional id-variable (stage 1 model)
id2	Optional id-variable (stage 2 model)
all	If TRUE return additional output (naive estimates)
formula	optional formula specifying non-linear relation
std.err	If FALSE calculations of standard errors will be skipped
...	Additional arguments to lower level functions

**Examples**

```

m <- lvm(c(x1,x2,x3)~f1,f1~z,
         c(y1,y2,y3)~f2,f2~f1+z)
latent(m) <- ~f1+f2
d <- simulate(m,100,p=c("f2,f2"=2,"f1,f1"=0.5),seed=1)

## Full MLE
ee <- estimate(m,d)

## Manual two-stage
## Not run:
m1 <- lvm(c(x1,x2,x3)~f1,f1~z); latent(m1) <- ~f1
e1 <- estimate(m1,d)
pp1 <- predict(e1,f1~x1+x2+x3)

d$u1 <- pp1[,]
d$u2 <- pp1[,]^2+attr(pp1,"cond.var")[1]
m2 <- lvm(c(y1,y2,y3)~eta,c(y1,eta)~u1+u2+z); latent(m2) <- ~eta
e2 <- estimate(m2,d)

## End(Not run)

## Two-stage
m1 <- lvm(c(x1,x2,x3)~f1,f1~z); latent(m1) <- ~f1
m2 <- lvm(c(y1,y2,y3)~eta,c(y1,eta)~u1+u2+z); latent(m2) <- ~eta
pred <- function(mu,var,data,...)
  cbind("u1"=mu[,1],"u2"=mu[,1]^2+var[1])
(mm <- twostage(m1,model2=m2,data=d,predict.fun=pred))

if (interactive()) {
  pf <- function(p) p["eta"]+p["eta~u1"]*u + p["eta~u2"]*u^2
  plot(mm,f=pf,data=data.frame(u=seq(-2,2,length.out=100)),lwd=2)
}

## Reduce test timing

```

```

## Splines
f <- function(x) cos(2*x)+x+-0.25*x^2
m <- lvm(x1+x2+x3~eta1, y1+y2+y3~eta2, latent=~eta1+eta2)
functional(m, eta2~eta1) <- f
d <- sim(m,500,seed=1,latent=TRUE)
m1 <- lvm(x1+x2+x3~eta1,latent=~eta1)
m2 <- lvm(y1+y2+y3~eta2,latent=~eta2)
mm <- twostage(m1,m2,formula=eta2~eta1,type="spline")
if (interactive()) plot(mm)

nonlinear(m2,type="quadratic") <- eta2~eta1
a <- twostage(m1,m2,data=d)
if (interactive()) plot(a)

kn <- c(-1,0,1)
nonlinear(m2,type="spline",knots=kn) <- eta2~eta1
a <- twostage(m1,m2,data=d)
x <- seq(-3,3,by=0.1)
y <- predict(a, newdata=data.frame(eta1=x))

if (interactive()) {
  plot(eta2~eta1, data=d)
  lines(x,y, col="red", lwd=5)

  p <- estimate(a,f=function(p) predict(a,p=p,newdata=x))$coefmat
  plot(eta2~eta1, data=d)
  lines(x,p[,1], col="red", lwd=5)
  confband(x,lower=p[,3],upper=p[,4],center=p[,1], polygon=TRUE, col=Col(2,0.2))

  l1 <- lm(eta2~splines::ns(eta1,knots=kn),data=d)
  p1 <- predict(l1,newdata=data.frame(eta1=x),interval="confidence")
  lines(x,p1[,1],col="green",lwd=5)
  confband(x,lower=p1[,2],upper=p1[,3],center=p1[,1], polygon=TRUE, col=Col(3,0.2))
}
## Reduce test timing

## Not run: ## Reduce timing
## Cross-validation example
ma <- lvm(c(x1,x2,x3)~u,latent=~u)
ms <- functional(ma, y~u, value=function(x) -.4*x^2)
d <- sim(ms,500)#,seed=1)
ea <- estimate(ma,d)

mb <- lvm()
mb1 <- nonlinear(mb,type="linear",y~u)
mb2 <- nonlinear(mb,type="quadratic",y~u)
mb3 <- nonlinear(mb,type="spline",knots=c(-3,-1,0,1,3),y~u)
mb4 <- nonlinear(mb,type="spline",knots=c(-3,-2,-1,0,1,2,3),y~u)
ff <- lapply(list(mb1,mb2,mb3,mb4),
  function(m) function(data,...) twostage(ma,m,data=data,st.derr=FALSE))
a <- cv(ff,data=d,rep=1)
a

```

```
## End(Not run)
```

---

twostageCV	<i>Cross-validated two-stage estimator</i>
------------	--

---

### Description

Cross-validated two-stage estimator for non-linear SEM

### Usage

```
twostageCV(
  model1,
  model2,
  data,
  control1 = list(trace = 0),
  control2 = list(trace = 0),
  knots.boundary,
  nmix = 1:4,
  df = 1:9,
  fix = TRUE,
  std.err = TRUE,
  nfolds = 5,
  rep = 1,
  messages = 0,
  ...
)
```

### Arguments

model1	model 1 (exposure measurement error model)
model2	model 2
data	data.frame
control1	optimization parameters for model 1
control2	optimization parameters for model 1
knots.boundary	boundary points for natural cubic spline basis
nmix	number of mixture components
df	spline degrees of freedom
fix	automatically fix parameters for identification (TRUE)
std.err	calculation of standard errors (TRUE)
nfolds	Number of folds (cross-validation)
rep	Number of repeats of cross-validation
messages	print information (>0)
...	additional arguments to lower

## Examples

```

## Reduce Ex.Timings##'
m1 <- lvm( x1+x2+x3 ~ u, latent= ~u)
m2 <- lvm( y ~ 1 )
m <- functional(merge(m1,m2), y ~ u, value=function(x) sin(x)+x)
distribution(m, ~u1) <- uniform.lvm(-6,6)
d <- sim(m,n=500,seed=1)
nonlinear(m2) <- y~u1
if (requireNamespace('mets', quietly=TRUE)) {
  set.seed(1)
  val <- twostageCV(m1, m2, data=d, std.err=FALSE, df=2:6, nmix=1:2,
                    nfolds=2)
  val
}

```

---

vars

---

*Extract variable names from latent variable model*


---

## Description

Extract exogenous variables (predictors), endogenous variables (outcomes), latent variables (random effects), manifest (observed) variables from a lvm object.

## Usage

```

vars(x,...)

endogenous(x,...)

exogenous(x,...)

manifest(x,...)

latent(x,...)

## S3 replacement method for class 'lvm'
exogenous(x, xfree = TRUE,...) <- value

## S3 method for class 'lvm'
exogenous(x,variable,latent=FALSE,index=TRUE,...)

## S3 replacement method for class 'lvm'
latent(x,clear=FALSE,...) <- value

```

**Arguments**

x	lvm-object
...	Additional arguments to be passed to the low level functions
variable	list of variables to alter
latent	Logical defining whether latent variables without parents should be included in the result
index	For internal use only
clear	Logical indicating whether to add or remove latent variable status
xfree	For internal use only
value	Formula or character vector of variable names.

**Details**

`vars` returns all variables of the `lvm-object` including manifest and latent variables. Similarly `manifest` and `latent` returns the observed resp. latent variables of the model. `exogenous` returns all manifest variables without parents, e.g. covariates in the model, however the argument `latent=TRUE` can be used to also include latent variables without parents in the result. Pr. default lava will not include the parameters of the exogenous variables in the optimisation routine during estimation (likelihood of the remaining observed variables conditional on the covariates), however this behaviour can be altered via the assignment function `exogenous<-` telling lava which subset of (valid) variables to condition on. Finally `latent` returns a vector with the names of the latent variables in `x`. The assignment function `latent<-` can be used to change the latent status of variables in the model.

**Value**

Vector of variable names.

**Author(s)**

Klaus K. Holst

**See Also**

[endogenous](#), [manifest](#), [latent](#), [exogenous](#), [vars](#)

**Examples**

```
g <- lvm(eta1 ~ x1+x2)
regression(g) <- c(y1,y2,y3) ~ eta1
latent(g) <- ~eta1
endogenous(g)
exogenous(g)
identical(latent(g), setdiff(vars(g),manifest(g)))
```

---

vec	<i>vec operator</i>
-----	---------------------

---

**Description**

vec operator

**Usage**

```
vec(x, matrix = FALSE, sep = ".", ...)
```

**Arguments**

x	Array
matrix	If TRUE a row vector (matrix) is returned
sep	Seperator
...	Additional arguments

**Details**

Convert array into vector

**Author(s)**

Klaus Holst

---

wait	<i>Wait for user input (keyboard or mouse)</i>
------	--

---

**Description**

Wait for user input (keyboard or mouse)

**Usage**

```
wait()
```

**Author(s)**

Klaus K. Holst

---

`wkm`*Weighted K-means*

---

**Description**

Weighted K-means via Lloyd's algorithm

**Usage**

```
wkm(  
  x,  
  mu,  
  data,  
  weights = rep(1, NROW(x)),  
  iter.max = 20,  
  n.start = 5,  
  init = "kmpp",  
  ...  
)
```

**Arguments**

<code>x</code>	Data (or formula)
<code>mu</code>	Initial centers (or number centers chosen randomly among <code>x</code> )
<code>data</code>	optional data frame
<code>weights</code>	Optional weights
<code>iter.max</code>	Max number of iterations
<code>n.start</code>	Number of restarts
<code>init</code>	method to create initial centres (default <code>kmeans++</code> )
<code>...</code>	Additional arguments to lower level functions

**Author(s)**

Klaus K. Holst

---

wrapvec	<i>Wrap vector</i>
---------	--------------------

---

**Description**

Wrap vector

**Usage**

```
wrapvec(x, delta = 0L, ...)
```

**Arguments**

x	Vector or integer
delta	Shift
...	Additional parameters

**Examples**

```
wrapvec(5,2)
```

---

zibreg	<i>Regression model for binomial data with unkown group of immortals</i>
--------	--

---

**Description**

Regression model for binomial data with unkown group of immortals (zero-inflated binomial regression)

**Usage**

```
zibreg(
  formula,
  formula.p = ~1,
  data,
  family = stats::binomial(),
  offset = NULL,
  start,
  var = "hessian",
  ...
)
```

**Arguments**

formula	Formula specifying
formula.p	Formula for model of disease prevalence
data	data frame
family	Distribution family (see the help page family)
offset	Optional offset
start	Optional starting values
var	Type of variance (robust, expected, hessian, outer)
...	Additional arguments to lower level functions

**Author(s)**

Klaus K. Holst

**Examples**

```
## Simulation
n <- 2e3
x <- runif(n,0,20)
age <- runif(n,10,30)
z0 <- rnorm(n,mean=-1+0.05*age)
z <- cut(z0,breaks=c(-Inf,-1,0,1,Inf))
p0 <- lava:::expit(model.matrix(~z+age) %>% c(-.4, -.4, 0.2, 2, -0.05))
y <- (runif(n)<lava:::ticol(-1+0.25*x-0*age))*1
u <- runif(n)<p0
y[u==0] <- 0
d <- data.frame(y=y,x=x,u=u*1,z=z,age=age)
head(d)

## Estimation
e0 <- zibreg(y~x*z,~1+z+age,data=d)
e <- zibreg(y~x,~1+z+age,data=d)
compare(e,e0)
e
PD(e0,intercept=c(1,3),slope=c(2,6))

B <- rbind(c(1,0,0,0,20),
           c(1,1,0,0,20),
           c(1,0,1,0,20),
           c(1,0,0,1,20))
prev <- summary(e,pr.contrast=B)$prevalence

x <- seq(0,100,length.out=100)
newdata <- expand.grid(x=x,age=20,z=levels(d$z))
fit <- predict(e,newdata=newdata)
plot(0,0,type="n",xlim=c(0,101),ylim=c(0,1),xlab="x",ylab="Probability(Event)")
count <- 0
for (i in levels(newdata$z)) {
  count <- count+1
}
```

```
lines(x, fit[which(newdata$z==i)], col="darkblue", lty=count)
}
abline(h=prev[3:4,1], lty=3:4, col="gray")
abline(h=prev[3:4,2], lty=3:4, col="lightgray")
abline(h=prev[3:4,3], lty=3:4, col="lightgray")
legend("topleft", levels(d$z), col="darkblue", lty=seq_len(length(levels(d$z))))
```

---

%++%

*Concatenation operator*

---

## Description

For matrices a block-diagonal matrix is created. For all other data types the operator is a wrapper of `paste`.

## Usage

```
x %++% y
```

## Arguments

x	First object
y	Second object of same class

## Details

Concatenation operator

## Author(s)

Klaus K. Holst

## See Also

`blockdiag`, [paste](#), [cat](#),

## Examples

```
## Block diagonal
matrix(rnorm(25),5)%++%matrix(rnorm(25),5)
## String concatenation
"Hello "%++%" World"
## Function composition
f <- log %++% exp
f(2)
```

---

`%ni%`*Matching operator (x not in y) oposed to the %in%-operator (x in y)*

---

**Description**

Matching operator

**Usage**`x %ni% y`**Arguments**

<code>x</code>	vector
<code>y</code>	vector of same type as <code>x</code>

**Value**

A logical vector.

**Author(s)**

Klaus K. Holst

**See Also**[match](#)**Examples**`1:10 %ni% c(1,5,10)`

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