Package 'FKF'

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Author David Luethi, Philipp Erb, Simon Otziger	
Maintainer Paul Smith <paul@waternumbers.co.uk></paul@waternumbers.co.uk>	
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Description This is a fast and flexible implementation of the Kalman filter, which can deal with NAs. It is entirely written in C and relies fully on linear algebra subroutines contained in BLAS and LAPACK. Due to the speed of the filter, the fitting of high-dimensional linear state space models to large datasets becomes possible. This package also contains a plot function for the visualization of the state vector and graphical diagnostics of the residuals.	
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fkf	,
Index	1

fkf Fast Kalman filter

Description

This function allows for fast and flexible Kalman filtering. Both, the measurement and transition equation may be multivariate and parameters are allowed to be time-varying. In addition "NA"-values in the observations are supported. fkf wraps the C-function FKF which fully relies on linear algebra subroutines contained in BLAS and LAPACK.

Usage

```
fkf(a0, P0, dt, ct, Tt, Zt, HHt, GGt, yt, check.input = TRUE)
```

Arguments

a0	A vector giving the initial value/estimation of the state variable.
P0	A matrix giving the variance of a0.
dt	A matrix giving the intercept of the transition equation (see Details).
ct	A matrix giving the intercept of the measurement equation (see Details).
Tt	An array giving the factor of the transition equation (see Details).
Zt	An array giving the factor of the measurement equation (see Details).
HHt	An array giving the variance of the innovations of the transition equation (see Details).
GGt	An array giving the variance of the disturbances of the measurement equation (see Details).
yt	A matrix containing the observations. "NA"-values are allowed (see Details).
check.input	A logical stating whether the input shall be checked for consistency ("storage.mode", "class", and dimensionality, see Details). This should always be 'TRUE' unless the performance becomes crucial and correctness of the arguments concerning dimensions, class and storage.mode is ensured.

Details

State space form

The following notation is closest to the one of Koopman et al. The state space model is represented by the transition equation and the measurement equation. Let m be the dimension of the state variable, d be the dimension of the observations, and n the number of observations. The transition equation and the measurement equation are given by

$$\alpha_{t+1} = d_t + T_t \cdot \alpha_t + H_t \cdot \eta_t$$
$$y_t = c_t + Z_t \cdot \alpha_t + G_t \cdot \epsilon_t,$$

where η_t and ϵ_t are iid $N(0, I_m)$ and iid $N(0, I_d)$, respectively, and α_t denotes the state variable. The parameters admit the following dimensions:

$$\begin{array}{lll} a_t in R^m & d_t in R^m & eta_t in R^m \\ T_t \in R^{m \times m} & H_t \in R^{m \times m} \\ y_t \in R^d & c_t \in R^d & \epsilon_t \in R^d \\ Z_t \in R^{d \times m} & G_t \in R^{d \times d} \end{array}$$

Note that fkf takes as input HHt and GGt which corresponds to H_tH_t' and G_tG_t' .

Iteration:

Let i be the loop variable. The filter iterations are implemented the following way (in case of no NA's):

```
Initialization: if(i == 1){ at[, i] = a0 Pt[,, i] = P0 }

Updating equations:
vt[, i] = yt[, i] - ct[, i] - Zt[,,i] %*% at[, i]
Ft[,, i] = Zt[,, i] %*% Pt[,, i] %*% t(Zt[,, i]) + GGt[,, i]
Kt[,, i] = Pt[,, i] %*% t(Zt[,, i]) %*% solve(Ft[,, i])
att[, i] = at[, i] + Kt[,, i] %*% vt[, i]
Ptt[, i] = Pt[,, i] - Pt[,, i] %*% t(Zt[,, i]) %*% t(Kt[,, i])

Prediction equations:
at[, i + 1] = dt[, i] + Tt[,, i] %*% att[, i]
Pt[,, i + 1] = Tt[,, i] %*% Ptt[,, i] %*% t(Tt[,, i]) + HHt[,, i]

Next iteration:
i <- i + 1
goto "Updating equations".</pre>
```

NA-values:

NA-values in the observation matrix yt are supported. If particular observations yt[,i] contain NAs, the NA-values are removed and the measurement equation is adjusted accordingly. When the full vector yt[,i] is missing the Kalman filter reduces to a prediction step.

Parameters:

The parameters can either be constant or deterministic time-varying. Assume the number of observations is n (i.e. $y=(y_t)_{t=1,...,n}, y_t=(y_{t1},...,y_{td})$). Then, the parameters admit the following classes and dimensions:

```
\begin{array}{ll} {\rm dt} & {\rm either~a~}m\times n~{\rm (time\mbox{-}varying)}~{\rm or~a~}m\times 1~{\rm (constant)}~{\rm matrix}. \\ {\rm Tt} & {\rm either~a~}m\times m\times n~{\rm or~a~}m\times m\times 1~{\rm array}. \\ {\rm HHt} & {\rm either~a~}m\times m\times n~{\rm or~a~}m\times m\times 1~{\rm array}. \\ {\rm ct} & {\rm either~a~}d\times n~{\rm or~a~}d\times 1~{\rm matrix}. \\ {\rm Zt} & {\rm either~a~}d\times m\times n~{\rm or~a~}d\times m\times 1~{\rm array}. \\ {\rm GGt} & {\rm either~a~}d\times d\times n~{\rm or~a~}d\times d\times 1~{\rm array}. \\ {\rm yt} & {\rm a~}d\times n~{\rm matrix}. \\ \end{array}
```

If check.input is TRUE each argument will be checked for correctness of the dimensionality, storage mode, and class. check.input should always be TRUE unless the performance becomes crucial and correctness of the arguments concerning dimensions, class, and storage.mode is ensured. Note that the class of the arguments if of importance. For instance, to check whether a parameter is constant the dim attribute is accessed. If, e.g., Zt is a constant, it could be a $d \times d$ -matrix. But the third dimension (i.e. dim(Zt)[3]) is needed to check for constancy. This requires Zt to be an

 $d \times d \times 1$ -array.

BLAS and LAPACK routines used:

The R function fkf basically wraps the C-function FKF, which entirely relies on linear algebra subroutines provided by BLAS and LAPACK. The following functions are used:

BLAS: dcopy, dgemm, daxpy. LAPACK: dpotri, dpotrf.

FKF is called through the .Call interface. Internally, FKF extracts the dimensions, allocates memory, and initializes the R-objects to be returned. FKF subsequently calls cfkf which performs the Kalman filtering.

The only critical part is to compute the inverse of F_t and the determinant of F_t . If the inverse can not be computed, the filter stops and returns the corresponding message in status (see **Value**). If the computation of the determinant fails, the filter will continue, but the log-likelihood (element logLik) will be "NA".

The inverse is computed in two steps: First, the Cholesky factorization of F_t is calculated by dpotrf. Second, dpotri calculates the inverse based on the output of dpotrf.

The determinant of F_t is computed using again the Cholesky decomposition.

Value

An S3-object of class "fkf", which is a list with the following elements:

```
att A m \times n-matrix containing the filtered state variables, i.e. a_{t|t} = E(\alpha_t|y_t).
```

at A $m \times (n+1)$ -matrix containing the predicted state variables, i.e. $a_t = E(\alpha_t | y_{t-1})$.

Ptt A $m \times m \times n$ -array containing the variance of att, i.e. $P_{t|t} = var(\alpha_t|y_t)$.

Pt A $m \times m \times (n+1)$ -array containing the variances of at, i.e. $P_t = var(\alpha_t|y_{t-1})$.

vt A $d \times n$ -matrix of the prediction errors given by $v_t = y_t - c_t - Z_t a_t$.

Ft A $d \times d \times n$ -array which contains the variances of vt, i.e. $F_t = var(v_t)$.

Kt A $m \times d \times n$ -array containing the "Kalman gain" (ambiguity, see calculation above).

logLik The log-likelihood.

status A vector which contains the status of LAPACK's dpotri and dpotrf. (0,0) means successful exit.

sys.time The time elapsed as an object of class "proc_time".

The first element of both at and Pt is filled with the function arguments a0 and P0, and the last, i.e.

the (n + 1)-th, element of at and Pt contains the predictions

$$at[, n+1] = E(\alpha_{n+1}|y_n)$$
 and $Pt[, n+1] = var(\alpha_{n+1}|y_n)$.

Usage

```
fkf(a0, P0, dt, ct, Tt, Zt, HHt, GGt, yt, check.input = TRUE)
```

References

Harvey, Andrew C. (1990). Forecasting, Structural Time Series Models and the Kalman Filter. Cambridge University Press.

Hamilton, James D. (1994). *Time Series Analysis*. Princeton University Press.

Koopman, S. J., Shephard, N., Doornik, J. A. (1999). *Statistical algorithms for models in state space using SsfPack* 2.2. Econometrics Journal, Royal Economic Society, vol. 2(1), pages 107-160.

See Also

plot to visualize and analyze fkf-objects, KalmanRun from the stats package, function dlmFilter from package dlm.

Examples

```
## <---->
## Example 1: ARMA(2, 1) model estimation.
## <----->
## This example shows how to fit an ARMA(2, 1) model using this Kalman
## filter implementation (see also stats' makeARIMA and KalmanRun).
n <- 1000
## Set the AR parameters
ar1 <- 0.6
ar2 < -0.2
ma1 <- -0.2
sigma <- sqrt(0.2)
## Sample from an ARMA(2, 1) process
a \leftarrow arima.sim(model = list(ar = c(ar1, ar2), ma = ma1), n = n,
              innov = rnorm(n) * sigma)
## Create a state space representation out of the four ARMA parameters
arma21ss <- function(ar1, ar2, ma1, sigma) {</pre>
   Tt \leftarrow matrix(c(ar1, ar2, 1, 0), ncol = 2)
   Zt \leftarrow matrix(c(1, 0), ncol = 2)
   ct <- matrix(0)
   dt \leftarrow matrix(0, nrow = 2)
   GGt <- matrix(0)</pre>
   H \leftarrow matrix(c(1, ma1), nrow = 2) * sigma
   HHt <- H %*% t(H)
   a0 < -c(0, 0)
   P0 \leftarrow matrix(1e6, nrow = 2, ncol = 2)
    return(list(a0 = a0, P0 = P0, ct = ct, dt = dt, Zt = Zt, Tt = Tt, GGt = GGt,
               HHt = HHt)
}
## The objective function passed to 'optim'
objective <- function(theta, yt) {</pre>
   sp <- arma21ss(theta["ar1"], theta["ar2"], theta["ma1"], theta["sigma"])</pre>
   ans \leftarrow fkf(a0 = sp$a0, P0 = sp$P0, dt = sp$dt, ct = sp$ct, Tt = sp$Tt,
              Zt = sp$Zt, HHt = sp$HHt, GGt = sp$GGt, yt = yt)
    return(-ans$logLik)
}
theta <- c(ar = c(0, 0), ma1 = 0, sigma = 1)
```

```
fit <- optim(theta, objective, yt = rbind(a), hessian = TRUE)
## Confidence intervals
rbind(fit$par - qnorm(0.975) * sqrt(diag(solve(fit$hessian))),
      fit$par + qnorm(0.975) * sqrt(diag(solve(fit$hessian))))
## Filter the series with estimated parameter values
sp <- arma21ss(fit$par["ar1"], fit$par["ar2"], fit$par["ma1"], fit$par["sigma"])</pre>
ans \leftarrow fkf(a0 = sp$a0, P0 = sp$P0, dt = sp$dt, ct = sp$ct, Tt = sp$Tt,
          Zt = sp$Zt, HHt = sp$HHt, GGt = sp$GGt, yt = rbind(a))
## Compare the prediction with the realization
plot(ans, at.idx = 1, att.idx = NA, CI = NA)
lines(a, lty = "dotted")
## Compare the filtered series with the realization
plot(ans, at.idx = NA, att.idx = 1, CI = NA)
lines(a, lty = "dotted")
## Check whether the residuals are Gaussian
plot(ans, type = "resid.qq")
## Check for linear serial dependence through 'acf'
plot(ans, type = "acf")
## <----->
## Example 2: Local level model for the Nile's annual flow.
## Transition equation:
## alpha[t+1] = alpha[t] + eta[t], eta[t] \sim N(0, HHt)
## Measurement equation:
## y[t] = alpha[t] + eps[t], eps[t] \sim N(0, GGt)
y <- Nile
y[c(3, 10)] \leftarrow NA + NA values can be handled
## Set constant parameters:
dt <- ct <- matrix(0)</pre>
Zt <- Tt <- matrix(1)</pre>
a0 <- y[1]
                   # Estimation of the first year flow
P0 <- matrix(100) # Variance of 'a0'
## Estimate parameters:
fit.fkf <- optim(c(HHt = var(y, na.rm = TRUE) * .5,
                  GGt = var(y, na.rm = TRUE) * .5),
                fn = function(par, ...)
                -fkf(HHt = matrix(par[1]), GGt = matrix(par[2]), ...)$logLik,
                yt = rbind(y), a0 = a0, P0 = P0, dt = dt, ct = ct,
                Zt = Zt, Tt = Tt)
## Filter Nile data with estimated parameters:
```

plot.fkf 7

plot.fkf

Plotting fkf objects

Description

Plotting method for objects of class fkf. This function provides tools for graphical analysis of the Kalman filter output: Visualization of the state vector, QQ-plot of the individual residuals, QQ-plot of the Mahalanobis distance, auto- as well as crosscorrelation function of the residuals.

Usage

```
## S3 method for class 'fkf'
plot(x, type = c("state", "resid.qq", "qqchisq", "acf"),
  CI = 0.95, at.idx = 1:nrow(x$at), att.idx = 1:nrow(x$att), ...)
```

Arguments

X	The output of fkf.
type	A string stating what shall be plotted (see Details).
CI	The confidence interval in case type $==$ "state". Set CI to NA if no confidence interval shall be plotted.
at.idx	An vector giving the indexes of the predicted state variables which shall be plotted if type $==$ "state".
att.idx	An vector giving the indexes of the filtered state variables which shall be plotted if type $==$ "state".
	Arguments passed to either plot, qqnorm, qqplot or acf.

8 plot.fkf

Details

The argument type states what shall be plotted. type must partially match one of the following:

state The state variables are plotted. By the arguments at.idx and att.idx, the user can specify which of the predicted (a_t) and filtered $(a_{t|t})$ state variables will be drawn.

resid.gq Draws a QQ-plot for each residual-series invt.

qqchisq A Chi-Squared QQ-plot will be drawn to graphically test for multivariate normality of the residuals based on the Mahalanobis distance.

acf Creates a pairs plot with the autocorrelation function (acf) on the diagonal panels and the crosscorrelation function (ccf) of the residuals on the off-diagnoal panels.

Value

Invisibly returns an list with components:

```
distance The Mahalanobis distance of the residuals as a vector of length n. std.resid The standardized residuals as an d \times n-matrix. It should hold that std.resid_{ij} iid \sim N_d(0, I),
```

where d denotes the dimension of the data and n the number of observations.

usage

```
plot(x, type = c("state", "resid.qq", "qqchisq", "acf"),CI = 0.95, at.idx = 1:nrow(x$at), att.idx = 1:
```

See Also

fkf

Examples

```
## Example 3: Local level model for the treering data
## Transition equation:
## alpha[t+1] = alpha[t] + eta[t], eta[t] \sim N(0, HHt)
## Measurement equation:
## y[t] = alpha[t] + eps[t], eps[t] \sim N(0, GGt)
y <- treering
y[c(3, 10)] \leftarrow NA + NA  values can be handled
## Set constant parameters:
dt <- ct <- matrix(0)</pre>
Zt \leftarrow Tt \leftarrow array(1,c(1,1,1))
a0 <- y[1]
                      # Estimation of the first width
P0 <- matrix(100) # Variance of 'a0'
## Estimate parameters:
fit.fkf <- optim(c(HHt = var(y, na.rm = TRUE) * .5,
                    GGt = var(y, na.rm = TRUE) * .5),
```

plot.fkf 9

Index

```
*Topic algebra
fkf, 2
*Topic hplot
plot.fkf, 7
*Topic models
fkf, 2
*Topic multivariate
fkf, 2
acf, 7, 8
ccf, 8
fkf, 2, 7, 8
KalmanRun, 5
plot, 5, 7
plot.fkf, 7
qqnorm, 7
qqplot, 7
```