# An R Companion to James Hamilton's "Time Series Analysis"

with R

Robert Bell, Matthieu Stigler

#### Foreword

RcompHam94 is an R package that implements many of the worked examples in Time Series Analysis (Hamilton, 1994) as well as providing access to the code and datasets used. In many cases RcompHam94 provides both simplified implmentations "from scratch" to allow the reader to explore the underlying logic and calculations, and more realistic implementations that make use of the large body of contributed packages in the Comprehensive R Archive Network (CRAN). Thus readers who have cut their teeth on the textbook can use this package as a stepping stone to doing their own analysis and/or research. Readers looking for additional introductory treatment of facilities available in CRAN can explore other excellent introductions such as http://cran.r-project.org/doc/contrib/Farnsworth-EconometricsInR.pdf and http://cran.r-project.org/web/packages/AER/AER.pdf (Kleiber and Zeileis, 2008).

We assume the reader has downloaded the R language, and package "RcompHam94" from http://www.r-project.org/ and has read "An Introduction to R" available here http://cran.r-project.org/doc/manuals/R-intro.html and also available as a PDF from the "Help" menu of the R package.

To load the package, just use:

```
library("RcompHam94")
```

Code shown in this document (and some not shown for brevity) can be executed using the R "demo" function. For a list of available demos, use:

```
demo(package = "RcompHam94")
```

To invoke a specific demo, say the demo called "p112", use:

```
demo(topic = "p112", package = "RcompHam94")
```

In general the demos are written so that the results of individual calculations can be examined after the fact by examining variables containing the results of those calculations.

Page references in the body of this document refer to  $\mathtt{Time}$   $\mathtt{Series}$   $\mathtt{Analy-sis}$ .

## Contents

1	Linear Difference Equations				
	1.1	Dynamic Multipliers for First Order Difference Equations $\ \ldots \ \ldots$	4		
	1.2	Comparing Transitory Versus Permanent Changes	5		
	1.3	Dynamic Multipliers for Second Order Difference Equations	6		
<b>2</b>	Stationary ARMA Processes				
	2.1	Autocorrelations for AR and MA Processes	7		
	2.2	R Facilities for ARMA Autocorrelations	9		
	2.3	Autocorrelations as a Function of the Moving Average Parameter	10		
	2.4	Realizations of ARMA Processes	11		
	2.5	R Facilities for simulating ARMA process	12		
3	Sample Autocorrelations and Partial Autocorrelations				
	3.1	A Box Jenkins Example	13		
	3.2	R Facilities for Sample Autocorrelations	15		
4	Spectral Analysis				
5	The Kalman Filter				
	5.1	Kalman Filtering Example Applied to Detecting Business Cycles	19		
	5.2	R facilities for Kalman Filtering	22		
6	Generalized Method of Moments				
	6.1	Classical Method of Moments	24		
	6.2	Generalized Method of Moments	24		
	6.3	R Facilities for Generalized Method of Moments	27		
7	Models of Nonstationary Time Series				
	7.1	Fractional Integration	27		
8	Univariate Processes with Unit Roots				
	8.1	Preamble	28		
	8.2	Dickey Fuller Tests for Unit Roots	29		
	8.3	Analyzing GNP data	32		
	8.4	Using Phillips Perron Tests	34		
	8.5	Augmented Dickey Fuller Tests	36		
	8.6	Example 17.10 - Bayesian Test of Autoregressive Coefficient	39		
	8.7	Determining Lag Length	39		
	8.8	R Facilities for Testing Unit Roots	40		

9	Cointegration				
	9.1	Testing Cointegration when the Cointegrating Vector is Known .	40		
	9.2	Estimating the Cointegrating Vector	48		
	9.3	Testing Hypotheses About the Cointegrating Vector	53		
10	Full-Information Maximum Likelihood Analysis of Cointegrated				
	Syst	ems	<b>55</b>		
	10.1	An Application of the Johansen Approach to the PPP data $$	55		
	10.2	Likelihood Ratio Tests on the Cointegration Vector	58		
11	Time Series Models of Heteroskedasticity				
	11.1	Preamble	60		
	11.2	Application of ARCH Models to US Fed Funds Data	62		
	11.3	R Facilities For GARCH models	66		
12	Mod	deling Time Series with Changes in Regime	66		
	12.1	Modeling Changes in Regime	66		
$R\epsilon$	fere	nces	69		

#### 1 Linear Difference Equations

```
> library("RcompHam94", lib.loc = ".")
```

# 1.1 Dynamic Multipliers for First Order Difference Equations

Page 3 describes calculations for dynamic multipliers for first order difference equations. An example of these calculations in action is given on page 4. A simple method to calculate dynamic multipliers is to simulate the difference equation calculating forward based on an initial shock at time t=1, assuming the value of y at time 0 is 0. R indexes arrays starting at 1 instead of 0, so subscripts are one more than the convention used in the text, meaning that the shock will be said to occur at time 2.

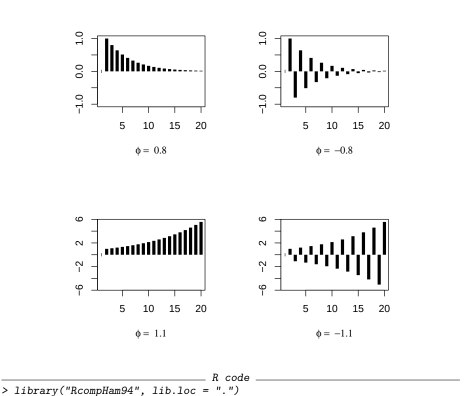
In the examples shown on page 4 there are actually four different equations being simulated, so we will use a matrix, rather than a vector, to store the results.

```
Phis <- c(0.8, -0.8, 1.1, -1.1)
> y <- array(dim = c(T, length(phis)))
> y[1, ] <- rep(0, length(phis))
> for (j in 2:T) y[j, ] <- phis * y[j - 1, ] + w[j]</pre>
```

We can check this calculation against the closed form expression on page 3.

```
output
[1] 1.00000000 0.80000000 0.64000000 0.51200000 0.40960000 0.32768000
[7] 0.26214400 0.20971520 0.16777216 0.13421773 0.10737418 0.08589935
[13] 0.06871948 0.05497558 0.04398047 0.03518437 0.02814750 0.02251800
[19] 0.01801440
```

Finally we can plot the results using a histogram plot reproducing figure 1.1.

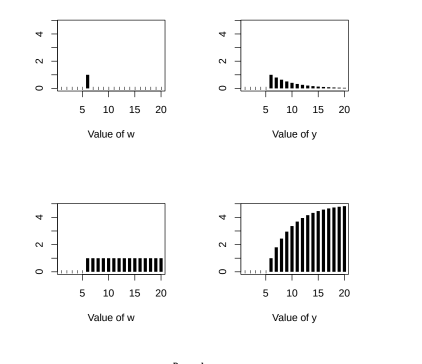


#### 1.2 Comparing Transitory Versus Permanent Changes

The above example examined the effect changing  $\phi$  on the dynamic multiplier. Pages 5 and 6 describe what happens when the permanence of the change is varied with a fixed multiplier, i.e. while leaving  $\phi$  unchanged.

```
R code
> phi <- 0.8
> T <- 20
> w <- 1 * cbind(1:T == 6, 1:T >= 6)
> y <- array(dim = c(T, 2))
> y[1:5, ] <- 0
> for (j in 6:T) y[j, ] <- phi * y[j - 1, ] + w[j, ]</pre>
```

The results can be plotted reproducing figures 1.2 and 1.3.



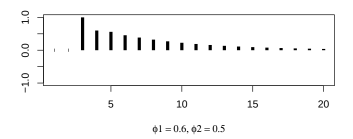
> library("RcompHam94", lib.loc = ".")

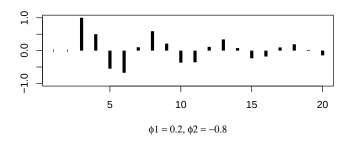
# 1.3 Dynamic Multipliers for Second Order Difference Equations

Finally we use similar techniques to calculate the effects of an impulse on a second order system. Here each column of phi represents the coefficients of a second order system.

```
R code
> T <- 20
> w <- 1 * (1:20 == 3)
> y <- array(dim = c(T, 2))
> y[1:2, ] <- 0
> phi <- array(c(0.6, 0.2, 0.5, -0.8), c(2, 2))
> for (j in 3:T) y[j, ] <- apply(X = phi * y[(j - 1):(j - 2), ],
+ MARGIN = 2, FUN = sum) + w[j]</pre>
```

The results can be plotted reproducing figure 1.4.





### 2 Stationary ARMA Processes

#### 2.1 Autocorrelations for AR and MA Processes

Pages 50 to 59 describe the calculation of autocorrelation functions of AR and MA processes. Following the expressions in the text we can calculate results using separate formulae for white noise, moving average, and autoregressive processes.

```
R code \longrightarrow 7 <- 20 

> specifications <- list(list(label = "White Noise", MA = vector(mode = "numeric"), AR = vector(mode = "numeric")), list(label = "MA(1)", MA = c(0.8), AR = vector(mode = "numeric")), list(label = "MA(4)", MA = c(-0.6, 0.5, -0.5, 0.3), AR = vector(mode = "numeric")), list(label = "AR(1) with 0.8", MA = vector(mode = "numeric"), AR = c(0.8)), list(label = "AR(1) with -0.8",
```

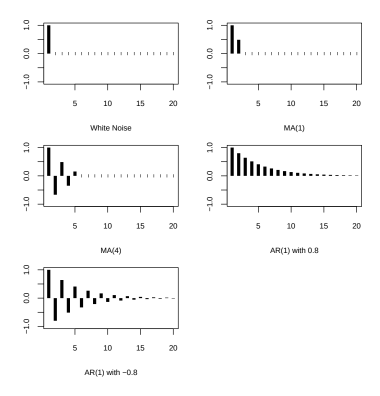
```
+ MA = vector(mode = "numeric"), AR = c(-0.8)))
> sigmasq < -1
```

White noise calculations are described on bottom of page 47 and the top of page 48

```
> specifications[[1]]$rho <- c(1, rep(0, T-1))
```

Moving average calculations are described on page 51.

Autocorrelation calculations are described on page 59



#### 2.2 R Facilities for ARMA Autocorrelations

Function ARMAacf can be used to calculate autocorrelations for an arbitrary ARMA process.

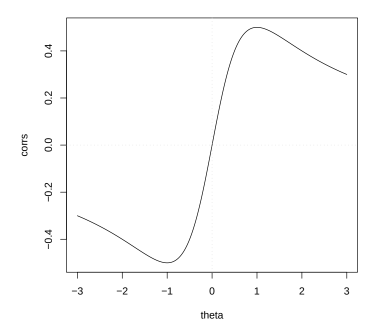
```
R code
> g3 <- ARMAacf(ar = numeric(0), ma = specifications[[3]]$MA, lag.max = T,
     pacf = FALSE)
> print(specifications[[3]]$rho)
                                 output
[1]
     1.0000000 -0.6666667
                            0.4871795 -0.3487179
                                                  0.1538462
                                                             0.000000
[7]
     0.0000000
                0.0000000
                            0.0000000 0.0000000
                                                             0.000000
                                                  0.0000000
[13]
     0.0000000
                 0.000000
                            0.0000000 0.0000000
                                                  0.000000
                                                             0.000000
[19]
     0.0000000
                0.0000000
                                R code -
> print(g3)
                    1
                                          3
1.0000000 -0.6666667
                       0.4871795 -0.3487179 0.1538462
                                                        0.0000000
                                                                   0.0000000
                                         10
                                                               12
                                                                          13
                                                    11
```

```
14
                        16
                                  17
                                           18
R code
> g4 <- ARMAacf(ar = specifications[[4]]$AR, ma = numeric(0), lag.max = T -
    1, pacf = FALSE)
> print(specifications[[4]]$rho)
                           output
[1] 1.00000000 0.80000000 0.64000000 0.51200000 0.40960000 0.32768000
[7] 0.26214400 0.20971520 0.16777216 0.13421773 0.10737418 0.08589935
[13] 0.06871948 0.05497558 0.04398047 0.03518437 0.02814750 0.02251800
[19] 0.01801440 0.01441152
                          _{-} R code _{-}
> print(g4)
1.00000000 0.80000000 0.64000000 0.51200000 0.40960000 0.32768000 0.26214400
                                  10
0.20971520 0.16777216 0.13421773 0.10737418 0.08589935 0.06871948 0.05497558
      14
               15
                        16
                                  17
                                           18
0.04398047 0.03518437 0.02814750 0.02251800 0.01801440 0.01441152
                           R code
> library("RcompHam94", lib.loc = ".")
```

### 2.3 Autocorrelations as a Function of the Moving Average Parameter

Figure 3.2 is easily generated from the formula for autocorrelations of an MA(1) process.

```
R code
> theta <- (-300:300) * 0.01
> corrs <- theta/(1 + theta^2)
> plot(theta, corrs, type = "l")
> grid(nx = 2, ny = 2)
```



```
> library("RcompHam94", lib.loc = ".")
```

#### 2.4 Realizations of ARMA Processes

Pages 55 shows some realizations of AR processes. We will assume the innovations are drawn from a standard normal distribution.

```
R code

> specifications <- list(list(label = "f = 0", MA = vector(mode = "numeric"),

+ AR = vector(mode = "numeric")), list(label = "f = .5", MA = vector(mode = "numeric"),

+ AR = c(0.5)), list(label = "f = .9", MA = vector(mode = "numeric"),

+ AR = c(0.9)))

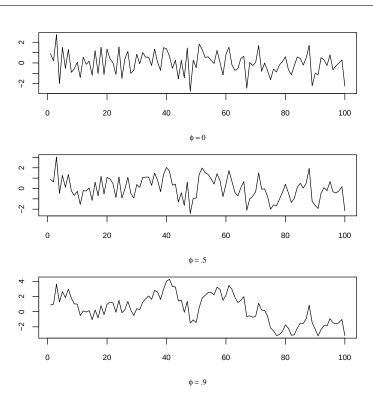
> T <- 100

> epsilon <- rnorm(T, 0, 1)
```

These can be calculated by iterating forward on the defining equations.

```
R code
> simulate.forward <- function(specification, epsilon) {
+ T <- length(epsilon)
+ AR <- specification$AR
+ MA <- specification$MA
```

```
+ presample <- rep(0, max(length(AR), length(MA)))
+ epsilon <- c(presample, epsilon)
+ Y <- vector(mode = "numeric", length = T + length(presample))
+ Y[1:length(presample)] <- 0
+ for (i in (length(presample) + 1):(T + length(presample))) Y[i] <- epsilon[[i]] +
+ ifelse(length(AR) > 0, t(AR) %*% Y[(i - 1):(i - length(AR))],
+ 0) + ifelse(length(MA) > 0, t(MA) %*% epsilon[(i -
+ 1):(i - length(MA))], 0)
+ Y[(length(presample) + 1):(T + length(presample))]
+ }
> for (i in 1:length(specifications)) specifications[[i]]$Y <- simulate.forward(specifications[[i]],
+ epsilon)</pre>
```



#### 2.5 R Facilities for simulating ARMA process

Function "simulate.forward" is a special case of capabilities provided by the function arima.sim in package stats, as the following code verifies.

```
R code

> for (specification in specifications) {

+ AR <- specification$AR

+ MA <- specification$MA
```

```
shift <- max(length(AR), length(MA))</pre>
     Y \leftarrow arima.sim(model = list(order = c(length(AR), 0, length(MA)),
         ar = AR, ma = MA), n = T, innov = epsilon[1:T], n.start = max(shift,
         1), start.innov = rep(0, max(shift, 1)))
     print(specification$Y[1:10])
     print(Y[1:10])
+ }
                              output
[1] 0.8770062 0.1957487 2.7218685 -1.9778921 1.4930980 -0.5158801
[7] 1.3022081 -0.9023128 -0.5719365 0.0748756
[1] 0.8770062 0.1957487 2.7218685 -1.9778921 1.4930980 -0.5158801
[7] 1.3022081 -0.9023128 -0.5719365 0.0748756
 [1] 0.8770062 0.6342518 3.0389944 -0.4583949
                                             1.2639006 0.1160702
[7] 1.3602432 -0.2221912 -0.6830321 -0.2666404
[7] 1.3602432 -0.2221912 -0.6830321 -0.2666404
[1] 0.8770062 0.9850542 3.6084173 1.2696835 2.6358132 1.8563518 2.9729247
[8] 1.7733195 1.0240510 0.9965215
[1] 0.8770062 0.9850542 3.6084173 1.2696835 2.6358132 1.8563518 2.9729247
[8] 1.7733195 1.0240510 0.9965215
```

# 3 Sample Autocorrelations and Partial Autocorrelations

```
R code
> library("RcompHam94", lib.loc = ".")
```

#### 3.1 A Box Jenkins Example

Example 4.1 from page 112 illustrates the Box-Jenkins approach based on autocorrelations. Here the data series is log changes of seasonally adjusted real US GNP from 1947 to 1988, available by simple transformations of the data in object "gnp1996". The data is prepared by selecting quarterly date from as shown, then computing the log of differences.

```
R code
> data(gnp1996, package = "RcompHam94")
> selection <- subset(gnp1996, Quarter >= "1947-01-01" & Quarter <=
+ "1988-10-01")
> y <- diff(log(selection$GNPH))
```

Page 110 shows how to compute sample autocorrelations - we will generate the first 20 to be used in plotting the results below.

```
R code
> max.lags <- 20
> T <- length(y)
> threshold <- 2/sqrt(T)
> gammas <- vector(mode = "numeric", length = max.lags + 1)
> gammas[[1]] <- 1/T * t(y - mean(y)) %*% (y - mean(y))
> for (j in 1:max.lags) gammas[j + 1] <- 1/T * t((y - mean(y))[(j + 1):T]) %*% (y - mean(y))[1:(T - j)]
> rhos <- gammas/gammas[[1]]</pre>
```

Page 111 shows how to compute sample partial autocorrelations.

A plot of the outputs reproducing figure 4.2 is shown below. The source code is provided in the demo.

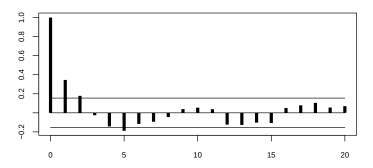


Figure 4.2(a) Sample autocorrelations

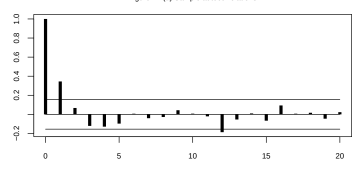


Figure 4.2(b) Sample partial autocorrelations

#### 3.2 R Facilities for Sample Autocorrelations

Function acf from R package "stats" performs the same function as acf, as we can readily confirm.

```
R code
> acf.correlation <- acf(y, lag.max = max.lags, type = "correlation",
+ plot = FALSE, demean = TRUE)</pre>
```

> print(as.vector(acf.correlation\$acf))

[19] 0.10451845 0.05540046 0.07001701

```
> print(rhos) R code ______
```

\_\_\_\_\_ output \_\_\_\_\_ [1] 1.00000000 0.34509475 0.17817758 -0.02537843 -0.14230681 -0.18827409

 $[7] \ -0.11613672 \ -0.09335581 \ -0.04441490 \ \ 0.03902657 \ \ 0.05412612 \ \ 0.03788102$ 

<sup>[7] -0.11613672 -0.09335581 -0.04441490 0.03902657 0.05412612 0.03788102</sup> 

<sup>[13] -0.12386994 -0.12725888 -0.10256196 -0.10719806 0.05022865 0.07874423</sup> 

```
[13] -0.12386994 -0.12725888 -0.10256196 -0.10719806 0.05022865 0.07874423
[19] 0.10451845 0.05540046 0.07001701
                                R code -
> acf.partial <- acf(y, lag.max = max.lags, type = "partial", plot = FALSE,
     demean = TRUE)
> print(as.vector(acf.partial$acf))
                               output
[1] 0.345094750 0.067075208 -0.120748043 -0.128609341 -0.096659383
 [6] 0.006935269 -0.040052970 -0.027544630 0.043507786 0.007543470
[11] -0.020592065 -0.186352407 -0.053599417 0.009939122 -0.066137883
     0.093638650 0.007111983 0.016895000 -0.045185857 0.023227306
                               _ R code _
> print(alphas)
                               output
     0.345094750
                  0.067075208 -0.120748043 -0.128609341 -0.096659383
[6] 0.006935269 -0.040052970 -0.027544630 0.043507786 0.007543470
[11] -0.020592065 -0.186352407 -0.053599417
                                            0.009939122 -0.066137883
Г16Т
    0.093638650 0.007111983 0.016895000 -0.045185857 0.023227306
```

### 4 Spectral Analysis

```
R code ______
> library("RcompHam94", lib.loc = ".")
```

Pages 167 to 170 give an example of the uses of spectral analysis, as applied to US Industrial Production from January 1947 to November 1989, available in data source "indprod". We will analyze the actual raw data, as well as one month and one year log changes.

```
R code

> data(indprod, package = "Ham94")

> selection <- subset(indprod, Month >= "1947-01-01" & Month <= 
+ "1989-11-01")

> raw.data <- selection$IPMFG6

> logdiff.data <- 100 * diff(log(raw.data), lag = 1)

> yeardiff.data <- 100 * diff(log(raw.data), lag = 12)
```

For plotting purposes, generate frequencies at regular intervals as show on page 159. The first spectrum uses unsmoothed estimates, the last two use a Bartlett kernel.

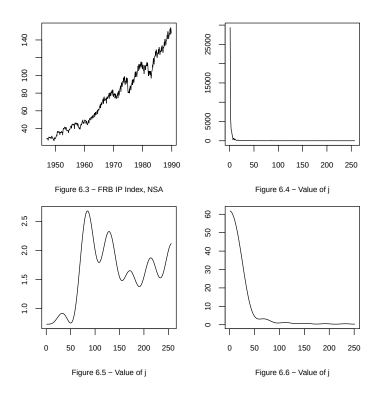
We show this in two ways:

- Step by step function (page 17)
- Built-in function (page 18)

#### Step by step function

```
_{-} R code _{-}
> s.Y.omega <- function(omega, gammas, params) {
     1/(2 * pi) * (gammas[[1]] + 2 * as.numeric(t(gammas[-1]) %*%
         cos(1:(length(gammas) - 1) * omega)))
+ }
> s.Y.omega.Bartlett <- function(omega, gammas, params) {
      1)) * gammas[2:(params + 1)]) %*% cos(1:params * omega)))
+ }
> generate.plot.data <- function(values, estimator, params) {</pre>
     T <- length(values)</pre>
     acf.covariance <- acf(values, lag.max = T - 1, type = "covariance",</pre>
         plot = FALSE, demean = TRUE)
     sapply(2 * pi/T * 1:((T - 1)/2), estimator, as.vector(acf.covariance$acf),
         params)
+ }
> raw.s.Y.omega <- generate.plot.data(raw.data, s.Y.omega, NULL)
\verb|> logdiff.s.Y.omega <- generate.plot.data(logdiff.data, s.Y.omega.Bartlett,
     12)
> yeardiff.s.Y.omega <- generate.plot.data(yeardiff.data, s.Y.omega.Bartlett,
```

The resulting output is shown below.



**Built-in function** We use here the function spectrum:

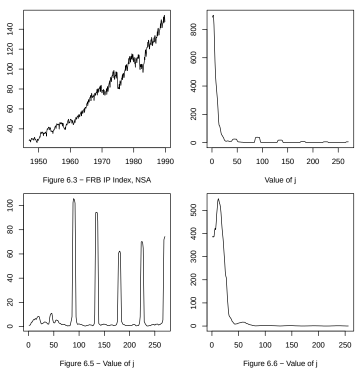
R code
> args(spectrum)

\_\_\_\_\_\_ output
function (x, ..., method = c("pgram", "ar"))

NULL

R code
> sp <- spectrum(raw.data, plot = FALSE, span = 10)
> x <- 100 \* diff(log(raw.data))
> sp2 <- spectrum(x, span = 6, plot = FALSE)
> x12 <- 100 \* diff(log(raw.data), lag = 12)
> sp3 <- spectrum(x12, span = 20, plot = FALSE)</pre>





#### 5 The Kalman Filter

```
R code 
> library("RcompHam94", lib.loc = ".")
```

# 5.1 Kalman Filtering Example Applied to Detecting Business Cycles

Page 376 describes an application of Kalman filtering to business cycles by James Stock and Mark Watson.

This can be implemented in two steps. The first is to implement the Kalman algorithm as described in the text. The following function follows the notation in Chapter 13.

```
R code

> kalman <- function(H, R, F, x, A, y, Q, xi.1.0, P.1.0) {

+ T <- dim(x)[[2]]

+ P.t.t_1 <- array(dim = c(dim(P.1.0), T + 1))

+ P.t.t_1[, , 1] <- P.1.0

+ P.t.t <- array(dim = c(dim(P.1.0), T))
```

```
K.t \leftarrow array(dim = c(dim(H), T))
      xi.t.t_1 \leftarrow array(dim = c(length(xi.1.0), T + 1))
      xi.t.t_1[, 1] <- xi.1.0
      xi.t.t \leftarrow array(dim = c(length(xi.1.0), T))
      L <- 0
      for (tt in 1:T) {
          V \leftarrow solve(t(H) %*% P.t.t_1[, , tt] %*% H + R)
          K.t[, , tt] \leftarrow P.t.t_1[, , tt] %*% H %*% V
          P.t.t[, , tt] \leftarrow P.t.t_1[, , tt] - K.t[, , tt] %*% t(H) %*%
              P.t.t_1[, , tt]
          P.t.t_1[, , tt + 1] \leftarrow F %*% P.t.t[, , tt] %*% t(F) +
          w \leftarrow y[, tt] - t(A) %*% x[, tt] - t(H) %*% xi.t.t_1[,
              tt]
          xi.t.t[, tt] \leftarrow xi.t.t_1[, tt] + K.t[, , tt] %*% w
          xi.t.t_1[, tt + 1] <- F %*% xi.t.t[, tt]
          L \leftarrow L - 1/2 * dim(y)[[1]] * log(2 * pi) + 1/2 * log(det(V)) -
              1/2 * t(w) %*% V %*% w
      }
      xi.t.T \leftarrow array(dim = c(length(xi.1.0), T))
      xi.t.T[, T] \leftarrow xi.t.t[, T]
      P.t.T \leftarrow array(dim = c(dim(P.1.0), T))
      P.t.T[, , T] \leftarrow P.t.t[, , T]
      for (tt in (T-1):1) {
          +
          xi.t.T[, tt] \leftarrow xi.t.t[, tt] + Jt %*% (xi.t.T[, tt +
              1] - xi.t.t_1[, tt + 1])
          P.t.T[, , tt] <- P.t.t[, , tt] + Jt %*% (P.t.T[, , tt +
+
              1] - P.t.t_1[, , tt + 1]) %*% t(Jt)
      list(xi.t.t = xi.t.t, xi.t.t_1 = xi.t.t_1, P.t.t = P.t.t,
          P.t.t_1 = P.t.t_1, K.t = K.t, log.likelihood = L, xi.t.T = xi.t.T,
          P.t.T = P.t.T)
+ }
```

The second is to specify the state space model as described on pp376-377 and estimate the parameters via maximum likelihood. Data for this analysis is consumption and income data form dataset "coninc" in log differences.

```
> data(coninc, package = "Ham94")
> YGR <- diff(log(coninc$GYD82))</pre>
```

```
> CGR <- diff(log(coninc$GC82))
> y <- t(cbind(YGR - mean(YGR), CGR - mean(CGR)))</pre>
```

The following helper function converts the parameters from a vector of labeled components into the correct inputs for the filter as shown in equations [13.1.28], [13.1.29], and [13.1.30].

The objective function is the log.likelihood obtained from the Kalman iteration.

```
R code
> objective <- function(THETA, y) {

+ params <- theta.y.to.params(THETA, y)

+ kalman(params$H, params$R, params$F, params$x, params$A,

+ y, params$Q, params$xi.1.0, params$P.1.0)$log.likelihood

+ }

> optimizer.results <- optim(par = THETA, fn = objective, gr = NULL,

+ y = y, control = list(trace = 0))</pre>
```

Finally calculate the smoothed results based on the ML estimated parameters.

```
R code

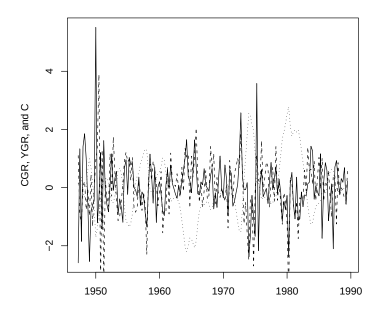
> params <- theta.y.to.params(optimizer.results$par, y)

> smoothed.results <- kalman(params$H, params$R, params$F, params$x,

+ params$A, y, params$Q, params$xi.1.0, params$P.1.0)

> smoothed.data <- smoothed.results$xi.t.T[1, ]
```

The results of the smoothed inference are shown below.



#### 5.2 R facilities for Kalman Filtering

There are several different packages in R for Kalman filtering, some that provide univariate support, others multivariate support. For example, package FKF is a fast implementation, but there are others. One key aspect of using such packages is specifying an interface to allow for time varying inputs, and providing results under those conditions. Some packages use caller supplied functions, others check for dimensions of (up to three dimensional) arrays, etc.

For example, a simple implementation of the example on page 382 using function "kalman" above might look like:

```
R code
> sigmasq <- 2
> params <- list(F = array(c(0, 1, 0, 0), c(2, 2)), Q = diag(c(sigmasq,
+ 0)), H = array(c(1, 0.8), c(2, 1)), R = array(0, c(1, 1)),
+ A = array(0.5, c(1, 1)), x = 1 %0% rep(1, 5), y = 1 %0% c(1,
+ seq(0.5, 4)), xi.1.0 = c(0, 0))
> params <- c(params, list(P.1.0 = array(solve(diag(length(params$xi.1.0)^2) -
+ params$F %x% params$F, as.vector(params$Q)), c(length(params$xi.1.0),
+ length(params$xi.1.0))))
> myResults <- kalman(params$H, params$R, params$F, params$x, params$A,
+ params$y, params$Q, params$xi.1.0, params$P.1.0)</pre>
```

We can perform the some operations using package FKF with a slight alteration of the function arguments. In particular, many of the arguments using an outer product as a quick way to convert them into a structure of one additional dimension, with the length of the additional dimension being 1. This is a convenient calling convention to specifying a \*non\* time varying parameter. If the parameter \*were\* time varying then the full extra dimension would be used. For example, the F matrix can be time varying in FKF (called Tt). A call exploiting this would then have a vector of two dimensional F matrices, one for each time index, i.e. a three dimensional array. If F is not time varying, (as in the case of the simple example above) then a three dimensional array with the third dimension being of length 1 is used.

```
R code
 fkfResults <- FKF::fkf(a0 = params$xi.1.0, P0 = params$P.1.0,</pre>
      dt = rep(0, length(params$xi.1.0)) %o% 1, Tt = params$F %o%
          1, HHt = params$Q %o% 1, ct = t(params$A) %*% params$x,
      Zt = t(params$H) \%0\% 1, GGt = params$R \%0\% 1, yt = params$y,
      check.input = TRUE)
The results can be confirmed by examing the output:
                                  R code -
> print(myResults$xi.t.t)
                                   output
                                            [,4]
          [,1]
                      [,2]
                                  [,3]
                                                     [,5]
[1,] 0.3048780 -0.1951600 1.02502699 1.100137 2.031900
                0.2439500 -0.03128374 1.124828 1.210125
                                  R code -
> print(fkfResults$att)
                                   output
          [,1]
                      [,2]
                                            [,4]
                                  [,3]
[1,] 0.3048780 -0.1951600 1.02502699 1.100137 2.031900
[2,] 0.2439024 0.2439500 -0.03128374 1.124828 1.210125
```

#### 6 Generalized Method of Moments

```
> library("RcompHam94", lib.loc = ".")
```

#### 6.1 Classical Method of Moments

Pages 409-410 gives a simple example of estimating the degrees of freedom of a standard t distribution. To illustrate, first generate a sample of 500 observations from a t distribution with 10 degrees of freedom.

```
_ R code _
> Y <- rt(500, 10)
Then maximize the sum of logs of a t density evaluated on the sample points.
                                  R code -
> objective <- function(nu, Y) {</pre>
      -sum(log(dt(Y, df = nu)))
+ }
> classical.results <- optimize(interval = c(1, 30), f = objective,
      Y = Y
> mu2 <- mean(Y^2)
> nu <- 2 * mu2/(mu2 - 1)
> print(classical.results)
                                 _ output _
$minimum
[1] 7.040553
$objective
[1] 780.6777
                                 _ R code _
> print(nu)
```

#### 6.2 Generalized Method of Moments

[1] 7.19724

Using the sample sample, we can estimate the degrees of freedom using GMM. To this end define a function following the GMM recipe in the text.

 $\_$  output  $\_$ 

```
R code
> compute.estimates <- function(Y, h, interval) {
+     g <- function(Y, THETA) {
+     apply(X = apply(X = Y, MARGIN = 1, FUN = h, THETA = THETA),
+     MARGIN = 1, FUN = mean)
+  }</pre>
```

```
objective <- function(THETA, Y, W) {
          g.value <- g(Y, THETA)</pre>
+
          t(g.value) %*% W %*% g.value
      r <- length(h(Y[1, ], interval[[1]]))</pre>
      a <- length(interval[[1]])</pre>
      T <- dim(Y)[[1]]
      stage.1.results <- optimize(interval = interval, f = objective,</pre>
          Y = Y, W = diag(r)
      temp <- apply(X = Y, MARGIN = 1, FUN = h, THETA = stage.1.results$objective)
      S \leftarrow 1/T * temp %*% t(temp)
      stage.2.results <- optimize(interval = interval, f = objective,</pre>
          Y = Y, W = solve(S)
      J.test \leftarrow 1 - pchisq(T * stage.2.results$objective, r - a)
      list(stage.1.results = stage.1.results, stage.2.results = stage.2.results,
          overidentifying = J.test)
+ }
```

Using this function is then a matter of specifying an appropriate function h to define an observation of the set of moments being targeted.

```
R code
> h <- function(Yt, THETA) {
+    nu <- THETA
+    c(Yt^2 - nu/(nu - 2), Yt^4 - 3 * nu^2/((nu - 2) * (nu - 4)))
+ }
> estimates <- compute.estimates(Y %o% 1, h, interval = c(5, 30))
> print(estimates)
```

```
output ____
```

```
$stage.1.results
```

\$stage.1.results\$minimum

[1] 7.564196

\$stage.1.results\$objective

[,1]

[1,] 0.0006450114

\$stage.2.results

\$stage.2.results\$minimum

[1] 7.727134

\$stage.2.results\$objective

A second example estimates the shape parameter of a two-sided gamma distribution.

```
_ R code -
> Yg <- rgamma(500, 10) * sign(runif(500, -1, 1))
> hg <- function(Yt, THETA) {</pre>
      k <- THETA
     nu <- k
     mu <- k
     sigma <- k
     skew <- 2/sqrt(k)
     kurt <- 6/k
      c(Yt^2 - sigma - mu^2, Yt^4 - (kurt * (sigma^2) + 3) - 4 *
          (skew * sigma^1.5) * mu - 6 * sigma * mu^2 - mu^4)
+ }
> gestimates <- compute.estimates(Yg %o% 1, hg, interval = c(5,
      30))
> print(gestimates)
                                _ output _
$stage.1.results
$stage.1.results$minimum
[1] 10.02213
$stage.1.results$objective
           [,1]
[1,] 0.08514715
$stage.2.results
$stage.2.results$minimum
[1] 10.02392
```

#### 6.3 R Facilities for Generalized Method of Moments

 $\operatorname{TBD}$ 

### 7 Models of Nonstationary Time Series

R code
> library("RcompHam94", lib.loc = ".")

#### 7.1 Fractional Integration

This example uses package fracdiff to compute the exponent of fractional integration as described on pp448-449. Data is US GDP and Treasury Yields.

	_ R code
> data(gnptbill, package = "Ham9	
<pre>&gt; print(fdGPH(gnptbill\$GNP))</pre>	
	output
\$d	
[1] 0.9588756	
\$sd.as	
[1] 0.2427173	
\$sd.reg	
[1] 0.04061276	
	_ R code
> print(fdGPH(gnptbill\$TBILL))	- It code
	output
\$d	output
[1] 0.9511594	

```
$sd.as
[1] 0.2427173

$sd.reg
[1] 0.227921
```

#### 8 Univariate Processes with Unit Roots

#### 8.1 Preamble

This section uses a few utility functions that follow procedures in the test for testing hypotheses about unit roots. First is the Newey West estimator described by [10.5.10] and [10.5.15].

Next are the Dickey Fuller stats described in [17.4.7] and [17.4.9], with an optional correction for serial correlation defined in [17.7.35] and [17.7.38].

```
}
<environment: namespace:RcompHam94>
The Phillips Perron stats are defined by [17.6.8] and [17.6.12]
                                 _ R code _
> print(Phillips.Perron)
                                  output -
function (T, rho, sigma.rho, s, lambda.hat.sq, gamma0)
    list(T = T, rho = rho, sigma.rho = sigma.rho, s.sq = s^2,
        lambda.hat.sq = lambda.hat.sq, gamma0 = gamma0, rho.stat = T *
            (rho - 1) - 1/2 * (T * sigma.rho/s)^2 * (lambda.hat.sq -
            gamma0), t.stat = (gamma0/lambda.hat.sq)^0.5 * (rho -
            1)/sigma.rho - 1/2 * (lambda.hat.sq - gamma0) * T *
            sigma.rho/s/(lambda.hat.sq^0.5))
}
<environment: namespace:RcompHam94>
Finally the Wald form of an F test as defined by [8.1.32].
                                 R code _
> print(Wald.F.Test)
                                _ output -
function (R, b, r, s2, XtX_1)
    v <- R %*% b - r
    as.numeric(t(v) %*% solve(s2 * R %*% XtX_1 %*% t(R)) %*%
        v/dim(R)[[1]])
<environment: namespace:RcompHam94>
```

#### 8.2 Dickey Fuller Tests for Unit Roots

Page 489 describes the analysis of nominal three month U.S. Treasury yield data from dataset gnptbill, shown below.

```
______ R code _______
> data(gnptbill, package = "Ham94")
> tbill.data <- data.frame(yt = gnptbill$TBILL[-1], yt_1 = gnptbill$TBILL[-length(gnptbill$TBILL)])
```

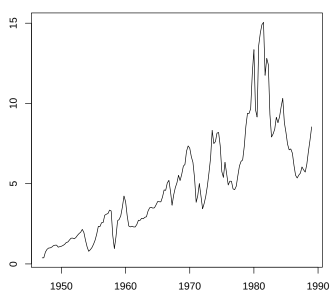


Figure 17.2 – Nominal Interest Rate

The regression model is shown in [17.4.13], and the results are shown below.

[1] 0.9969357

```
[1] 0.01059183
$zeta
numeric(0)
$rho.stat
[1] -0.5147943
$t.stat
[1] -0.2893034
A similar analysis is described on page 494 , but a constant is included in the
regression model [17.4.37].
> case2.lms <- summary(lm(yt ~ 1 + yt_1, tbill.data))</pre>
> case2.DF <- Dickey.Fuller(T = length(tbill.data$yt), rho = case2.lms$coefficients[["yt_1",
      "Estimate"]], sigma.rho = case2.lms$coefficients[["yt_1",
      "Std. Error"]])
> print(case2.lms$coefficients)
                                __ output .
             Estimate Std. Error t value
(Intercept) 0.2105899 0.11212302 1.878204 6.210685e-02
            0.9669104 0.01913305 50.536135 1.013453e-102
                                __ R code _
> print(case2.DF)
                                 _ output _
[1] 168
$rho
[1] 0.9669104
$sigma.rho
[1] 0.01913305
$zeta
numeric(0)
```

\$sigma.rho

```
$rho.stat
[1] -5.559061

$t.stat
[1] -1.729450
```

Example 17.5 describes how to test the joint hypothesis that the trend coefficient is 0 and the autoregressive coefficient is 1.

```
R code

> F <- Wald.F.Test(R = diag(2), b = case2.lms$coefficients[, "Estimate"],

+ r = c(0, 1), s2 = case2.lms$sigma^2, XtX_1 = case2.lms$cov.unscaled)

> print(F)

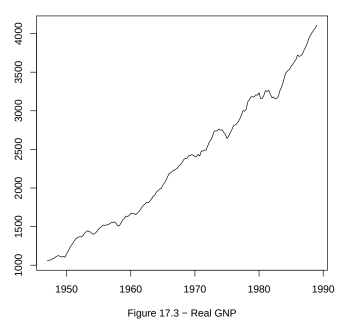
output

[1] 1.806307
```

#### 8.3 Analyzing GNP data

A similar analysis can be conducted on log real GNP data described beginning on page 501, shown below.

```
R code
> logGNP <- 100 * log(gnptbill$GNP)
> gnp.data <- data.frame(tt = seq(1, length(gnptbill$GNP) - 1),
+ yt = logGNP[-1], yt_1 = logGNP[-length(gnptbill$GNP)])</pre>
```



The regression

model here incorporates a time trend, based on the shape of the GDP graph

```
_ R code _
> case4.lms <- summary(lm(yt ~ 1 + yt_1 + tt, gnp.data))</pre>
> case4.DF <- Dickey.Fuller(T = length(gnp.data$yt), rho = case4.lms$coefficients[["yt_1",
     "Estimate"]], sigma.rho = case4.lms$coefficients[["yt_1",
     "Std. Error"]])
> print(case4.lms$coefficients)
                           output -
            Estimate Std. Error t value
                                          Pr(>|t|)
(Intercept) 27.26477184 13.54992552 2.012171 4.582876e-02
          yt_1
          tt
                        ____ R code _
> print(case4.DF)
                          _ output _
$Т
[1] 168
```

\$rho

[1] 0.962522

```
$sigma.rho
[1] 0.01930452
$zeta
numeric(0)
$rho.stat
[1] -6.296298
$t.stat
[1] -1.941409
                                 _ R code
> F <- Wald.F.Test(R = cbind(rep(0, 2), diag(2)), b = case4.lms$coefficients[,
      "Estimate"], r = c(1, 0), s2 = case4.lms\$sigma^2, XtX_1 = case4.lms\$cov.unscaled)
> print(F)
                               __ output _
[1] 2.442251
      Using Phillips Perron Tests
Examples 17.6 and 17.7 reanalyze the case 2 and case 4 regressions above using
the Phillips Perron tests as shown on pages 511-513.
                                 R code
> case2.PP <- Phillips.Perron(T = length(case2.lms$residuals),</pre>
      rho = case2.lms$coefficients[["yt_1", "Estimate"]], sigma.rho = case2.lms$coefficients[["yt_1"
          "Std. Error"]], s = case2.lms$sigma, lambda.hat.sq = as.numeric(Newey.West(case2.lms$resid
          1, 4)), gamma0 = mean(case2.lms$residuals^2))
> print(case2.lms$coefficients)
```

> print(case2.PP)

yt\_1

\$T [1] 168 \_ output

\_ R code -

\_ output -

0.9669104 0.01913305 50.536135 1.013453e-102

Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.2105899 0.11212302 1.878204 6.210685e-02

```
$rho
 [1] 0.9669104
 $sigma.rho
 [1] 0.01913305
 $s.sq
  [1] 0.6375998
 $lambda.hat.sq
  [1] 0.6880069
 $gamma0
  [1] 0.6300093
 $rho.stat
 [1] -6.028975
 $t.stat
 [1] -1.795686
                                                                                                                                                                                                                      _ R code _
> case4.PP <- Phillips.Perron(T = length(case4.lms$residuals),
                                        \label{local_constraint} \begin{subarray}{ll} $r$ ho = case 4.1 ms $coefficients [["yt\_1", "Estimate"]], sigma.rho = case 4.1 ms $coefficients [["yt\_1", "Yt\_1"]], sigma.rho = case 4.1 ms $coefficients
                                                                   "Std. \ Error"]], \ s = case 4. \\ lms \\ sigma, \ lambda. \\ hat. \\ sq = as. \\ numeric (Newey. \\ West (case 4. \\ lms \\ sresident \\ lms \\ sresident \\ lms \\ l
                                                                  1, 4)), gamma0 = mean(case4.lms$residuals^2))
 > print(case4.lms$coefficients)
                                                                                                                                                                                                                      output -
                                                                                                   Estimate Std. Error
                                                                                                                                                                                                                                                                                                                                                  Pr(>|t|)
                                                                                                                                                                                                                                                     t value
 (Intercept) 27.26477184 13.54992552 2.012171 4.582876e-02
yt_1
                                                                                     _ R code _
 > print(case4.PP)
                                                                                                                                                                                                                 _ output _
 $Т
  [1] 168
```

\$rho

```
[1] 0.962522

$sigma.rho
[1] 0.01930452

$s.sq
[1] 1.156270

$lambda.hat.sq
[1] 2.117173

$gamma0
[1] 1.135623

$rho.stat
[1] -10.76066

$t.stat
[1] -2.439143
```

#### 8.5 Augmented Dickey Fuller Tests

Example 17.8 illustrates incorporates the use of lagged regressors to (putatively) eliminate serial correlation in the residuals. The function "embed" is useful for creating lagged regressors.

```
R code
> tbill.data <- list(it = gnptbill$TBILL[-1:-5], delta.it_ = embed(diff(gnptbill$TBILL[-length(gnptb
      4), it_1 = gnptbill$TBILL[c(-1:-4, -(length(gnptbill$TBILL):length(gnptbill$TBILL)))])
> tbill.lms <- summary(lm(it ~ delta.it_ + 1 + it_1, tbill.data))</pre>
> tbill.adf <- Dickey.Fuller(T = length(gnptbill$TBILL) - 5, rho = tbill.lms$coefficients[["it_1",
      "Estimate"]], sigma.rho = tbill.lms$coefficients[["it_1",
      "Std. Error"]], zeta = tbill.lms$coefficients[paste("delta.it",
      1:4, sep = "_"), "Estimate"])
> print(tbill.lms$coefficients)
                               _ output .
             Estimate Std. Error
                                                 Pr(>|t|)
                                   t value
(Intercept) 0.1954328 0.10863764 1.798942 7.393646e-02
delta.it_1 0.3346654 0.07882340 4.245762 3.705074e-05
delta.it_2 -0.3879736 0.08082096 -4.800408 3.643800e-06
```

delta.it\_3 0.2761332 0.07998276 3.452409 7.130684e-04

```
delta.it_4 -0.1067090 0.07944645 -1.343156 1.811475e-01
             0.9690445 0.01860387 52.088332 2.094220e-101
it_1
                                _ R code _
> print(tbill.adf)
                              ____ output _
$Т
[1] 164
$rho
[1] 0.9690445
$sigma.rho
[1] 0.01860387
$zeta
delta.it_1 delta.it_2 delta.it_3 delta.it_4
0.3346654 -0.3879736  0.2761332 -0.1067090
$rho.stat
[1] -5.74363
$t.stat
[1] -1.663928
The next test checks whether or not the farthest lag is different from zero, i.e.
whether or not the right number of lags are included in the equation.
                                  R code -
> print(tbill.lms$coefficients[["delta.it_4", "t value"]])
                                 _ output _
[1] -1.343156
Example 17.9 performs a similar analysis for the GNP data.
                                 _ R code _
> gnp.data <- list(yt = logGNP[-1:-5], delta.yt_ = embed(diff(logGNP[-length(logGNP)]),
      4), yt_1 = logGNP[c(-1:-4, -(length(logGNP):length(logGNP)))],
      t = 6:length(logGNP))
> gnp.lms <- summary(lm(yt ~ delta.yt_ + 1 + yt_1 + t, gnp.data))
> gnp.adf <- Dickey.Fuller(T = length(logGNP) - 5, rho = gnp.lms$coefficients[["yt_1",
      "Estimate"]], sigma.rho = gnp.lms$coefficients[["yt_1", "Std. Error"]],
```

```
zeta = gnp.lms$coefficients[paste("delta.yt", 1:4, sep = "_"),
         "Estimate"])
> F \leftarrow Wald.F.Test(R = cbind(rep(0, 2) \%0\% rep(0, 5), diag(2)),
     b = gnp.lms$coefficients[, "Estimate"], r = c(1, 0), s2 = gnp.lms$sigma^2,
     XtX_1 = gnp.lms$cov.unscaled)
> print(gnp.lms$coefficients)
                             output _
             Estimate Std. Error t value
                                               Pr(>|t|)
(Intercept) 35.91807717 13.57200191 2.6464834 8.961726e-03
delta.yt_1 0.32908487 0.07769385 4.2356619 3.869829e-05
delta.yt_2 0.20856825 0.08128118 2.5660092 1.122316e-02
delta.yt_3 -0.08424648 0.08182895 -1.0295437 3.048077e-01
delta.yt_4 -0.07453301 0.07879621 -0.9458959 3.456552e-01
yt_1
           0.03783123 \quad 0.01521561 \quad 2.4863440 \ 1.395295 e-02
                            __ R code _
> print(gnp.adf)
                             _ output _
[1] 164
$rho
[1] 0.9496901
$sigma.rho
[1] 0.01938565
$zeta
delta.yt_1 delta.yt_2 delta.yt_3 delta.yt_4
$rho.stat
[1] -13.28363
$t.stat
[1] -2.595211
                            __ R code __
> print(F)
```

	output
[1] 3.743228	

# 8.6 Example 17.10 - Bayesian Test of Autoregressive Coefficient

Page 532 describes a test on the autoregressive coefficient that weights prior probabilities.

### 8.7 Determining Lag Length

Page 530 describes an iterative process to determine the correct lag length. This is easily expressed in terms of the structures used above.

#### 8.8 R Facilities for Testing Unit Roots

TBD

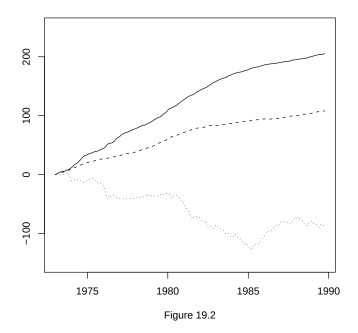
## 9 Cointegration

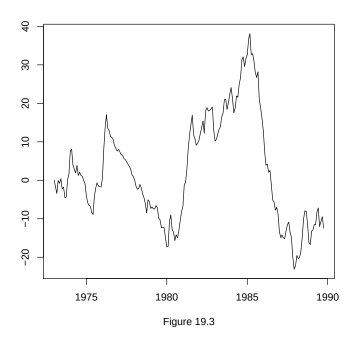
```
R code ______ R code ______
```

# 9.1 Testing Cointegration when the Cointegrating Vector is Known

Section 19.2, beginning on page 582 describes cointegration testing of purchasing power parity between Italian lire and US dollars. The data used is 100 times log monthly price levels and spot nominal and real exchange rates, normalized to a value of zero at the start of the series.

```
R code
> data(ppp, package = "Ham94")
> selection <- subset(ppp, Month >= "1973-01-01" & Month <= "1989-10-01")
> ppp.data <- data.frame(Month = selection$Month, pstar = 100 *
+ log(selection$PC6IT/selection$PC6IT[[1]]), p = 100 * log(selection$PZUNEW/selection$PZUNEW[[1]])
+ ner = -100 * log(selection$EXRITL/selection$EXRITL[[1]]))
> ppp.data[["rer"]] <- ppp.data$p - ppp.data$p ropp.data$pstar</pre>
```





To save time define a simple utility function to perform augmented Dickey Fuller analysis according to the conventions in the text.

```
R code
> do.DF <- function(series, lag) {</pre>
                       T <- length(series)</pre>
                       df.lms \leftarrow summary(lm(yt \sim yt_1 + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(lag + tt + delta.yt_ + 1, list(yt = series[-1:-(la
                                       1)], delta.yt_= embed(diff(series[-T]), lag), yt_1 = series[c(-1:-lag, yt_1)]
                                       -(T:T), tt = (lag + 2):T))
                       df.results <- Dickey.Fuller(T = length(series) - lag - 1,</pre>
                                     rho = df.lms$coefficients[["yt_1", "Estimate"]], sigma.rho = df.lms$coefficients[["yt_1",
                                                       "Std. Error"]], zeta = df.lms$coefficients[paste("delta.yt_",
                                                      1:lag, sep = ""), "Estimate"])
                       F \leftarrow Wald.F.Test(R = cbind(rep(0, 2), diag(2), rep(0, 2) %o%)
                                      rep(0, lag)), b = df.lms$coefficients[, "Estimate"],
                                      r = c(1, 0), s2 = df.lmssigma^2, XtX_1 = df.lms$cov.unscaled)
                       print(df.lms$coefficients)
                       print(df.results)
                       print(F)
 + }
```

Following the text, check each series with a Dickey Fuller test with a regression estimated with twelve lags.

```
R code
> for (series.name in c("p", "pstar", "ner", "rer")) do.DF(series = ppp.data[[series.name]],
+ lag = 12)
```

```
output -
                                                    Pr(>|t|)
               Estimate Std. Error
                                       t value
(Intercept) 0.136160926 0.085779070
                                     1.5873444 1.142502e-01
yt_1
            0.994004087 0.003067474 324.0464885 6.323397e-244
tt
            0.002927051 0.001766655
                                    1.6568325 9.935541e-02
            0.553397837 0.075217880
                                    7.3572644 7.109482e-12
delta.yt_1
delta.yt_2 -0.056908322 0.085440124 -0.6660609 5.062543e-01
delta.yt_3 0.070125117 0.084906900 0.8259060 4.099884e-01
delta.yt_4 0.060389596 0.081969953 0.7367284 4.622797e-01
delta.yt_5 -0.078232496 0.078488461 -0.9967388 3.202754e-01
delta.yt_6 -0.048376861 0.070721885 -0.6840437 4.948576e-01
delta.yt_7 0.165843348 0.068915448 2.4064757 1.715410e-02
delta.yt_8 -0.070207448 0.070014467 -1.0027563 3.173709e-01
delta.yt_9
            0.244644550 0.070161410
                                     3.4868819 6.187074e-04
delta.yt_10 -0.110047172 0.072579707 -1.5162251 1.312771e-01
delta.yt_11 0.117580628 0.072937432
                                     1.6120753 1.087579e-01
delta.yt_12  0.046702346  0.068650314  0.6802933  4.972230e-01
$Т
```

```
[1] 189
```

\$rho

[1] 0.994004

\$sigma.rho

[1] 0.003067474

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6
0.55339784 -0.05690832 0.07012512 0.06038960 -0.07823250 -0.04837686
delta.yt\_7 delta.yt\_8 delta.yt\_9 delta.yt\_10 delta.yt\_11 delta.yt\_12
0.16584335 -0.07020745 0.24464455 -0.11004717 0.11758063 0.04670235

\$rho.stat

[1] -10.78352

\$t.stat

[1] -1.954675

#### [1] 2.412933

Estimate Std. Error t value Pr(>|t|) (Intercept) 0.768007976 0.253071035 3.0347526 2.776788e-03 yt\_1 0.999456707 0.004116999 242.7633949 3.768702e-222 tt -0.002406065 0.004989081 -0.4822662 6.302229e-01 delta.yt\_1 0.420701728 0.076110499 5.5275124 1.170691e-07 delta.yt\_2 -0.011592127 0.081521266 -0.1421976 8.870885e-01 delta.yt\_3 0.013439685 0.080162382 0.1676558 8.670488e-01 delta.yt\_4 0.077206365 0.080125530 0.9635676 3.366000e-01 delta.yt\_5 -0.036494296 0.080087139 -0.4556824 6.491866e-01 delta.yt\_6 0.145282237 0.078670504 1.8467180 6.648647e-02 delta.yt\_7 -0.099118088 0.078839877 -1.2572075 2.103634e-01 delta.yt\_8 0.046717520 0.078598766 0.5943798 5.530301e-01 delta.yt\_9 -0.049982364 0.078111841 -0.6398820 5.230909e-01 delta.yt\_10 -0.034638353 0.078168372 -0.4431249 6.582258e-01 delta.yt\_11 0.075555037 0.077993666 0.9687330 3.340230e-01 delta.yt\_12 0.021863739 0.073346671 0.2980877 7.659919e-01 \$Т

\$rho

[1] 189

[1] 0.9994567

#### \$sigma.rho

[1] 0.004116999

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6 0.42070173 -0.01159213 0.01343968 0.07720637 -0.03649430 0.14528224 delta.yt\_7 delta.yt\_8 delta.yt\_9 delta.yt\_10 delta.yt\_11 delta.yt\_12 -0.09911809 0.04671752 -0.04998236 -0.03463835 0.07555504 0.02186374

#### \$rho.stat

[1] -0.2382095

#### \$t.stat

[1] -0.1319633

#### [1] 4.249956

Estimate Std. Error t value Pr(>|t|) (Intercept) -0.389337356 0.413800921 -0.94088084 3.480703e-01 yt\_1 0.982941298 0.010766440 91.29678192 6.506909e-149 -0.007384125 0.006883901 -1.07266573 2.849066e-01 tt delta.yt\_1 0.348829755 0.074439036 4.68611329 5.595654e-06 delta.yt\_2 -0.025567401 0.079110764 -0.32318485 7.469433e-01 delta.yt\_3 0.002617322 0.078947706 0.03315261 9.735909e-01 delta.yt\_4 0.011689457 0.080007934 0.14610372 8.840086e-01 delta.yt\_5 0.099314112 0.079948258 1.24222983 2.158234e-01 delta.yt\_6 0.001387289 0.080819939 0.01716518 9.863245e-01 delta.yt\_7 0.063205400 0.080614348 0.78404653 4.340788e-01 delta.yt\_8 0.117223384 0.080560981 1.45508883 1.474464e-01 delta.yt\_9 -0.061127657 0.080788556 -0.75663757 4.502903e-01 delta.yt\_10 0.081739596 0.080696462 1.01292665 3.125017e-01 delta.yt\_11 0.037261364 0.080646524 0.46203311 6.446347e-01 delta.yt\_12 -0.030363466 0.076740775 -0.39566275 6.928385e-01 \$Т

#### [1] 189

#### \$rho

[1] 0.9829413

#### \$sigma.rho

[1] 0.01076644

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6
0.348829755 -0.025567401 0.002617322 0.011689457 0.099314112 0.001387289
delta.yt\_7 delta.yt\_8 delta.yt\_9 delta.yt\_10 delta.yt\_11 delta.yt\_12
0.063205400 0.117223384 -0.061127657 0.081739596 0.037261364 -0.030363466

#### \$rho.stat

[1] -9.112996

#### \$t.stat

[1] -1.584433

#### Γ17 1.489674

Estimate Std. Error t value Pr(>|t|) (Intercept) 0.0532014210 0.390557357 0.13621923 8.918054e-01 0.9712932573 0.014145189 68.66597772 5.679805e-128 yt\_1 -0.0004612496 0.003237185 -0.14248477 8.868620e-01 tt delta.yt\_1 0.3178370194 0.074163266 4.28563944 3.010943e-05 delta.yt\_2 -0.0149166870 0.078078854 -0.19104644 8.487119e-01 delta.yt\_3 0.0127973250 0.077727723 0.16464299 8.694161e-01 delta.yt\_4 0.0224258044 0.078676900 0.28503671 7.759550e-01 delta.yt\_5 0.0845155831 0.078339518 1.07883716 2.821536e-01 delta.yt\_6 -0.0030653274 0.079071534 -0.03876651 9.691210e-01 delta.yt\_7 0.0299137752 0.078750797 0.37985362 7.045173e-01 delta.yt\_8 0.0824197050 0.078641636 1.04804158 2.960730e-01 delta.yt\_9 -0.0478615036 0.078647910 -0.60855405 5.436137e-01 delta.yt\_10 0.0755667133 0.078405880 0.96378886 3.364893e-01 delta.yt\_11 0.0504082264 0.078279945 0.64394816 5.204570e-01 delta.yt\_12 -0.0124704308 0.075997755 -0.16408946 8.698512e-01 \$Т

#### \$rho

[1] 189

[1] 0.9712933

#### \$sigma.rho

[1] 0.01414519

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6
0.317837019 -0.014916687 0.012797325 0.022425804 0.084515583 -0.003065327
delta.yt\_7 delta.yt\_8 delta.yt\_9 delta.yt\_10 delta.yt\_11 delta.yt\_12

```
$rho.stat
[1] -13.48204
$t.stat
[1] -2.029435
[1] 2.078078
Now check the real exchange rate with a Phillips Perron test
                                _ R code _
> pp.lms <- summary(lm(zt ~ zt_1 + 1, data.frame(zt = ppp.data$rer[-1],
      zt_1 = ppp.data$rer[-length(ppp.data$rer)])))
> PP.results <- Phillips.Perron(T = length(pp.lms$residuals), rho = pp.lms$coefficients[["zt_1",
      "Estimate"]], sigma.rho = pp.lms$coefficients[["zt_1", "Std. Error"]],
      s = pp.lmssigma, lambda.hat.sq = as.numeric(Newey.West(pp.lms$residuals %o%)
          1, 12)), gamma0 = mean(pp.lms$residuals^2))
> print(pp.lms$coefficients)
                                _ output _
              Estimate Std. Error t value
                                                   Pr(>|t|)
(Intercept) -0.0297931 0.17835718 -0.1670418 8.675068e-01
             0.9865420 0.01275287 77.3584248 1.854719e-150
zt_1
                             ____ R code __
> print(PP.results)
                                oxdot output oxdot
$Т
[1] 201
$rho
[1] 0.986542
$sigma.rho
[1] 0.01275287
$s.sq
[1] 6.205887
```

 $0.029913775 \quad 0.082419705 \quad -0.047861504 \quad 0.075566713 \quad 0.050408226 \quad -0.012470431$ 

\$lambda.hat.sq

```
[1] 13.03064

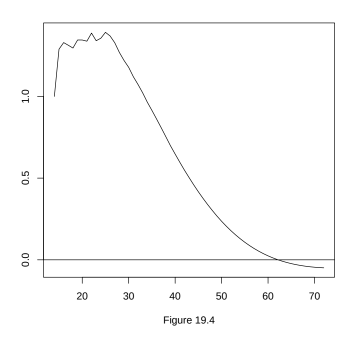
$gamma0
[1] 6.144137

$rho.stat
[1] -6.35068

$t.stat
[1] -1.706128
```

Estimating the impulse response function gives a sense of the persistence of deviations from PPP.

```
R code
> ar.results <- ar(ppp.data$rer, aic = FALSE, order.max = 13, method = "ols",
+ demean = TRUE)
> tt <- seq(1, 72)
> start.innov <- rep(0, 13)
> et <- c(start.innov, 1, rep(0, length(tt) - 14))
> arima.sim.output <- arima.sim(list(order = c(13, 0, 0), ar = ar.results$ar),
+ n = length(tt), innov = et, n.start = length(start.innov),
+ start.innov = start.innov)
> irf <- as.vector(arima.sim.output)</pre>
```



#### 9.2 Estimating the Cointegrating Vector

Page 598 shows an example of the Phillips Ouliaris Hansen procedure for estimating a cointegrating vector.

R code \_\_\_\_\_

0.05134848 0.012045369 4.262923 3.114337e-05 0.53004097 0.006708385 79.011705 3.148050e-152

pstar

```
> print(poh.residual.lms$coefficients)
                                 output
     Estimate Std. Error t value
                                      Pr(>|t|)
u_1 0.9833108 0.01171956 83.90338 7.71577e-158
                                 _ R code _
> print(POH.results)
                                 _ output _
$Т
[1] 201
$rho
[1] 0.9833108
$sigma.rho
[1] 0.01171956
$s.sq
[1] 0.1630028
$lambda.hat.sq
[1] 0.4082242
$gamma0
[1] 0.1621919
$rho.stat
[1] -7.542281
$t.stat
[1] -2.020981
A second example performs a similar analysis on quarterly US consumption and
income data from 1947Q1 to 1989Q4.
> data(coninc, package = "Ham94")
R code -
> selection <- subset(coninc, Quarter >= "1947-01-01" & Quarter <=
```

> coninc.data <- data.frame(Quarter = selection\$Quarter, cons = 100 \*
+ log(selection\$GC82), inc = 100 \* log(selection\$GYD82))</pre>

"1989-07-01")



Test individual

series for unit root status using Dickey Fuller.

```
R code
> for (series.name in c("inc", "cons")) do.DF(series = coninc.data[[series.name]],
+ lag = 6)
```

```
output
                                               Pr(>|t|)
              Estimate
                       Std. Error
                                    t value
(Intercept) 20.336729221 15.04162460 1.35203010 1.783352e-01
yt_1
           tt
           0.023796844 0.01985318 1.19864142 2.324968e-01
delta.yt_1 -0.006528755 0.08092856 -0.08067307 9.358060e-01
delta.yt_2 -0.035846316 0.08025935 -0.44663103 6.557649e-01
delta.yt_3 0.102128545 0.07758036 1.31642276 1.899755e-01
delta.yt_4 -0.187536343 0.07699406 -2.43572477 1.599577e-02
delta.yt_5 -0.037187883 0.07813842 -0.47592314 6.347992e-01
delta.yt_6
           0.027855951 0.07662877 0.36351818 7.167132e-01
$Т
[1] 164
```

#### \$rho

[1] 0.970585

#### \$sigma.rho

[1] 0.02306293

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6 -0.006528755 -0.035846316 0.102128545 -0.187536343 -0.037187883 0.027855951

#### \$rho.stat

[1] -4.242382

#### \$t.stat

[1] -1.275428

#### [1] 1.132134

Estimate Std. Error t value Pr(>|t|)
(Intercept) 29.46860131 15.19248322 1.9396830 5.423391e-02
yt\_1 0.95552168 0.02360001 40.4881863 2.508405e-84
tt 0.03721088 0.02006161 1.8548306 6.552012e-02
delta.yt\_1 0.03624864 0.07979877 0.4542506 6.502840e-01
delta.yt\_2 0.25964745 0.07935028 3.2721680 1.315743e-03
delta.yt\_3 0.06273192 0.08172798 0.7675697 4.439106e-01
delta.yt\_4 -0.05234112 0.08122252 -0.6444163 5.202580e-01
delta.yt\_5 -0.04791625 0.07956524 -0.6022260 5.479037e-01
delta.yt\_6 -0.06782142 0.07919698 -0.8563637 3.931186e-01
\$T
[1] 164

#### \$rho

[1] 0.9555217

#### \$sigma.rho

[1] 0.02360001

#### \$zeta

delta.yt\_1 delta.yt\_2 delta.yt\_3 delta.yt\_4 delta.yt\_5 delta.yt\_6 0.03624864 0.25964745 0.06273192 -0.05234112 -0.04791625 -0.06782142

#### \$rho.stat

[1] -9.011597

#### \$t.stat

[1] -1.884673

Estimate cointegration vector, then check for unit root status of the residual using Phillips Perron.

```
> poh.cointegration.lm <- lm(cons ~ 1 + inc, coninc.data)</pre>
> poh.residual.lms <- summary(lm(u ~ 0 + u_1, data.frame(u = poh.cointegration.lm$residuals[-1],
                  u_1 = poh.cointegration.lm$residuals[-length(poh.cointegration.lm$residuals)])))
> POH.results <- Phillips.Perron(T = length(poh.residual.lms$residuals),
                 rho = poh.residual.lms$coefficients[["u_1", "Estimate"]],
                 sigma.rho = poh.residual.lms$coefficients[["u_1", "Std. Error"]],
                  s = poh.residual.lms\$sigma, lambda.hat.sq = as.numeric(Newey.West(poh.residual.lms\$residuals % lambda.hat.sq = lambda.hat.
                             1, 6)), gamma0 = mean(poh.residual.lms$residuals^2))
> print(summary(poh.cointegration.lm)$coefficients)
                                                                                             _ output -
                                     Estimate Std. Error
                                                                                                              t value
                                                                                                                                                    Pr(>|t|)
                                                                                                    0.2840347 7.767315e-01
(Intercept) 0.6675807 2.350348907
                                  0.9864943 0.003217444 306.6080542 5.567137e-234
inc
                                                                                               R code -
> print(poh.residual.lms$coefficients)
                                                                                            _ output -
              Estimate Std. Error t value
                                                                                                              Pr(>|t|)
u_1 0.7818542 0.04788553 16.32757 1.402076e-36
                                                                                            _ R code _
> print(POH.results)
                                                                                            _ output _
$Т
[1] 170
$rho
[1] 0.7818542
$sigma.rho
[1] 0.04788553
$s.sq
[1] 1.22395
```

```
$lambda.hat.sq
[1] 1.030594

$gamma0
[1] 1.216750

$rho.stat
[1] -32.04525

$t.stat
[1] -4.27529
```

#### 9.3 Testing Hypotheses About the Cointegrating Vector

Page 608-612 illustrate a technique that uses leads and lags to produce a stationary vector for hypothesis testing.

```
R code
> T <- length(coninc.data$Quarter)
> lead.lag.data <- list(ct = coninc.data$cons[c(-1:-5, -((T - 3):T))],
+ yt = coninc.data$inc[c(-1:-5, -((T - 3):T))], delta.yt = diff(coninc.data$inc[c(-1:-4,
+ -((T - 3):T))]), delta.yt_ = embed(diff(coninc.data$inc[-((T - 4):T)]), 4), delta.yt. = embed(diff(coninc.data$inc[-1:-5])[(T - 4):T)], 4)[(T - 9):1, ], tt = 6:(T - 4))</pre>
```

The regression is estimated with both no trend and trend, and the corrected t-stat is calculated.

```
R code
> no.trend.lm <- lm(ct ~ 1 + yt + delta.yt. + delta.yt + delta.yt_,
+ lead.lag.data)
> trend.lm <- lm(ct ~ 1 + yt + tt + delta.yt. + delta.yt + delta.yt_,
+ lead.lag.data)
> for (model in list(no.trend.lm, trend.lm)) {
+ lags <- 2
+ cms <- summary(model)
+ T <- length(cms$residuals)
+ cfs <- cms$coefficients
+ t.rho <- (cfs[["yt", "Estimate"]] - 1)/cfs[["yt", "Std. Error"]]
+ rms <- summary(lm(u ~ 0 + u_, list(u = cms$residuals[-c(1:lags)],
+ u_ = embed(cms$residuals[-T], lags))))
+ sigma1.hat.sq <- mean(rms$residuals^2)</pre>
```

```
1:lags, sep = ""), "Estimate"]))
+
     t.a <- t.rho * cms$sigma/lambda.11
     print(cfs)
     print(rms$coefficients)
     print(T)
     print(cms$sigma)
     print(t.rho)
     print(sigma1.hat.sq)
     print(lambda.11)
     print(t.a)
+ }
                            output -
             Estimate Std. Error
                                  t value
                                              Pr(>|t|)
(Intercept) -4.51922906 2.340224673 -1.9311091 5.534290e-02
           0.99215853 0.003063317 323.8837231 1.617626e-216
           0.48592391 0.115704789 4.1996871 4.551158e-05
delta.yt.1
delta.yt.2 0.26411856 0.114892015 2.2988418 2.288546e-02
delta.yt.3 0.28614193 0.115594505 2.4753939 1.441397e-02
delta.yt.4 0.14530952 0.118799555 1.2231487 2.231790e-01
         -0.24036007 0.117415901 -2.0470828 4.238356e-02
delta.yt
delta.yt_1 -0.01101143 0.113899420 -0.0966768 9.231113e-01
delta.yt_2 0.06969114 0.111505773 0.6250003 5.329142e-01
delta.yt_3 0.04055551 0.111155199 0.3648548 7.157303e-01
delta.yt_4 0.02150153 0.110083985 0.1953193 8.454056e-01
    Estimate Std. Error t value
                                 Pr(>|t|)
u_1 0.7179687 0.07722647 9.296924 1.127578e-16
u_2 0.2057401 0.07684783 2.677241 8.207043e-03
[1] 162
[1] 1.516006
[1] -2.559799
[1] 0.3809180
[1] 8.089864
[1] -0.4796954
              Estimate Std. Error
                                   t value
                                             Pr(>|t|)
(Intercept) 198.87166510 15.01478288 13.2450577 5.215628e-27
            уt
tt
            delta.yt.1
delta.yt.2
            0.15407283 0.07749787 1.9880910 4.862147e-02
```

delta.yt.3

```
delta.yt.4
        -0.05124600 0.07998305 -0.6407108 5.226882e-01
delta.yt
delta.yt_1
         0.12737594  0.07708222  1.6524685  1.005308e-01
         delta.yt_2
delta.yt_3
         delta.yt_4
         Estimate Std. Error t value
                         Pr(>|t|)
u_1 0.6871713 0.07786238 8.825460 1.937474e-15
u_2 0.1291820 0.07666487 1.685022 9.395837e-02
[1] 162
[1] 1.017016
[1] -13.90793
[1] 0.3439489
[1] 3.193478
[1] -4.429212
```

# 10 Full-Information Maximum Likelihood Analysis of Cointegrated Systems

```
R code
> library("RcompHam94", lib.loc = ".")
```

# 10.1 An Application of the Johansen Approach to the PPP data

Section 20.3 reanalyzes the data used in Chapter 19 using the FIML approach.

```
R code
> data(ppp, package = "Ham94")
> selection <- subset(ppp, Month >= "1973-01-01" & Month <= "1989-10-01")
> ppp.data <- data.frame(pstar = 100 * log(selection$PC6IT/selection$PC6IT[[1]]),
+ p = 100 * log(selection$PZUNEW/selection$PZUNEW[[1]]), ner = -100 *
+ log(selection$EXRITL/selection$EXRITL[[1]]))
> y <- as.matrix(ppp.data)</pre>
```

First conduct the auxiliary regressions. Given that the right hand sides consists of lagged values of the changes in y for both [20.2.4] and [20.2.5], construct a regression with both lagged y and lagged changes of y as left hand side.

```
R code ______
> delta.y <- diff(y)
> lags <- 12</pre>
```

Now calculate the canonical correlations according to [20.2.6], [20.2.7], [20.2.8], and calculate eigenvalues according to [20.2.9], and log likelihood as in [20.2.10]. Note that u is T rows by n columns so that ut is the t-th row of matrix u, so only a single inner product, rather than sum of outer products, is needed.

```
R code
> SigmaUU <- 1/T * t(u) %*% u
> SigmaVV <- 1/T * t(v) %*% v
> SigmaUV <- 1/T * t(u) %*% v
> eigen.results <- eigen(solve(SigmaVV) %*% t(SigmaUV) %*% solve(SigmaUU) %*%
      SigmaUV)
> lambda <- eigen.results$values
> LRT <- -T * sum(log(1 - lambda))
> print(SigmaUU)
                                 output -
               Response pstar Response p Response ner
                   0.17931504 0.01531134
                                            0.02715177
Response pstar
                   0.01531134  0.04341512  -0.03267373
Response p
Response ner
                   0.02715177 -0.03267373
                                            4.60842626
                                _ R code _
> print(SigmaVV)
                                 output
               Response pstar Response p Response ner
                    1503.5545
Response pstar
                                794.7041
                                            -697.4981
Response p
                     794.7041
                                421.5535
                                             -365.1883
Response ner
                    -697.4981
                               -365.1883
                                             414.1322
                                 _ R code _
> print(SigmaUV)
```

-		output		
	Response pstar	Response p	Response ner	
Response pstar	-3.5787320	-1.7958934	1.5095381	
Response p	-0.8602478	-0.4969721	0.5243431	
Response ner	-3.1461173	-2.0636489	-2.2685853	
		R code		
> print(lambda	)			
[1] 0.12002316	0.05077020 0.03	3174158		
		R code		
> print(T * log	g(1 - lambda))			
	0.045504	output		
[1] -24.165480	-9.847724 -6	. 096434		
		R code		
<pre>&gt; print(LRT)</pre>				
[1] 40.10964		output		
[1] 40.10904				
			rst cointegrating vector unity for the first coefficient	
		R code		
> ahat1 <- eig	en.results\$vecto	ors[, 1]		
> ahat1.tilde	<- ahat1/sqrt(t	(ahat1)	SigmaVV %*% ahat1)	
> ahat1.normal	<- ahat1/ahat1	[[1]]		
<pre>&gt; print(ahat1)</pre>				
		output		
[1] -0.4888515	1 0.87144476 -0	0.04010268		
> print(ahat1.		R code		
	,			
	0 0.79841545 -0	output		
		- /		
> print(ahat1.		R code		
[1] 1.0000000	0 -1.78263694 (	output 0.08203448		

#### 10.2 Likelihood Ratio Tests on the Cointegration Vector

Page 649 shows how to conduct hypothesis tests on the cointegration vector. The follow code implements [20.3.10] - [20.3.14] and subsequent calculations.

```
R code
> D = cbind(c(1, 0, 0), c(0, 0, 1))
> SigmaVV.tilde <- t(D) %*% SigmaVV %*% D</pre>
> SigmaUV.tilde <- SigmaUV %*% D
> eigen.results <- eigen(solve(SigmaVV.tilde) %*% t(SigmaUV.tilde) %*%
      solve(SigmaUU) %*% SigmaUV.tilde)
> lambda.tilde <- eigen.results$values
> h <- 1
> LRT <- T * sum(log(1 - lambda[1:h])) + T * sum(log(1 - lambda.tilde[1:h]))
> ahat1.normal.tilde <- eigen.results$vectors[, 1]/eigen.results$vectors[,
      1][[1]]
> print(SigmaVV.tilde)
                                  _ output _
           [,1]
                     [,2]
[1,] 1503.5545 -697.4981
[2,] -697.4981 414.1322
                                   _ R code _
> print(SigmaUV.tilde)
                                   output -
                      [,1]
                                  [,2]
Response pstar -3.5787320 1.5095381
                -0.8602478 0.5243431
Response p
Response ner
                -3.1461173 -2.2685853
                                   R code _
> print(lambda.tilde)
                                   _{\scriptscriptstyle -} output _{\scriptscriptstyle -}
[1] 0.05828948 0.03295258
                                    R code
> print(T * log(1 - lambda.tilde))
                                  _ output _
[1] -11.350839
                -6.332964
```

	R code	
> print(LRT)		
	au+mu+	
[1] 12.81464	output	
<pre>&gt; print(ahat1.normal.tilde)</pre>	R code	
	output	
[1] 1.000000 1.012463		
Page 650 shows a second examp	ble.	
	R code	
> h <- 1		
> D = c(1, -1, -1) %0% 1		
> SigmaVV.tilde <- t(D) %*% S	igmaVV %*% D	
> SigmaUV.tilde <- SigmaUV %*	% D	
> eigen.results <- eigen(solv	e(SigmaVV.tilde) %*% t(Sigma	aUV.tilde) %*%
+ solve(SigmaUU) %*% Sigm	aUV.tilde)	
> lambda.tilde <- eigen.resul	ts\$values	
> LRT <t *="" -="" lam<="" sum(log(1="" td=""><td></td><td>- lambda.tilde[1:h]))</td></t>		- lambda.tilde[1:h]))
> print(SigmaVV.tilde)	-	
	output	
[,1]		
[1,] 1414.452		
> print(SigmaUV.tilde)	R code	
	output	
[,1]		
Response pstar -3.2923768		
Response p -0.8876187		
Response ner 1.1861170		
	R code	
> print(lambda.tilde)		
	output	
[1] 0 04912925	output	

## 11 Time Series Models of Heteroskedasticity

```
R code ______
> library("RcompHam94", lib.loc = ".")
```

#### 11.1 Preamble

Page 658 and forward provide examples of ARCH models. Several utility functions are needed for these examples. The function "arch.fitted.values" calculates the value of ht given the conditional information set YT and a parameter vector THETA as described on page 660, [21.1.17] to [21.1.20].

Function "arch.standard.errors" calculates values for standard errors according to the description on page 663, particularly equations [21.1.25], and also using [21.1.21] for the estimate of the outer product estimate of the information matrix.

```
R code
> arch.standard.errors <- function(THETA, YT) {</pre>
      x \leftarrow YT$x
      v <- YT$v
      k < -dim(x)[[2]]
      alpha <- THETA[grep("alpha.*", names(THETA))]</pre>
      zeta <- THETA["zeta"]</pre>
      m <- length(alpha)</pre>
      T \leftarrow length(y) - m
      a \leftarrow k + 1 + m
      fv <- arch.fitted.values(THETA, YT)</pre>
      h <- fv$h
      u2 <- fv$u^2
      S \leftarrow array(0, c(a, a))
      D \leftarrow array(0, c(a, a))
      for (tt in (m + 1):length(y)) {
           temp <- c(t(alpha) %*% ((u2[(tt - 1):(tt - m)] %o% rep(1,
               k)) * x[(tt - 1):(tt - m), ]), c(1, u[(tt - 1):(tt -
               m)]^2))
           st \leftarrow (u2[tt] - h[tt])/(2 * h[tt]^2) * temp + c(u2[tt]/h[tt] *
               x[tt, ], rep(0, a - k))
           S \leftarrow S + 1/T * st %*% t(st)
           D \leftarrow D + 1/T * (1/(2 * h[tt]^2) * temp %*% t(temp) +
               rbind(cbind(1/h[tt] * x[tt, ] %*% t(x[tt, ]), array(0,
                    c(k, a - k))), array(0, c(a - k, a))))
       diag(1/T * solve(D) %*% S %*% solve(D))^0.5
+ }
```

The following two helper functions calculate the likelihood values under different distributional assumptions. The normal likelihood is calculated according to [21.1.20], the scaled t according to [21.1.24].

```
R code
> arch.normal <- function(THETA, YT) {

+ fv <- arch.fitted.values(THETA, YT)
+ m <- length(THETA[grep("alpha.*", names(THETA))])
+ h <- fv$h[-1:-m]
+ u <- fv$u[-1:-m]
+ -1/2 * (length(h) * log(2 * pi) - sum(log(h)) - sum(u^2/h))
+ }
> arch.scaled.t <- function(THETA, YT) {
+ fv <- arch.fitted.values(THETA, YT)</pre>
```

```
+ m <- length(THETA[grep("alpha.*", names(THETA))])
+ h <- fv$h[-1:-m]
+ u <- fv$u[-1:-m]
+ nu <- THETA[grep("nu", names(THETA))]
+ result <- length(h) * log(gamma((nu + 1)/2)/(sqrt(pi) * gamma(nu/2)) *
+ (nu - 2)^-0.5) - 1/2 * sum(log(h)) - (nu + 1)/2 * sum(log(1 + u^2/(h * (nu - 2))))
+ }</pre>
```

GMM estimates are calculated according to the recipe in Chapter 14, notably equations [14.1.7] and [14.1.10]. Functions h and S are specified by the caller.

```
R code
> GMM.estimates <- function(YT, h, THETA, S) {
      g <- function(YT, THETA) {</pre>
          apply(X = apply(X = YT, MARGIN = 1, FUN = h, THETA = THETA),
              MARGIN = 1, FUN = mean)
      objective <- function(THETA, YT, W) {
          g.value <- g(YT, THETA)</pre>
          as.numeric(t(g.value) %*% W %*% g.value)
      r \leftarrow length(h(YT[1, ], THETA))
      a <- length(THETA)
      stage.1.results \leftarrow optim(par = THETA, fn = objective, gr = NULL,
          YT = YT, W = diag(r)
      temp <- t(apply(X = YT, MARGIN = 1, FUN = h, THETA = stage.1.results$par))
      ST <- S(temp)
      stage.2.results <- optim(par = stage.1.results$par, fn = objective,
          gr = NULL, YT = YT, W = solve(ST))
      list(stage.1.results = stage.1.results, stage.2.results = stage.2.results)
+ }
```

#### 11.2 Application of ARCH Models to US Fed Funds Data

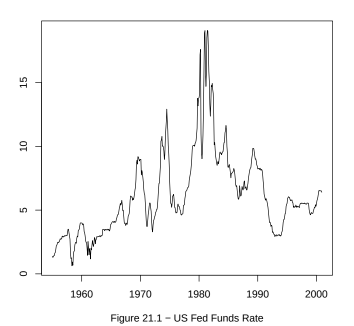
The dataset for these examples is the US Fed Funds Rate, monthly between Jan 1955 and December 2000, shown below.

```
R code

> data(fedfunds, package = "Ham94")

> selection <- subset(fedfunds, Month >= "1955-01-01" & Month <= 
+ "2000-12-01")

> y <- selection$FFED
```



A first step is

to characterize the autocorrelation structure of the squared residuals. These two regressions show that a second order AR process seems to fit the data pretty well.

```
R code
> y.lm <- lm(y ~1 + y_1, list(y = y[-1], y_1 = y[-length(y)])
> u <- y.lm$residuals
> u2.lm <- lm(u2 ~ 1 + u2_lag, list(u2 = u[-1:-4]^2, u2_lag = embed(u[-length(u)]^2,
      4)))
> F34 \leftarrow Wald.F.Test(R = cbind(rep(0, 2) \%0\% rep(0, 3), diag(2)),
      b = u2.lm$coefficients, r = c(0, 0), s2 = summary(u2.lm)$sigma^2,
      XtX_1 = summary(u2.lm)$cov.unscaled)
> F34.sig <- 1 - pf(F34, 2, length(u2.lm$residuals) - u2.lm$rank)
> F234 \leftarrow Wald.F.Test(R = cbind(rep(0, 3) %o% rep(0, 2), diag(3)),
      b = u2.lm$coefficients, r = c(0, 0, 0), s2 = summary(u2.lm)$sigma^2,
      XtX_1 = summary(u2.lm)$cov.unscaled)
> F234.sig <- 1 - pf(F234, 3, length(u2.lm$residuals) - u2.lm$rank)
> accept.arch <- pchisq(length(u2.lm$residuals) * summary(u2.lm)$r.squared,
      4)
> print(F34)
                                 output
[1] 0.8225742
```

	R с	ode	
<pre>&gt; print(F34.sig)</pre>			
[1] 0.439847	out <sub>]</sub>	out	
	R co	ode	
> print(F234)			
[1] 11.88167	out	out	
[1] 11.00107			
	R с	ode	
> print(F234.sig)			
[1] 1.513714e-07	out <sub>]</sub>	out	
1.513/14e-0/			
	R C	nde	
> print(accept.arch)		Jue	
	out;	out	
[1] 1			
Next we use a mayir	num likelihood esti	mation to estima	te the parameters for
the second order equa			the parameters for
the second order equi	acion appaining nor	mar circis.	
> YT <- list(y = y[-		ode	1)
> THETA $\leftarrow$ c(beta =	_		
+ $alpha = c(0.1)$		, 2000 var (y . 1	myrobradarb,,
> optimizer.results		ETA, fn = arch.n	ormal, gr = NULL,
+ $YT = YT$			
> print(optimizer.re	esults\$par)		
	out	out	
beta.(Intercept)	beta.y_1	zeta	alpha1
0.25226382	0.94858488	0.02734929	0.95530391
alpha2			
0.29858866			
	R co	ode	
> se <- arch.standar			T)
> print(se)			

Now use GMM to estimate the same parameters following page 664. The initial values for the regression coefficients are derived from the (homoskedastic) regression above, as is the presample variance. The estimator for S assumes no correlation at leads and lags.

```
_{-} R code _{-}
> h <- function(wt, THETA) {
      beta <- THETA[grep("beta.*", names(THETA))]</pre>
      zeta <- THETA["zeta"]</pre>
      alpha <- THETA[grep("alpha.*", names(THETA))]</pre>
      m <- length(alpha)</pre>
      k <- length(beta)
      yt <- wt[grep("yt.*", names(wt))]</pre>
      xt <- wt[grep("xt.*", names(wt))]</pre>
      ylagt <- wt[grep("ylagt.*", names(wt))]</pre>
      xlagt <- t(array(wt[grep("xlagt.*", names(wt))], c(k, m)))</pre>
      ut <- yt - t(xt) %*% beta
      zt <- c(1, (ylagt - t(xlagt) %*% beta)^2)</pre>
      c(ut * xt, (ut^2 - t(zt) %*% c(zeta, alpha)) * zt)
+ }
> S.estimator <- function(ht) {
      1/dim(ht)[[1]] * t(ht) %*% ht
+ }
> THETA <- c(beta = y.lm$coefficients, zeta = var(y.lm$residuals),
      alpha = c(0.1, 0.1)
> m <- length(THETA[grep("alpha.*", names(THETA))])</pre>
> T <- length(YT$y) - m
> w \leftarrow as.matrix(data.frame(yt = YT$y[-1:-m], xt = YT$x[-1:-m]
      ], ylagt = embed(YT$y[-(T + m)], m), xlagt = embed(YT$x[-(T + m)])
      m), ], m)))
> estimates <- GMM.estimates(YT = w, h = h, THETA = THETA, S.estimator)
> print(estimates$stage.1.results$par)
                                  _ output _
beta.(Intercept)
                           beta.y_1
                                                                  alpha1
                                                  zeta
      0.05788674
                         0.98955937
                                           0.32491651
                                                              0.01073606
          alpha2
      0.02105476
```

R code

	ou	tput		
beta.(Intercept)	beta.y_1	zeta	alpha1	
0.02579794	0.99791508	-0.17911928	0.01239927	
alpha2				
0.07770754				

#### 11.3 R Facilities For GARCH models

TBD

## 12 Modeling Time Series with Changes in Regime

```
R code ______
> library("RcompHam94", lib.loc = ".")
```

#### 12.1 Modeling Changes in Regime

Page 697 describes an example of the application of Markov switching models to US GNP from 1951Q1 to 1984Q4.

```
R code

> data(gnpdata, package = "Ham94")

> selection <- subset(gnpdata, Quarter >= "1951-01-01" & Quarter <= 
+ "1984-04-01")

> d <- selection$Quarter[-1]

> g <- diff(100 * log(selection$GNP), lag = 1, differences = 1)
```

The actual implementation uses the technique of collapsing multi-period states into a single state, p691, p698. During the maximum likelihood estimation process the state probabilities will change, but the layout of the matrix is still the same. The following code fragment precalculates the transition matrix structure with the five possible values, then uses a separate 5 element lookup vector to populate it.

The bulk of the work is done by the following function, based on the algorithm in section 22.4. Ergodic probabilities are defined as on page 684, including equation [22.2.26]. The loop uses equations [22.4.24], [22.4.2], [22.4.5], [22.4.8], [22.4.7], [22.4.6] and [22.4.14].

```
R code
> infer.regimes <- function(THETA, YT) {</pre>
      phi <- THETA[grep("phi*", names(THETA))]</pre>
      mu <- THETA[grep("mu*", names(THETA))]</pre>
      sigma <- THETA["sigma"]</pre>
      p11star <- THETA["p11star"]</pre>
      p22star <- THETA["p22star"]</pre>
      T <- length(YT)
      tp <- c(0, p11star, 1 - p22star, 1 - p11star, p22star)</pre>
      P <- array(tp[transit], c(nstates, nstates))</pre>
      A <- rbind(diag(nstates) - P, rep(1, nstates))
      ergodic.pi <- (solve(t(A) %*% A) %*% t(A))[, nstates + 1]
      xi.t.t <- ergodic.pi %o% rep(1, nlags)
      xi.t.t_1 <- cbind(xi.t.t, ergodic.pi)</pre>
      log.likelihood <- 0</pre>
      for (tt in (nlags + 1):T) {
          residuals <- as.vector(((rep(1, nstates) %o% YT[tt:(tt -
               nlags)]) - array(mu[lagstate], c(nstates, nlags +
               1))) %*% c(1, -phi))
          eta.t <- dnorm(residuals, mean = 0, sd = sigma)
          fp <- eta.t * xi.t.t_1[, tt - 1]</pre>
          fpt <- sum(fp)</pre>
          xi.t.t <- cbind(xi.t.t, fp/fpt)</pre>
          log.likelihood <- log.likelihood + log(fpt)</pre>
          xi.t.t_1 \leftarrow cbind(xi.t.t_1, P %*% xi.t.t[, tt])
      }
      xi.t.T <- xi.t.t[, T] %0% 1
      for (tt in (T - 1):1) xi.t.T <- cbind(xi.t.t[, tt] * (t(P) %*%
           (xi.t.T[, 1]/xi.t.t_1[, tt + 1])), xi.t.T)
      list(log.likelihood = log.likelihood, xi.t.t = xi.t.t, xi.t.T = xi.t.T)
+ }
```

Initial values of the parameters for transition probabilities are set from historical averages. The phi and sigma values are obtained from a (non-state) regression of change in GDP on 4 of its own lags.

Now we are in a position to optimize, then calculated the smoothed probabilities from the optimal parameters.

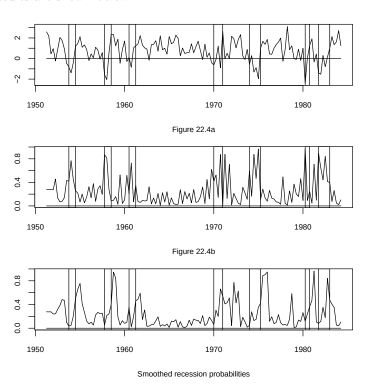
```
R code
> objective <- function(THETA, YT) {</pre>
      -infer.regimes(THETA, YT)$log.likelihood
+ }
> optimizer.results <- optim(par = THETA, hessian = TRUE, fn = objective,
      gr = NULL, YT = g)
> se <- diag(solve(optimizer.results$hessian))^0.5
> print(optimizer.results$par)
                                  output
                  p22star
                                                             phi1
                                                                          phi2
    p11star
0.869651020
             0.657920015 1.095327317 -0.198544833 0.311107386 0.092829514
        phi3
                     phi4
                                 sigma
-0.125038400 -0.007166502 0.872625052
                                 – R code –
> print(se)
                                 output .
              p22star
                                        mu2
                                                                         phi3
  p11star
                             mu1
                                                   phi1
                                                              phi2
0.13323951 0.04404274 0.23169921
                                        NaN 0.08762475 0.10748667 0.09374541
      phi4
                sigma
0.08826466
                  NaN
                                 R code -
> regimes <- infer.regimes(optimizer.results$par, g)
> recession.probability <- as.vector((1:nstates > nstates/2) %*%
```

smoothed.recession.probability <- as.vector((1:nstates > nstates/2) %\*%

regimes\$xi.t.t)

regimes\$xi.t.T)

The results are shown below.



# References

Hamilton, J. (1994): Time series analysis. Princeton University Press.

Kleiber, C., and A. Žeileis (2008): Applied Econometrics with R. Springer-Verlag, New York.