## ASTA

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## 1 Concepts, terminology and examples

### 1.1 Plotting, manipulation, trends and seasonality

- A time series is a collection of the same variable measured at different points in time. We will always assume the data is observed at equidistant points in time (i.e. same time difference between consecutive observations).
- To get appropriate summaries and plots of the data we must tell R that the data is a time series, which is called a ts object in $R$. A built-in example is the dataset AirPassengers which we will abbreviate to AP to save typing:

```
AP <- AirPassengers
plot(AP, ylab = "Bookings (1000s)")
```



- The generic function plot automatically detected that AP was a time series (ts object) and plotted the correct time on the horizontal axis (by calling plot.ts automatically).
- We can aggregate the time series over each year (sums the values by default, so we add FUN = mean to get mean number of monthly bookings for each year), which clearly shows an increasing trend in the number of passenger bookings:

```
APyear <- aggregate(AP, FUN = mean)
plot(APyear)
```



Such a systematic change in a time series that is not periodic is known as the trend of the series.

- The repeating (periodic) pattern within each year seen in the original data is known as seasonality or seasonal variation. The function cycle indicates which cycle (aka season) each observation belongs to (in this case 1-12). A boxplot of the number of bookings for each cycle component (month) clearly shows the seasonality:

```
cyc <- cycle(AP)
cyc <- factor(cyc, labels = month.abb)
boxplot(AP ~ cyc)
```



- If we only want to consider a subset of the data the function window is used:

```
APJun55Dec56 <- window(AP, start = c(1955, 6), end = c(1956, 12))
APJun55Dec56
```



- The window function can also sub-sample the time series according to the frequency:

```
APAug <- window(AP, start = c(1949, 8), end = end(AP), freq = TRUE)
```


### 1.2 Stochastic trend

- The trend for the passenger bookings data was very clear from the data (as well as the seasonal pattern), and a reasonable explanation may be increasing population, increasing prosperity and technological advances making tickets cheaper.
- In other situations the behaviour of the trend may be less clear and we will need to handle it differently. For the quarterly exchange rate between GBP and NZD the picture is less clear:

```
www <- "https://asta.math.aau.dk/eng/static/datasets?file=pounds_nz.dat"
exchange_data <- read.table(www, header = TRUE)
exchange <- ts(exchange_data, start = 1991, freq = 4)
plot(exchange)
```



## Time

- In general it is dangerous to extrapolate the trend, and this is even more so the case for a series with a stochastic trend, which this example may have.
- Especially for financial time series stochastic trends with abrupt changes to the trend are common.


### 1.3 Multiple series

- Monthly time series from Jan. 1958 to Dec. 1990 of supply of three goods in Australia:
- Electricity (Giga Watt hours)
- Beer (Mega liters)
- Chocolate (tonnes)

```
CBEdata <- read.table("https://asta.math.aau.dk/eng/static/datasets?file=cbe.dat",
            header = TRUE)
CBE <- ts(CBEdata, start = 1958, freq = 12)
plot(CBE)
```


## CBE



- For overlapping time series with the same frequency we can get the data for the overlapping period like this:

APCBE <- ts.intersect (AP, CBE)
plot(APCBE)


- In the realm of time series people often find spurious correlations and mistake them for causal relationships. E.g. the number of passenger bookings in USA looks very similar to electricity consumption in Australia and there is a strong correlation:

```
cor(APCBE)
```

| \#\# | AP | CBE.choc | CBE.beer | CBE.elec |
| :--- | ---: | ---: | ---: | ---: |
| \#\# AP | 1.0000000 | 0.6375808 | -0.4434747 | 0.8841668 |
| \#\# CBE.choc | 0.6375808 | 1.0000000 | -0.6204467 | 0.7644756 |
| \#\# CBE.beer | -0.4434747 | -0.6204467 | 1.0000000 | -0.3120179 |
| \#\# CBE.elec | 0.8841668 | 0.7644756 | -0.3120179 | 1.0000000 |

However, this does not imply that US air passengers really influence the electricity consumption in Australia (or vice versa)! Time series with similar trends and seasonality will typically show strong correlation even though they are unrelated. Better explanations may be due to similar population growth, seasonal patterns, etc.

### 1.4 Decomposition

### 1.4.1 Decomposition - trend term

- It can be useful to decompose the time series $x_{t}$ into the trend $m_{t}$, the seasonal variation $s_{t}$, and an error term $z_{t}$. Here we will focus on an additive decomposition $x_{t}=m_{t}+s_{t}+z_{t}$, but multiplicative models $x_{t}=m_{t} \cdot s_{t} \cdot z_{t}$ are also available.
- To estimate the trend a simple solution is to use a moving average by averaging the preceding and following $L$ values relative to the current point in time, where $L$ is chosen to average out the season effect. So if the season effect is weekly we choose $L=3$ such that

$$
\hat{m}_{t}=\left(x_{t-3}+x_{t-2}+x_{t-1}+x_{t}+x_{t+1}+x_{t+2}+x_{t+3}\right) / 7
$$

E.g. if the current time $t$ is a Tuesday the moving average averages today's value with the ones from the preceding Sat., Sun. and Mon. and the following Wed., Thu. and Fri. For an even season length like 12 months we have to do something else if we want an equal amount of past and future in the moving average, and we use half of the value six months in the past/future:

$$
\hat{m}_{t}=\frac{\frac{1}{2} x_{t-6}+x_{t-5}+x_{t-4}+\cdots+x_{0}+\cdots+x_{t+4}+x_{t+5}+\frac{1}{2} x_{t+6}}{12}
$$

E.g., the moving average for Jul. 1958 uses half the value in Jan. 1958 and half the value in Jan. 1959 and all the 11 values Feb.-Dec. 1958:

```
choc <- CBE[, "choc"]
i <- 7
(choc[i-6]/2 + sum(choc[(i-5):(i+5)]) + choc[i+6]/2)/12
```

```
## [1] 2413.625
```

```
## [1] 2413.625
```

This computation can be done for all time points (except the first six and last six) using decompose, which returns a list with a component trend (and other things to be discussed shortly):

```
choc_decomp <- decompose(choc)
choc_trend <- choc_decomp$trend
window(choc_trend, end = c(1958, 8))
\begin{tabular}{lrrrrrrr} 
\#\# & Jan & Feb & Mar & Apr & May & Jun & Jul \\
\#\# & 1958 & NA & NA & NA & NA & NA & NA 2413.625 \\
\#\# & Aug & & & & & \\
\#\# & 1958 & 2409.500 & & & & &
\end{tabular}
```

- Note that the procedure automatically chooses the appropriate size of the moving average based on the frequency of the time series, which was set to 12 in this case.


### 1.4.2 Decomposition - seasonal term

- To estimate the seasonal effect we subtract the trend and average for each period in the season (e.g. each month):

```
choc_no_trend <- choc - choc_trend
plot(choc_no_trend)
```



```
choc_jan <- mean(window(choc_no_trend, st = \(c(1958,1)\), freq \(=\) TRUE), na.rm \(=\) TRUE) choc_jan
\#\# [1] -2450.538
choc_may <- mean(window(choc_no_trend, st \(=c(1958,5)\), freq \(=\) TRUE), na.rm \(=\) TRUE) choc_may
\#\# [1] 1171.215
```

It appears that chocolate consumption typically is much higher in May than in January. The season effects are also calculated by decompose and stored in the list as seasonal:

```
plot(choc_decomp$seasonal)
```



### 1.4.3 Decomposition - random term

- The entire decomposition can also be plotted:
plot (choc_decomp)


## Decomposition of additive time series



- Here the random term is simply given by $z_{t}=x_{t}-m_{t}-s_{t}$, and this can be accessed through the random entry of the decomposition output choc_decomp\$random.


## 2 Stationarity and autocorrelation

- Consider the excess chocolate consumption data after adjusting for trend and season (and omitting the first six and last six entries which are NA):

```
choc_rand <- na.omit(choc_decomp$random)
plot(choc_rand)
```



This is a single realization of something we may think of as a random experiment. If things had been different (different weather, different chocolate advertisements, different economic fluctuations, ...) we could imagine another series.

This hypothetical scenario applies to any time series $x_{t}$ where we imagine our dataset is somehow randomly selected among an entire ensemble of possible time series.

- Calculating the mean of all such hypothetical series would give us the mean function $\mu(t)=E\left(x_{t}\right)$ for all $t$. Since we already adjusted for trend and season we expect a mean value around zero at each time point, and we might assume $\mu(t)=\mu$ for all $t$, where $\mu$ is a fixed constant (zero in this case). If this is truly the case we say the series is stationary in the mean (first order stationary). In general we can estimate $\mu$ by the sample mean as we have been used to:

$$
\hat{\mu}=\bar{x}=\frac{1}{n} \sum_{t=1}^{n} x_{t}
$$

- The variance (square of std. dev.) is in general also a function of time $\sigma^{2}(t)=E\left[\left(x_{t}-\mu(t)\right)^{2}\right]$. If we assume that we also have stationarity of the variance, $\sigma^{2}(t)=\sigma^{2}$, we can estimate $\sigma^{2}$ by the sample variance

$$
\hat{\sigma^{2}}=\frac{1}{n-1} \sum_{t=1}^{n}\left(x_{t}-\bar{x}\right)^{2} .
$$

- Now consider a process that is stationary in the mean and variance. The variables at different times may be correlated and the process is called second order stationary if the covariance only depends on the number of time steps between the variables, $\operatorname{Cov}\left(x_{t}, x_{t+k}\right)=\gamma(k)$ for all $t$, and $\gamma$ is called the autocovariance function. The number of time steps $k$ is called the lag. The autocovariance can be estimated from data:

$$
\hat{\gamma}(k)=c_{k}=\frac{1}{n} \sum_{t=1}^{n-k}\left(x_{t}-\bar{x}\right)\left(x_{t+k}-\bar{x}\right)
$$

- The more useful autocorrelation function (acf) is the normalized version that takes values between -1 and $+1: \rho(k)=\gamma(k) / \sigma^{2}$, which is estimated based on the estimate of the autocovariance above: $\hat{\rho}(k)=r_{k}=\frac{c_{k}}{c_{0}}$.
- The autocorrelation at lag 0 is always equal to $1, \rho(0)=1$ and $r_{0}=1$.
- For lag 1 it is the correlation between today's value $x_{t}$ and tomorrows value $x_{t+1}$. If we look directly at the chocolate data this correlation is very strong (simply because there is an increasing trend, so a high value is typically followed by a high value):

```
i <- 1:(length(choc)-1)
plot(choc[i], choc[i+1])
abline(lsfit(choc[i], choc[i+1]))
```


$\operatorname{cor}(\operatorname{choc}[i], \operatorname{choc}[i+1])$
\#\# [1] 0.8025919

- After removing the trend and seasonality there isn't really any dependence on the preceding value:

```
i <- 1:(length(choc_rand)-1)
plot(choc_rand[i], choc_rand[i+1])
abline(lsfit(choc_rand[i], choc_rand[i+1]))
```



```
cor(choc_rand[i], choc_rand[i+1])
## [1] -0.02278652
```


### 2.1 Correlogram (empirical acf)

- To calculate the empirical autocorrelation at many lags we use acf which produces a plot by default:

```
choc_acf <- acf(choc)
```


## Series choc



This is called the correlogram.

- It is possible to calculate the correlogram for any time series (also non-stationary). However, the theoretical properties only make sense for a stationary process.
- The correlogram value at lag $k, r_{k}$ is an estimate of the true autocorrelation $\rho(k)$.
- In particular if $\rho(k)=0$ then $r_{k}$ will usually not be exactly zero. Rather it is approximately normally distributed with a mean close to zero and standard deviation $1 / \sqrt{n}$, where $n$ is the length of the series. Based on this approximate significance bands are drawn at $\pm 1.96 / \sqrt{n}$.
- The significance bands are only valid for a single lag! Even if $\rho(k)=0$ for all lags $k=1,2, \ldots$ we expect $1 / 20$ of the $r_{k}$ 's to fall outside the line. Furthermore the estimates are heavily correlated so if one falls outside it is also more likely that the one next does.
- It is typical to look at a few specific lags. E.g. lag 1 for the immediate past and lag 12 if a season of length 12 is expected. A spike at lag 12 indicates that there is a seasonal pattern that has not been adequately accounted for. It appears the seasonal adjustment for the chocolate data hasn't been perfect since there is a somewhat large autocorrelation at lag 12 (1 year):

```
acf(choc_rand)
```


## Series choc_rand



### 2.2 Typical patterns in correlograms

It can be illustrative to consider a few specific examples of correlograms:

- Completely random:

```
x <- rnorm(100)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(x)
acf(x, main = "", lag.max = 30)
```




- Trend only:

```
x <- 1:100
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(x)
acf(x, main = "", lag.max = 30)
```



- Sinusoidal wave:

```
x <- sin((1:100)*2*pi/10)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(x)
acf(x, main = "", lag.max = 100)
```




- Repeated pattern:

```
x0 <- c(3,5,2,1,4)
x <- rep(x0, 20)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(x)
acf(x, main = "", lag.max = 30)
```



### 2.3 Partial autocorrelation function

- If a time series has a strong correlation with the values lagged by one time step it most likely also has a strong correlation with the values lagged by two time steps: When today's value is highly influenced by yesterdays, then yesterdays will be highly influenced by the day before, and thus the lag 2 value influences the current value through the lag 1 value.
- The partial autocorrelation (pacf) at lag 2 is the correlation between the time series and the lag 2 values after controlling for the lag 1 values.
- This is analogous to the multiple regression setup

$$
x_{t}=\beta_{0}+\beta_{1} x_{t-1}+\beta_{2} x_{t-2}+\epsilon
$$

where $\beta_{2}$ is a partial effect of the corresponding predictor $\left(x_{t-2}\right)$ when controlling for the other predictor ( $x_{t-1}$ ).

- In general the pacf at lag $k$ is the correlation of $x_{t}$ and $x_{t-k}$ after controlling for the intermediate variables $x_{t-1}, \ldots, x_{t-k-1}$.
- A more detailed description of partial correlation in the multiple regression setup is in Section 11.7 of Agresti (2013).
- For a strong deterministic trend the pacf only shows strong correlation in the first lag and after controlling for this the following lags show no extra correlation:

```
x <- 1:100
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(x)
pacf(x, main = "")
```




## 3 Models for time series data

There are two fundamentally different model classes for time series data.

- Discrete time stochastic processes
- Continuous time stochastic processes

Next lecture will focus on discrete time models and in the remainder we will touch upon continuous time models.

### 3.1 Continuous time stochastic processes

In this setup we see the underlying $X_{t}$ as a continuous function of $t$ for $t$ in some interval $[0, T]$.

In principle we imagine that there are infinitely many data points, simply because there are infinitely many time points between 0 and $T$. Of course this is never true: In practice we will always only have finitely many data points.

But it makes sense to believe that the real data actually contains all the data points. We are just not able to measure them (and to store them in a computer).

With a model for all datapoints, we are - through simulation - able to describe the behaviour of data. Also between the observations.

### 3.2 Wiener process

A key example of a process in continuous time will be the so-called Wiener process.
Three simulated realizations (black, blue and red) of this process can be seen here

```
## Warning in rgl.init(initValue, onlyNULL): RGL: unable to open X11 display
## Warning: 'rgl_init' failed, running with rgl.useNULL = TRUE
## Warning: .onUnload failed in unloadNamespace() for 'rgl', details:
## call: fun(...)
## error: object 'rgl_quit' not found
## Package 'Sim.DiffProc', version 4.0
## browseVignettes('Sim.DiffProc') for more informations.
```



A Wiener process has the following properties:

- It starts in 0: $W_{0}=0$.
- It has independent increments: For $0<s<t$ it holds that $W_{t}-W_{s}$ is independent of everything that has happened up to time $s$, that is $W_{u}$ for all $u \leq s$.
- It has normally distributed increments: For $0<s<t$ it holds that the increment $W_{t}-W_{s}$ is normally distributed with variance $t-s$ :

$$
W_{t}-W_{s} \sim \operatorname{Normal}\left(\mu=0, \sigma^{2}=t-s\right)
$$

The intuition of this process is that it somehow changes direction all the time: How the process changes after time $s$ will be independent of what has happened before time $s$. So whether the process should increase or decrease after $s$ will not be affected by how much it was increasing or decreasing before.

This gives the very bumpy behaviour over time.

### 3.3 Stochastic differential equations

A common way to define a continuous time stochastic process model is through a stochastic differential equation (SDE) which we will turn to shortly, but before doing so we will recall some basic things about ordinary differential equations.

Suppose $f$ is an unknown differentiable function and $f(0)=1$. Recall the mathematical description of a differential equation

$$
\frac{d f(t)}{d t}=-4 f(t)
$$

With the condition $f(0)=1$ this equation has the solution

$$
f(t)=\exp (-4 t)
$$



With a slightly unusual notation we can rewrite this as

$$
d f(t)=-4 \cdot f(t) d t
$$

This equation has the following (hopefully intuitive) interpretation:

- When time increases by a small amount $d t$ the value of $f$ changes (approximately) by $-4 f(t) d t$.

So when $t$ is increased, then $f$ is decreased. And the decrease is determined by the value of $f(t)$. That is why $f$ decreases slower and slower, when $t$ is increased.
We say that the function has a drift towards zero, and this drift is determined by the value of the function.

### 3.4 Stochastic differential equations

It will probably never be true that data behaves exactly like the exponentially decreasing curve on the previous slide.

Instead we will consider a model, where some random noise from a Wiener process has been added. Two different (black/blue) simulated realizations can be seen below


The type of process that is simulated above is described formally by the equation

$$
d X_{t}=-4 X_{t} d t+0.1 d W_{t}
$$

This is called a Stochastic Differential Equation (SDE), and the processes simulated above are called solutions of the stochastic differential equation.

The SDE $d X_{t}=-4 X_{t} d t+0.1 d W_{t}$ has two terms:

- $-4 X_{t} d t$ is the drift term.
- $0.1 d W_{t}$ is the diffusion term.

The intuition behind this notation is very similar to the intuition in the equation $d f(t)=-4 \cdot f(t) d t$ for an ordinary differential equation. When the time is increased by the small amount $d t$, then the process $X_{t}$ is increased by $-4 X_{t} d t$ AND by how much the process $0.1 W_{t}$ has increased on the time interval $[t, t+d t]$.

So this process has a drift towards zero, but it is also pushed in a random direction (either up or down) by the Wiener process (more precisely, the process $0.1 W_{t}$ )

## 4 Basic stochastic models

### 4.1 White noise

Suppose $y_{t}$ is a time series and we have a model which gives us predictions $\hat{y}_{t}$. Then the residuals are $x_{t}=y_{t}-\hat{y}_{t}$ for each time point $t=1, \ldots, n$. If the model has correctly captured the trend, seasonality and any autocorrelation in the original process $y_{t}$ then the correlogram (acf) of the residual process $x_{t}$ should show no obvious patterns. Such a residual process with no autocorrelation or other structure is called white noise.

### 4.1.1 Definition and properties of white noise

A time series $w_{t}, t=1, \ldots, n$ is white noise if the variables $w_{1}, w_{2}, \ldots, w_{n}$ are independent and identically distributed with a mean of zero.

From the definition it follows that white noise is a second order stationary process since the variance function $\sigma^{2}(t)=\sigma^{2}$ is the same constant for all $t$ and the autocovariance is $\operatorname{Cov}\left(w_{t}, w_{t+k}\right)=0$ for all $k \neq 0$ which does not depend on $t$. We summarize this as:

$$
\begin{gathered}
\mu=0 \\
\gamma(k)=\operatorname{Cov}\left(w_{t}, w_{t+k}\right)=\left\{\begin{array}{l}
\sigma^{2} \text { for } k=0 \\
0 \text { for } k \neq 0
\end{array}\right. \\
\rho(k)=\left\{\begin{array}{l}
1 \text { for } k=0 \\
0 \text { for } k \neq 0
\end{array}\right.
\end{gathered}
$$

Often we will also assume the distribution of each $w_{t}$ is Gaussian (i.e. $\left.w_{t} \sim \operatorname{norm}(0, \sigma)\right)$ and then we call it Gaussian white noise.

### 4.1.2 Simulation of white noise

To understand how white noise behaves we can simulate it with R and plot both the series and the autocorrelation:

```
w <- rnorm(100, mean = 0, sd = 1)
par(mfrow = c(2,1), mar = c(4,4,0,0))
ts.plot(w)
acf(w)
```



It is a good idea to repeat this simulation and plot a few times to appreciate the variablility of the results.

### 4.2 Random walk

A time series $x_{t}$ is called a random walk if

$$
x_{t}=x_{t-1}+w_{t}
$$

where $w_{t}$ is a white noise series. Using $x_{t-1}=x_{t-2}+w_{t-1}$ we get

$$
x_{t}=x_{t-2}+w_{t-1}+w_{t}
$$

Substituting for $x_{t-2}$ we get

$$
x_{t}=x_{t-3}+w_{t-2}+w_{t-1}+w_{t}
$$

Continuing this way we would get an infinite sum of white noise

$$
x_{t}=w_{t}+w_{t-1}+w_{t-2}+w_{t-3}+\ldots
$$

However, we will assume we have a fixed starting point $x_{0}=0$ such that

$$
x_{t}=w_{1}+w_{2}+\cdots+w_{t}
$$

### 4.2.1 Properties of random walk

A random walk $x_{t}$ has a constant mean function

$$
\mu(t)=0
$$

since the random walk at time $t$ is a sum of $t$ white noise terms that all have mean zero.
However, the variance function

$$
\sigma^{2}(t)=t \cdot \sigma^{2}
$$

clearly depends on the time $t$, so the process is not stationary. The variance function is derived from the general fact that for independent random variables, $y_{1}$ and $y_{2}$, the variance of the sum is

$$
\operatorname{Var}\left(y_{1}+y_{2}\right)=\operatorname{Var}\left(y_{1}\right)+\operatorname{Var}\left(y_{2}\right) .
$$

Thus,

$$
\operatorname{Var}\left(x_{t}\right)=\operatorname{Var}\left(w_{1}+w_{2}+\cdots+w_{t}\right)=\sigma^{2}+\sigma^{2}+\cdots+\sigma^{2}=t \sigma^{2}
$$

The non-stationary autocovariance function is

$$
\operatorname{Cov}\left(x_{t}, x_{t+k}\right)=t \sigma^{2}, \quad k=0,1, \ldots
$$

which only depends on how many white noise terms $x_{t}$ and $x_{t+k}$ have in common $(t)$ and not how far they are separated $(k)$.

By combining the two results we obtain the non-stationary autocorrelation function

$$
\operatorname{Cor}\left(x_{t}, x_{t+k}\right)=\frac{\operatorname{Cov}\left(x_{t}, x_{t+k}\right)}{\sqrt{\operatorname{Var}\left(x_{t}\right) \operatorname{Var}\left(x_{t+k}\right)}}=\frac{t \sigma^{2}}{\sqrt{t \sigma^{2}(t+k) \sigma^{2}}}=\frac{1}{\sqrt{1+k / t}}
$$

When $t$ is large compared to $k$ we have very high correlation (close to one) and even though the process is not stationary we expect the correlogram of a reasonably long random walk to show very slow decay.

### 4.2.2 Simulation of random walk

We already know how to simulate Gaussian white noise (with rnorm) and the random walk is just a cumulative sum of white noise:

```
w <- rnorm(1000)
rw <- cumsum(w)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(rw)
acf(rw, lag.max = 100)
```



### 4.2.3 Differencing

The slowly decaying acf for random walk is a classical sign of non-stationarity, indicating there may be some kind of trend. In this case there is no real trend, since the theoretical mean is constant zero, but we refer to the apparent trend which seems to change directions unpredictiably as a stochastic trend.

If a time series shows these signs of non-stationarity we can try to study the time series of differences and see if that is stationary and easier to understand:

$$
\nabla x_{t}=x_{t}-x_{t-1}
$$

Since we assume/define $x_{0}=0$ we get

$$
\begin{gathered}
\nabla x_{1}=x_{1} \\
\nabla x_{2}=x_{2}-x_{1} \\
\nabla x_{3}=x_{3}-x_{2}
\end{gathered}
$$

etc.
Specifically when we difference a random walk we recover the white noise series $\nabla x_{t}=w_{t}$ :

```
diffrw <- diff(rw)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
ts.plot(diffrw)
acf(diffrw, lag.max = 30)
```



In general if a time series needs to be differenced to become stationary we say that the series is integrated (of order 1).

### 4.2.4 Example: Exchange rate

We have previously studied the exchange rate from GBP to NZD and observed what looked like an unpredictable stochastic trend and we would like to see if it could reasonably be described as a random walk.

```
wWW <- "https://asta.math.aau.dk/eng/static/datasets?file=pounds_nz.dat"
exchange_data <- read.table(www, header = TRUE)
exchange <- ts(exchange_data, start = 1991, freq = 4)
plot(exchange)
```



To this end we difference the series and see if the difference looks like white noise:

```
diffexchange <- diff(exchange)
par(mfrow = c(2,1), mar = c(4,4,0.5,0.5))
plot(diffexchange)
acf(diffexchange)
```



The first order difference looks reasonably stationary, so the original exchange rate series could be considered integrated of order 1 . However, there is an indication of significant autocorrelation at lag 1 , so a random walk might not be a completely satisfactory model for this dataset.

## 5 Auto-regressive (AR) models

### 5.1 Auto-regressive model of order 1: AR(1)

A significant auto-correlation at lag 1 means that $x_{t}$ and $x_{t-1}$ are correlated so the previous value $x_{t-1}$ can be used to predict the current value $x_{t}$. This is the idea behind an auto-regressive model of order one AR(1):

$$
x_{t}=\alpha_{1} x_{t-1}+w_{t}
$$

where $w_{t}$ is white noise and the auto-regressive coefficient $\alpha_{1}$ is a parameter to be estimated from data. The model is only stationary if $-1<\alpha_{1}<1$ such that the dependence of the past decreases with time.

### 5.1.1 Properties of AR(1) models

For a stationary $\operatorname{AR}(1)$ model with $-1<\alpha_{1}<1$ it can be shown that

- $\mu(t)=0$
- $\operatorname{Var}\left(x_{t}\right)=\sigma^{2}(t)=\sigma^{2} /\left(1-\alpha_{1}^{2}\right)$
- $\gamma(k)=\alpha_{1}^{k} \sigma^{2} /\left(1-\alpha_{1}^{2}\right)$
- $\rho(k)=\alpha_{1}^{k}$

Furthermore, the partial autocorrelation of an $\operatorname{AR}(1)$ model is zero for all lags $k \geq 2$.
Below are the theoretical correlograms for the following $\mathrm{AR}(1)$ models:

- Model 1: $x_{t}=0.7 x_{t-1}+w_{t}$
- Model 2: $x_{t}=-0.7 x_{t-1}+w_{t}$
$\mathrm{k}<-0: 10$
$\operatorname{par}($ mfrow $=c(2,1), \operatorname{mar}=c(4,4,0.5,0.5))$
plot (k, $0.7^{\wedge} \mathrm{k}$, type $=$ "h", xlab = "")
plot $\left(k,(-0.7)^{\wedge} k\right.$, type $\left.=" h ", y l i m=c(-1,1)\right)$
abline(h=0)



### 5.1.2 Simulation of AR(1) models

$R$ has a built-in function arima.sim to simulate $\operatorname{AR}(1)$ and other more complicated models called ARMA and ARIMA. It needs the model (i.e. the autoregressive coefficient $\alpha_{1}$ ) and the desired number of time steps n. To simulate 200 time steps of $\operatorname{AR}(1)$ with $\alpha_{1}=0.7$ we do:

```
x <- arima.sim(model = list(ar = 0.7), n = 200)
plot(x)
```



```
acf(x)
points(k, 0.7^k, col = "red")
```


## Series x



Here we have compared the empirical correlogram with the theoretical values of the model.

### 5.1.3 Fitted AR(1) models

To estimate the autoregressive parameter $\alpha_{1}$ we use the function ar:

```
fit <- ar(x, order.max = 1)
```

The resulting object contains the value of the estimated parameter and a bunch of other information. In this case the input data are artificial so we know we should ideally get a value close to 0.7 :
fit\$ar
\#\# [1] 0.7009574

An estimate of the variance of the estimate $\hat{\alpha}_{1}$ is given in fit\$asy.var.coef (the estimated std. error is the square root of this):

```
fit$asy.var.coef
## [,1]
## [1,] 0.002568983
se <- sqrt(fit$asy.var.coef)
se
```

\#\# [,1]
\#\# [1,] 0.05068514
ci <- c(fit\$ar - 2*se, fit\$ar + 2*se)
ci
\#\# [1] 0.59958710 .8023277

There are several different methods that can be used to estimate the model, but we ignore the details of this. However, first step in the estimation is to subtract the mean $\bar{x}$ of the time series before doing anything else, so the model that is fitted is actually:

$$
x_{t}-\bar{x}=\alpha_{1} \cdot\left(x_{t-1}-\bar{x}\right)+w_{t}
$$

To predict the value of $x_{t}$ given $x_{t-1}$ we use that $w_{t}$ is white noise so we expect it to be zero on average:

$$
\hat{x}_{t}-\bar{x}=\hat{\alpha}_{1} \cdot\left(x_{t-1}-\bar{x}\right)
$$

So the predictions are given by

$$
\hat{x}_{t}=\bar{x}+\hat{\alpha}_{1} \cdot\left(x_{t-1}-\bar{x}\right), \quad t \geq 2
$$

Given the predictions we can estimate the model errors as usual by the model residuals:

$$
\hat{w}_{t}=x_{t}-\hat{x}_{t}, \quad t \geq 2
$$

If we believe the model describes the dataset well the residuals should look like a sample of white noise:

```
res <- na.omit(fit$resid)
par(mfrow = c(2,1), mar = c(4,4,1,1))
plot(res)
acf(res)
```



Time


This naturally looks good for this artificial dataset.

### 5.1.4 $\mathrm{AR}(1)$ model fitted to exchange rate

A random walk is an example of a $\operatorname{AR}(1)$ model with $\alpha_{1}=1$, and it is non-stationary. This didn't provide an ideal fit for the exchange rate dataset, so we might suggest a stationary $\operatorname{AR}(1)$ model with $\alpha_{1}$ as a parameter to be estimated from data:

```
fitexchange <- ar(exchange, order.max = 1)
fitexchange$ar
## [1] 0.890261
resexchange <- na.omit(fitexchange$resid)
par(mfrow = c(2,1), mar = c(4,4,1,1))
plot(resexchange)
acf(resexchange)
```



This does not appear to really provide a better fit than the random walk model proposed earlier.
An alternative would be to propose a $\mathrm{AR}(1)$ model for the differenced time series $\nabla x_{t}=x_{t}-x_{t-1}$ :

```
dexchange <- diff(exchange)
fitdiff <- ar(dexchange, order.max = 1)
fitdiff$ar
## [1] 0.3451507
resdiff <- na.omit(fitdiff$resid)
par(mfrow = c(2,1), mar = c(4,4,1,1))
plot(resdiff)
acf(resdiff)
```




## Lag

### 5.1.5 Prediction from AR(1) model

We can use a fitted $\mathrm{AR}(1)$ model to predict future values of a time series. If the last observed time point is $t$ then we predict $x_{t+1}$ using the equation given previously:

$$
\hat{x}_{t+1}=\bar{x}+\hat{\alpha}_{1} \cdot\left(x_{t}-\bar{x}\right)
$$

If we want to predict $x_{t+2}$ we use

$$
\hat{x}_{t+2}=\bar{x}+\hat{\alpha}_{1} \cdot\left(\hat{x}_{t+1}-\bar{x}\right) .
$$

And we can continue this way. Prediction is performed by predict in R. E.g. for the $\mathrm{AR}(1)$ model fitted to the exchange rate data the last observation is in third quarter of 2000 . If we want to predict 1 year ahead to third quarter of 2001 (probably a bad idea due to the stochastic trend):

```
pred1 <- predict(fitexchange, n.ahead = 4)
pred1
## $pred
## Qtr1 Qtr2 Qtr3 Qtr4
## 2000 3.453332
## 2001 3.384188 3.322631 3.267830
##
## $se
## Qtr1 Qtr2 Qtr3 Qtr4
## 2000 0.1767767
## 2001 0.2366805 0.2750411 0.3020027
```

Note how the prediction returns both the predicted value and a standard error for this value. So we predict that the exchange rate in third quarter of 2001 would be within $3.27 \pm 0.6$ with approximately $95 \%$ probability.
We can plot a prediction and approximate $95 \%$ pointwise prediction intervals with ts.plot (where we use a 10 year prediction - which is a very bad idea - to see how it behaves in the long run):

```
pred10 <- predict(fitexchange, n.ahead = 40)
lower10 <- pred10$pred-2*pred10$se
upper10 <- pred10$pred+2*pred10$se
ts.plot(exchange, pred10$pred, lower10, upper10, lty = c(1,2,3,3))
```



### 5.2 Auto-regressive models of higher order

The first order auto-regressive model can be generalised to higher order by adding more lagged terms to explain the current value $x_{t}$. An $\mathrm{AR}(\mathrm{p})$ process is

$$
x_{t}=\alpha_{1} x_{t-1}+\alpha_{2} x_{t-2}+\cdots+\alpha_{p} x_{t-p}+w_{t}
$$

where $w_{t}$ is white noise and $\alpha_{1}, \alpha_{2}, \ldots, \alpha_{p}$ are parameters to be estimated from data.
The parameters cannot be chosen arbitrarily if we want the model to be stationary. To check that a given $\mathrm{AR}(\mathrm{p})$ model is stationary we must find all the roots of the characteristic equation

$$
1-\alpha_{1} z-\alpha_{2} z^{2}-\cdots-\alpha_{p} z^{p}=0
$$

and check that the absolute value of each root is greater than 1.

### 5.2.1 Estimation of AR(p) models

For an $\mathrm{AR}(\mathrm{p})$ model there are typically two things we need to estimate:

1. The maximal non-zero lag $p$ in the model.
2. The autoregressive coefficients/parameters $\alpha_{1}, \ldots, \alpha_{p}$.

For the first point we note the following theoretical property of $A R(p)$ processes:
For an $\mathrm{AR}(\mathrm{p})$ process the theoretical value of the partial autocorrelation at lag $k$ is $\alpha_{k}$. Thus, for all lags greater than $p$ the theoretical partial autocorrelation is zero. So heuristically we can choose the maximal lag $p$ of an $\mathrm{AR}(\mathrm{p})$ process by looking at when the estimated partial autocorrelation function is close to zero.

In practice it can be hard to say exactly when the pacf is sufficienty close to zero and more advanced methods are often used. The function ar in R uses AIC (Akaike's Information Criterion) to automatically select the value for $p$.

Once the order is chosen and the estimates $\hat{\alpha}_{1}, \ldots, \hat{\alpha}_{p}$ are found the corresponding standard errors can be found as the square root of the diagonal of the matrix stored as asy.var.coef in the fitted model object.

### 5.2.2 Example of AR(p) model

We use an example of monthly global temperatures expressed as anomalies from the monthly average in 1961-1990. We reduce the dataset to the yearly mean temperature, and fit an $A R(p)$ model to this. A good fit would indicate that the higher temperatures over the last decade could be explained by a purely stochastic process which just has dependence on the temperature anomalies from previous year and eventually might as well start decreasing again. However, this does not mean that there is no climate crisis! There is lots of scientific evidence of this based on much more complicated models and more detailed data.

```
global_data <- scan("https://asta.math.aau.dk/eng/static/datasets?file=global.dat")
global_monthly <- ts(global_data, st = c(1856,1), end = c(2005,12), freq = 12)
global <- aggregate(global_monthly, FUN = mean)
plot(global)
```



```
globalfit <- ar(global, order.max = 10)
globalfit
##
## Call:
## ar(x = global, order.max = 10)
##
## Coefficients:
## 
## 0.6825 0.0032 0.0672 0.1730
##
## Order selected 4 sigma^2 estimated as 0.01371
globalresid <- na.omit(globalfit$resid)
par(mfrow = c(2,1), mar = c(4,4,1,1))
plot(globalresid)
acf(globalresid, lag.max = 30)
```



## 6 Moving average models

Another class of models are moving average (MA) models. An moving average process of order $q$, MA(q), is defined by

$$
x_{t}=w_{t}+\beta_{1} w_{t-1}+\beta_{2} w_{t-2}+\cdots+\beta_{q} w_{t-q}
$$

where $w_{t}$ is a white noise process with mean zero and variance $\sigma_{w}^{2}$ and $\beta_{1}, \beta_{2}, \ldots, \beta_{q}$ are parameters to be estimated.

Since a moving average process is a finite sum of stationary white noise terms it is itself stationary and therefore the mean and variance is time-invariant (same constant mean and variance for all $t$ ):

- Mean $\mu(t)=0$
- Variance $\sigma^{2}(t)=\sigma_{w}^{2}\left(1+\beta_{1}^{2}+\beta_{2}^{2}+\cdots+\beta_{q}^{2}\right)$

The autocorrelation function, for $k \geq 0$, is

$$
\rho(k)= \begin{cases}1 & k=0 \\ \sum_{i=0}^{q-k} \beta_{i} \beta_{i+k} / \sum_{i=0}^{q} \beta_{i}^{2} & k=1,2, \ldots, q \\ 0 & k>q\end{cases}
$$

where $\beta_{0}=1$.

### 6.1 Simulation of MA(q) processes

To simulate a MA(q) process we just need the white noise process $w_{t}$ and then transform it using the MA coefficients. If we e.g. want to simulate a model with $\beta_{1}=-0.7, \beta_{2}=0.5$, and $\beta_{3}=-0.2$ we can use arima.sim:
xsim <- arima.sim(list(ma $=c(-.7, .5,-.2)), \mathrm{n}=200)$ plot(xsim)


The theoretical autocorrelations are in this case:

$$
\begin{gathered}
\rho(1)=\frac{1 \cdot(-0.7)+(-0.7) \cdot 0.5+0.5 \cdot(-0.2)}{1+(-0.7)^{2}+0.5^{2}+(-0.2)^{2}}=-0.65 \\
\rho(2)=\frac{1 \cdot 0.5+(-0.7) \cdot(-0.2)}{1+(-0.7)^{2}+0.5^{2}+(-0.2)^{2}}=0.36 \\
\rho(3)=\frac{1 \cdot(-0.2)}{1+(-0.7)^{2}+0.5^{2}+(-0.2)^{2}}=-0.11
\end{gathered}
$$

acf(xsim)
points(1:3, c(-.65, .36, -.11), col = "red")

## Series xsim



### 6.1.1 Estimation of MA(q) models

To estimate the parameters of a MA(q) model we use arima:

```
xfit <- arima(xsim, order = c(0,0,3))
xfit
##
## Call:
## arima(x = xsim, order = c(0, 0, 3))
##
## Coefficients:
\begin{tabular}{lrrrr} 
\#\# & ma1 & ma2 & ma3 & intercept \\
\#\# & -0.5879 & 0.4464 & -0.1359 & 0.0301 \\
\#\# s.e. & 0.0734 & 0.0950 & 0.0906 & 0.0500
\end{tabular}
##
## sigma^2 estimated as 0.9567: log likelihood = -279.62, aic = 569.25
```

The function arima does not include automatic selection of the order of the model so this has to be chosen beforehand or selected by comparing several proposed models using the AIC.

## 7 Mixed models: Auto-regressive moving average models

A time series $x_{t}$ follows an auto-regressive moving average (ARMA) process of order $(p, q)$, denoted $A R M A(p, q)$, if

$$
x_{t}=\alpha_{1} x_{t-1}+\alpha_{2} x_{t-2}+\cdots+\alpha_{p} x_{t-p}+w_{t}+\beta_{1} w_{t-1}+\beta_{2} w_{t-2}+\cdots+\beta_{q} w_{t-q}
$$

where $w_{t}$ is a white noise process and $\alpha_{1}, \alpha_{2}, \ldots, \alpha_{p}, \beta_{1}, \beta_{2}, \ldots, \beta_{q}$ are parameters to be estimated.
We can simulate an ARMA model with arima.sim. E.g. an ARMA(1,1) model:

```
xarma <- arima.sim(model = list(ar = -0.6, ma = 0.5), n = 200)
plot(xarma)
```



Estimation is done with arima as before.

### 7.0.1 Example with exchange rate data

For the exchange rate data we may e.g. suggest either a $\mathrm{AR}(1)$, $\mathrm{MA}(1)$ or $\operatorname{ARMA}(1,1)$ model. We can compare fitted model using AIC (smaller is better):

```
exchange_ar <- arima(exchange, order = c(1,0,0))
AIC(exchange_ar)
## [1] -37.40417
exchange_ma <- arima(exchange, order = c(0,0,1))
AIC(exchange_ma)
```

```
exchange_arma <- arima(exchange, order = c(1,0,1))
AIC(exchange_arma)
## [1] -42.27357
exchange_arma
##
## Call:
## arima(x = exchange, order = c(1, 0, 1))
##
## Coefficients:
\begin{tabular}{lrrr} 
\#\# & ar1 & ma1 & intercept \\
\#\# & 0.8925 & 0.5319 & 2.9597 \\
\#\# s.e. & 0.0759 & 0.2021 & 0.2435
\end{tabular}
##
## sigma^2 estimated as 0.01505: log likelihood = 25.14, aic = -42.27
par(mfrow = c(2,1), mar = c(4,4,1,1))
resid_arma <- na.omit(exchange_arma$residuals)
plot(resid_arma)
acf(resid_arma)
```



Time


