# R Package diagram: visualising simple graphs, flowcharts, and webs

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#### Abstract

This document describes how to use the **diagram** package (Soetaert 2009a) for plotting small networks, flow charts, and (food) webs.

Together with R-package shape (Soetaert 2009b) this package has been written to produce the figures of the book (Soetaert and Herman 2009b).

The electrical network symbols were added to produce a figure of the book (Soetaert, Cash, and Mazzia 2012)

Keywords: diagram, food web, flow chart, arrows, R.

# 1. Introduction

There are three ways in which package **diagram** can be used:

- function plotmat takes as input a matrix with transition coefficients or interaction strengths. It plots the corresponding network consisting of (labeled) boxes (the components) connected by arrows. Each arrow is labeled with the value of the coefficients.
- function plotweb takes as input a matrix with (flow) values, and plots a web. Here the components are connected by arrows whose thickness is determined by the value of the coefficients.
- Flowcharts can be made by adding separate objects (textboxes) to the figure and connecting these with arrows.

Three datasets have been included:

- Rigaweb, the planktonic food web of the Gulf of Riga (Donali, Olli, Heiskanen, and Andersen 1999).
- Takapotoweb, the Takapoto atoll planktonic food web (Niquil, Jackson, Legendre, and Delesalle 1998).
- Teasel, the transition matrix describing the population dynamics of Teasel, a European perennial weed ((Caswell 2001; Soetaert and Herman 2009b).

The food webs were generated using R packages **LIM** and **limSolve** (Soetaert, Van den Meersche, and van Oevelen 2009; Soetaert and van Oevelen 2009) which contain functions to read and solve food web problems respectively.

# 2. plotmat - plotting networks based on matrix input

This is the quickest method of plotting a network. The network is specified in a matrix, which gives the magnitudes of the links (from columns to rows).

The position of the elements (boxes) is specified by argument pos. Thus, setting pos=c(1,2,1) indicates that the 4 elements will be arranged in three equidistant rows; on the first row one element, on the second row two elements and on the third row one element.

# 2.1. Simple examples

Below are some simple examples of the use of plotmat. In the first graph - four simple boxes are put; no arrows drawn

The second graph contains round boxes with arrows, labeled "flow"

The third graph has diamond-shaped boxes including self-arrows.

The fourth graph has hexagonal-shaped boxes, with curved arrows. The arrows are enlarged and the arrowhead pointing from box 2 to 4 is colored red.

```
> par(mar = c(1, 1, 1, 1), mfrow = c(2, 2))
> #
> #
> names <- c("A", "B", "C", "D")
> M <- matrix(nrow = 4, ncol = 4, byrow = TRUE, data = 0)
> plotmat(M, pos = c(1, 2, 1), name = names, lwd = 1,
          box.lwd = 2, cex.txt = 0.8, box.size = 0.1,
          box.type = "square", box.prop = 0.5)
+
> #
> M[2, 1] <- M[3, 1] <- M[4, 2] <- M[4, 3] <- "flow"
> plotmat(M, pos = c(1, 2, 1), curve = 0, name = names, lwd = 1,
          box.lwd = 2, cex.txt = 0.8, box.type = "circle", box.prop = 1.0)
> #
> #
> diag(M) <- "self"
> plotmat(M, pos = c(2, 2), curve = 0, name = names, lwd = 1, box.lwd = 2,
          cex.txt = 0.8, self.cex = 0.5, self.shiftx = c(-0.1, 0.1, -0.1, 0.1),
          box.type = "diamond", box.prop = 0.5)
> M <- matrix(nrow = 4, ncol = 4, data = 0)
> M[2, 1] <- 1; M[4, 2] <- 2; M[3, 4] <- 3; M[1, 3] <- 4
> Col <- M
> Col[] <- "black"
> Col[4, 2] <- "darkred"
> pp < -plotmat(M, pos = c(1, 2, 1), curve = 0.2, name = names, lwd = 1,
                box.lwd = 2, cex.txt = 0.8, arr.type = "triangle",
```

```
+ box.size = 0.1, box.type = "hexa", box.prop = 0.25,
+ arr.col = Col, arr.len = 1)
> mtext(outer = TRUE, side = 3, line = -1.5, cex = 1.5, "plotmat")
> #
> par(mfrow = c(1, 1))
```

The contents of pp shows the position of the various items.

> pp

#### \$arr

```
ArrowX
                                                  ArrowY
                                                             TextX
 row col
             Angle Value
                                 rad
                                                                        TextY
   2
           53.1301
                       1 0.08333333 0.3085298 0.7169284 0.2843333 0.7346667
2
        2 -53.1301
                       2 0.08333333 0.3077450 0.2841193 0.2843333 0.2653333
                       4 0.08333333 0.6918629 0.7164048 0.7156667 0.7346667
        3 -53.1301
3
    1
   3
        4 53.1301
                       3 0.08333333 0.6910769 0.2825485 0.7156667 0.2653333
```

#### \$comp

```
x y
[1,] 0.50 0.8333333
[2,] 0.25 0.5000000
[3,] 0.75 0.5000000
[4,] 0.50 0.1666667
```

#### \$radii

```
x y
[1,] 0.1 0.025
[2,] 0.1 0.025
[3,] 0.1 0.025
[4,] 0.1 0.025
```

# \$rect

```
xleft ybot xright ytop
[1,] 0.40 0.8083333 0.60 0.8583333
[2,] 0.15 0.4750000 0.35 0.5250000
[3,] 0.65 0.4750000 0.85 0.5250000
[4,] 0.40 0.1416667 0.60 0.1916667
```

#### 2.2. a schematic representation of an ecosystem model

In the example below, first the main components and arrows are drawn (plotmat), and the output of this function written in list pp. This contains, a.o. the positions of the components (boxes), arrows, etc.. It is used to draw an arrow from the middle of the arrow connecting fish and zooplankton ("ZOO") to detritus. Function straightarrow (see below) is used to draw this arrow.

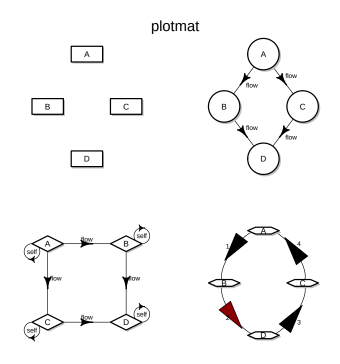


Figure 1: Four simple examples of plotmat

```
> names <- c("PHYTO", "NH3", "ZOO", "DETRITUS", "BotDET", "FISH")
> M <- matrix(nrow = 6, ncol = 6, byrow = TRUE, data = c(
     p n z
                d
                     b
                         f
      0, 1, 0,
                0,
                    0,
                         0, #p
      0, 0, 4, 10, 11,
                         0, #n
      2, 0, 0, 0,
                    0, 0, #z
      8, 0, 13, 0,
                    0, 12, #d
      9, 0, 0, 7,
                    0, 0, #b
                0,
      0, 0, 5,
                    0,
      ))
> #
> pp <- plotmat(M, pos = c(1, 2, 1, 2), curve = 0, name = names,
                lwd = 1, box.lwd = 2, cex.txt = 0.8,
                box.type = "square", box.prop = 0.5, arr.type = "triangle",
                arr.pos = 0.4, shadow.size = 0.01, prefix = "f",
                main = "NPZZDD model")
> #
           <- pp$comp[names=="PHYTO"]</pre>
> phyto
           <- pp$comp[names=="Z00"]</pre>
> zoo
> nh3
           <- pp$comp[names=="NH3"]</pre>
> detritus <- pp$comp[names=="DETRITUS"]</pre>
> fish
           <- pp$comp[names=="FISH"]</pre>
> #
```

#### NPZZDD model

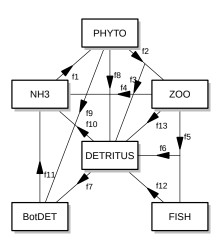


Figure 2: An NPZZDD model

```
> # flow5->detritus
> #
> m2 <- 0.5*(zoo+fish)
> m1 <- detritus
> m1[1] <- m1[1] + pp$radii[4,1]</pre>
> mid <- straightarrow (to = m1, from = m2, arr.type = "triangle",
                         arr.pos = 0.4, lwd = 1)
> text(mid[1], mid[2]+0.03, "f6", cex = 0.8)
> # flow2->detritus
> #
> m2 <- 0.5*(zoo+phyto)
> m1 <- detritus
> m1[1] <-m1[1] + pp$radii[3,1]*0.2
> m1[2] <-m1[2] + pp$radii[3,2]</pre>
> mid <- straightarrow (to = m1, from = m2, arr.type = "triangle",</pre>
                         arr.pos = 0.3, lwd = 1)
> text(mid[1]-0.01, mid[2]+0.03, "f3", cex = 0.8)
```

# 2.3. Plotting a transition matrix

The next example uses formulae to label the arrows <sup>1</sup>. This is done by passing a data.frame rather than a matrix to function plotmat

```
> # Create population matrix
> #
                   <- 6
> Numgenerations
> DiffMat <- matrix(data = 0, nrow = Numgenerations, ncol = Numgenerations)
> AA <- as.data.frame(DiffMat)
> AA[[1,4]] <- "f[3]"
> AA[[1,5]] <- "f[4]"
> AA[[1,6]] <- "f[5]"
> #
> AA[[2,1]] <- "s[list(0,1)]"
> AA[[3,2]] <- "s[list(1,2)]"
> AA[[4,3]] <- "s[list(2,3)]"
> AA[[5,4]] <- "s[list(3,4)]"
> AA[[6,5]] <- "s[list(4,5)]"
> #
> name <- c(expression(Age[0]), expression(Age[1]), expression(Age[2]),</pre>
             expression(Age[3]), expression(Age[4]), expression(Age[5]))
> #
> plotmat(A = AA, pos = 6, curve = 0.7, name = name, lwd = 2,
          arr.len = 0.6, arr.width = 0.25, my = -0.2,
          box.size = 0.05, arr.type = "triangle", dtext = 0.95,
          main = "Age-structured population model 1")
```

#### 2.4. Another transition matrix

The data set Teasel contains the transition matrix of the population dynamics model of teasel (Dipsacus sylvestris), a European perennial weed, (Caswell 2001; Soetaert and Herman 2009b)

# > Teasel

```
DS 1yr DS 2yr R small R medium R large
          0.000
                  0.00
                          0.000
                                   0.000
                                           0.000 322.380
DS 1yr
                                                    0.000
DS 2yr
          0.966
                  0.00
                          0.000
                                   0.000
                                           0.000
R small
          0.013
                  0.01
                          0.125
                                   0.000
                                           0.000
                                                    3.448
R medium 0.007
                  0.00
                          0.125
                                   0.238
                                           0.000 30.170
R large
          0.008
                  0.00
                          0.038
                                   0.245
                                           0.167
                                                    0.862
          0.000
                  0.00
                          0.000
                                   0.023
                                           0.750
                                                    0.000
```

This dataset is plotted using curved arrows; we specify the curvature in a matrix called curves.

<sup>&</sup>lt;sup>1</sup>This is now possible thanks to Yvonnick Noel, Univ. Rennes, France

#### Age-structured population model 1

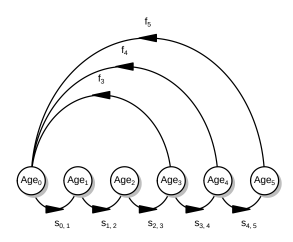


Figure 3: A transition matrix

# 3. plotweb - plotting webs based on matrix input

Given a matrix containing flow values (from rows to columns), function plotweb will plot a web. The elements are positioned on a circle, and connected by arrows; the magnitude of web flows determines the thickness of the arrow.

This function is less flexible than plotmat, although it does allow to color the arrows differently.

```
> BB <- matrix(nrow = 20, ncol = 20, 1:20)
> diag(BB) <- 0
```

# **Teasel population model**

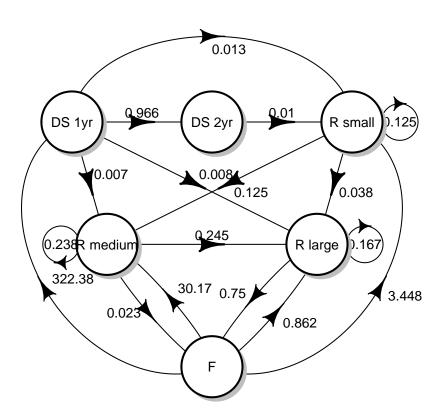


Figure 4: The Teasel data set

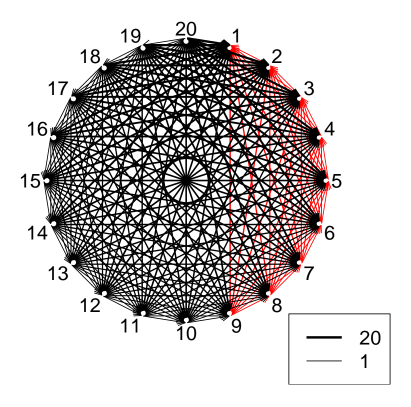


Figure 5: Plotweb

```
> Col <- BB
> Col[] <- "black"
> Col[BB<10]<- "red"
> plotweb(BB, legend = TRUE, maxarrow = 3, arr.col = Col)
> par(mfrow = c(1, 1))
```

# 3.1. Foodwebs

Dataset Rigaweb ((Donali et al. 1999) contains flow values for the food web of the Gulf of Riga planktonic system.

# > Rigaweb

	P1	P2	В	N	Z	D	DOC	C02
P1	0.0000	0.0000	0.0000	4.12297	10.49431	0.000000	1.565910	17.22501
P2	0.0000	0.0000	0.0000	0.00000	16.79755	4.457164	2.723090	29.95399

```
0.0000 0.0000
                           0.0000 9.44000 0.00000 0.000000 0.000000 244.99223
В
N
            0.0000 0.0000
                           13.40297
                           0.0000 0.00000 0.00000 3.183226 3.963226
Ζ
            0.0000 0.0000
                                                                30.19580
D
            0.0000 0.0000
                           0.0000 0.00000 12.34039 0.000000 0.000000
                                                                 0.00000
            0.0000 0.0000 261.1822 0.00000 0.00000 0.000000 0.000000
DOC
                                                                 0.00000
C<sub>02</sub>
            31.3182 54.4618
                           0.00000
Sedimentation 0.0000 0.0000
                           0.00000
            Sedimentation
P1
                    0.10
P2
                    0.34
В
                    0.00
N
                    0.00
Z
                   0.78
D
                   13.92
DOC
                    0.00
C<sub>0</sub>2
                    0.00
Sedimentation
                   0.00
> plotweb(Rigaweb, main = "Gulf of Riga food web",
        sub = "mgC/m3/d", val = TRUE)
```

# 4. functions to create flow charts

The various functions are given in table  $(1)^2$ .

The code below generates a flow chart

```
> par(mar = c(1, 1, 1, 1))
> openplotmat()
> elpos \leftarrow coordinates (c(1, 1, 2, 4))
> fromto <- matrix(ncol = 2, byrow = TRUE,
                    data = c(1, 2, 2, 3, 2, 4, 4, 7, 4, 8))
> nr
         <- nrow(fromto)
> arrpos <- matrix(ncol = 2, nrow = nr)</pre>
> for (i in 1:nr)
   arrpos[i, ] <- straightarrow (to = elpos[fromto[i, 2], ],</pre>
                                 from = elpos[fromto[i, 1], ],
                                 1wd = 2, arr.pos = 0.6, arr.length = 0.5)
                                     lab = "start",
                                                               box.col = "green",
> textellipse(elpos[1,], 0.1,
              shadow.col = "darkgreen", shadow.size = 0.005, cex = 1.5)
+
             (elpos[2,], 0.15, 0.05, lab = "found term?",
                                                               box.col = "blue",
> textrect
              shadow.col = "darkblue", shadow.size = 0.005, cex = 1.5)
             (elpos[4,], 0.15, 0.05, lab = "related?",
> textrect
                                                               box.col = "blue",
              shadow.col = "darkblue", shadow.size = 0.005, cex = 1.5)
> textellipse(elpos[3,], 0.1, 0.1, lab = c("other","term"), box.col = "orange",
```

<sup>&</sup>lt;sup>2</sup>textparallel was implemented by Michael Folkes, Canada

Karline Soetaert

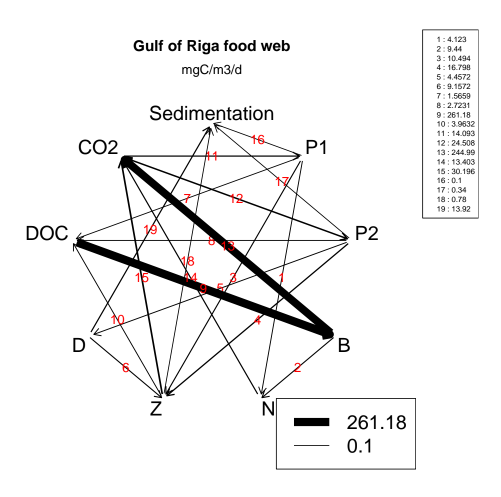


Figure 6: The Gulf of Riga data set

Table 1: Summary of flowchart functions

Function	Description
openplotmat	creates an empty plot
coordinates	calculates coordinates of elements, neatly arranged in rows/columns
bentarrow	adds 2-segmented arrow between two points
$\operatorname{curvedarrow}$	adds curved arrow between two points
segmentarrow	adds 3-segmented arrow between two points
selfarrow	adds a circular self-pointing arrow
splitarrow	adds a branched arrow between several points
straightarrow	adds straight arrow between two points
treearrow	adds dendrogram-like branched arrow between several points
shadowbox	adds a box with a shadow to a plot
textdiamond	adds lines of text in a diamond-shaped box to a plot
textellipse	adds lines of text in a ellipse-shaped box to a plot
textempty	adds lines of text on a colored background to a plot
texthexa	adds lines of text in a hexagonal box to a plot
textmulti	adds lines of text in a multigonal box to a plot
textparallel	adds lines of text in a parallelogram to a plot
textplain	adds lines of text to a plot
textrect	adds lines of text in a rectangular-shaped box to a plot
textround	adds lines of text in a rounded box to a plot

The different types of text boxes are generated with the following code:

```
> openplotmat(main = "textbox shapes")
> rx <- 0.1
> ry <- 0.05
> pos <- coordinates(c(1, 1, 1, 1, 1, 1, 1, 1, 1, 1), mx = -0.2)
> textdiamond(mid = pos[1,], radx = rx, rady = ry, lab = LETTERS[1],
+ cex = 2, shadow.col = "lightblue")
```

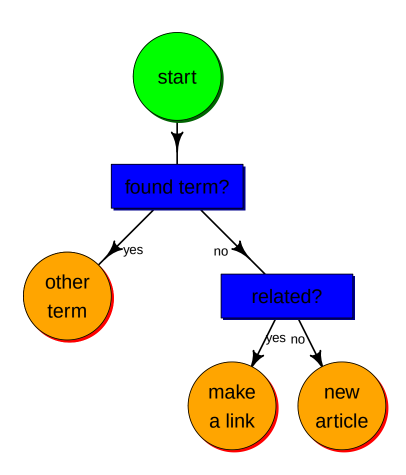


Figure 7: A flow chart

```
> textellipse(mid = pos[2,], radx = rx, rady = ry, lab = LETTERS[2],
              cex = 2, shadow.col = "blue")
> texthexa(mid = pos[3,], radx = rx, rady = ry, lab = LETTERS[3],
           cex = 2, shadow.col = "darkblue")
> textmulti(mid = pos[4,], nr = 7, radx = rx, rady = ry, lab = LETTERS[4],
           cex = 2, shadow.col = "red")
> textrect(mid = pos[5,], radx = rx, rady = ry, lab = LETTERS[5],
           cex = 2, shadow.col = "darkred")
> textround(mid = pos[6,], radx = rx, rady = ry, lab = LETTERS[6],
           cex = 2, shadow.col = "black")
> textparallel(mid = pos[7,], radx = rx, rady = ry, lab = LETTERS[7],
           cex = 2, theta = 40, shadow.col = "black")
> textempty(mid = pos[8,], lab = LETTERS[8], cex = 2, box.col = "yellow")
> pos[ ,1] <- pos[ ,1] + 0.5
> text(pos[ ,1],pos[ ,2], c("textdiamond", "textellipse", "texthexa",
                    "textmulti", "textrect", "textround",
                    "textparallel", "textempty"))
The different types of arrows are generated with the following code:
> par(mar = c(1, 1, 1, 1))
> openplotmat(main = "Arrowtypes")
> elpos <- coordinates (c(1, 2, 1), mx = 0.1, my = -0.1)
> curvedarrow(from = elpos[1, ], to = elpos[2, ], curve = -0.5,
              1ty = 2, 1col = 2)
> straightarrow(from = elpos[1, ], to = elpos[2, ], lty = 3, lcol = 3)
> segmentarrow (from = elpos[1, ], to = elpos[2, ], lty = 1, lcol = 1)
               (from = elpos[2:3, ], to = elpos[4, ], lty = 4, lcol = 4)
> treearrow
> bentarrow (from = elpos[3, ], to = elpos[3, ]-c(0.1, 0.1),
             arr.pos = 1, lty = 5, lcol = 5)
> bentarrow(from = elpos[1, ], to = elpos[3, ], lty = 5, lcol = 5)
> selfarrow(pos = elpos[3, ], path = "R", lty = 6, curve = 0.075, lcol = 6)
> splitarrow(from = elpos[1, ], to = elpos[2:3, ], lty = 1,
             lwd = 1, dd = 0.7, arr.side = 1:2, lcol = 7)
> for ( i in 1:4)
    textrect (elpos[i, ], 0.05, 0.05, lab = i, cex = 1.5)
> legend("topright", lty = 1:7, legend = c("segmentarrow",
         "curvedarrow", "straightarrow", "treearrow", "bentarrow",
```

# 5. functions to draw electrical networks

"selfarrow", "splitarrow"), lwd = c(rep(2, 6), 1), col = 1:7)

Since version 1.6, it is possible to use diagram to draw electrical networks. Below I give an example of a small transistor circuit.

```
> layoutmat <- matrix(data = c(rep(1, 12), 2, 3, 4, 5),
+ nrow = 4, ncol = 4, byrow = TRUE)
```

# textbox shapes

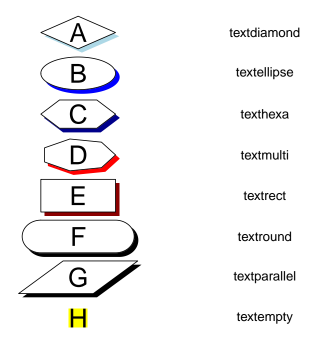


Figure 8: The text boxes

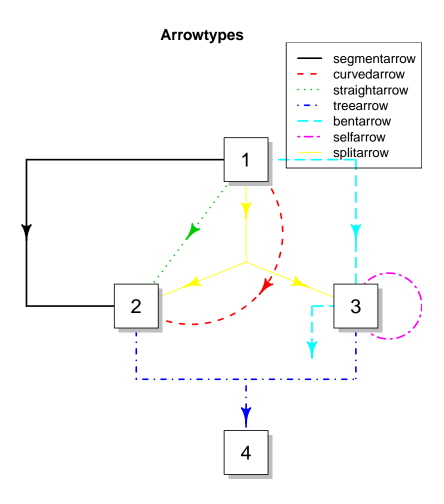


Figure 9: The arrow types

```
> nf <- layout(layoutmat, respect = FALSE)
> par(lwd = 1.5)
> par(mar = c(0, 0, 2, 0))
> emptyplot(main = "transistor Amplifier", asp = FALSE,
           vlim = c(-0.1, 1), xlim = c(-0.1, 1.1)
> x1 <- 0; x2 <- 0.2; x3 <- 0.4; x4 <- 0.6; x5 <- 0.8; x6 <- 1
> y1 <- 0.05; y2 <- 0.4; y3 <- 0.5; y4 <- 0.6; y5 <- 0.95
> x23 <- (x2 + x3)/2
> x56 < - (x5 + x6)/2
> lines(c(x2, x6, x6, x2, x2, x1, x1, x23, x3, x3),
        c(y1, y1, y5, y5, y1, y1, y3, y3, y4, y5))
> lines(c(x23, x3, x3),
                                  c(y3, y2, y1))
> lines(c(x3, x4, x4),
                                  c(y2, y2, y1))
> lines(c(x3, x5, x5),
                                   c(y4, y4, y1))
> en.Amplifier(c(x23, y3), r = 0.035)
> en.Signal(c(x1, 0.2), lab = expression("U"["in"]))
> en.Signal(c(x6, y2), lab = expression("U"["b"]))
> straightarrow(c(x1 - 0.05, 0.23), c(x1 - 0.05, 0.17),
                arr.pos =1, arr.type = "triangle", lwd = 1)
> straightarrow(c(x6 + 0.05, y2 + 0.03), c(x6 + 0.05, y2 - 0.03),
                arr.pos = 1, arr.type = "triangle", lwd = 1)
> en.Node(c(x1, y3), lab = "u1")
> en.Node(c(x2, y3), lab = "u2")
> en.Node(c(x3, y2), lab = "u3", pos = 1.5)
> en.Node(c(x3, y4), lab = "u4", pos = 2.5)
> en.Node(c(x5, y4), lab = "u5")
> en.Capacitator(c(0.5*(x1 + x2),y3), lab = "C1", vert = FALSE)
> en.Capacitator(c(x4, y4), lab = "C3", vert = FALSE)
> en.Capacitator(c(x4, 0.5*(y1+y2)), lab = "C2", vert = TRUE)
> en.Resistor(c(x1, y2), lab = "R0")
> en.Resistor(c(x2, 0.5*(y1+y2)), lab = "R1")
> en.Resistor(c(x2, 0.5*(y4+y5)), lab = "R2")
> en.Resistor(c(x3, 0.5*(y4+y5)), lab = "R4")
> en.Resistor(c(x3, 0.5*(y1+y2)), lab = "R3")
> en.Resistor(c(x5, 0.5*(y1+y2)), lab = "R5")
> en.Ground(c(1.0, 0.05))
> par(mar=c(2, 2, 2, 2))
> emptyplot(main = "transistor")
> lines(c(0.1, 0.5, 0.9), c(0.5, 0.5, 0.9))
> lines(c(0.5, 0.9), c(0.5, 0.1))
> lines(c(0.5, 0.5), c(0.4, 0.6))
> text(0.2, 0.4, "Gate", font = 3)
> text(0.8, 0.9, "Drain", font = 3,adj = 1)
> text(0.8, 0.1, "Source", font = 3,adj = 1)
> en.Amplifier(c(0.5, 0.5), r = 0.15)
> box(col = "grey")
> emptyplot(main = "capacitator")
```

```
> straightarrow(c(0.5, 0.9), c(0.5, 0.1),
                arr.pos = 0.3, arr.length = 0.25, arr.type = "triangle")
> en.Capacitator(c(0.5, 0.5), width = 0.075, length = 0.5, vert = TRUE)
> text(0.4, 0.65, "i", font = 3, cex = 2)
> straightarrow(c(0.8, 0.3), c(0.8, 0.77), arr.pos = 1,
                arr.length = 0.25, arr.type = "triangle", lwd = 1)
> text(0.925, 0.65, "v", font = 3, cex = 2)
> text(0.15, 0.5, "C", font = 3, cex = 2)
> box(col = "grey")
> emptyplot(main = "resistor")
> straightarrow(c(0.5, 0.9), c(0.5, 0.1), arr.pos = 0.2,
                arr.length = 0.25, arr.type = "triangle", lwd = 1)
> text(0.4, 0.85, "i", font = 3, cex = 2)
> en.Resistor(c(0.5, 0.5), width = 0.25, length = 0.35)
> straightarrow(c(0.8, 0.3), c(0.8, 0.77), arr.pos = 1,
                arr.length = 0.25, arr.type = "triangle", lwd = 1)
> text(0.925, 0.65, "v", font = 3, cex = 2)
> text(0.5, 0.5, "R", font = 3, cex = 2)
> box(col = "grey")
> emptyplot(main = "voltage source")
> lines(c(0.5, 0.5), c(0.1, 0.9))
> en.Signal(c(0.5, 0.5), r = 0.15)
> straightarrow(c(0.8, 0.3), c(0.8, 0.77), arr.pos = 1,
                arr.length = 0.25, arr.type = "triangle", lwd = 1)
> text(0.925, 0.65, "v", font = 3, cex = 2)
> box(col = "grey")
```

This vignette was created using Sweave (Leisch 2002).

The package is on CRAN, the R-archive website ((R Development Core Team 2008))

More examples can be found in the demo's of package ecolMod (Soetaert and Herman 2009a)

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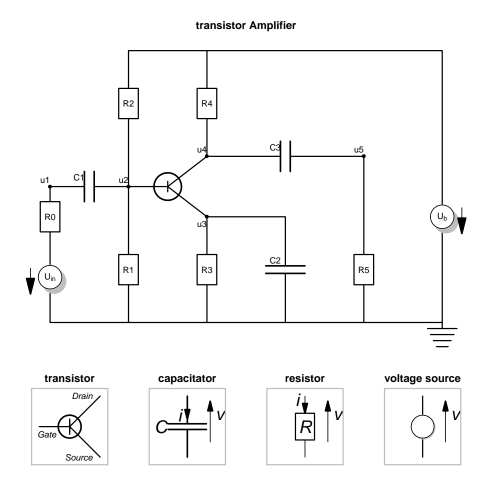


Figure 10: Drawing an electrical network with package diagram  $\,$ 

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