Package Vignette for DirichletReg: JSS Code

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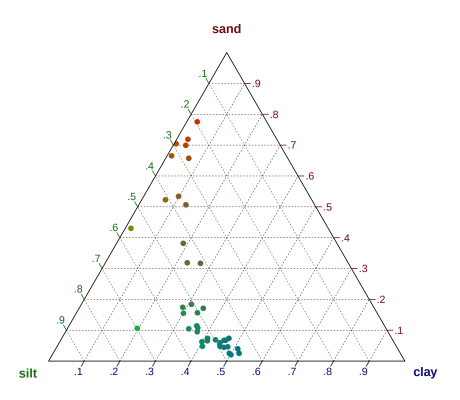
Abstract

This package vignette contains the full code from the JSS article. This document was generated using R 2.13.2 (R Development Core Team 2011) and DirichletReg 0.3.0.

Keywords: DirichletReg package, Dirichlet regression.

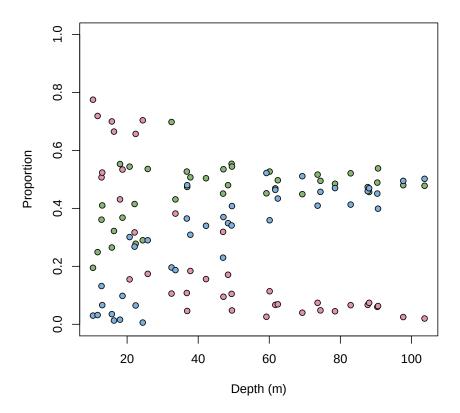
```
> library(DirichletReg)
> head(ArcticLake)
  sand silt clay depth
1 0.775 0.195 0.030 10.4
2 0.719 0.249 0.032 11.7
3 0.507 0.361 0.132 12.8
4 0.522 0.409 0.066 13.0
5 0.700 0.265 0.035 15.7
6 0.665 0.322 0.013 16.3
> AL <- DR_data(ArcticLake[, 1:3])</pre>
> AL
This object contains compositional data with 3 dimensions.
Number of observations: 39
* The data were normalized.
To access the data, use the function getdata().
> head(getdata(AL), width = 15, height = 10)
       sand
                 silt
                           clay
1 0.7750000 0.1950000 0.0300000
2 0.7190000 0.2490000 0.0320000
3 0.5070000 0.3610000 0.1320000
4 0.5235707 0.4102307 0.0661986
5 0.7000000 0.2650000 0.0350000
6 0.6650000 0.3220000 0.0130000
```

> plot(AL, reset_par = FALSE)



```
> plot(rep(ArcticLake$depth, 3), unlist(getdata(AL)), pch = 21,
+ bg = rep(rainbow_hcl(3), each = 39), xlab = "Depth (m)",
+ ylab = "Proportion", ylim = 0:1)
```

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```
> lake1 <- DirichReg(AL ~ depth, ArcticLake)
> summary(lake1)
```

Call:

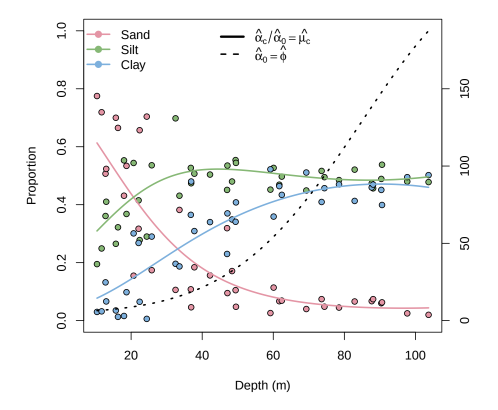
DirichReg(formula = AL ~ depth, data = ArcticLake)

Standardized Residuals:

```
Min 1Q Median 3Q Max sand -1.7724 -0.8319 0.0120 1.3435 2.2862 silt -1.0863 -0.5343 -0.1279 0.2892 1.4493 clay -2.0868 -0.7516 -0.0012 0.4391 1.9660
```

```
Beta-Coefficients for variable no. 3: clay
           Estimate Std. Error z-Value p-Value
depth 0.064302 0.005738 11.21 < 2e-16 ***
       ______
Signif. codes: `***' < .001, `**' < 0.01, `*' < 0.05, `.' < 0.1
Log-likelihood: 101.4 on 6 df (102+2 iterations)
Link: Log
Parametrization: common
> lake2 <- DirichReg(AL ~ depth + I(depth^2), ArcticLake)</pre>
> anova(lake1, lake2)
Analysis of Deviance Table
Model 1:
DirichReg(formula = AL ~ depth, data = ArcticLake)
DirichReg(formula = AL ~ depth + I(depth^2), data = ArcticLake)
         Deviance N. par Difference df p-value
Model 1 -202.7393 6
Model 2 -217.9937 9 15.25441 3 0.001611655
> lake2
Call:
DirichReg(formula = AL ~ depth + I(depth^2), data = ArcticLake)
using the common parametrization
Log-likelihood: 109 on 9 df (99+2 iterations)
Coefficients for variable no. 1: sand
(Intercept) depth I(depth^2)
 1.4361967 -0.0072383 0.0001324
Coefficients for variable no. 2: silt
(Intercept) depth I(depth^2)
 -0.0259705 0.0717450 -0.0002679
Coefficients for variable no. 3: clay
(Intercept) depth I(depth^2)
-1.7931487 0.1107906 -0.0004872
> plot(rep(ArcticLake$depth, 3), unlist(getdata(AL)), pch = 21,
     bg = rep(rainbow_hcl(3), each = 39), xlab = "Depth (m)",
     ylab = "Proportion", ylim = 0:1, main = "Sediment Composition in an Arctic Lake")
> Xnew <- data.frame(depth = seq(min(ArcticLake$depth), max(ArcticLake$depth),
    length.out = 100))
```

Sediment Composition in an Arctic Lake



```
> AL <- DR_data(ArcticLake[, 1:3])
> dd <- range(ArcticLake$depth)
> X <- data.frame(depth = seq(dd[1], dd[2], length.out = 200))
> pp <- predict(DirichReg(AL ~ depth + I(depth^2), ArcticLake),
+ X)
> plot(AL, cex = 0.1, reset_par = FALSE)
> points(DirichletReg:::coord.trafo(AL$Y[, c(2, 3, 1)]), pch = 16,
+ cex = 0.5, col = gray(0.5))
> lines(DirichletReg:::coord.trafo(pp[, c(2, 3, 1)]), lwd = 3,
+ col = rainbow_hcl(2, l = 25)[2])
> Dols <- log(cbind(ArcticLake[, 2]/ArcticLake[, 1], ArcticLake[,</pre>
```

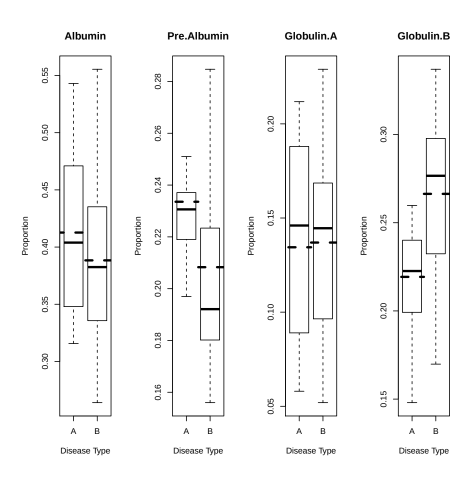
```
+ 3]/ArcticLake[, 1]))
> ols <- lm(Dols ~ depth + I(depth^2), ArcticLake)
> p2 <- predict(ols, X)
> p2m <- exp(cbind(0, p2[, 1], p2[, 2]))/rowSums(exp(cbind(0, p2[, + 1], p2[, 2])))
> lines(DirichletReg:::coord.trafo(p2m[, c(2, 3, 1)]), lwd = 3,  
+ col = rainbow_hcl(2, 1 = 25)[1], lty = "21")
```



```
> B <- DR_data(BloodSamples[1:30, 1:4])</pre>
> blood1 <- DirichReg(B ~ Disease | phi ~ 1, BloodSamples)</pre>
> blood2 <- DirichReg(B ~ Disease | phi ~ Disease, BloodSamples)</pre>
> anova(blood1, blood2)
Analysis of Deviance Table
Model 1:
DirichReg(formula = B ~ Disease | phi ~ 1, data = BloodSamples)
DirichReg(formula = B ~ Disease | phi ~ Disease, data = BloodSamples)
           Deviance
                       N. par
                                Difference
                                              df
                                                     p-value
          -303.8560
Model 1
Model 2
          -304.6147
                            8
                                 0.7586655
                                               1
                                                   0.3837465
> summary(blood1)
```

Call:

```
DirichReg(formula = B ~ Disease | phi ~ 1, data = BloodSamples)
Standardized Residuals:
            Min 1Q Median 3Q
Albumin -2.1310 -0.9307 -0.1234 0.8149 2.8429
Pre.Albumin -1.0687 -0.4054 -0.0789 0.1947 1.5691
Globulin.A -2.0503 -1.0392 0.1938 0.7927 2.2393
Globulin.B -1.8176 -0.5347 0.1488 0.5115 1.3284
MEAN MODELS:
______
Coefficients for variable no. 1: Albumin
- variable omitted (reference category) -
Coefficients for variable no. 2: Pre.Albumin
         Estimate Std. Error z-Value p-Value
DiseaseB -0.05761 0.11575 -0.498 0.619
______
Coefficients for variable no. 3: Globulin.A
        Estimate Std. Error z-Value p-Value
(Intercept) -1.11639 0.09935 -11.237 <2e-16 ***
DiseaseB 0.07002 0.13604 0.515 0.607
Coefficients for variable no. 4: Globulin.B
        Estimate Std. Error z-Value p-Value
DiseaseB 0.25192 0.11300 2.229 0.0258 *
______
PRECISION MODEL:
     Estimate Std. Error z-Value p-Value
(Intercept) 4.2227 0.1475 28.64 <2e-16 ***
Signif. codes: `***' < .001, `**' < 0.01, `*' < 0.05, `.' < 0.1
Log-likelihood: 151.9 on 7 df (43+2 iterations)
Links: Logit (Means) and Log (Precision)
Parametrization: alternative
> par(mfrow = c(1, 4))
> for (i in 1:4) {
    boxplot(B$Y[, i] ~ BloodSamples$Disease[1:30], main = paste(names(BloodSamples)[i]),
       xlab = "Disease Type", ylab = "Proportion")
    segments(c(-5, 1.5), unique(fitted(blood2)[, i]), c(1.5, 1.5)
       5), unique(fitted(blood2)[, i]), 1 \text{wd} = 3, 1 \text{ty} = 2)
+ }
```



```
> alpha <- predict(blood2, data.frame(Disease = factor(c("A", "B"))),</pre>
\verb|> L <- sapply(1:2, function(i) ddirichlet(DR_data(BloodSamples[31:36,
      1:4])$Y, unlist(alpha[i, ])))
> LP <- L/rowSums(L)
> dimnames(LP) \leftarrow list(paste("C", 1:6), c("A", "B"))
> print(round(LP * 100, 1), print.gap = 2)
              В
        Α
     59.4
           40.6
C 1
C 2
     43.2
          56.8
С 3
     38.4
           61.6
C 4
     43.8
           56.2
C 5
     36.6
           63.4
     70.2
           29.8
> B2 <- DR_data(BloodSamples[, c(1, 2, 4)])
> plot(B2, cex = 0.001, reset_par = FALSE)
> div.col <- diverge_hcl(100)</pre>
> temp <- (alpha/rowSums(alpha))[, c(2, 4, 1)]
> points(DirichletReg:::coord.trafo(temp/rowSums(temp)), pch = 22,
      bg = div.col[c(1, 100)], cex = 1, lwd = 0.25)
> temp <- B2$Y[1:30, c(2, 3, 1)]
> points(DirichletReg:::coord.trafo(temp/rowSums(temp)), pch = 21,
```

```
+ bg = (div.col[c(1, 100)])[BloodSamples$Disease[1:30]], cex = 0.5,

+ lwd = 0.25)

> temp < -B2\$Y[31:36, c(2, 3, 1)]

> points(DirichletReg:::coord.trafo(temp/rowSums(temp)), pch = 21,

+ bg = div.col[round(100 * LP[, 2], 0)], cex = 0.5, lwd = 0.5)

> legend("topleft", bty = "n", legend = c("Disease A", "Disease B",

+ NA, "Expected Values"), pch = c(21, 21, NA, 22), pt.bg = c(div.col[c(1, 100)], NA, "white"))
```

- Disease A
- Disease B


```
Deviance N. par
                            Difference
                                         df
                                               p-value
         -133.4682
Model 1
                        8
Model 2
        -131.8037
                         7
                              1.664453
                                             0.1970031
> a <- ReadingSkills$accuracy</pre>
> logit_a <- log(a/(1 - a))
> rlr <- lm(logit_a ~ dyslexia * iq, ReadingSkills)</pre>
> summary(rlr)
Call:
lm(formula = logit_a ~ dyslexia * iq, data = ReadingSkills)
Residuals:
    Min
             1Q Median
-2.66405 -0.37966 0.03687 0.40887 2.50345
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept)
              2.8067 0.2822 9.944 2.27e-12 ***
                         0.4517 -5.338 4.01e-06 ***
              -2.4113
dyslexiayes
iq
              0.7823
                         0.2992 2.615 0.0125 *
dyslexiayes:iq -0.8457
                         0.4510 -1.875 0.0681 .
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.2 on 40 degrees of freedom
Multiple R-squared: 0.6151,
                          Adjusted R-squared: 0.5862
F-statistic: 21.31 on 3 and 40 DF, p-value: 2.083e-08
> summary(rs2)
Call:
DirichReg(formula = acc ~ dyslexia * iq | phi ~ dyslexia + iq,
data = ReadingSkills)
Standardized Residuals:
            Min 1Q Median
                                      3Q
1 - data -1.5661 -0.8204 -0.5112 0.5211 3.4334
         -3.4334 -0.5211 0.5112 0.8204 1.5661
data
MEAN MODELS:
Coefficients for variable no. 1: 1 - data
- variable omitted (reference category) -
Coefficients for variable no. 2: data
             Estimate Std. Error z-Value p-Value
(Intercept)
              1.8649 0.2991 6.235 4.52e-10 ***
              -1.4833
                         0.3029 -4.897 9.74e-07 ***
dyslexiayes
iq
```

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```
Estimate Std. Error z-Value p-Value
(Intercept) 1.5579 0.3336 4.670 3.01e-06 ***
dyslexiayes 3.4931 0.5880 5.941 2.83e-09 ***
iq 1.2291 0.4596 2.674 0.00749 **
```

Signif. codes: `***' < .001, `**' < 0.01, `*' < 0.05, `.' < 0.1

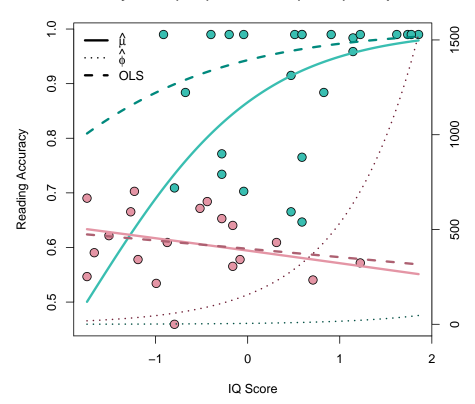
Log-likelihood: 65.9 on 7 df (39+1 iterations) Links: Logit (Means) and Log (Precision)

Parametrization: alternative

PRECISION MODEL:

```
> g.ind <- as.numeric(ReadingSkills$dyslexia)</pre>
> plot(accuracy ~ iq, ReadingSkills, pch = 21, bg = rainbow_hcl(2)[3 -
      g.ind], cex = 1.5, main = "Dyslexic (Red) vs. Control (Green) Group",
      xlab = "IQ Score", ylab = "Reading Accuracy")
> x <- seq(min(ReadingSkills$iq), max(ReadingSkills$iq), length.out = 200)
> n \leftarrow length(x)
> X <- data.frame(dyslexia = rep(c("yes", "no"), each = n), iq = c(x,
> pv <- predict(rs2, X, TRUE, TRUE, TRUE)
> lines(x, pv$mu[-(1:n), 2], col = rainbow_hcl(2)[2], lwd = 3)
> lines(x, pv$mu[1:n, 2], col = rainbow_hcl(2)[1], lwd = 3)
> olsN <- 1/(1 + exp(-predict(rlr, X[-(1:n), ])))
> olsD <- 1/(1 + exp(-predict(rlr, X[1:n, ])))</pre>
> lines(x, olsD, col = rainbow_hcl(2, 1 = 50)[1], lwd = 3, lty = 2)
> lines(x, olsN, col = rainbow_hcl(2, 1 = 50)[2], lwd = 3, lty = 2)
> par(new = TRUE)
> plot(x, pv$phi[-(1:n)], col = rainbow_hcl(2, 1 = 25)[2], lty = 3,
      type = "1", ylim = c(0, \max(pv\$phi)), axes = F, ann = F,
> lines(x, pv$phi[1:n], col = rainbow_hcl(2, 1 = 25)[1], lty = 3,
      type = "1", lwd = 2)
+
> axis(4)
> legend("topleft", legend = c(expression(hat(mu)), expression(hat(phi)),
      "OLS"), 1 \text{ty} = c(1, 3, 2), 1 \text{wd} = c(3, 2, 3), b \text{ty} = "n"
```

Dyslexic (Red) vs. Control (Green) Group



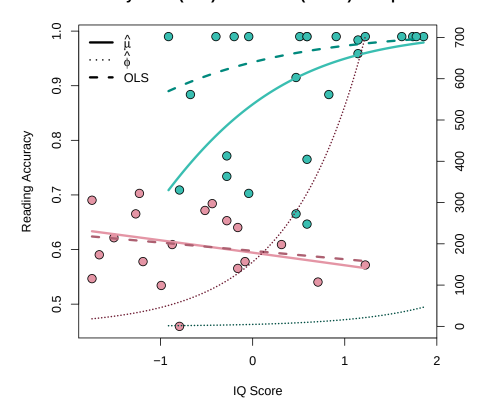
```
> gcol <- rainbow_hcl(2)[3 - as.numeric(ReadingSkills$dyslexia)]</pre>
> tmt <- c(-3, 3)
> par(mfrow = c(3, 2))
 qqnorm(residuals(rlr, "pearson"), ylim = tmt, xlim = tmt, pch = 21,
      bg = gcol, main = "Normal QQPlot: OLS Residuals", cex = 0.75,
      1wd = 0.5)
> abline(0, 1, lwd = 2)
> qqline(residuals(rlr, "pearson"), lty = 2)
 qqnorm(residuals(rs2, "standardized")[, 2], ylim = tmt, xlim = tmt,
      pch = 21, bg = gcol, main = "Normal QQPlot: DirichReg Residuals",
      cex = 0.75, 1wd = 0.5)
> abline(0, 1, lwd = 2)
> qqline(residuals(rs2, "standardized")[, 2], 1ty = 2)
> plot(ReadingSkills$iq, residuals(rlr, "pearson"), pch = 21, bg = gcol,
      ylim = c(-3, 3), main = "OLS Residuals", xlab = "IQ", ylab = "Pearson Residuals",
      cex = 0.75, 1wd = 0.5)
> abline(h = 0, lty = 2)
> lines(smooth.spline(ReadingSkills$iq, residuals(rlr, "pearson")))
 plot(ReadingSkills$iq, residuals(rs2, "standardized")[, 2], pch = 21,
      bg = gcol, ylim = c(-3, 3), main = "DirichReg Residuals",
      xlab = "IQ", ylab = "Standardized Residuals", cex = 0.75,
      1wd = 0.5)
> abline(h = 0, lty = 2)
> lines(smooth.spline(ReadingSkills$iq, residuals(rs2, "standardized")[,
```

```
2]))
  plot(fitted(rlr), residuals(rlr, "pearson"), pch = 21, bg = gcol,
       ylim = c(-3, 3), main = "OLS Residuals", xlab = "Fitted",
       ylab = "Pearson Residuals", cex = 0.75, lwd = 0.5)
> abline(h = 0, lty = 2)
> lines(smooth.spline(fitted(rlr), residuals(rlr, "pearson")))
  plot(fitted(rs2)[, 2], residuals(rs2, "standardized")[, 2], pch = 21,
       bg = gcol, ylim = c(-3, 3), main = "DirichReg Residuals",
       xlab = "Fitted", ylab = "Standardized Residuals", cex = 0.75,
       1wd = 0.5)
> abline(h = 0, lty = 2)
> lines(smooth.spline(fitted(rs2)[, 2], residuals(rs2, "standardized")[,
       Normal Q-Q-Plot: OLS Residuals
                                           Normal Q-Q-Plot: DirichReg Residuals
Sample Quantiles
                                      Sample Quantiles
   ī
                    0
                                             -3
                                                 -2
       -3
              Theoretical Quantiles
                                                    Theoretical Quantiles
              OLS Residuals
                                                  DirichReg Residuals
                                      Standardized Residuals
Pearson Residuals
   6
                   0
                                                          0
            -1
                   IQ
                                                          IQ
              OLS Residuals
                                                  DirichReg Residuals
                                      Standardized Residuals
Pearson Residuals
   T
   က
                                               0.6
                                                     0.7
                  2
                                                           0.8
                                                                 0.9
                   Fitted
                                                         Fitted
> g.ind <- as.numeric(ReadingSkills$dyslexia)</pre>
> g1 <- g.ind == 1
> g2 <- g.ind != 1
> plot(accuracy ~ iq, ReadingSkills, pch = 21, bg = rainbow_hcl(2)[3 -
       g.ind], cex = 1.5, main = "Dyslexic (Red) vs. Control (Green) Group",
       xlab = "IQ Score", ylab = "Reading Accuracy", xlim = range(ReadingSkills$iq))
> x1 <- seq(min(ReadingSkills$iq[g1]), max(ReadingSkills$iq[g1]),
       length.out = 200)
> x2 <- seq(min(ReadingSkills$iq[g2]), max(ReadingSkills$iq[g2]),
```

length.out = 200)

```
> n \leftarrow length(x1)
> X1 <- data.frame(dyslexia = factor(rep(0, n), levels = 0:1, labels = c("no",
      "yes")), iq = x1)
> X2 <- data.frame(dyslexia = factor(rep(1, n), levels = 0:1, labels = c("no",
      "yes")), iq = x2)
> pv1 <- predict(rs2, X1, TRUE, TRUE, TRUE)
> pv2 <- predict(rs2, X2, TRUE, TRUE, TRUE)
> lines(x1, pv1$mu[, 2], col = rainbow_hcl(2)[2], lwd = 3)
> lines(x2, pv2$mu[, 2], col = rainbow_hcl(2)[1], lwd = 3)
> olsN <- 1/(1 + exp(-predict(rlr, X1)))</pre>
> olsD <- 1/(1 + exp(-predict(rlr, X2)))
> lines(x2, olsD, col = rainbow_hcl(2, l = 50)[1], lwd = 3, lty = 2)
> lines(x1, olsN, col = rainbow_hcl(2, 1 = 50)[2], lwd = 3, lty = 2)
> par(new = TRUE)
> plot(x1, pv1$phi, col = rainbow_hcl(2, 1 = 25)[2], lty = "11",
      type = "1", ylim = c(0, max(pv2\$phi)), axes = F, ann = F,
      lwd = 2, xlim = range(ReadingSkills$iq))
> lines(x2, pv2$phi, col = rainbow_hcl(2, 1 = 25)[1], lty = "11",
      type = "1", 1wd = 2)
> axis(4)
> legend("topleft", legend = c(expression(hat(mu)), expression(hat(phi)),
      "OLS"), 1 \text{ty} = c(1, 3, 2), 1 \text{wd} = c(3, 2, 3), 6 \text{ty} = "n"
```

Dyslexic (Red) vs. Control (Green) Group



References

R Development Core Team (2011). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL http://www.R-project.org/.