

CORRELATED BINARY DATA

ANIKO SZABO

ABSTRACT. We define a class for describing data from toxicology experiments and implement fitting of a variety of existing models and trend tests.

"../R/CorrBin-package.R" 1≡

```
#'Nonparameterics for Correlated Binary and Multinomial Data
#
#This package implements nonparametric methods for analysing exchangeable
#binary and multinomial data with variable cluster sizes with emphasis on trend testing. The
#input should specify the treatment group, cluster-size, and the number of
#responses (i.e. the number of cluster elements with the outcome of interest)
#for each cluster.
#
#'\tabular{ll}{
#' Package: \tab CorrBin\cr
#' Type: \tab Package\cr
#' Version: \tab 1.5\cr
#' Date: \tab 2014-12-18\cr
#' License: \tab GPL 2\cr
#' LazyLoad: \tab yes\cr
#'}
#'\itemize{ \item The \code{\link{CBData}} and \code{\link{read.CBData}}
#functions create a 'CBData' object used by the analysis functions.
#'\item \code{\link{ran.CBData}} can be used to generate random data with
#prespecified mean response and within-cluster correlation.
#'\item \code{\link{mc.test.chisq}} tests the assumption of marginal compatibility
#underlying all the methods, while \code{\link{mc.est}} estimates the
#distribution of the number of responses under marginal compatibility.
#'\item Finally, \code{\link{trend.test}} performs three different tests for trend
#along the treatment groups. }
#
#'\@name CorrBin-package
#'\@aliases CorrBin-package CorrBin
#'\@docType package
#'\@author Aniko Szabo
#
#'\@maintainer Aniko Szabo <aszabo@mcw.edu>
#'\@references Szabo A, George EO. (2009) On the Use of Stochastic Ordering to
#Test for Trend with Clustered Binary Data. \emph{Biometrika}
#
#'\@Stefanescu, C. & Turnbull, B. W. (2003) Likelihood inference for exchangeable
#binary data with varying cluster sizes. \emph{Biometrics}, 59, 18-24
#
#'\@Pang, Z. & Kuk, A. (2007) Test of marginal compatibility and smoothing
#methods for exchangeable binary data with unequal cluster sizes.
#\emph{Biometrics}, 63, 218-227
#'\@keywords package nonparametric
```



```

if (!is.data.frame(x)) stop("x has to be a data frame")
nms <- names(x)
process.var <- function(var){
  if (is.character(var)){
    if (var %in% nms) res <- x[[var]]
    else stop(paste("Variable '", var, "' not found"))
  }
  else {
    if (is.numeric(var)){
      if (var %in% seq(along=nms)) res <- x[[var]]
      else stop(paste("Column", var, " not found"))
    }
    else stop(paste("Invalid variable specification:",var))
  }
}
trtvar <- factor(process.var(trt))
csvar <- process.var(clustersize)
nrespvar <- process.var(nresp)
if (is.null(freq)) freqvar <- rep(1, nrow(x))
else freqvar <- process.var(freq)

d <- data.frame(Trt=trtvar, ClusterSize=csvar, NResp=nrespvar, Freq=freqvar)
d <- aggregate(d$Freq, list(Trt=d$Trt, ClusterSize=d$ClusterSize, NResp=d$NResp),sum)
names(d)[4] <- "Freq"
d$ClusterSize <- as.numeric(as.character(d$ClusterSize))
d$NResp <- as.numeric(as.character(d$NResp))
class(d) <- c("CBData", "data.frame")
d}
◇

```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

Defines: `CBdata.data.frame` Never used.

The `read.CBData` function reads in clustered binary data from a tab-delimited text file. The first column should give the treatment group, the second the size of the cluster, the third the number of responses in the cluster. Optionally, a fourth column could give the number of times the given combination occurs in the data.

```
"../R/CBData.R" 4a≡
```

```
#'Read data from external file into a CBData object
#
#'#A convenience function to read data from specially structured file directly
#'#into a \code{CBData} object.
#
#'#@export
#'#@param file name of file with data. The first column should contain the
#'#treatment group, the second the size of the cluster, the third the number of
#'#responses in the cluster. Optionally, a fourth column could give the number
#'#of times the given combination occurs in the data.
#'#@param with.freq logical indicator of whether a frequency variable is present
#'#in the file
#'#@param ... additional arguments passed to \code{\link[utils]{read.table}}
#'#@return a \code{CBData} object
#'#@author Aniko Szabo
#'#@seealso \code{\link{CBData}}
#'#@keywords IO file
#
◇
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

```
"../R/CBData.R" 4b≡
```

```
read.CBData <- function(file, with.freq=TRUE, ...){
  d <- read.table(file, col.names=c("Trt","ClusterSize","NResp", if (with.freq) "Freq"), ...)
  if (!with.freq) d$Freq <- 1
  d <- aggregate(d$Freq, list(Trt=d$Trt, ClusterSize=d$ClusterSize, NResp=d$NResp),sum)
  names(d)[4] <- "Freq"
  d$ClusterSize <- as.numeric(as.character(d$ClusterSize))
  d$NResp <- as.numeric(as.character(d$NResp))
  d <- CBData(d, "Trt", "ClusterSize", "NResp", "Freq")
  d}
◇
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

Defines: `read.CBdata` Never used.

The `[.CMDData]` function defines subsetting of `CMDData` objects. If the subsetting is only affecting the rows, then the appropriate attributes are preserved, and the unused levels of `Trt` are dropped. Otherwise the returned object does not have a `CMDData` class anymore.

```
"../R/CBData.R" 5a≡
```

```
#'Extract from a CBData/CMDData object
#
#The extracting syntax works as for \code{\link{[.data.frame]}}, and in general the returned object is no
#However if the columns are not modified, then the result is still a \code{CBData}/\code{CMDData} object
# and the unused levels of treatment groups dropped.
#
# @param x \code{CMDData} object.
# @param i numeric, row index of extracted values
# @param j numeric, column index of extracted values
# @param drop logical. If TRUE the result is coerced to the lowest possible dimension.
# The default is the same as for \code{\link{[.data.frame]}}: to drop if only one column is left, but not
# @return a \code{CBData}/\code{CMDData} object
# @author Aniko Szabo
# @seealso \code{CBData}/, \code{\link{CMDData}}
# @keywords manip
# @name Extract
#
NULL
```

◇
File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

```
"../R/CBData.R" 5b≡
```

```
# @rdname Extract
# @export

"[.CBData" <- function(x, i, j, drop){
  res <- NextMethod("[")
  if (NCOL(res) == ncol(x)){
    res <- "[.data.frame"(x, i, )
    res$Trt <- droplevels(res$Trt)
    res
  }
  else {
    class(res) <- setdiff(class(res), "CBData")
  }
  res
}
◇
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
Defines: `[.CBData` Never used.

`unwrap.CBData` is a utility function that reformats a `CBData` object so that each row is one observation (instead of one cluster). A new ‘ID’ variable is added to indicate clusters. It is first defined as a generic function to allow generalization.

```
"../R/CBData.R" 5c≡
```

```
#'Unwrap a clustered object
#
#\code{unwrap.CBData} is a utility function that reformats a CBData object so
#that each row is one observation (instead of one or more clusters). A new
#‘ID’ variable is added to indicate clusters. This form can be useful for
#setting up the data for a different package.
#
```

```

#'@aliases unwrap unwrap.CBData
#'@export
#'@param object a \link{CBData} object
#'@param \dots other potential arguments; not currently used
#'@return For \code{unwrap.CBData}: a data frame with one row for each cluster element (having a binary
#'outcome) with the following standardized column names
#'@return \item{Trt}{factor, the treatment group}
#'@return \item{ClusterSize}{numeric, the cluster size}
#'@return \item{ID}{factor, each level representing a different cluster}
#'@return \item{Resp}{numeric with 0/1 values, giving the response of the cluster
#'element}
#'author Aniko Szabo
#'keywords manip
#'examples
#'
#'data(shelltox)
#'ush <- unwrap(shelltox)
#'head(ush)
#'
◇

```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

Uses: `unwrap.CBData` [6a](#).

"../R/CBData.R" 6a≡

```

unwrap <- function(object,...) UseMethod("unwrap")

#'@rdname unwrap
#'@method unwrap CBData
unwrap.CBData <- function(object,...){
  freqs <- rep(1:nrow(object), object$Freq)
  cb1 <- object[freqs,]
  cb1$Freq <- NULL
  cb1$ID <- factor(1:nrow(cb1))
  pos.idx <- rep(1:nrow(cb1), cb1$NResp)
  cb.pos <- cb1[pos.idx,]
  cb.pos$Resp <- 1
  cb.pos$NResp <- NULL
  neg.idx <- rep(1:nrow(cb1), cb1$ClusterSize-cb1$NResp)
  cb.neg <- cb1[neg.idx,]
  cb.neg$Resp <- 0
  cb.neg$NResp <- NULL
  res <- rbind(cb.pos, cb.neg)
  res[order(res$ID),]
}
◇

```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

Defines: `unwrap.CBData` [5c](#), [8b](#).

2. RAO-SCOTT ADJUSTED COCHRAN-ARMITAGE TEST

The RS-adjusted CA test for trend is based on design-effect adjustment.

"../R/CBData.R" 6b≡

```

#'Rao-Scott trend test

```

```
#'
#' \code{RS.trend.test} implements the Rao-Scott adjusted Cochran-Armitage test
#' for linear increasing trend with correlated data.
#'
#' The test is based on calculating a \dfn{design effect} for each cluster by
#' dividing the observed variability by the one expected under independence. The
#' number of responses and the cluster size are then divided by the design
#' effect, and a Cochran-Armitage type test statistic is computed based on these
#' adjusted values.
#'
#' The implementation aims for testing for \emph{increasing} trend, and a
#' one-sided p-value is reported. The test statistic is asymptotically normally
#' distributed, and a two-sided p-value can be easily computed if needed.
#'
#' @export
#' @param cbdata a \code{\link{CBData}} object
#' @return A list with components
#' @return \item{statistic}{numeric, the value of the test statistic}
#' @return \item{p.val}{numeric, asymptotic one-sided p-value of the test}
#' @author Aniko Szabo, aszabo@mcw.edu
#' @seealso \code{\link{SO.trend.test}}, \code{\link{GEE.trend.test}} for
#' alternative tests; \code{\link{CBData}} for constructing a CBData object.
#' @references Rao, J. N. K. & Scott, A. J. A (1992) Simple Method for the
#' Analysis of Clustered Data \emph{Biometrics}, 48, 577-586.
#' @keywords htest nonparametric
#' @examples
#'
#' data(shelltox)
#' RS.trend.test(shelltox)
#'
#'
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

Uses: [GEE.trend.test](#) [8b](#), [RS.trend.test](#) [7](#).

"../R/CBData.R" 7≡

```
RS.trend.test <- function(cbdata){
  dat2 <- cbdata[rep(1:nrow(cbdata), cbdata$Freq),] #each row is one sample
  dat2$Trt <- factor(dat2$Trt) #remove unused levels
  x.i <- pmax(tapply(dat2$NResp, dat2$Trt, sum), 0.5) #"continuity" adjustment to avoid RS=NaN
  n.i <- tapply(dat2$ClusterSize, dat2$Trt, sum)
  m.i <- table(dat2$Trt)
  p.i.hat <- x.i/n.i
  r.ij <- dat2$NResp - dat2$ClusterSize*p.i.hat[dat2$Trt]
  v.i <- m.i/(m.i-1)/n.i^2*tapply(r.ij^2, dat2$Trt, sum)
  d.i <- n.i * v.i / (p.i.hat*(1-p.i.hat)) #design effect
  x.i.new <- x.i/d.i
  n.i.new <- n.i/d.i
  p.hat <- sum(x.i.new)/sum(n.i.new)

  scores <- (1:nlevels(dat2$Trt))-1
  mean.score <- sum(scores*n.i.new)/sum(n.i.new)
  var.scores <- sum(n.i.new*(scores-mean.score)^2)
  RS <- (sum(x.i.new*scores) - p.hat*sum(n.i.new*scores)) /
    sqrt(p.hat*(1-p.hat)*var.scores)
  p.val <- pnorm(RS, lower.tail=FALSE)
  list(statistic=RS, p.val=p.val)
}
```

◇
 File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Defines: `RS.trend.test` [6b](#), [8a](#).

3. GEE BASED TEST

"../R/CBData.R" [8a](#)≡

```
#'GEE-based trend test
#
#'\code{GEE.trend.test} implements a GEE based test for linear increasing trend
# for correlated binary data.
#
# 'The actual work is performed by the \code{\link[geepack]{geese}} function of
# the \code{geepack} library. This function only provides a convenient wrapper
# to obtain the results in the same format as \code{\link{RS.trend.test}} and
# \code{\link{SO.trend.test}}.
#
# 'The implementation aims for testing for \emph{increasing} trend, and a
# one-sided p-value is reported. The test statistic is asymptotically normally
# distributed, and a two-sided p-value can be easily computed if needed.
#
# @export
# @param cbdata a \code{\link{CBData}} object
# @param scale.method character string specifying the assumption about the
# change in the overdispersion (scale) parameter across the treatment groups:
# "fixed" - constant scale parameter (default); "trend" - linear trend for the
# log of the scale parameter; "all" - separate scale parameter for each group.
# @return A list with components
# @return \item{statistic}{numeric, the value of the test statistic}
# @return \item{p.val}{numeric, asymptotic one-sided p-value of the test}
# @author Aniko Szabo, aszabo@mcw.edu
# @seealso \code{\link{RS.trend.test}}, \code{\link{SO.trend.test}} for
# alternative tests; \code{\link{CBData}} for constructing a CBData object.
# @keywords htest models
# @examples
#
# data(shelltox)
# GEE.trend.test(shelltox, "trend")
#
```

◇
 File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Uses: `GEE.trend.test` [8b](#), `RS.trend.test` [7](#).

"../R/CBData.R" [8b](#)≡

```
GEE.trend.test <- function(cbdata, scale.method=c("fixed", "trend", "all")){
  require(geepack)
  ucb <- unwrap.CBData(cbdata)
  scale.method <- match.arg(scale.method)
  if (scale.method=="fixed") {
    geemod <- geese(Resp~unclass(Trt), id=ucb$ID, scale.fix=FALSE, data=ucb,
                    family=binomial, corstr="exch") }
  else if (scale.method=="trend"){
    geemod <- geese(Resp~unclass(Trt), sformula=~unclass(Trt), id=ucb$ID, data=ucb,
                    family=binomial, sca.link="log", corstr="exch")}
```



```

else if (scale.method=="all"){
  geemod <- geese(Resp~unclass(Trt), id=ucb$ID, sformula=~Trt, data=ucb,
    family=binomial, sca.link="log", corstr="exch") }
geesum <- summary(geemod)
testres <- geesum$mean[2,"estimate"]/geesum$mean[2,"san.se"]
p <- pnorm(testres, lower.tail=FALSE)
list(statistic=testres, p.val=p)
}

```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Defines: `GEE.trend.test` [6b](#), [8a](#).
 Uses: `unwrap.CBData` [6a](#).

4. GENERATING RANDOM DATA

`ran.CBData` generates a random `CBData` object from a given two-parameter distribution. `sample.sizes` is a dataset with variables `Trt`, `ClusterSize` and `Freq` giving the number of clusters to be generated for each `Trt/ClusterSize` combination. `p.gen.fun` and `rho.gen.fun` are functions that generate the parameter values for each treatment group ($g = 1$ corresponds to the lowest group, $g = 2$ to the second, etc). `pdf.fun` is a function(`p`, `rho`, `n`) generating the pdf of the number of responses given the two parameters `p` and `rho`, and the cluster size `n`.

"../R/CBData.R" 9≡

```

#'Generate random correlated binary data
#'
#'\code{ran.mc.CBData} generates a random \code{\link{CBData}} object from a
#'\code{given two-parameter distribution.
#'\code{dfn{p.gen.fun}} and \code{dfn{rho.gen.fun}} are functions that generate the
#'\code{parameter values for each treatment group; \code{dfn{pdf.fun}} is a function
#'\code{generating the pdf of the number of responses given the two parameters
#'\code{dfn{p}} and \code{dfn{rho}}, and the cluster size \code{dfn{n}}.
#'\code{p.gen.fun} and \code{rho.gen.fun} expect the parameter value of 1 to
#'\code{represent the first group, 2 - the second group, etc.
#'\code{@export
#'\code{@param sample.sizes} a dataset with variables \code{Trt}, \code{ClusterSize} and \code{Freq} giving
#'\code{the number of clusters to be generated for each Trt/ClusterSize combination.
#'\code{@param p.gen.fun} a function of one parameter that generates the value of the
#'\code{first parameter of \code{pdf.fun}} (\code{p}) given the group number.
#'\code{@param rho.gen.fun} a function of one parameter that generates the value of
#'\code{the second parameter of \code{pdf.fun}} (\code{rho}) given the group number.
#'\code{@param pdf.fun} a function of three parameters (\code{p}, \code{rho}, \code{n}) giving the
#'\code{PDF of the number of responses in a cluster given the two parameters
#'\code{(\code{p}, \code{rho})}, and the cluster size (\code{n}). Functions implementing two
#'\code{common distributions: the beta-binomial (\code{\link{betabin.pdf}})} and
#'\code{q-power (\code{\link{qpower.pdf}})} are provided in the package.
#'\code{@return} a \code{CBData} object with randomly generated number of responses with
#'\code{sample sizes specified in the call.
#'\code{@author} Aniko Szabo, aszabo@mcw.edu
#'\code{@seealso} \code{\link{betabin.pdf}} and \code{\link{qpower.pdf}}
#'\code{@keywords} distribution
#'\code{@examples
#'\code{

```

```
#' set.seed(3486)
#' ss <- expand.grid(Trt=0:3, ClusterSize=5, Freq=4)
#' #Trt is converted to a factor
#' rd <- ran.CBData(ss, p.gen.fun=function(g)0.2+0.1*g)
#' rd
#'
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Uses: `ran.CBData` [10](#).

"../R/CBData.R" 10≡

```
ran.CBData <- function(sample.sizes, p.gen.fun=function(g)0.3,
                        rho.gen.fun=function(g)0.2, pdf.fun=qpower.pdf){
  ran.gen <- function(d){
    # d is subset(sample.sizes, Trt==trt, ClusterSize==cs)
    cs <- d$ClusterSize[1]
    trt <- unclass(d$Trt)[1]
    n <- d$Freq[1]
    p <- p.gen.fun(trt)
    rho <- rho.gen.fun(trt)
    probs <- pdf.fun(p, rho, cs)
    tmp <- rmultinom(n=1, size=n, prob=probs)[,1]
    cbind(Freq=tmp, NResp=0:cs, ClusterSize=d$ClusterSize, Trt=d$Trt)}

  sst <- if (is.factor(sample.sizes$Trt)) sample.sizes$Trt else factor(sample.sizes$Trt)
  a <- by(sample.sizes, list(Trt=sst, ClusterSize=sample.sizes$ClusterSize), ran.gen)
  a <- data.frame(do.call(rbind, a))
  a$Trt <- factor(a$Trt, labels=levels(sst))
  a <- a[a$Freq>0, ]
  class(a) <- c("CBData", "data.frame")
  a
}
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Defines: `ran.CBData` [1](#), [9](#), [11a](#).

4.1. Parametric pdf generating functions. `betabin.pdf` and `qpower.pdf` provide two classic distributions – beta-binomial and q-power – for generating correlated binary data. Either can be used in `ran.CBData`.

```
"../R/CBData.R" 11a≡
```

```
#'Parametric distributions for correlated binary data
#
#'\code{qpwr.pdf} and \code{betabin.pdf} calculate the probability
#distribution function for the number of responses in a cluster of the q-power
#and beta-binomial distributions, respectively.
#
#The pdf of the q-power distribution is \deqn{P(X=x) =
#\frac{\sum_{k=0}^x (-1)^k \binom{x}{k} q^{(n-x+k)\gamma}}{\sum_{k=0}^n (-1)^k \binom{n}{k} q^{(n-k)\gamma}}, where
#C(n,x) = \sum_{k=0}^x (-1)^k \binom{n}{k} q^{(n-k)\gamma}, where \gamma =
#\frac{q^{2\gamma} - q^2}{q(1-q)}.}
#
#The pdf of the beta-binomial distribution is \deqn{P(X=x) = \binom{n}{x}
#\frac{B(\alpha+x, n+\beta-x)}{B(\alpha, \beta)} P(X=x) = C(n,x)
#B(a+x, n+b-x)/B(a,b), where \alpha =
#p \frac{1-\rho}{\rho}, and \beta =
#(1-p) \frac{1-\rho}{\rho}.
#
# @export
# @rdname pdf
# @aliases qpwr.pdf betabin.pdf
# @param p numeric, the probability of success.
# @param rho numeric between 0 and 1 inclusive, the within-cluster correlation.
# @param n integer, cluster size.
# @return a numeric vector of length \eqn{n+1} giving the value of \eqn{P(X=x)}
# for \eqn{x=0, \ldots, n}.
# @author Aniko Szabo, aszabo@mcw.edu
# @seealso \code{\link{ran.CBData}} for generating an entire dataset using
# these functions
# @references Kuk, A. A (2004) litter-based approach to risk assessment in
# developmental toxicity studies via a power family of completely monotone
# functions \emph{Applied Statistics}, 52, 51-61.
#
# Williams, D. A. (1975) The Analysis of Binary Responses from Toxicological
# Experiments Involving Reproduction and Teratogenicity \emph{Biometrics}, 31,
# 949-952.
# @keywords distribution
# @examples
#
# the distributions have quite different shapes
# with q-power assigning more weight to the "all affected" event than other distributions
# plot(0:10, betabin.pdf(0.3, 0.4, 10), type="o", ylim=c(0,0.34),
#      ylab="Density", xlab="Number of responses out of 10")
# lines(0:10, qpwr.pdf(0.3, 0.4, 10), type="o", col="red")
#
#
#
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).
 Uses: [ran.CBData](#) [10](#).

```
"../R/CBData.R" 11b≡
```

```
betabin.pdf <- function(p, rho, n){
  a <- p*(1/rho-1)
  b <- (1-p)*(1/rho-1)
  idx <- 0:n
  res <- choose(n, idx)*beta(a+idx, b+n-idx)/beta(a,b)
  res
}
```

}

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).

"../R/CBData.R" 12≡

```
#'@rdname pdf
#'@export
qpower.pdf <- function(p, rho, n){
  .q <- 1-p
  gamm <- log2(log(.q^2+rho*.q*(1-.q))/log(.q))
  res <- numeric(n+1)
  for (y in 0:n){
    idx <- 0:y
    res[y+1] <- choose(n,y) * sum( (-1)^idx * choose(y,idx) * .q^((n-y+idx)^gamm))
  }
  res <- pmax(pmin(res,1),0) #to account for numerical imprecision
  res
}
```

File defined by [2ab](#), [4ab](#), [5abc](#), [6ab](#), [7](#), [8ab](#), [9](#), [10](#), [11ab](#), [12](#).