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Single-Source Capture-Recapture Models With singleRcapture

Piotr Chlebicki

Maciej Beresewicz

Stockholm University

Poznań University of Economics and Business Statistical Office in Poznań

Abstract

Estimating population size is an important issue in official statistics, social sciences and natural sciences. One way to approach this problem is to use capture-recapture methods, which can be classified according to the number of sources used, the main distinction being between methods based on one source and those based on two or more sources. In this presentation we will introduce the **singleRcapture** R package for fitting SSCR models. The package implements state-of-the-art models as well as some new models proposed by the authors (e.g. extensions of zero-truncated one-inflated and one-inflated zero-truncated models). The software is intended for users interested in estimating the size of populations, particularly those that are difficult to reach or for which information is available from only one source and dual/multiple system estimation cannot be used.

Keywords: population size estimation, truncated distributuons, count regression models, R.

1. Introduction

1.1. Literature review

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The subject of this workshop is the **singleRcapture** package and its lightweight extension that allows for integration with other R packages called **singleRcaptureExtra**.

The package is available on CRAN: CRAN.R-project.org/package=singleRcapture while the extension is available on: https://github.com/ncn-foreigners/singleRcaptureExtra. The **singleRcapture** package is an R language package that focuses on implementing state of the art methods for frequentist point and interval estimation of size of closed populations in single-source capture-recapture (SSCR) setting (e.g. estimation of the population size of irregular migrants at set time point in a given area).

The beginning of inference in single source capture-recapture dates back to the seminal van der Heijden, Bustami, Cruyff, Engbersen, and van Houwelingen (2003) paper in which the zero truncated poisson model was applied to study the size of population of irregular migrants in fours cities in Netherlands.

1.2. How do we estimate population size with only one register? The basics of SSCR

Let Y_k represent the number of times k-th unit was observed in source data. Clearly, we don not know how often $Y_k = 0$ and to find the total population size N we need to estimate it. In general, we assume that conditional distribution of Y_k given a~vector of covariates \boldsymbol{x}_k follows some version of zero truncated count data distribution. Knowing the parameters of the distribution we may estimate the population size using Horwitz-Thompson type estimator:

$$\hat{N} = \sum_{k=1}^{N} \frac{I_k}{\mathbb{P}[Y_k > 0 | \boldsymbol{X}_k]} = \sum_{k=1}^{N_{obs}} \frac{1}{\mathbb{P}[Y_k > 0 | \boldsymbol{X}_k]},$$

where $I_k := \mathcal{I}_{\mathbb{N}}(Y_k)$, and maximum likelihood estimate of N is obtained after substituting regression estimates for $\mathbb{P}[Y_k > 0 | \boldsymbol{x}_k]$ into the equation above. Most of the methods relate to poisson processes.

The analytic variance estimation is then done by computing two parts of the decomposition due to the law of total variance:

$$\operatorname{var}[\hat{N}] = \mathbb{E}\left[\operatorname{var}\left[\hat{N}|I_1,\dots,I_n\right]\right] + \operatorname{var}\left[\mathbb{E}[\hat{N}|I_1,\dots,I_n]\right],\tag{1}$$

where the first addend is by the multivariate δ method seen to be:

$$\mathbb{E}\left[\operatorname{var}\left[\hat{N}|I_{1},\ldots,I_{n}\right]\right] = \left.\left(\frac{\partial(N|I_{1},\ldots,I_{N})}{\partial\boldsymbol{\beta}}\right)^{T}\operatorname{cov}\left[\boldsymbol{\beta}\right]\left(\frac{\partial(N|I_{1},\ldots,I_{N})}{\partial\boldsymbol{\beta}}\right)\right|_{\boldsymbol{\beta}=\hat{\boldsymbol{\beta}}},\tag{2}$$

while the later part of the decomposition in (1) is under the assumption of independence of I_k 's and after some omitted simplifications one sees that this is optimally estimated via:

$$\operatorname{var}\left(\mathbb{E}(\hat{N}|I_1,\dots,I_n)\right) = \operatorname{var}\left(\sum_{k=1}^N \frac{I_k}{\mathbb{P}(Y_k > 0)}\right)$$

$$\approx \sum_{k=1}^{N_{obs}} \frac{1 - \mathbb{P}(Y_k > 0)}{\mathbb{P}(Y_k > 0)^2},$$
(3)

which forms the basis of confidence interval creation. Confidence intervals are usually constructed under the assumption of (asymptotic) normality of \hat{N} or asymptotic normality of $\ln(\hat{N}-N)$ (or log normality of \hat{N}). The latter of which is an attempt to address a common

criticism of student type confidence intervals in SSCR, that is a possibly skewed distribution of \hat{N} , and results in the confidence interval of the form (for confidence level of α):

$$\left(N_{obs} + \frac{\hat{N} - N_{obs}}{G}, N_{obs} + \left(\hat{N} - N_{obs}\right)G\right),\,$$

where:

$$G = \exp\left(z\left(1 - \frac{\alpha}{2}\right)\sqrt{\ln\left(1 + \frac{\widehat{\text{Var}}(\hat{N})}{\left(\hat{N} - N_{obs}\right)^2}\right)}\right).$$

Existing implementations

There are some packages implementing zero truncated count data models such as **VGAM** and **countreg** and they can be integrated within the **singleRcapture** ecosystem by the lightweight extention **singleRcaptureExtra**.

2. Basic usage

2.1. The estimatePopsize function

The main function that **singleRcapture** is built around is **estimatePopsize**. The leading design principle was to make using **estimatePopsize** as close to standard **stats::glm** as possible. The most important arguments are:

- formula the main formula (i.e for the Poisson λ parameter),
- data the data.frame (or data.frame coercible) object,
- model either a function a string or a family class object specifying which model should be used possible values are listed in documentation. The supplied argument should have the form model = "ztpoisson", model = ztpoisson or model = ztpoisson(lambdaLink = "log") the third way is the only one where the user may (but doesn't have to) select a link function.
- method numerical method used to fit regression IRLS or optim,
- popVar a method for estimating variance of \hat{N} and confidence interval creation (either bootstrap, analytic or skipping the estimation entirely),
- controlMethod, controlModel, controlPopVar control parameters for numerical fitting, specifying additional formulas (inflation, dispersion) and population size estimation respectively. We will tackle these arguments separately,
- offset a matrix of offset values with number of columns matching the number of distribution parameters providing offset values to each of linear predictors.

With the formula, data, model being the three arguments which must be provided in estimatePopsize syntax.

Example with R code

The package should be installed from CRAN https://cran.r-project.org/package=singleRcapture with the usual code:

```
R> install.packages("singleRcapture")
```

To showcase the main function let us recreate the zero truncated Poisson model from van der Heijden et al. (2003) on the same data included in the package under the name netherlandsimmigrant:

```
R> library(singleRcapture)
R> head(netherlandsimmigrant)
```

	capture	gender	age		${\tt reason}$		${\tt nation}$
1	1	${\tt male}$	<40yrs	Other	reason	North	${\tt Africa}$
2	1	${\tt male}$	<40yrs	Other	reason	North	${\tt Africa}$
3	1	${\tt male}$	<40yrs	Other	reason	North	Africa
4	1	${\tt male}$	<40yrs	Other	reason		Asia
5	1	${\tt male}$	<40yrs	Other	reason		Asia
6	2	male	<40yrs	Other	reason	North	Africa

This data set contains information about immigrants in four cities (Amsterdam, Rotterdam, The Hague and Utrecht) in Netherlands that have been staying in the country illegally in 1995 and have appeared in police records that year. The number of times each individual appeared in the records is included in the capture variable with the available covariates being gender, age, reason, nation being respectively the persons gender and age, reason for being captured and region of the world from which each person comes:

R> summary(netherlandsimmigrant)

capture	gender	age	reason			
Min. :1.000	female: 398	<40yrs:1769	Illegal stay: 259			
1st Qu.:1.000	male :1482	>40yrs: 111	Other reason:1621			
Median :1.000						
Mean :1.162						
3rd Qu.:1.000						
Max. :6.000						
	nation					
American and Australia: 173						
Asia	: 284					
North Africa	:1023					
Rest of Africa	: 243					
Surinam	: 64					
Turkey	: 93					

The basic syntax is indeed vary similar to that of glm with the output of the summary method being also quite simmilar except for the additional results of the population size estimates:

```
R> basicModel <- estimatePopsize(</pre>
     formula = capture ~ gender + age + nation,
     model = ztpoisson(),
     data = netherlandsimmigrant
+ )
R> summary(basicModel)
Call:
estimatePopsize.default(formula = capture ~ gender + age + nation,
     data = netherlandsimmigrant, model = ztpoisson())
Pearson Residuals:
      Min.
              1st Qu.
                            Median
                                           Mean
                                                    3rd Qu.
-0.486442 -0.486442 -0.298080 0.002093 -0.209444 13.910844
Coefficients:
_____
For linear predictors associated with: lambda
                         Estimate Std. Error z value P(>|z|)
(Intercept)
                         -1.3411 0.2149 -6.241 4.35e-10 ***
                         gendermale
age>40yrs
nationAsia

      nationNorth Africa
      0.1900
      0.1940
      0.979
      0.327398

      nationRest of Africa
      -0.9106
      0.3008
      -3.027
      0.002468
      **

      nationSurinam
      -2.3364
      1.0136
      -2.305
      0.021159
      *

      nationTurkey
      -1.6754
      0.6028
      -2.779
      0.005445
      **

Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
AIC: 1712.901
BIC: 1757.213
Residual deviance: 1128.553
Log-likelihood: -848.4504 on 1872 Degrees of freedom
Number of iterations: 8
______
Population size estimation results:
Point estimate 12690.35
Observed proportion: 14.8% (N obs = 1880)
Std. Error 2808.169
95% CI for the population size:
            lowerBound upperBound
```

```
normal 7186.444 18194.26
logNormal 8431.275 19718.32
95% CI for the share of observed population:
lowerBound upperBound
normal 10.332927 26.16037
logNormal 9.534281 22.29793
```

One point which we should make while analysing this data set is that there is a disproportionate number of individuals who were observed only once (see table bellow):

R> table(netherlandsimmigrant\$capture)

```
1 2 3 4 5 6
1645 183 37 13 1 1
```

Since there is a reasonable suspicion that the act of observing a unit in the dataset may led to undesirable consequences from the point of view of the subject of the observation (here possible deportation, detainment or similar). For those reason one should

Warning in estimatePopsize.default(formula = capture ~ nation, model = oiztgeom(omegaLink NOTE: Second derivative test failing does not

necessarily mean that the maximum of score function that was found numericaly is invalid since $R^{\hat{}}k$ is not a bounded space.

Additionally in one inflated and hurdle models second derivative test often fails even on

Warning in estimatePopsize.default(formula = capture ~ nation, model = oiztgeom(omegaLink = "cloglog"), : Switching from observed information matrix to Fisher information matrix because hessian of log-likelihood is not negative define.

```
R> summary(modelInflated)
```

```
Call:
estimatePopsize.default(formula = capture ~ nation, data = netherlandsimmigrant,
     model = oiztgeom(omegaLink = "cloglog"), popVar = "bootstrap",
     controlModel = controlModel(omegaFormula = ~gender + age),
     controlPopVar = controlPopVar(bootType = "semiparametric"))
Pearson Residuals:
     Min. 1st Qu. Median
                                   Mean 3rd Qu.
-0.41643 -0.41643 -0.30127 0.00314 -0.18323 13.88376
Coefficients:
______
For linear predictors associated with: lambda
                       Estimate Std. Error z value P(>|z|)
                        -1.2552 0.2149 -5.840 5.22e-09 ***
(Intercept)
nationAsia

      nationNorth Africa
      0.2057
      0.1838
      1.119
      0.26309

      nationRest of Africa
      -0.6692
      0.2548
      -2.627
      0.00862 **

      nationSurinam
      -1.5205
      0.6271
      -2.425
      0.01532 *

      nationTurkey
      -1.1888
      0.4343
      -2.737
      0.00619 **

_____
For linear predictors associated with: omega
              Estimate Std. Error z value P(>|z|)
(Intercept) -1.4577 0.3884 -3.753 0.000175 *** gendermale -0.8738 0.3602 -2.426 0.015267 *
age>40yrs
               1.1745
                           0.5423 2.166 0.030326 *
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
AIC: 1677.125
BIC: 1726.976
Residual deviance: 941.5416
Log-likelihood: -829.5625 on 3751 Degrees of freedom
Number of iterations: 10
_____
Population size estimation results:
Point estimate 6699.953
Observed proportion: 28.1% (N obs = 1880)
Boostrap sample skewness: 1.621389
O skewness is expected for normally distributed variable
Bootstrap Std. Error 1719.353
95% CI for the population size:
lowerBound upperBound
```

5001.409 11415.969

```
95% CI for the share of observed population:
lowerBound upperBound
16.46816 37.58941
```

The implementation

Methods

```
R> (popEst <- popSizeEst(basicModel))</pre>
```

Point estimate: 12690.35

Variance: 7885812

95% confidence intervals:

lowerBound upperBound normal 7186.444 18194.26 logNormal 8431.275 19718.32

the popEst object is of the popSizeEstResults class and list type and contains the following fields:

- pointEstimate, variance numerics containing point estimate and variance of this estimate.
- confidenceInterval a data.frame with confidence intervals.
- boot If bootstrap was performed a numeric vector containing the \hat{N} values from the bootstrap, a character vector with value "No bootstrap performed" otherwise.
- control a controlPopVar object with controls used to obtained the object.

```
R> dfb <- dfbeta(basicModel)
R> apply(dfb, 2, quantile)
```

```
(Intercept)
                     gendermale
                                    age>40yrs
                                                 nationAsia nationNorth Africa
0%
    -0.0099087523 -0.0905349877 -0.0200100688 -9.555875e-02
                                                                 -9.660498e-02
25% -0.0015325874 -0.0007770049 0.0001792919 -5.288544e-04
                                                                 -8.417624e-04
50%
     0.0001906118 -0.0002829978 0.0003789034 6.642632e-05
                                                                 -1.768274e-04
75%
     0.0005208531 0.0010171840 0.0006909682 1.199821e-04
                                                                  8.674555e-05
100% 0.0866193890 0.0221346456 0.1600608785 1.799137e-01
                                                                  3.125955e-02
    nationRest of Africa nationSurinam nationTurkey
0%
           -9.449682e-02 -9.313964e-02 -9.619821e-02
25%
           -2.436010e-04 -6.616693e-05 -2.199799e-04
            2.984337e-05 1.969480e-05 7.918067e-05
50%
75%
            8.278833e-05 3.543883e-05 1.427684e-04
            1.097872e-01 9.933829e-01 3.209798e-01
100%
```

R> dfp <- dfpopsize(basicModel, dfbeta = dfb)
R> summary(dfp)

2.2. Marginal frequencies

A popular method of testing the model fit in single source capture-recapture studies is comparing the fitted marginal frequencies $\sum_{j=1}^{N_{obs}} \hat{\mathbb{P}}\left[Y_j = k|\boldsymbol{x}_j, Y_j > 0\right]$ with the observed marginal

frequencies $\sum_{j=1}^{N} \mathcal{I}_{\{k\}}(Y_k) = \sum_{j=1}^{N_{obs}} \mathcal{I}_{\{k\}}(Y_k)$ for $k \geq 1$. If a fitted model bears sufficient resem-

blance to the real data collection process these quantities should be quite close and both G and χ^2 tests may be employed in order to test the statistical significance of the discrepancy with the following **singleRcapture** syntax:

```
R> margFreq <- marginalFreq(basicModel)
R> summary(margFreq, df = 1, drop15 = "group")
```

Test for Goodness of fit of a regression model:

```
Test statistics df P(>X^2)
Chi-squared test 50.06 1 1.5e-12
G-test 34.31 1 4.7e-09
```

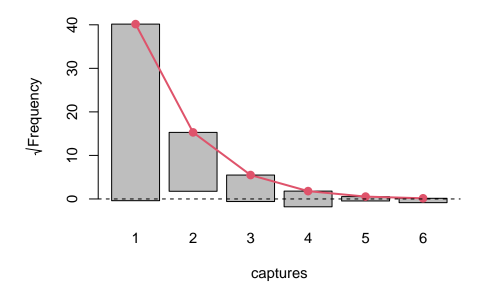
```
Cells with fitted frequencies of < 5 have been grouped Names of cells used in calculating test(s) statistic: 1 2 3
```

where the drop15 argument is used to indicate how to handle the cells with less than 5 fitted observations, note however that currently there is no continuity correction.

2.3. Plots

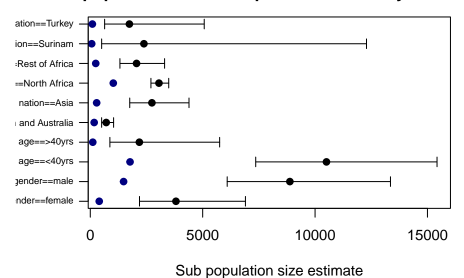
The singleRStaticCountData class has a plot method implementing several types of quick demonstrative plots such as the rootogram Kleiber and Zeileis (2016) for comparing the fitted and marginal frequencies which we can get with the syntax:

```
R> plot(basicModel, plotType = "rootogram")
```



R> plot(basicModel, plotType = "strata")

idence intervals and point estimates for specified sub popserved population sizes are presented as navy coloured



The full list of plot types along with the list of optional arguments which may be passed from the call to the plot method down to base R and graphics functions is listed in the help file:

R> ?plot.singleRStaticCountData

3. Detailed information

3.1. Fitting method

As previously showcased the **singleRcapture** package supports modelling (linear) dependence on covariates of all parameters. To that end a modified IRLS algorithm is employed, full details are available in Yee (2015). In order to employ the algorithm a modified model matrix is created $X_{\rm vlm}$ at call to estimatePopsize. In the context of the models implemented in singleRcapture this matrix can be written as:

$$\boldsymbol{X}_{vlm} = \begin{pmatrix} \boldsymbol{X}_1 & \boldsymbol{0} & \dots & \boldsymbol{0} \\ \boldsymbol{0} & \boldsymbol{X}_2 & \dots & \boldsymbol{0} \\ \vdots & \vdots & \ddots & \vdots \\ \boldsymbol{0} & \boldsymbol{0} & \dots & \boldsymbol{X}_p \end{pmatrix}$$
(4)

where each X_i corresponds to a model matrix associated with user specified formula.

In the context of multi-parameter families we have a matrix of linear predictors η instead of a vector, with the number of columns matching the number of parameters in the distribution.

"Weights" are then modified to be information matrices $\mathbb{E}\left[-\frac{\partial^2 \ell}{\partial \boldsymbol{\eta}_{(k)}^T \partial \boldsymbol{\eta}_{(k)}}\right]$ where $\boldsymbol{\eta}_{(k)}$ is the k'th row of $\boldsymbol{\eta}$, while in the usual IRLS they are scalars $\mathbb{E}\left[-\frac{\partial^2 \ell}{\partial \boldsymbol{\eta}_k^2}\right]$ which is often just $-\frac{\partial^2 \ell}{\partial \boldsymbol{\eta}^2}$.

- 1. Initialize with iter $\leftarrow 1, \eta \leftarrow \text{start}, W \leftarrow I, \ell \leftarrow \ell(\beta)$.
- 2. Store values from the previous step: $\ell_- \leftarrow \ell, W_- \leftarrow W, \beta_- \leftarrow \beta$ (the last assignment is omitted during the first iteration), and assign values in current iteration $\boldsymbol{\eta} \leftarrow \boldsymbol{X}_{\text{vlm}}\boldsymbol{\beta} + \boldsymbol{o}, \boldsymbol{W}_{(k)} \leftarrow \mathbb{E}\left[-\frac{\partial^2 \ell}{\partial \boldsymbol{\eta}_{(k)}^T \partial \boldsymbol{\eta}_{(k)}}\right], Z \leftarrow \boldsymbol{\eta}_{(k)} + \frac{\partial \ell}{\partial \boldsymbol{\eta}_{(k)}} \boldsymbol{W}_{(k)}^{-1} - \boldsymbol{o}_{(k)}.$
- 3. Assign current coefficient value: $\beta \leftarrow (X_{\text{vlm}}WX_{\text{vlm}})^{-1}X_{\text{vlm}}WZ$.
- 4. If $\ell(\beta) < \ell(\beta_-)$ try selecting the smallest value h such that for $\beta_h \leftarrow 2^{-h} \left(\beta + \beta_-\right)$ the inequality $\ell(\beta_h) > \ell(\beta_-)$ holds if this is successful $\beta \leftarrow \beta_h$ else stop the algorithm.
- 5. If convergence is achieved or iter is higher than maxiter end algorithm, else iter← 1+iter and return to step 2.

3.2. The estimatePopsizeFit function

```
R> X <- matrix(data = 0, nrow = 2 * NROW(farmsubmission), ncol = 7)
R> X[1:NROW(farmsubmission), 1:4] <- model.matrix(
+ ~ 1 + log_size + log_distance + C_TYPE,
+ farmsubmission
+ )
```

```
R > X[-(1:NROW(farmsubmission)), 5:7] <- X[1:NROW(farmsubmission), c(1, 3, 4)]
R> # this attribute tells the function which elements of the design matrix
R> # correspond to which linear predictor
R > attr(X, "hwm") <- c(4, 3)
R> start <- glm.fit(# get starting points
   y = farmsubmission$TOTAL_SUB,
    x = X[1:NROW(farmsubmission), 1:4],
    family = poisson()
+ )$coefficients
R> res <- estimatePopsizeFit(</pre>
                = farmsubmission$TOTAL_SUB,
   Χ
                = X,
               = "IRLS",
  method
  priorWeights = 1,
               = ztoigeom(),
+ family
  control = controlMethod(silent = TRUE),
+ coefStart = c(start, 0, 0, 0),
  etaStart = matrix(X %*% c(start, 0, 0, 0), ncol = 2),
offset = cbind(rep(0. NROW(farmsubmission)).
  offset
               = cbind(rep(0, NROW(farmsubmission)),
                         rep(0, NROW(farmsubmission)))
+ )# extract results
R> 11 <- ztoigeom() $makeMinusLogLike(y = farmsubmission $TOTAL_SUB, X = X)
R> print(c(res$beta, -ll(res$beta), res$iter))
[1] -2.784523e+00 6.170270e-01 -6.455925e-02 5.346108e-01 -3.174491e+00
[6] 1.280589e-01 -1.086452e+00 -1.727876e+04 1.500000e+01
R> # Compare with optim call
R> res2 <- estimatePopsizeFit(</pre>
+ y = farmsubmission$TOTAL_SUB,
+ X = X,
+ method = "optim",
+ priorWeights = 1,
  family = ztoigeom(),
  coefStart = c(start, 0, 0, 0),
    control = controlMethod(silent = TRUE),
  offset = cbind(rep(0, NROW(farmsubmission)), rep(0, NROW(farmsubmission)))
+ )# extract results
R> c(res2$beta, -11(res2$beta), res2$iter)
-2.640779e+00 6.258275e-01 -8.293688e-02 5.324707e-01 -1.243731e-01
                                               function
                                                             gradient
-1.629884e-01 -1.105502e+00 -1.728034e+04 1.002000e+03
```

3.3. Avaiable models

The full list of implemented models in **singleRcapture** along with the expressions for probability density functions and point estimates is found in the collective help file for all family functions:

R> ?ztpoisson

Here we limit ourselves to just listing the family functions:

• Zero-truncated and zero-one-truncated Poisson, geometric, NB type II regression where the untruncated distribution is parameterized as:

$$\mathbb{P}[Y = y | \lambda, \alpha] = \frac{\Gamma(y + \alpha^{-1})}{\Gamma(\alpha^{-1}) y!} \left(\frac{\alpha^{-1}}{\alpha^{-1} + \lambda}\right)^{\alpha^{-1}} \left(\frac{\lambda}{\lambda + \alpha^{-1}}\right)^{y}.$$

• Zero-truncated one-inflated (ztoi) modifications distributions where the new probability \mathbb{P}^* measure is defined in terms of count data measure \mathbb{P} with support on $\mathbb{N} \cup \{0\}$ as:

$$\mathbb{P}^*[Y = y] = \begin{cases} \mathbb{P}[Y = 0] & y = 0, \\ \omega (1 - \mathbb{P}[Y = 0]) + (1 - \omega) \mathbb{P}[Y = 1] & y = 1, \\ (1 - \omega) \mathbb{P}[Y = y] & y > 1, \end{cases}$$
$$\mathbb{P}^*[Y = y | Y > 0] = \omega \mathcal{I}_{f11}(y) + (1 - \omega) \mathbb{P}[Y = y | Y > 0].$$

• One-inflated zero-truncated (oizt) modifications distributions where the new probability \mathbb{P}^* measure is defined as:

$$\begin{split} \mathbb{P}^*[Y = y] &= \omega \mathcal{I}_{\{1\}}(y) + (1 - \omega) \mathbb{P}[Y = y], \\ \mathbb{P}^*[Y = y | Y > 0] &= \omega \frac{\mathcal{I}_{\{1\}}(y)}{1 - (1 - \omega) \mathbb{P}[Y = 0]} + (1 - \omega) \frac{\mathbb{P}[Y = y]}{1 - (1 - \omega) \mathbb{P}[Y = 0]}. \end{split}$$

• Generalized Chao's and Zelterman's estimators via logistic regression on variable Z defined as Z=1 if Y=2 and Z=0 if Y=1 with $Z\sim b(p)$ where $\mathrm{logit}(p)=\ln(\lambda/2)$ for poisson parameter λ ,

$$\hat{N} = N_{obs} + \sum_{k=1}^{f_1 + f_2} \left(2 \exp\left(\boldsymbol{x}_k \hat{\boldsymbol{\beta}}\right) + 2 \exp\left(2\boldsymbol{x}_k \hat{\boldsymbol{\beta}}\right) \right)^{-1}, \quad \text{(Chao's estimator)}$$

$$\hat{N} = \sum_{k=1}^{N_{obs}} \left(1 - \exp\left(-2 \exp\left(\boldsymbol{x}_k \hat{\boldsymbol{\beta}}\right)\right) \right)^{-1}. \quad \text{(Zelterman's estimator)}$$

• Alternative approaches to modelling one-inflation that mimic hurdle models where the first type zero truncated hurdle model (ztHurdle) is defined as:

$$\mathbb{P}^*[Y=y] = \begin{cases} \frac{\mathbb{P}[Y=0]}{1-\mathbb{P}[Y=1]} & y=0, \\ \pi(1-\mathbb{P}[Y=1]) & y=1, \\ (1-\pi)\frac{\mathbb{P}[Y=y]}{1-\mathbb{P}[Y=1]} & y>1, \end{cases}$$

$$\mathbb{P}^*[Y=y|Y>0] = \pi \mathcal{I}_{\{1\}}(y) + (1-\pi)\mathcal{I}_{\mathbb{N}\backslash\{1\}}(y) \frac{\mathbb{P}[Y=y]}{1-\mathbb{P}[Y=0] - \mathbb{P}[Y=1]}$$

• The Hurdle zero truncarted (Hurdlezt) is defined as:

$$\begin{split} \mathbb{P}^*[Y=y] &= \begin{cases} \pi & y=1,\\ (1-\pi)\frac{\mathbb{P}[Y=y]}{1-\mathbb{P}[Y=1]} & y \neq 1, \end{cases} \\ \mathbb{P}^*[Y=y|Y>0] &= \begin{cases} \pi\frac{1-\mathbb{P}[Y=1]}{1-\mathbb{P}[Y=0]-\mathbb{P}[Y=1]} & y=1,\\ (1-\pi)\frac{\mathbb{P}[Y=y]}{1-\mathbb{P}[Y=0]-\mathbb{P}[Y=1]} & y>1. \end{cases} \end{split}$$

Takeaways of different models

- The dispersion parameter in nb is often interpreted as indicating unobserved heterogeneity
- Geometric is the light version of that
- inflated models model inflation
- Hurdle models can also model deflation as well as both inflation and deflation simultaneously so they are more flexible
- By contrast the interpretation of the ω inflation parameter is more convenient that the interpretation of the π probability parameter.

3.4. Structure of a family function

• makeMinusLogLike - A factory function for creating the:

$$\ell(oldsymbol{eta}), rac{\partial \ell}{\partial oldsymbol{eta}}, rac{\partial^2 \ell}{\partial oldsymbol{eta}^T \partial oldsymbol{eta}}$$

functions from y vector and X_{vlm} the argument deriv with possible values in c(0, 1, 2) provides which derivative to return with the default 0 being just the minus log-likelihood.

- links List with link functions.
- mu.eta, variance Functions of linear predictors that return expected value and variance. There is a 'type' argument with 2 possible values "trunc" and "nontrunc" that specifies whether to return $\mathbb{E}[Y|Y>0]$, var[Y|Y>0] or $\mathbb{E}[Y]$, var[Y] respectively, also the deriv argument with values in c(0, 1, 2) is used for indicating the derivative with respect to the linear predictors with is used for providing standard error in predict method.
- family Character that specifies name of the model.
- valideta, validmu For now only returns true. In near future will be used to check whether applied linear predictors are valid (i.e. are transformed into some elements of parameter space the subjected to inverse link function).

- funcZ, Wfun Functions that create pseudo residuals and working weights used in IRLS algorithm.
- devResids Function that given the linear predictors prior weights vector and response vector returns deviance residuals.
- pointEst, popVar Functions that given prior weights linear predictors and in the later case also estimation of $cov(\hat{\beta})$ and X_{vlm} matrix return point estimate for population size and analytic estimation of its variance. There is a additional boolean parameter contr in the former function that if set to true returns contribution of each unit.
- etaNames Names of linear predictors.
- densityFunction A function that given linear predictors returns value of PMF at values x. Additional argument type specifies whether to return $\mathbb{P}[Y|Y>0]$ or $\mathbb{P}[Y]$.
- simulate A function that generates values of dependent vector given linear predictors.
- getStart Expression for generating starting points.

3.5. Bootstrap algorithms

There are three types of bootstrap algorithms which the user may specify in controlPopVar controls with bootType argument which has three possible values "parametric", "semiparametric", "nonparametric" with the nonparametric being bootstrap being the usual bootstrap algorithm which as argued in Norris and Pollock (1996) and Zwane and Van der Heijden (2003). The idea of semiparametric bootstrap is to modify the usual bootstrap to include the additional uncertainty due to the sample size being a random variable. This type of bootstrap can be in short described as:

- 1. Draw the sample size $N'_{obs} \sim \text{Be}\left(N', \frac{N' N_{obs}}{N'}\right)$, where $N' = \lfloor \hat{N} \rfloor + b \left(\lfloor \hat{N} \rfloor \hat{N}\right)$.
- 2. Draw N'_{obs} units from the data uniformly without replacement.
- 3. Obtain new population size estimate using bootstrap data.
- 4. Repeat 1-3 B times.

In other words we first draw the sample size and then the sample conditional on the sample size. Note that in using semi-parametric bootstrap one implicitly assumes that the population size estimate \hat{N} is accurate. The last implemented bootstrap type is the parametric algorithm which in short first draws the finite population of size $\approx \hat{N}$ from the superpopulation model and then samples from this population according to the selected model:

- 1. Draw the number of covariates equal to $\lfloor \hat{N} \rfloor + b \left(\lfloor \hat{N} \rfloor \hat{N} \right)$ proportional to the estimated contribution $(\mathbb{P}\left[Y_k > 0 | \boldsymbol{x}_k \right])^{-1}$ with replacement.
- 2. Using the fitted model and regression coefficients $\hat{\beta}$ draw for each covariate the Y value from the corresponding probability measure on $\mathbb{N} \cup \{0\}$.
- 3. Truncate units with drawn Y value equal to 0.

- 4. Obtain population size estimate based on the truncated data.
- 5. Repeat 1-4 B times.

Note however that for this type of algorithm to result in consistent standard error estimates it is imperative that the estimated model for the entire superpopulation probability space is consistent which may be much less realistic than semiparametric bootstrap. The parametric bootstrap algorithm is the default in **singleRcapture**.

Additional arguments accepted by the contorlPopVar function which are relevant to bootstrap are:

- alpha, B significance level and number of bootstrap samples to be performed respectively with 0.05 and 500 being the default options.
- cores number of process cores to use in bootstrap (1 by default) parallel computing is done via **doParallel**, **foreach**, **parallel** packages.
- keepbootStat logical value indicating whether to keep a vector of statistics produced by bootstrap.
- traceBootstrapSize, bootstrapVisualTrace logical values indicating whether sample and population size should be tracked (FALSE by default) these work only when cores = 1.
- fittingMethod, bootstrapFitcontrol fitting method (by default the same as used in the original call) and control parameters (controlMethod) for model fitting in bootstrap.

4. Integration with the VGAM, countreg packages

The singleRcaptureExtra extensions allows for converting objects created by vglm, vgam, countreg functions from packages VGAM, countreg to a singleRStaticCountData via the respective estimatePopsize methods for their classes. The help files for all the methods and all the controll functions are accessed by:

```
R> ?estimatePopsize.vgam
```

R> ?controlEstPopVgam

Using the fitted zerotrunc, vglm, vgam class objects in population size estimation such as the one additive models with smooth terms for dataset from Böhning, Vidal-Diez, Lerdsuwansri, Viwatwongkasem, and Arnold (2013):

```
R> library(VGAM)
```

Loading required package: stats4

Loading required package: splines

```
R> library(singleRcaptureExtra)
R> modelVgam <- vgam(</pre>
+ TOTAL\_SUB \sim (s(log\_size, df = 3) + s(log\_distance, df = 2)) / C\_TYPE,
   data = farmsubmission,
   family = posnegbinomial(lsize = negloglink)
+ )
can be a complished with the following syntax simple syntax:
R> modelVgamPop <- estimatePopsize(modelVgam)</pre>
Compare with a simmilar linear model from base singleRcapture:
R> modelBase <- estimatePopsize(</pre>
   TOTAL_SUB ~ (log_size + log_distance) * C_TYPE,
   data = farmsubmission,
  model = ztnegbin()
+ )
R> summary(modelBase)
Call:
estimatePopsize.default(formula = TOTAL_SUB ~ (log_size + log_distance) *
   C_TYPE, data = farmsubmission, model = ztnegbin())
Pearson Residuals:
    Min. 1st Qu.
                     Median
                                Mean
                                      3rd Qu.
-0.729357 -0.317558 -0.152482 0.000609 0.148985 6.604269
Coefficients:
For linear predictors associated with: lambda
                     Estimate Std. Error z value P(>|z|)
(Intercept)
                     -1.77609 0.45894 -3.870 0.000109 ***
                       0.49391 0.02521 19.594 < 2e-16 ***
log_size
                     log_distance
C_TYPEDairy
                      -1.68591 0.55327 -3.047 0.002310 **
log_size:C_TYPEDairy
                      log_distance:C_TYPEDairy 0.08568 0.04874 1.758 0.078762 .
_____
For linear predictors associated with: alpha
          Estimate Std. Error z value P(>|z|)
(Intercept) 0.57673 0.07267 7.936 2.09e-15 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
AIC: 34481.99
```

```
BIC: 34533.76
Residual deviance: 17611.16
Log-likelihood: -17233.99 on 24065 Degrees of freedom
Number of iterations: 9
Population size estimation results:
Point estimate 38877
Observed proportion: 31% (N obs = 12036)
Std. Error 1749.448
95% CI for the population size:
         lowerBound upperBound
normal
           35448.14
                      42305.85
logNormal
           35661.32
                      42530.37
95% CI for the share of observed population:
         lowerBound upperBound
normal
           28.44996
                      33.95382
logNormal
           28.29978
                      33.75085
R> summary(modelVgamPop)
Call:
estimatePopsize.vgam(formula = modelVgam)
______
Population size estimation results:
Point estimate 37760.01
Observed proportion: 31.9% (N obs = 12036)
Std. Error 1630.429
95% CI for the population size:
         lowerBound upperBound
normal
           34564.42
                      40955.59
           34757.77
logNormal
                      41158.93
95% CI for the share of observed population:
         lowerBound upperBound
           29.38793
                    34.82193
normal
           29.24274
                      34.62823
logNormal
-----
-- Summary of foreign object --
Call:
vgam(formula = TOTAL_SUB ~ (s(log_size, df = 3) + s(log_distance,
   df = 2))/C_TYPE, family = posnegbinomial(lsize = negloglink),
   data = farmsubmission)
```

```
Names of additive predictors: loglink(munb), negloglink(size)
```

Dispersion Parameter for posnegbinomial family: 1

Log-likelihood: -17214.62 on 24063.17 degrees of freedom

Number of Fisher scoring iterations: 11

DF for Terms and Approximate Chi-squares for Nonparametric Effects

```
Df Npar Df Npar Chisq
(Intercept):1
(Intercept):2
                                                       1
                                                       1
s(log_size, df = 3)
                                                             1.8
                                                                      51.949
s(\log_{distance}, df = 2)
                                                             1.0
                                                                       3.503
s(log_size, df = 3):s(log_distance, df = 2):C_TYPE 2
                                                        P(Chi)
(Intercept):1
(Intercept):2
s(log_size, df = 3)
                                                      0.000000
s(\log_{distance}, df = 2)
                                                      0.063835
s(log_size, df = 3):s(log_distance, df = 2):C_TYPE
```

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A. Implementing custom singleRcapture family function

Suppose we want to implement a very specific zero truncated family function in the **singleRcapture** which corresponds to the following "untruncated" distribution:

$$\mathbb{P}[Y = y | \lambda, \pi] = \begin{cases}
1 - \frac{1}{2}\lambda - \frac{1}{2}\pi & \text{when: } y = 0 \\
\frac{1}{2}\pi & \text{when: } y = 1 \\
\frac{1}{2}\lambda & \text{when: } y = 2,
\end{cases}$$
(5)

with $\lambda, \pi \in (0,1)$ being dependent on covariates. The following would be one way of implementing it, with lambda, pi in the code meaning $\frac{1}{2}\lambda, \frac{1}{2}\pi$ in the equation above:

```
R> myFamilyFunction <- function(lambdaLink = c("logit", "cloglog", "probit"), + piLink = c("logit", "cloglog", "probit"), + ...) {
```

```
if (missing(lambdaLink)) lambdaLink <- "logit"</pre>
+
    if (missing(piLink))
                                  piLink <- "logit"</pre>
    links <- list()</pre>
    attr(links, "linkNames") <- c(lambdaLink, piLink)</pre>
+
    lambdaLink <- switch(lambdaLink,</pre>
      "logit"
               = singleRcapture:::singleRinternallogitLink,
+
      "cloglog" = singleRcapture:::singleRinternalcloglogLink,
      "probit" = singleRcapture:::singleRinternalprobitLink
    )
+
   piLink <- switch(piLink,</pre>
      "logit" = singleRcapture:::singleRinternallogitLink,
      "cloglog" = singleRcapture:::singleRinternalcloglogLink,
      "probit" = singleRcapture:::singleRinternalprobitLink
    )
+
+
    links[1:2] <- c(lambdaLink, piLink)</pre>
   mu.eta <- function(eta, type = "trunc", deriv = FALSE, ...) {</pre>
                    piLink(eta[, 2], inverse = TRUE) / 2
             <-
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
+
+
      if (!deriv) {
        switch (type,
          "nontrunc" = pi + 2 * lambda,
          "trunc" = 1 + lambda / (pi + lambda)
        )
+
      } else {
+
        # Only necessary if one wishes to use standard errors in predict method
+
        switch (type,
          "nontrunc" = {
            matrix(c(2, 1) * c(
+
              lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2,
                  piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2
+
            ), ncol = 2)
+
          },
+
          "trunc" = {
            matrix(c(
              pi / (pi + lambda) ^ 2,
              -lambda / (pi + lambda) ^ 2
+
+
              lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2,
                  piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2
            ), ncol = 2)
```

```
)
      }
    }
    variance <- function(eta, type = "nontrunc", ...) {</pre>
+
                     piLink(eta[, 2], inverse = TRUE) / 2
+
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
      switch (type,
      "nontrunc" = pi * (1 - pi) + 4 * lambda * (1 - lambda - pi),
      "trunc" = lambda * (1 - lambda) / (pi + lambda)
+
      )
    }
+
+
    Wfun <- function(prior, y, eta, ...) {
                     piLink(eta[, 2], inverse = TRUE) / 2
             <-
+
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
      G01 <- ((lambda + pi) ^ (-2)) * piLink(eta[, 2], inverse = TRUE, deriv = 1) *
+
        lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) * prior / 4
      G00 <- ((lambda + pi) ^ (-2)) - (pi ^ (-2)) - lambda / ((lambda + pi) * (pi ^ 2
      G00 \leftarrow G00 * prior * (piLink(eta[, 2], inverse = TRUE, deriv = 1) ^ 2) / 4
+
+
+
      G11 <- ((lambda + pi) ^ (-2)) - (((lambda + pi) * lambda) ^ -1)
      G11 \leftarrow G11 * prior * (lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) ^ 2) / 4
+
      matrix(
        -c(G11, # lambda)
           GO1, # mixed
+
+
           GO1, # mixed
           G00 # pi
        ),
        dimnames = list(rownames(eta), c("lambda", "mixed", "mixed", "pi")),
+
        ncol = 4
+
      )
    }
+
+
    funcZ <- function(eta, weight, y, prior, ...) {</pre>
                     piLink(eta[, 2], inverse = TRUE) / 2
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
+
      weight <- weight / prior</pre>
                           - ((lambda + pi) ^ -1)
      GO \leftarrow (2 - y) / pi
      G1 \leftarrow (y - 1) / lambda - ((lambda + pi) ^ -1)
```

```
G1 <- G1 * lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2
                      piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2
      GO <- GO *
      uMatrix \leftarrow matrix(c(G1, G0), ncol = 2)
+
      weight <- lapply(X = 1:nrow(weight), FUN = function (x) {</pre>
        matrix(as.numeric(weight[x, ]), ncol = 2)
+
+
      })
      pseudoResid <- sapply(X = 1:length(weight), FUN = function (x) {</pre>
        #xx <- chol2inv(chol(weight[[x]])) # less computationally demanding</pre>
        xx <- solve(weight[[x]]) # more stable</pre>
        xx %*% uMatrix[x, ]
+
+
      7)
+
      pseudoResid <- t(pseudoResid)</pre>
      dimnames(pseudoResid) <- dimnames(eta)</pre>
+
      pseudoResid
+
    }
+
    minusLogLike <- function(y, X, offset,</pre>
                               weight
                                         = 1,
+
                               NbyK
                                         = FALSE,
                               vectorDer = FALSE,
                               deriv
                                         = 0.
                               ...) {
      y <- as.numeric(y)</pre>
      if (is.null(weight)) {
        weight <- 1
      }
+
      if (missing(offset)) {
+
        offset \leftarrow cbind(rep(0, NROW(X) / 2), rep(0, NROW(X) / 2))
      }
      if (!(deriv %in% c(0, 1, 2))) stop("Only score function and derivatives up to 2
+
      deriv <- deriv + 1 # to make it conform to how switch in R works, i.e. indexing
+
      switch (deriv,
+
+
        function(beta) {
          eta <- matrix(as.matrix(X) %*% beta, ncol = 2) + offset
                         piLink(eta[, 2], inverse = TRUE) / 2
          lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
          -sum(weight * ((2 - y) * log(pi) + (y - 1) * log(lambda) - log(pi + lambda)
        },
+
        function(beta) {
          eta <- matrix(as.matrix(X) %*% beta, ncol = 2) + offset
                         piLink(eta[, 2], inverse = TRUE) / 2
          lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
```

```
GO \leftarrow (2 - y) / pi - ((lambda + pi) ^ -1)
          G1 \leftarrow (y - 1) / lambda - ((lambda + pi) ^ -1)
          G1 <- G1 * weight * lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2
                                   piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2
          GO <- GO * weight *
+
          if (NbyK) {
+
            XX <- 1: (attr(X, "hwm")[1])
            return(cbind(as.data.frame(X[1:nrow(eta), XX]) * G1, as.data.frame(X[-(1:
          7
          if (vectorDer) {
+
            return(cbind(G1, G0))
+
+
          as.numeric(c(G1, G0) \%*\% X)
        },
+
        function (beta) {
          lambdaPredNumber <- attr(X, "hwm")[1]</pre>
+
          eta <- matrix(as.matrix(X) %*% beta, ncol = 2) + offset
          рi
                         piLink(eta[, 2], inverse = TRUE) / 2
                 <-
          lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
          res <- matrix(nrow = length(beta), ncol = length(beta),
+
+
                         dimnames = list(names(beta), names(beta)))
          # pi^2 derivative
          dpi <- (2 - y) / pi - (lambda + pi) ^ -1
          G00 \leftarrow ((lambda + pi) ^ (-2)) - (2 - y) / (pi ^ 2)
+
          G00 \leftarrow t(as.data.frame(X[-(1:(nrow(X) / 2)), -(1:lambdaPredNumber)] *
+
          (G00 * ((piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2) ^ 2) +
          dpi * piLink(eta[, 2], inverse = TRUE, deriv = 2) / 2) * weight)) %*%
          as.matrix(X[-(1:(nrow(X) / 2)), -(1:lambdaPredNumber)])
          # mixed derivative
          G01 \leftarrow (lambda + pi) ^ (-2)
+
+
          G01 \leftarrow t(as.data.frame(X[1:(nrow(X) / 2), 1:lambdaPredNumber]) *
          GO1 * (lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2) *
          (piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2) * weight) %*%
          as.matrix(X[-(1:(nrow(X) / 2)), -(1:lambdaPredNumber)])
          # lambda^2 derivative
          G11 \leftarrow ((lambda + pi) ^ (-2)) - (y - 1) / (lambda ^ 2)
          dlambda \leftarrow (y - 1) / lambda - ((lambda + pi) ^ -1)
          G11 <- t(as.data.frame(X[1:(nrow(X) / 2), 1:lambdaPredNumber] *</pre>
          (G11 * ((lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2) ^ 2) +
```

```
dlambda * lambdaLink(eta[, 1], inverse = TRUE, deriv = 2) / 2) * weight)) %
          X[1:(nrow(X) / 2), 1:lambdaPredNumber]
          res[-(1:lambdaPredNumber), -(1:lambdaPredNumber)] <- G00</pre>
          res[1:lambdaPredNumber, 1:lambdaPredNumber] <- G11
          res[1:lambdaPredNumber, -(1:lambdaPredNumber)] <- t(G01)</pre>
          res[-(1:lambdaPredNumber), 1:lambdaPredNumber] <- G01</pre>
          res
        }
+
      )
    }
+
+
+
    validmu <- function(mu) {</pre>
      (sum(!is.finite(mu)) == 0) \&\& all(0 < mu) \&\& all(2 > mu)
    }
+
    # this is optional
    devResids <- function(y, eta, wt, ...) {</pre>
    }
    pointEst <- function (pw, eta, contr = FALSE, ...) {</pre>
                     piLink(eta[, 2], inverse = TRUE) / 2
              <-
+
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
      N <- pw / (lambda + pi)</pre>
      if(!contr) {
        N \leftarrow sum(N)
      }
+
      N
+
    }
    popVar <- function (pw, eta, cov, Xvlm, ...) {</pre>
                     piLink(eta[, 2], inverse = TRUE) / 2
+
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
      bigTheta1 <- -pw / (pi + lambda) ^ 2 # w.r to pi
      bigTheta1 <- bigTheta1 * piLink(eta[, 2], inverse = TRUE, deriv = 1) / 2</pre>
+
      bigTheta2 <- -pw / (pi + lambda) ^ 2 # w.r to lambda
      bigTheta2 <- bigTheta2 * lambdaLink(eta[, 1], inverse = TRUE, deriv = 1) / 2# w
+
      bigTheta <- t(c(bigTheta2, bigTheta1) %*% Xvlm)</pre>
      f1 <- t(bigTheta) %*% as.matrix(cov) %*% bigTheta
+
      f2 <- sum(pw * (1 - pi - lambda) / ((pi + lambda) ^ 2))
```

```
f1 + f2
+
    dFun <- function (x, eta, type = c("trunc", "nontrunc")) {
      if (missing(type)) type <- "trunc"</pre>
+
              <-
                     piLink(eta[, 2], inverse = TRUE) / 2
      рi
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
+
+
      switch (type,
        "trunc" = {
           (pi * as.numeric(x == 1) + lambda * as.numeric(x == 2)) / (pi + lambda)
+
        },
+
        "nontrunc" = {
+
           (1 - pi - lambda) * as.numeric(x == 0) +
          pi * as.numeric(x == 1) + lambda * as.numeric(x == 2)
+
      )
+
    }
    simulate <- function(n, eta, lower = 0, upper = Inf) {</pre>
      рi
                      piLink(eta[, 2], inverse = TRUE) / 2
              <-
+
      lambda <- lambdaLink(eta[, 1], inverse = TRUE) / 2</pre>
+
      CDF <- function(x) {</pre>
        ifelse(x == Inf, 1,
+
+
        ifelse(x < 0, 0,
        ifelse(x < 1, 1 - pi - lambda,
        ifelse(x < 2, 1 - lambda, 1))))
      }
      1b <- CDF(lower)</pre>
+
      ub <- CDF(upper)</pre>
+
      p_u <- stats::runif(n, lb, ub)</pre>
      sims \leftarrow rep(0, n)
      cond <- CDF(sims) <= p_u</pre>
      while (any(cond)) {
+
        sims[cond] <- sims[cond] + 1</pre>
        cond <- CDF(sims) <= p_u</pre>
+
      }
+
+
      sims
    }
+
    getStart <- expression(</pre>
      if (method == "IRLS") {
+
+
        etaStart <- cbind(</pre>
           family$links[[1]](mean(observed == 2) * (1 + 0 * (observed == 2))), # lambd
           family$links[[2]](mean(observed == 1) * (1 + 0 * (observed == 1))) # pi
        ) + offset
      } else if (method == "optim") {
```

```
init <- c(
          family$links[[1]](weighted.mean(observed == 2, priorWeights) * 1 + .0001),
          family$links[[2]](weighted.mean(observed == 1, priorWeights) * 1 + .0001)
        if (attr(terms, "intercept")) {
+
          coefStart <- c(init[1], rep(0, attr(Xvlm, "hwm")[1] - 1))</pre>
+
+
          coefStart <- rep(init[1] / attr(Xvlm, "hwm")[1], attr(Xvlm, "hwm")[1])</pre>
        if ("(Intercept):pi" %in% colnames(Xvlm)) {
          coefStart <- c(coefStart, init[2], rep(0, attr(Xvlm, "hwm")[2] - 1))</pre>
          coefStart <- c(coefStart, rep(init[2] / attr(Xvlm, "hwm")[2], attr(Xvlm, "h</pre>
+
+
+
     }
    )
+
+
    structure(
+
      list(
        makeMinusLogLike = minusLogLike,
        densityFunction = dFun,
        links
                  = links,
+
        mu.eta
                  = mu.eta,
        valideta = function (eta) {TRUE},
+
+
        variance = variance,
        Wfun
                  = Wfun,
        funcZ
                 = funcZ,
        devResids = devResids,
        validmu = validmu,
+
        pointEst = pointEst,
+
                = popVar,
        popVar
        family
                  = "myFamilyFunction",
        etaNames = c("lambda", "pi"),
        simulate = simulate,
+
        getStart = getStart,
+
        extraInfo = c(
                     = "pi / 2 + lambda",
+
          mean
                     = paste0("(pi / 2) * (1 - pi / 2) + 2 * lambda * (1 - lambda / 2
          variance
          popSizeEst = "(1 - (pi + lambda) / 2) ^ -1",
                     = "1 + lambda / (pi + lambda)",
          varianceTr = paste0("lambda * (1 - lambda / 2) / (pi + lambda)")
        )
+
      ),
      class = c("singleRfamily", "family")
+ }
```

A quick tests shows us that this implementation in fact works:

```
R> set.seed(123)
R> Y <- simulate(</pre>
     myFamilyFunction(lambdaLink = "logit", piLink = "logit"),
     nsim = 1000, eta = matrix(0, nrow = 1000, ncol = 2),
     truncated = FALSE
+ )
R> mm <- estimatePopsize(</pre>
     formula = Y \sim 1,
     data = data.frame(Y = Y[Y > 0]),
     model = myFamilyFunction(lambdaLink = "logit",
                             piLink = "logit"),
+
+
    controlPopVar = controlPopVar(
         # the usual observed information matrix is ill-suited for this distribution
         covType = "Fisher"
     )
+ )
R> summary(mm)
Call:
estimatePopsize.default(formula = Y ~ 1, data = data.frame(Y = Y[Y >
    0]), model = myFamilyFunction(lambdaLink = "logit", piLink = "logit"),
    controlPopVar = controlPopVar(covType = "Fisher"))
Pearson Residuals:
  Min. 1st Qu. Median Mean 3rd Qu.
-0.8198 -0.8198 0.8099 0.0000 0.8099 0.8099
Coefficients:
For linear predictors associated with: lambda
           Estimate Std. Error z value P(>|z|)
(Intercept) 0.01217
                      0.20253 0.06 0.952
-----
For linear predictors associated with: pi
           Estimate Std. Error z value P(>|z|)
AIC: 687.4249
BIC: 695.8259
Residual deviance: 0
Log-likelihood: -341.7124 on 984 Degrees of freedom
Number of iterations: 2
```

```
Population size estimation results:
Point estimate 986
Observed proportion: 50% (N obs = 493)
Std. Error 70.30092
95% CI for the population size:
          lowerBound upperBound
normal
            848.2127
                       1123.787
            866.3167
logNormal
                       1144.053
95% CI for the share of observed population:
          lowerBound upperBound
normal
            43.86951
                       58.12221
logNormal
            43.09241
                       56.90759
```

Where the link functions such as singleRcapture:::singleRinternalcloglogLink are just internal functions in singleRcapture that compute link functions their inverses and derivatives of both links and inverse link up to third order:

```
R> singleRcapture:::singleRinternalcloglogLink
```

```
function (x, inverse = FALSE, deriv = 0)
    deriv <- deriv + 1
    if (isFALSE(inverse)) {
        res <- switch(deriv, log(-log(1 - x)), -1/((1 - x) *
            log(1 - x)), -(1 + log(1 - x))/((x - 1)^2 * log(1 - x))
            x)^2, (2 * log(1 - x)^2 + 3 * log(1 - x) + 2)/(log(1 - x)^2)
            x)^3 * (x - 1)^3)
    }
    else {
        res <- switch(deriv, 1 - \exp(-\exp(x)), \exp(x - \exp(x)),
            (1 - \exp(x)) * \exp(x - \exp(x)), (\exp(2 * x) - 3 *
                exp(x) + 1) * exp(x - exp(x))
    }
    res
}
<bytecode: 0x123dab290>
<environment: namespace:singleRcapture>
```

one might of course include code for cumputing them manually.

References

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Affiliation:

Piotr Chlebicki Stockholm University Matematiska institutionen Albano hus 1 106 91 Stockholm, Sweden

E-mail: piotr.chlebicki@math.su.se

URL: https://github.com/Kertoo, https://www.su.se/profiles/pich3772

Maciej Beręsewicz Poznań University of Economics and Business Statistical Office in Poznań Poznań University of Economics and Business Department of Statistics Institute of Informatics and Quantitative Economics Al. Niepodległosci 10 61-875 Poznań, Poland

Statistical Office in Poznań ul. Wojska Polskiego 27/29 60-624 Poznań, Poland

E-mail: maciej.beresewicz@ue.poznan.pl

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