DRAFT: Analysis of the reporting styles using generalized ordered threshold models.

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Abstract

The *hopit* package provides R functions to fit and analyze ordered response data in the context of reporting styles. In this vignette we describe the formulation and fit of *hopit* models as well as functions used to analyse reporting styles.

1. Introduction

The *hopit* package provides R functions to fit and analyze ordered response data in the context of reporting styles.

The ordered response data classifies a measure of interest into ordered categories collected during a survey. If the dependent variable is a happiness then a respondent typically answers a question: "Taking all things together, would you say you are ...?" and have some response options e.g."very happy", "pretty happy", "not too happy", "very unhappy" (Liao, Fu, and Yi 2005). Similarly if interviewees are asked to evaluate their health in general (e.g. "Would you say your health is ...?") they may choose among several categories, e.g. very good, good, fair, bad, and very bad (Jürges 2007). In political sciences a respondent may be asked for an opinion about recent legislation (e.g. "Rate your feelings about the proposed legislation") and asked to choose among several categories "strongly oppose", "mildly oppose", "indifferent", "mildly support", "strongly support" (Greene and Hensher 2010). It is easy to imagine other multi-level ordinal variables that might by use during the survey and to which methodology described below could be applied with.

Practically, it is assumed that when responding to a survey question about their general happiness, health, feeling, attitude or other status, participants assess their true value of this unobserved continuous variable, and project it to a provided discrete scale. The thresholds that each individual uses to categorize their true status into a specific response option may be affected by the choice of a reference group, earlier life experiences, and cross-cultural differences in using scales, and thus, may differ across individuals depending on their gender, age, cultural background, education, and personality traits, among other factors.

From the modeling perspective, one of the main tasks is to compute this continuous estimate of individuals' underlying, latent variable based on several specific characteristics of the considered response (e.g. health variables or happiness variables) and accounting also for variations in reporting across socio-demographic and cultural groups. More specifically, to build the latent, underlying variable a generalized hierarchical ordered threshold model is fitted, which regresses the reported status/attitude/feeling on two sets of independent variables (Boes and Winkelmann 2006; Greene et al. 2014). When a dependent reported ordered variable is self-rated health status then the first set of variables - health variables - assesses individuals' specific aspects of health, and might include chronic conditions, mobility level, difficulties with a range of daily activities, performance on grip strength test, anthropometric measures, lifestyle behaviors, etc. Using the second set of independent variables (threshold variables), the model also adjusts for the differences across socio-demographic and cultural groups like cultural background, gender, age, education, etc. (King et al. 2004; Jürges 2007; but see Rebelo and Pereira 2014).

Once the model is fitted its estimates (latent variable and threshold coefficients) can be used to calculate the differences in reporting styles among groups of people having different contextual characteristics realized by calculation of differences between expected and reported ordinal response measures (Jürges 2007).

2. Generalized (hierarchical) ordered threshold model

Ordered threshold models are used with ordered categorical dependent variables. The generalized ordered threshold models (Ierza 1985; Boes and Winkelmann 2006; Greene et al. 2014) are an extension to the ordered threshold models (McKelvey and Zavoina 1975). In the latter models the thresholds are constant, whereas generalized models allows thresholds to be dependent on covariates. Greene and Hensher (2010) and Greene et al. (2014) pointed out that also thresholds must be ordered so that a model has a sense. This motivated Greene and coauthors to call this models **HOPIT**, which stands for hierarchical ordered probit models.

In the the self-rated health example, the response variable is self-rated health and latent variable h_i can depend on different health conditions and diseases (health variables X). Variables X are modeled with parallel regression assumption. According to the assumption, coefficients, which describe the relationship between lowest and all higher response categories, are the same as those coefficients, which describe the relationship between another (e.g. adjacent) lowest and the remaining higher response categories. In the considered case h_i is modeled as a linear function of X and their coefficients β :

$$h_i = \sum_{k=1}^K \beta_k X_{i,k} = X'\beta \tag{1}$$

where index $i \in 1...N$ is number of cases (e.g. respondents), X is in the form of model matrix, and K is number of columns in X. As described above, the categorization (response mechanism) of the latent variable h_i is modeled in terms of thresholds $\alpha_{i,j}$ assuming that thresholds of lower order are never greater than thresholds of higher orders:

$$\begin{cases} y_{i} = 1 \iff \alpha_{i,0} \leq h_{i} < \alpha_{i,1} \\ y_{i} = 2 \iff \alpha_{i,1} \leq h_{i} < \alpha_{i,2} \\ \cdots \\ y_{i} = j \iff \alpha_{i,j-1} \leq h_{i} < \alpha_{i,j} \\ \cdots \\ y_{i} = J \iff \alpha_{i,J-1} \leq h_{i} < \alpha_{i,J} \end{cases}$$

$$(2)$$

The thresholds (cut points, α) are modeled by threshold variables γ and intercepts λ . It is assumed that they model contextual characteristics of the respondent (e.g. country of origin, gender, age, etc.). Threshold variables are modeled without parallel regression assumption, thus each threshold is modeled by a variable independently (Boes and Winkelmann 2006; Greene et al. 2014).

Different parametrizations of thresholds exist (Greene et al. 2014; Rebelo and Pereira 2014; Jürges 2007). In the package, King et al. (2004) and Jürges (2007) parametrization is used, which assumes that:

$$\alpha_{i, j} = \begin{cases} -\infty & for \ j = 0\\ \lambda_1 + \sum_{m=1}^{M} \gamma_{1,m} Y_{i,m} & for \ j = 1\\ \alpha_{i, j-1} + exp(\lambda_j + \sum_{m=1}^{M} \gamma_{j,m} Y_{i,m}) & for \ J - 1 \ge j \ge 2\\ \infty & for \ j = J \end{cases}$$
(3)

The condition $y_i = j \iff \alpha_{j-1,i} \le h_i < \alpha_{j,i}$ can be easily expressed in terms of the probability, which leads to:

$$P(y_i = j) = P(\alpha_{j-1,i} \le h_i < \alpha_{j,i}), \tag{4}$$

hence

$$P(y_i = j) = \Phi(\alpha_{i, j} - h_i) - \Phi(\alpha_{i, j-1} - h_i), \tag{5}$$

where Φ is a distribution function (cdf, cumulative density function). For example, for probit regression it is standard normal cdf $\Phi(x) = \frac{1}{2} + \frac{1}{2} * erf\left(\frac{x}{\sqrt{2}}\right)$ whereas for logit regression it takes the form $\Phi(x) = \frac{1}{1+e^{-x}}$. In

reporting styles analyses the typical choice is the probit model. It simply assumes that h_i is affected by a random noise ϵ_i having standard normal distribution $\epsilon_i \sim \mathcal{N}(0,1)$.

Using all definitions presented above the log likelihood function can be constructed

$$\ln L = \sum_{i=1}^{N} \sum_{j=1}^{J} z_{i, j} \ln \left[\Phi(\alpha_{i, j} - h_{i}) - \Phi(\alpha_{i, j-1} - h_{i}) \right], \tag{6}$$

where $z_{i,j}$ is an indicator function defined as:

$$z_{i, j} = \begin{cases} 0 & \text{for } y_i = j\\ 1 & \text{for } y_i \neq j \end{cases}$$
 (7)

3. Analysis of reporting styles

The model estimates are used to determine reporting behavior, i.e in how the continuous latent variable is projected onto categorical response measure. Practically, it is done by comparing actual categorical ordered responses with theoretical ones that are adjusted for heterogeneity in reporting behaviors and are more comparable across individuals.

One of the first steps of the analysis is standardization of the latent variable to obtain latent index H_i .

$$H_i = 1 - \frac{h_i - \min_i h_i}{\max_i h_i - \min_i h_i} \tag{8}$$

In the self-rated health example H_i is a proxy for true underlying health of an individual, and varies from 0 representing the (model-based) worst health state to 1 representing the (model-based) best health in the sample.

The predicted latent variable h_i obtained from the model is also used to standardize latent variable coefficients. In the self-rated health example the standardized coefficients are called disability weights D_k (Jürges 2007) and are calculated for each health variable to provide information about the impact of a specific health measure on the latent index. The disability weight for a health variable is equal to the ratio of corresponding health coefficient and the difference between the lowest and highest values of predicted latent health.

$$D_k = \frac{\beta_k}{\max_i h_i - \min_i h_i} \tag{9}$$

While the latent index H_i is intend to reflect underlying health, happiness or other status across individuals, the standardized coefficients D_k , like disability weights, are computed for an average individual in the study population. The relation between H_i and D_k follows the equation:

$$H_i = C - \sum_{k=1}^K D_k X_{i,k}, \quad \text{where } C = \frac{\max_i h_i}{\max_i h_i - \min_i h_i}$$
 (10)

Reporting styles analysis is based on the reclassification of individuals into new response categories. The classification is based on calculated latent index H_i and is thus adjusted for inter-individual differences in reporting behavior. There are two methods of reclassification: (1) Jürges (2007) percentile method (see also Rebelo and Pereira 2014) and (2) reclassification based on estimated thresholds.

The Jürges' percentile method is based on on original distribution of categorical response variable. First for each category j an empirical distribution function is constructed.

$$\hat{F}(j) = \frac{1}{N} \sum_{i=1}^{N} \mathbf{1}_{y_i} \le j \tag{11}$$

Where 1 is indicator function taking 1 if the condition is true or 0 otherwise. The calculated cumulative frequencies of latent index H_i are used as percentiles (cut points), so each individual i can be reclassified into new response categories.

In the second case the reclassification is based on eq. (2), so each individual has its own, model-derived cut-points.

4. Installing and loading the package

The newest available version of the package is always available from GitHub. It can be installed using devtools package

```
library(devtools)
install_github("maciejdanko/hopit")
library(hopit)
```

In examples presented below we will use artificially generated data set.

```
data(healthsurvey)
head(healthsurvey)
```

```
##
     ID
            health diabetes obese IADL_problems hypertenssion high_cholesterol
##
  1
      1 Very good
                          no
                                 no
                                                 no
##
   2
      2
              Good
                                                               yes
                                                                                  yes
                          no
                                 no
                                                 no
## 3
      3
              Good
                         yes
                                 no
                                                 no
                                                                no
                                                                                   no
## 4
      4
              Good
                          no
                                 no
                                                 no
                                                                no
                                                                                   no
## 5
      5 Excellent
                          no
                                 no
                                                 no
                                                                no
                                                                                   no
##
   6
      6
              Good
                          no
                                 no
                                                 no
                                                               yes
                                                                                  yes
                                                     poor_mobility
##
     respiratory_problems heart_atack_or_stroke
                                                                     very_poor_grip
## 1
                         no
                                                  no
                                                                 no
## 2
                                                 yes
                                                                 no
## 3
                         no
                                                  no
                                                                 no
                                                                                  no
## 4
                         no
                                                  no
                                                                 no
                                                                                  no
## 5
                         no
                                                  no
                                                                 no
                                                                                  nο
## 6
                        yes
                                                  no
                                                                yes
                                                                                  no
##
     depression other_diseases
                                    sex ageclass education contHM country
## 1
                                    man [80,120)
                                                                 3.6
              no
                              yes
                                                       prim-
                                                                             Y
                                          [70,80)
                                                                 4.4
                                                                             Y
## 2
                                                       prim-
              no
                              yes
                                    man
## 3
                                          [50,60)
                                                                 4.5
                                                                             Х
              no
                               no
                                    man
                                                       prim-
                                          [60,70)
                                                                 5.1
                                                                             Y
## 4
                                                        sec+
             yes
                               no
                                    man
## 5
                               no woman [80,120)
                                                                 3.3
                                                                             Z
                                                       prim-
              no
                                                                             Y
                                    man [80,120)
## 6
                                                       prim-
                                                                 5.1
                              yes
##
          csw psu
                     ssu
## 1 2407.48
               YB
                    <NA>
## 2 1198.12
               YΒ
                    <NA>
      885.26
               XC
                  XCgis
## 4
      772.04
               YΑ
                    <NA>
## 5 1304.24
               ZB
                    <NA>
## 6
     917.16
               YD
                    <NA>
```

5. Fitting the model using the *hopit()* function

Generalized ordered probit model can be fitted using hopit() function. The function takes two kinds of formulas: (1) latent.formula that models the latent variable and (2) *thresh.formula that models thresholds.

```
model1<- hopit(latent.formula = health ~ hypertenssion + high_cholesterol +</pre>
                             heart_atack_or_stroke + poor_mobility + very_poor_grip +
                             depression + respiratory problems +
                             IADL_problems + obese + diabetes + other_diseases,
               thresh.formula = ~ sex + ageclass,
               decreasing.levels = TRUE,
               control=list(trace=FALSE),
               data = healthsurvey)
summary(model1)
## Formula (latent variables):
## health ~ hypertenssion + high_cholesterol + heart_atack_or_stroke +
##
       poor_mobility + very_poor_grip + depression + respiratory_problems +
       IADL_problems + obese + diabetes + other_diseases
##
## Formula (threshold variables): ~sex + ageclass
## Link: probit
## Number of cases: 10000
## Response levels: Excellent, Very good, Good, Fair, Poor
## Robust SE were used (sandwich estimator of varcov).
##
                            Estimate Std. Error z value Pr(>|z|)
##
                                                   7.76 8.4e-15 ***
## hypertenssionyes
                             0.19232
                                        0.02478
## high_cholesterolyes
                             0.09780
                                        0.02918
                                                   3.35
                                                          0.00080 ***
## heart_atack_or_strokeyes
                                                  10.81
                                                         < 2e-16 ***
                             0.34401
                                        0.03183
## poor_mobilityyes
                             0.72832
                                        0.03564
                                                  20.44 < 2e-16 ***
                             0.49720
                                        0.12299
                                                   4.04 5.3e-05 ***
## very_poor_gripyes
## depressionyes
                             0.25323
                                        0.02390
                                                  10.59
                                                          < 2e-16 ***
                                                  11.02 < 2e-16 ***
## respiratory_problemsyes
                             0.36777
                                        0.03337
## IADL_problemsyes
                                        0.03637
                                                  16.93 < 2e-16 ***
                             0.61579
## obeseyes
                             0.18991
                                        0.03295
                                                   5.76
                                                          8.3e-09 ***
## diabetesyes
                             0.33726
                                        0.04010
                                                   8.41 < 2e-16 ***
## other diseasesyes
                             0.33533
                                        0.02370
                                                  14.15 < 2e-16 ***
## (L).1|2
                            -0.09248
                                        0.03194
                                                  -2.90 0.00379 **
## (L).2|3
                            -0.26826
                                        0.03236
                                                   -8.29
                                                         < 2e-16 ***
## (L).3|4
                                                   2.59 0.00968 **
                             0.07514
                                        0.02905
## (L).4|5
                            -0.20346
                                        0.05222
                                                  -3.90 9.8e-05 ***
## (G).sexwoman.1|2
                             0.02373
                                        0.03015
                                                   0.79
                                                          0.43112
## (G).sexwoman.2|3
                                        0.03460
                                                   0.39
                             0.01366
                                                          0.69304
## (G).sexwoman.3|4
                                        0.02869
                                                   1.28 0.20192
                             0.03661
## (G).sexwoman.4|5
                             0.11848
                                        0.05039
                                                   2.35
                                                          0.01872 *
## (G).ageclass[60,70).1|2
                            -0.01835
                                        0.03383
                                                  -0.54
                                                          0.58763
## (G).ageclass[60,70).2|3
                             0.05336
                                        0.04068
                                                   1.31
                                                          0.18962
## (G).ageclass[60,70).3|4
                             0.06003
                                        0.03616
                                                   1.66 0.09693
## (G).ageclass[60,70).4|5
                             0.16842
                                        0.06492
                                                   2.59
                                                          0.00949 **
## (G).ageclass[70,80).1|2
                            -0.32157
                                        0.04391
                                                  -7.32
                                                          2.4e-13 ***
## (G).ageclass[70,80).2|3
                             0.17131
                                        0.04774
                                                   3.59
                                                          0.00033 ***
## (G).ageclass[70,80).3|4
                             0.19360
                                        0.03777
                                                   5.13 3.0e-07 ***
```

0.06653

3.49 0.00048 ***

0.23234

(G).ageclass[70,80).4|5

```
## (G).ageclass[80,120).1|2 -0.33134
                                         0.07274
                                                    -4.56 5.2e-06 ***
## (G).ageclass[80,120).2|3 0.14976
                                         0.07590
                                                     1.97 0.04848 *
## (G).ageclass[80,120).3|4 0.17851
                                                     3.55 0.00038 ***
                                         0.05025
## (G).ageclass[80,120).4|5 0.22378
                                                     2.92 0.00354 **
                                         0.07674
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Theta: 1
## Log-likelihood: -12945.98
## Deviance: 25891.96
## AIC: 25953.96
model1 contains 11 dichotomous health variables and two threshold variables. The fitted coefficient can be
accessed by coef() function
# extract parameters in a form of list
cm1 <- coef(model1, aslist = TRUE)</pre>
# types of returned coefficients
names(cm1)
## [1] "latent.params" "thresh.lambda" "thresh.gamma"
                                                         "logTheta"
# latent health variables
cm1$latent.params
##
           hypertenssionyes
                                  high_cholesterolyes heart_atack_or_strokeyes
##
                  0.1923166
                                             0.0978032
                                                                       0.3440052
##
           poor_mobilityyes
                                                                  depressionyes
                                    very_poor_gripyes
                  0.7283236
                                             0.4972017
                                                                       0.2532285
##
##
    respiratory_problemsyes
                                     IADL_problemsyes
                                                                        obeseves
                                                                       0.1899097
##
                  0.3677676
                                             0.6157910
                diabetesyes
                                    other_diseasesyes
##
##
                  0.3372606
                                             0.3353300
model1 can be further extended by adding country of origin to the treshold formula to control for cultural
differences.
model2<- hopit(latent.formula = health ~ hypertenssion + high_cholesterol +</pre>
                              heart_atack_or_stroke + poor_mobility +
                              very_poor_grip + depression + respiratory_problems +
                              IADL_problems + obese + diabetes + other_diseases,
               thresh.formula = ~ sex + ageclass + country,
               decreasing.levels = TRUE,
               control=list(trace=FALSE),
               data = healthsurvey)
The fit of both models can be compared using AIC() function:
AIC(model2, model1)
##
     model2
              model1
## 25154.19 25953.96
or using Likelihood Ratio Test (LRT) as models are nested:
anova(model2, model1)
## Full model:
## -- Formula (latent variables):
```

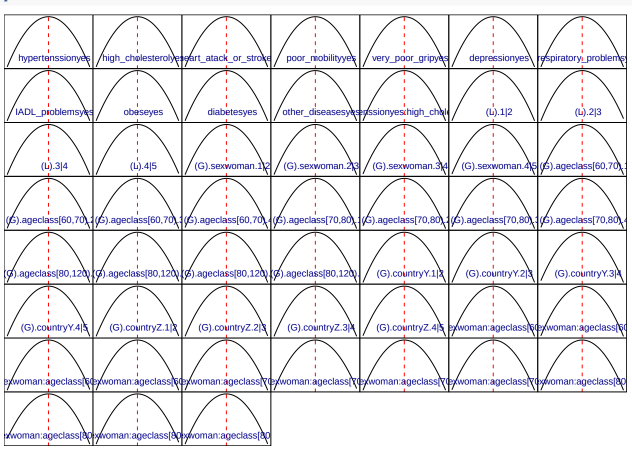
```
## health ~ hypertenssion + high_cholesterol + heart_atack_or_stroke +
       poor_mobility + very_poor_grip + depression + respiratory_problems +
##
       IADL problems + obese + diabetes + other diseases
## -- Formula (threshold variables): ~sex + ageclass + country
## -- Theta: FALSE
##
## Nested model:
## -- Formula (latent variables):
## health ~ hypertenssion + high_cholesterol + heart_atack_or_stroke +
       poor_mobility + very_poor_grip + depression + respiratory_problems +
       IADL_problems + obese + diabetes + other_diseases
## -- Formula (threshold variables): ~sex + ageclass
## -- Theta: FALSE
##
## Likelihood ratio test:
   Chi^2 df Pr(>Chi^2)
## 815.78 8
                  <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Both latent formula and thresh formula allow to specify interactions, like interaction between gender (sex)
and age (ageclass):
model3<- hopit(latent.formula = health ~ hypertenssion * high_cholesterol +</pre>
                             heart_atack_or_stroke + poor_mobility +
                             very_poor_grip + depression + respiratory_problems +
                             IADL_problems + obese + diabetes + other_diseases,
               thresh.formula = ~ sex * ageclass + country,
               decreasing.levels = TRUE,
               control=list(trace=FALSE),
               data = healthsurvey)
print(anova(model3,model2), short=TRUE)
##
## Likelihood ratio test:
   Chi^2 df Pr(>Chi^2)
## 28.923 13
                 0.00671 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
The hopit() function has also an option to include survey design using the survey package. The example
below fit a model using simple two level cluster sampling design
design <- svydesign(ids = ~ country + psu, weights = healthsurvey$csw, data = healthsurvey)
model2s<- hopit(latent.formula = health ~ hypertenssion + high_cholesterol +</pre>
                              heart atack or stroke + poor mobility +
                              very_poor_grip + depression + respiratory_problems +
                              IADL problems + obese + diabetes + other diseases,
                thresh.formula = ~ sex + ageclass + country,
                decreasing.levels = TRUE,
                design = design,
                control=list(trace=FALSE),
                data = healthsurvey)
```

Ignoring survey design could lead to biased results. Here, in presented examples it has however an minor importance, which is seen by comparing coefficients of latent variable for both models:

##	No survey design Has	survey design
## hypertenssionyes	0.18475332	0.18777955
## high_cholesterolyes	0.08972562	0.09368829
## heart_atack_or_strokeyes	0.34659838	0.34676962
## poor_mobilityyes	0.70346456	0.70603471
## very_poor_gripyes	0.51424418	0.54768793
## depressionyes	0.24998274	0.24922297
<pre>## respiratory_problemsyes</pre>	0.37863461	0.37984848
## IADL_problemsyes	0.59262343	0.60999389
## obeseyes	0.19041874	0.18874130
## diabetesyes	0.32839067	0.32328477
## other_diseasesyes	0.32936970	0.32876106

The fit accuracy of the model can be assessed using profile() function, which calculate and plot profile of the log likelihood function around fitted coefficient values.

profile(model3)



All parameters seem to be at arg.max (at optimum).

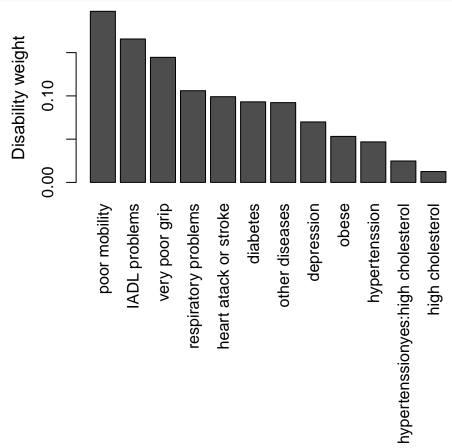
6. Analyses of the reporting styles using hopit package

Let's look at latent health variables of model2.

model3\$coef.ls\$latent.params ## hypertenssionyes high_cholesterolyes ## 0.16692322 0.04478911 ## heart_atack_or_strokeyes poor_mobilityyes ## 0.35296700 0.70450393 ## very_poor_gripyes depressionyes ## 0.51500758 0.24905559 ## respiratory_problemsyes IADL_problemsyes 0.37763441 0.59062305 ## ## obeseves diabetesyes 0.18939406 ## 0.33180729 ## other_diseasesyes hypertenssionyes:high_cholesterolyes ## 0.32861717 0.08822596

We can standardize them using Jürges' approach (Jürges 2007) to obtain disability weights. The standardization can be doen using standardizeCoef() function.

```
# A function that modifies coefficient names.
txtfun <- function(x) gsub('_',' ',substr(x,1,nchar(x)-3))
# Calcualte and plot disability weights
sc <- standardizeCoef(model3, plotf = TRUE, namesf = txtfun)</pre>
```

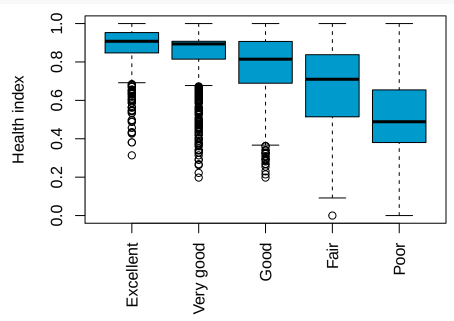


sc

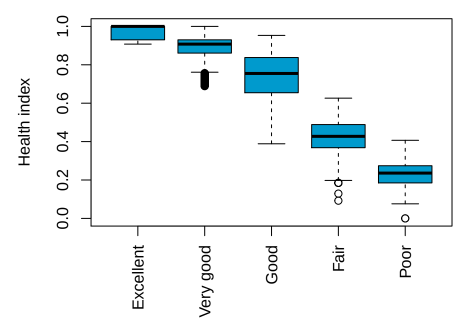
```
##
                                             [,1]
                                      0.19778802
## poor mobility
## IADL problems
                                      0.16581620
## very poor grip
                                      0.14458732
## respiratory problems
                                      0.10602008
## heart atack or stroke
                                      0.09909476
## diabetes
                                      0.09315421
## other diseases
                                      0.09225859
## depression
                                      0.06992184
## obese
                                      0.05317199
## hypertenssion
                                      0.04686335
## hypertenssionyes:high cholesterol 0.02476926
## high cholesterol
                                      0.01257445
```

The *namesf* argument is a function or a character vector that is used to rename the coefficent names. Here, it removes last 3 letters ("yes"), which is a reference level for each variable and exchanges "__" with spaces in variable names.

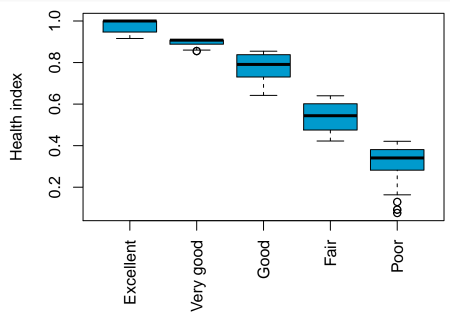
The latent index is simply calculated using *latentindex()* function.



The boxplot shows reported health status vs. health index. It is also possible to plot expected categorical health status on Y axis calcualted according to the eq. (2).

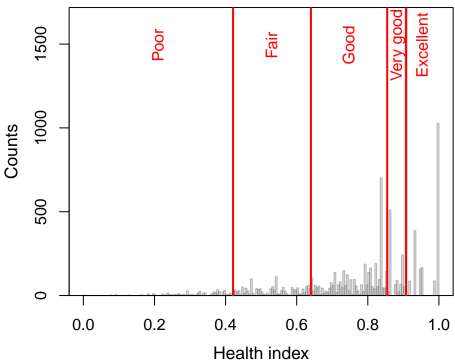


or according to Jürges' method:

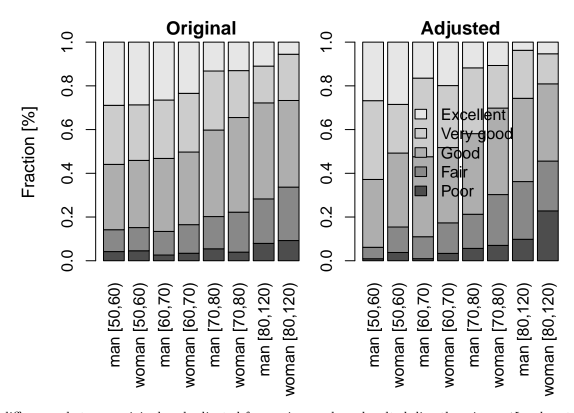


The central part of reporting styles analyses is to determine the cut-points used to calcualte adjusted health status for each individual. The calcualtion and ploting of cut-points is realized by getCutPoints() function.

z=getCutPoints(model=model3)



```
# Health index cut-points
z$cutpoints
##
       4.41%
                17.68%
                           52.34%
                                     77.63%
## 0.4210433 0.6400315 0.8545694 0.9077414
#Adjusted health levels for individuals: Jurges method
table(z$adjused.health.levels)
##
##
        Poor
                  Fair
                             Good Very good Excellent
##
         450
                  1347
                             3505
                                       2787
                                                  1909
#Adjusted health levels for individuals: Estimated model thresholds
table(model3$Ey_i)
##
## Excellent Very good
                             Good
                                       Fair
                                                  Poor
         737
                  4361
                             4029
                                         807
                                                    66
#Original health levels for individuals
table(model3$y_i)
##
## Excellent Very good
                                                  Poor
                             Good
                                       Fair
        2237
                  2529
                             3466
                                       1327
                                                   441
The analysis of health levels is done by getLevels() function
# Health levels for combination of age and gender, and pooled country of origin.
hl <- getLevels(model=model3, formula=~ sex + ageclass, data = healthsurvey,
                       sep=' ', plotf=TRUE, legbty = 'n')
```

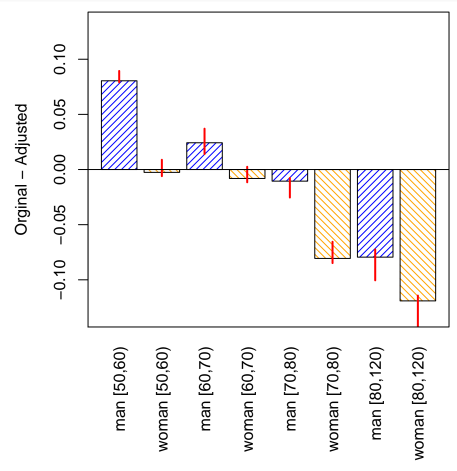


The differences between original and adjusted frequencies can be calcualted directly using getLevels output:

```
round(100*(hl$original - hl$adjusted),2)
##
##
                                      Good Very good Excellent
                       Poor
                               Fair
                                     -1.15
##
     man [50,60)
                       3.28
                               4.77
                                                -9.02
                                                            2.13
##
     woman [50,60)
                       0.77 - 1.03
                                     -3.09
                                                 3.13
                                                            0.21
##
     man [60,70)
                       1.66
                               0.77
                                     -3.19
                                                -9.26
                                                           10.03
##
     woman [60,70)
                       0.06 -0.87
                                     -1.25
                                                            3.55
                                                -1.50
     man [70,80)
                      -0.19
                             -0.86
##
                                      2.68
                                                -3.07
                                                            1.44
##
     woman [70,80)
                      -3.09 - 4.97
                                      3.75
                                                 1.97
                                                            2.34
##
     man [80,120)
                      -1.87
                             -6.07
                                      5.84
                                                -5.14
                                                            7.24
##
     woman [80,120) -13.55
                               1.64
                                      4.31
                                                 7.39
                                                            0.21
```

7. Bootstraping Confidence Intervals

The package offers functions to calculate confidence intervals for any measure derived from the model. As an example we show calculation of confidence intervals of the difference between original and adjusted frequencies of combined "Poor" + "Fair" healt categories.



The results show that men tend to over-report bad health at ages (50,60] and (50,70], wherease women at ages [70,80) and both sexes at ages (80, 120] under-report bad health.

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