Evaluate a set of hypotheses with GORIC or GORICA: ANOVA Example

Rebecca M. Kuiper

15 August 2024

Contents

Example 1: Lucas Data	1
Data preparation	1
Preparation for $GORIC(A)$	
GORIC examples	
GORICA examples	
Example 2: Berzonsky et al.	7
Data preparation	7
Preparation for GORIC(A)	
GORIC examples	
GORICA examples	
Example 3: Holubar	10
Data preparation	10
Preparation for $GORIC(A)$	11
Model selection using GORIC	
Example 4: Sesame	11
Calculate GORIC values and weights	12
Calculate GORIC values and weights for H1 and its complement	
Calculate GORIC values and weights for H2 and its complement	
Notes on weights	14
Note 1: GORIC weights versus GORICA weights	14
Note 2: complement in case of two hypotheses of interest	
Note 3: weights when using complement not per se higher	
Extra: Two methods to calculate the penalty	18
Note on not full row-rank	19

Example 1: Lucas Data

Data preparation

First, load the required libraries (after they have been installed). These libraries contain functions, such as goric, that will be used in the R code below. Each time you reopen R, you will have to load the required libraries.

```
## First, install the packages, if you have not done this already:
if (!require("psych")) install.packages("psych")
if (!require("restriktor")) install.packages("restriktor")

## Then, load the packages:
library(psych) # for the function describeBy
library(restriktor) # for the goric function

# If you want to use restriktor from github:
#if (!require("devtools")) install.packages("devtools")
#library(devtools)
#install_github("LeonardV/restriktor")
#library(restriktor) # for goric function
```

Second, it is necessary to load the data.

Notably, it is only possible to load the data if you are using the correct working directory (with both your R script and data file). The command getwd() shows you your current working directory. You can change the working directory to the one you prefer using the function setwd() by specifying the correct location between parentheses. Alternatively, in Rstudio, you can use the "Session" tab (on top) or you can use the "Files"-pane (on top of probably the right lower box of your Rstudio-screen, this pane is located next to the panes for "Plots", "Packages", "Help", and "Viewer").

If you open the data file Data_Lucas.txt in a text editor, you can see that the variable labels have been inserted (using quotes; i.e., "...") in the first line of the file, which is called a header. Therefore, you have to specify 'header = TRUE' when loading the data:

```
# Load the data
Lucas <- read.table("data/Data_Lucas.txt", header = TRUE)</pre>
```

Since a .txt file was loaded, R does not know the measurement levels of the variables and assumes all of them to be continuous, meaning that they are of interval or ratio type. Hence, especially when there are more than two groups, one has to tell R that the variable group is a factor by using the factor() function on the group variable (i.e., a grouping / categorical / nominal variable):

```
# Make the variable group a factor
Lucas$group <- factor(Lucas$group)</pre>
```

To inspect the first 6 rows of the dataset, use the head() function:

```
head(Lucas) # Look at first (6) rows of the data
```

```
group Influence
       1
               3.58
2
       1
              -0.15
3
               0.67
      1
4
               2.22
       1
               2.56
5
       1
       1
               1.70
```

To see a more detailed overview of the data via descriptive statistics split by group variable, use the describeBy() function with Lucas\$group set to be a grouping variable, as follows:

```
descrstat <- describeBy(Lucas$Influence, Lucas$group, mat = TRUE, digits = 3)
descrstat</pre>
```

```
item group1 vars n mean
                                 sd median trimmed
                                                                            skew kurtosis
                                                    mad
                                                           min max range
X11
                  1 30 2.329 1.860
                                    2.330
                                            2.243 2.009 -0.45 6.74
                                                                   7.19
                                                                          0.371
                                                                                   -0.719 0.340
                   1 30 1.328 1.149 1.320
X12
       2
              2
                                            1.272 1.231 -0.38 3.87 4.25
                                                                          0.318
                                                                                   -0.943 0.210
```

```
X13
       3
                   1 30 3.200 1.790 3.475
                                             3.243 1.824 -0.31 6.84 7.15 -0.149
                                                                                   -0.556 0.327
X14
       4
              4
                   1 30 2.231 1.450
                                    1.665
                                             2.099 1.171 0.47 5.12 4.65 0.724
                                                                                   -0.867 0.265
X15
      5
                   1 30 3.229 1.500 3.730
                                             3.358 1.460 -0.46 5.67 6.13 -0.630
                                                                                   -0.563 0.274
```

Preparation for GORIC(A)

ANOVA model

First, an R-object with unconstrained estimates is needed, that is, in this example, the five group means and one residual variance. The linear regression model using lm() function is specified as follows:

```
lm_fit_Lucas <- lm(Influence ~ group - 1, data = Lucas)</pre>
```

Note that:

- 1. y ~ group 1 instructs the function 1m (linear model) to regress the variable y on the variable group.
- 2. The 1 instructs the function 1m to drop the intercept and, therefore, estimate the means of each group, resulting, here, in five group means.

If the intercept is not dropped, 'y \sim group' would estimate an intercept – representing the mean of the reference group – and the mean differences between the other (here, four) groups and the reference group.

3. The results are collected in, what is called, an R-object; here, named lm_fit_Lucas.

It can be helpful to check the names used in this model, because these are needed when specifying the hypotheses:

```
names(coef(lm_fit_Lucas))
```

```
[1] "group1" "group2" "group3" "group4" "group5"
```

Note: Im object or coef() and vcov() When calculating the GORIC, the goric() function can use Im objects as input.

When calculating the GORICA, the goric() function can use lm or glm objects (even most lavaan objects) as input, but one also enter the (structural) parameter estimates and their covariance matrix. For most R-objects, the latter can be obtained using coef() and vcov(); e.g.:

```
est <- coef(lm_fit_Lucas)
VCOV <- vcov(lm_fit_Lucas)</pre>
```

This will be illustrated in an GORICA example below.

Set of hypotheses

To evaluate the hypotheses of interest, it is necessary to specify the restrictions in these hypotheses correctly:

- Within the restriktor() and goric() functions, it is possible to use the following operators: >, <, =, <=, >=, == (where the last three denote the same constraint as the first three).
- The goric() and the restriktor() functions can deal with:
 - pairwise restrictions separated by a semicolon; (e.g., "beta1 > beta2; beta2 > beta3") or an & sign (e.g., "beta1 > beta2 & beta2 > beta3").
 - combined restrictions consisting of more than one operator (e.g., "beta1 > beta2 > beta3").

Note that one should use the labels of the parameter estimates (in the example above: group1 – group5).

• One can also define hypothesis in terms of linear functions of parameters. For more details, see 'Extra possibility specification hypotheses' near the end of the goric tutorial called 'Tutorial_GORIC_restriktor_General' (https://github.com/rebeccakuiper/Tutorials).

Let us specify the following theory-based hypotheses:

```
H1 <- 'group5 = group3 > group1 > group2; group3 > group4 > group2'

# Note: H1 is not full row-rank (see below and the goric() tutorial for more details).

H2 <- 'group3 > group1 > group4 = group5 > group2'
```

To prevent from selecting a weak hypothesis, that is, a hypothesis not supported by the data, one should include a failsafe/safeguard hypothesis. This can be:

- the unconstrained hypothesis (which includes all possible hypotheses, thus including the one(s) of interest);
- the complement (which includes all other possible hypotheses, thus excluding the one(s) of interest),

where the first option is the default. Notably, currently, the complement can only be used for one hypothesis of interest. Therefore, the examples w.r.t the use of the complement only evaluate H1 (and not the whole set).

Seed values

In the calculation of the GORIC(A), an iterative process is used to calculate the penalty / complexity part. Therefore, one needs to set a seed value using set.seed(). This has two advantages:

- 1. Using the same seed value leads to the same penalty value every time this code is run. This is thus helpful to reproduce results.
- 2. Using different seed values, allows for a sensitivity check on the penalty value. If it is sensitive, then increase number of iterations used in calculation of the penalty (see below).

GORIC examples

The GORIC can be used for normal linear models, like ANOVA and regression models. The goric() function calculates the *GORIC* value by default (type = "goric"). To calculate the *GORICA* values, the argument type has to be set to gorica (type = "gorica", see example below).

Note: For (more) information regarding interpreting the GORIC(A) output, see 'Guidelines_output_GORIC' (https://github.com/rebeccakuiper/Tutorials).

Example 1.1a: Using the unconstrained as failsafe

```
set.seed(123) # Set seed value
output <- goric(lm_fit_Lucas, hypotheses = list(H1 = H1, H2 = H2))
#summary(output)
output</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	H1	-278.051	3.458	563.019	0.493	0.407	0.899
2	Н2	-281.761	3.136	569.794	0.012	0.561	0.030
3	unconstrained	-278.048	6.000	568.097	0.495	0.032	0.071

Ratio GORIC-weights:

	vs. H1	vs. H2	vs. unconstrained
H1	1.000	29.593	12.668
H2	0.034	1.000	0.428

```
unconstrained 0.079 2.336 1.000
round(output$ratio.gw, 2)
```

```
vs. H1 vs. H2 vs. unconstrained
H1 1.00 29.59 12.67
H2 0.03 1.00 0.43
unconstrained 0.08 2.34 1.00
```

It can be seen that the order-restricted hypothesis H_1 has > 1 times more support than H_u (the unconstrained hypothesis). Hence, H_1 is not a weak hypotheses and can thus be compared to the other (weak and non-weak) competing hypotheses:

 H_1 is 29.6 times more likely than H_2 .

Example 1.1b: Using the complement as failsafe

```
set.seed(123) # Set seed value
output_c <- goric(lm_fit_Lucas, hypotheses = list(H1), comparison = "complement")
#summary(output_c)
output_c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	H1	-278.051	3.458	563.019	0.499	0.903	0.903
2	complement	-278.048	5.691	567.479	0.501	0.097	0.097

The order-restricted hypothesis 'H1' has 9.30 times more support than its complement.

The order-restricted hypothesis H_1 has 9.3 times more support than its complement.

GORICA examples

The *GORICA* can be used for a broad range of models. Besides normal linear models (e.g., ANOVA and regression models) it can be applied also to logistic regression and SEM models and much more. To calculate the *GORICA* values, one should use type = "gorica".

Example 1.2a: Using the unconstrained as failsafe

```
set.seed(123) # Set seed value
output_gorica <- goric(lm_fit_Lucas, hypotheses = list(H1 = H1, H2 = H2), type = "gorica")
#summary(output_gorica)
output_gorica</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

	model	loglik	penalty	gorica	loglik.weights	penalty.weights	gorica.weights
1	H1	1.647	2.458	1.622	0.493	0.407	0.898
2	H2	-2.029	2.136	8.330	0.012	0.561	0.031
3	unconstrained	1.650	5.000	6.700	0.494	0.032	0.071

```
Ratio GORICA-weights:
```

```
vs. H1 vs. H2 vs. unconstrained
```

```
H1 1.000 28.622 12.669
H2 0.035 1.000 0.443
unconstrained 0.079 2.259 1.000
```

round(output_gorica\$ratio.gw, 2)

```
vs. H1 vs. H2 vs. unconstrained H1 1.00 28.62 12.67 H2 0.03 1.00 0.44 unconstrained 0.08 2.26 1.00
```

In Example 1, the same analysis is done with the GORIC, you can see that the (relative) weights are (about) the same for the GORIC and GORICA. The ratio of weights may differ (somewhat see a note at the end of this tutorial).

From this output, it can be seen that the order-restricted hypothesis H_1 has > 1 times more support than H_u (the unconstrained hypothesis). Hence, H_1 is not a weak hypotheses and can thus be compared to the other (weak and non-weak) competing hypotheses:

 H_1 is 28.6 times more likely than H_2 .

```
set.seed(123) # Set seed value
est <- coef(lm_fit_Lucas)
VCOV <- vcov(lm_fit_Lucas)
output_gorica_alt <- goric(est, VCOV = VCOV, hypotheses = list(H1 = H1, H2 = H2), type = "gorica", mix_"
#summary(output_gorica_alt)
output_gorica_alt</pre>
```

Alternative input

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

	model	loglik	penalty	gorica	loglik.weights	penalty.weights	gorica.weights
1	H1	1.647	2.457	1.619	0.493	0.406	0.898
2	Н2	-2.029	2.131	8.320	0.012	0.562	0.031
3	${\tt unconstrained}$	1.650	5.000	6.700	0.494	0.032	0.071

Ratio GORICA-weights:

	vs. H1	vs. H2	vs. unconstrained
H1	1.000	28.516	12.686
H2	0.035	1.000	0.445
unconstrained	0.079	2.248	1.000

round(output_gorica_alt\$ratio.gw, 2)

	vs.	H1	vs.	H2	vs.	${\tt unconstrained}$
H1	1	.00	28	. 52		12.69
H2	0	.04	1	.00		0.44
unconstrained	0	.08	2	. 25		1.00

Example 1.2b: Using the complement as failsafe

```
H1 <- 'group5 = group3 > group1 > group2; group3 > group4 > group2'
# Note: H1 is not full row-rank;
# for more details, see below and/or the goric tutorial.
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

```
gorica loglik.weights penalty.weights gorica.weights
        model
               loglik penalty
           H1
                1.647
                         2.458
                                 1.622
                                                  0.499
                                                                   0.903
                                                                                   0.903
1
                1.650
  complement
                         4.691
                                 6.082
                                                  0.501
                                                                   0.097
                                                                                   0.097
```

The order-restricted hypothesis 'H1' has 9.30 times more support than its complement.

The order-restricted hypothesis H_1 has 9.3 times more support than its complement. Notably, the weights are also now the same as for the GORIC (Example 2).

Example 2: Berzonsky et al.

Next, the R code to apply the GORIC(A) to the data of Berzonsky et al. is shown. The instruction on loading and preparing the data are similar to the ones from Lucas example (Example 1). Hence, for more comments and details, see that example.

Data preparation

```
# Read Data.
BerzEtAl <- read.table("data/Data_BerzEtAl.txt", header = TRUE)</pre>
BerzEtAl$group <- factor(BerzEtAl$group)</pre>
# this command tells R that group is a factor and not a continuous variable
# Inspect data
head(BerzEtAl)
  group Influence
        39.79709
1
      1
         26.58804
2
3
      1 36.04999
4
      1 35.92915
      1 27.02636
5
      1 31.08900
# Compute descriptive statistics for each group
descrip <- describeBy(BerzEtAl$Influence, BerzEtAl$group, mat = TRUE, digits = 3)</pre>
descrip
```

```
item group1 vars n mean
                                sd median trimmed
                                                           min
                                                                  max range
                                                                               skew kurtosis
                                                    mad
X11
      1
             1
                  1 15 32.00 4.461 33.508 31.934 3.768 25.055 39.797 14.742 -0.114
                                                                                      -1.353 1.152
X12
      2
                  1 15 21.00 4.627 21.426 20.976 3.795 13.179 29.138 15.959 -0.081
                                                                                      -1.007 1.195
X13
      3
             3
                       7.00 4.192 6.364
                                            6.853 3.498
                                                         1.456 14.453 12.996
                                                                                      -1.100 1.082
                  1 15
                                                                             0.440
X14
      4
             4
                  1 15 14.00 6.242 14.708
                                           14.259 7.264
                                                         0.906 23.724 22.818 -0.349
                                                                                      -0.809 1.612
X15
      5
             5
                  1 15 14.00 4.368 14.068
                                           14.195 3.532 5.194 20.266 15.072 -0.465
                                                                                      -0.746 1.128
X16
                  1 15 0.01 0.010 0.011
                                            0.009 0.010 -0.004 0.038 0.042 1.081
                                                                                       1.119 0.003
X17
                  1 15 0.10 0.093 0.083
                                           0.096 0.129 -0.026 0.284 0.310 0.179
                                                                                      -1.232 0.024
```

Preparation for GORIC(A)

```
# Using the R package lm
lm fit BerzEtAl <- lm(Influence ~ group-1, data=BerzEtAl)</pre>
# Check names used in model
names(coef(lm_fit_BerzEtAl))
[1] "group1" "group2" "group3" "group4" "group5" "group6" "group7" "group8"
# Specify restrictions using those names
# Set of hypotheses
H1 <- 'group1 > group2; group1 > group3; group1 > group4;
group5 > group6; group5 > group7; group5 > group8;
group1 > group5; group2 > group6; group3 > group7; group4 > group8;
group1 - group5 > group2 - group6;
group1 - group5 > group3 - group7;
group1 - group5 > group4 - group8;'
H2 <- 'group1 > group2; group1 > group3; group1 > group4;
group5 > group6; group5 > group7; group5 > group8;
group1 > group5; group2 > group6; group3 > group7; group4 > group8;
group1 - group5 > 2*(group2 - group6);
group1 - group5 > 2*(group3 - group7);
group1 - group5 > 2*(group4 - group8);'
GORIC examples
Example 2.1a: Using the unconstrained as failsafe
set.seed(123) # Set seed value
output B <- goric(lm fit BerzEtAl, hypotheses = list(H1 = H1, H2 = H2), mix weights = "boot")
#summary(output_B)
output B
restriktor (0.5-80): generalized order-restricted information criterion:
Results:
          model
                   loglik penalty
                                       goric loglik.weights penalty.weights goric.weights
                              6.084 668.772
                                                       0.229
                                                                                       0.846
1
              H1 -328.302
                                                                        0.489
```

```
Ratio GORIC-weights:
```

H2 -361.183

vs. H1 vs. H2 vs. unconstrained 5.48e+00 Н1 1.00e+00 1.92e+14 0.00e+00 1.00e+00 H2 0.00e+00 unconstrained 1.83e-01 3.51e+13 1.00e+00

round(output_B\$ratio.gw, 2)

3 unconstrained -327.087

2

vs. H1 vs. H2 vs. unconstrained

6.091 734.550

9.000 672.173

0.000

0.771

0.485

0.026

0.000

0.154

```
H1 1.00 1.920420e+14 5.48

H2 0.00 1.000000e+00 0.00

unconstrained 0.18 3.506845e+13 1.00
```

The order-restricted hypothesis H_1 has > 1 times more support than H_u (unconstrained). Hence, H_1 is not a weak hypotheses and can thus be compared to the other (weak and non-weak) competing hypotheses: H_1 is much more supported than H_2 .

Example 2.1b: Using the complement as failsafe

```
set.seed(123)
output_B_c <- goric(lm_fit_BerzEtAl, hypotheses = list(H1), comparison = "complement", mix_weights = "b
#summary(output_B_c)
output_B_c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	H1	-328.302	6.084	668.772	0.229	0.942	0.827
2	complement	-327.087	8.863	671.900	0.771	0.058	0.173

The order-restricted hypothesis 'H1' has 4.78 times more support than its complement.

The order-restricted hypothesis H_1 has 4.8 times more support than its complement.

GORICA examples

Example 2.2a: Using the unconstrained as failsafe

```
set.seed(123) # Set seed value
output_B_gorica <- goric(lm_fit_BerzEtAl, hypotheses = list(H1 = H1, H2 = H2), type = "gorica", mix_wei,
#summary(output_B_gorica)
output_B_gorica</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

	model	loglik	penalty	gorica	loglik.weights	penalty.weights	gorica.weights
1	H1	-8.396	5.084	26.959	0.241	0.489	0.854
2	Н2	-50.102	5.091	110.388	0.000	0.485	0.000
3	${\tt unconstrained}$	-7.250	8.000	30.499	0.759	0.026	0.146

Ratio GORICA-weights:

```
      vs. H1
      vs. H2
      vs. unconstrained

      H1
      1.00e+00
      1.31e+18
      5.87e+00

      H2
      0.00e+00
      1.00e+00
      0.00e+00

      unconstrained
      1.70e-01
      2.23e+17
      1.00e+00
```

```
round(output_B_gorica$ratio.gw, 2)
```

	vs. H1	vs. H2 vs.	unconstrained
H1	1.00 1.	.306966e+18	5.87
H2	0.00 1.	.000000e+00	0.00
unconstrained	0.17 2.	226433e+17	1.00

The order-restricted hypothesis H_1 has > 1 times more support than H_u (unconstrained). Hence, H_1 is not a weak hypotheses and can thus be compared to the other (weak and non-weak) competing hypotheses: H_1 is much more supported than H_2 .

Example 2.2b: Using the unconstrained as failsafe

```
set.seed(123)
output_B_gorica_c <- goric(lm_fit_BerzEtAl, hypotheses = list(H1), comparison = "complement", type = "g
#summary(output_B_gorica_c)
output_B_gorica_c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

```
        model
        loglik
        penalty
        gorica
        loglik.weights
        penalty.weights
        gorica.weights

        1
        H1
        -8.396
        5.084
        26.959
        0.241
        0.942
        0.837

        2
        complement
        -7.250
        7.863
        30.226
        0.759
        0.058
        0.163
```

The order-restricted hypothesis 'H1' has 5.12 times more support than its complement.

The order-restricted hypothesis H_1 has 5.1 times more support than its complement.

Example 3: Holubar

Data preparation

First, read in the Holubar dataset, and tell R that the variable gr (group) is a factor instead of a continuous variable (although it is not necessary because it consists of only two groups).

```
Holubar <- read.table("data/Data_Holubar.txt", header = TRUE) # load the data
Holubar$gr <- factor(Holubar$gr) # tell R that gr is a factor
```

If you want a more detailed overview of the data, also by means of descriptive statistics splitted by group, use head(Holubar)

```
item group1 vars n mean
                                sd median trimmed
                                                    mad
                                                           min
                                                                 max range
                                                                              skew kurtosis
X11
                   1 20 0.98 1.20
                                   1.177
                                            0.930 1.118 -0.719 3.617 4.335
                                                                             0.190
                                                                                     -0.778 0.268
X12
              2
                   1 27 0.02 1.88 0.169
       2
                                            0.111 1.951 -4.961 2.921 7.883 -0.517
                                                                                     -0.104 0.362
X13
       3
                   1 28 0.27 1.72 -0.099
                                            0.203 2.435 -2.276 3.796 6.073 0.292
                                                                                     -1.025 0.325
```

Preparation for GORIC(A)

ANOVA model

Then, fit an ANOVA-model by means of the lm() function (linear model) and directly check the names that are used in this model:

```
lm_fit_Holubar <- lm(at ~ gr - 1, data = Holubar)
names(coef(lm_fit_Holubar))
[1] "gr1" "gr2" "gr3"</pre>
```

Set of hypotheses

The following hypothesis will be evaluated:

```
H1 <- 'gr2 > gr1 > gr3'
```

Model selection using GORIC

Calculate the GORIC values and weights:

```
set.seed(123) # Set seed value
output_Hol <- goric(lm_fit_Holubar, hypotheses = list(H1), comparison = "complement")
#summary(output_Hol)
output_Hol</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
        model
        loglik
        penalty
        goric
        loglik.weights
        penalty.weights
        goric.weights

        1
        H1
        -144.981
        2.803
        295.569
        0.125
        0.710
        0.259

        2
        complement
        -143.038
        3.697
        293.469
        0.875
        0.290
        0.741
```

The order-restricted hypothesis 'H1' has 0.35 times more support than its complement.

Example 4: Sesame

```
# read in the sesame data from a text file
Sesame <- read.table("data/sesamesimANOVA.txt", header=TRUE)

# make viewcat a factor, that is, a categorical variable
Sesame$viewcat <- factor(Sesame$viewcat)

# Inspect data
head(Sesame)</pre>
```

```
# Check names used in model
names(coef(fit_Sesame))

[1] "viewcat1" "viewcat2" "viewcat4"

# Specify restrictions using those names

# Example hypotheses:
H1 <- 'viewcat1 = viewcat2 < viewcat3 < viewcat4'
H2 <- 'viewcat1 < viewcat2 < viewcat3 < viewcat4'
H3 <- 'viewcat1 = viewcat2 < viewcat3 = viewcat4'</pre>
```

Calculate GORIC values and weights

Here, we assume that the interest lies in H1 to H3. Since these do not cover all possibilities, the unconstrained hypotheses is included in the set.

```
set.seed(123)
goric_sesam <- goric(fit_Sesame, hypotheses = list(H1, H2, H3))
#summary(goric_sesam)
goric_sesam</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	H1	-918.485	2.810	1842.590	0.065	0.310	0.134
2	H2	-916.540	3.094	1839.269	0.457	0.233	0.704
3	НЗ	-919.644	2.500	1844.287	0.021	0.422	0.057
4	${\tt unconstrained}$	-916.540	5.000	1843.080	0.457	0.035	0.105

Note: Hypotheses 'H2' and 'unconstrained' have equal likelihood values (i.e., the hypotheses overlap). 'round(goric_sesam\$ratio.gw, 2)

```
vs. H1 vs. H2 vs. H3 vs. unconstrained
H1
               1.00 0.19
                           2.34
                                              1.28
Н2
                      1.00 12.29
                                              6.72
               5.26
НЗ
               0.43
                      0.08
                            1.00
                                              0.55
               0.78
                                              1.00
unconstrained
                      0.15
                             1.83
```

From the output, it is concluded that H2 is not a weak hypothesis (nor is H1). Thus, its support can be compared to that of the other hypotheses:

H2 is 5.3 times more supported than H1 and 12.3 times more than H3.

Hence, H2 is the preferred hypothesis (and has quite some evidence).

Calculate GORICA values and weights

```
set.seed(123)
gorica_sesam <- goric(fit_Sesame, hypotheses = list(H1, H2, H3), type = "gorica")
#summary(gorica_sesam)
gorica_sesam</pre>
```

 ${\tt restriktor}\ (0.5\text{--}80)\colon {\tt generalized}\ {\tt order-restricted}\ {\tt information}\ {\tt criterion}\ {\tt approximation}\colon$

Results:

	model	loglik	penalty	gorica	loglik.weights	penalty.weights	gorica.weights
1	H1	-7.052	1.810	17.725	0.066	0.310	0.136
2	Н2	-5.125	2.094	14.438	0.456	0.233	0.702
3	НЗ	-8.216	1.500	19.432	0.021	0.422	0.058
4	${\tt unconstrained}$	-5.125	4.000	18.249	0.456	0.035	0.104

Note: Hypotheses 'H2' and 'unconstrained' have equal likelihood values (i.e., the hypotheses overlap).
round(gorica_sesam\$ratio.gw, 2)

```
vs. H1 vs. H2 vs. H3 vs. unconstrained
H1
                1.00
                       0.19
                              2.35
                                                 1.30
H2
                5.17
                       1.00 12.15
                                                 6.72
                0.43
                                                 0.55
НЗ
                       0.08
                             1.00
                0.77
                       0.15
                              1.81
                                                 1.00
unconstrained
```

Calculate GORIC values and weights for H1 and its complement

Here, we assume that the interest lies only in H1. In that case, it should be evaluated against its complement.

```
set.seed(123)
goric_sesam_1c <- goric(fit_Sesame, hypotheses = list(H1), comparison = "complement")
#summary(goric_sesam_1c)
goric_sesam_1c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
model
                loglik penalty
                                           loglik.weights penalty.weights goric.weights
                                     goric
              -918.485
                          2.810 1842.590
                                                    0.125
                                                                      0.868
                                                                                     0.484
          H1
1
                                                    0.875
                                                                                     0.516
  complement
               -916.540
                          4.690 1842.460
                                                                      0.132
```

The order-restricted hypothesis 'H1' has 0.94 times more support than its complement.

From the output, it is concluded that the support for H1 is comparable to that of its complement, with a slight preference for the complement. Note that the complement has the highest (log) likelihood but also the highest complexity/penalty, when balancing fit and complexity this results in a slight preference for the complement of H1.

Calculate GORICA values and weights for H1 and its complement

```
set.seed(123)
gorica_sesam_1c <- goric(fit_Sesame, hypotheses = list(H1), type = "gorica", comparison = "complement")
#summary(gorica_sesam_1c)
gorica_sesam_1c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

```
        model
        loglik
        penalty
        gorica
        loglik.weights
        penalty.weights
        gorica.weights

        1
        H1
        -7.052
        1.810
        17.725
        0.127
        0.868
        0.488

        2
        complement
        -5.125
        3.690
        17.629
        0.873
        0.132
        0.512
```

The order-restricted hypothesis 'H1' has 0.95 times more support than its complement.

Calculate GORIC values and weights for H2 and its complement

Here, we assume that the interest lies only in H2. In that case, it should be evaluated against its complement.

```
set.seed(123)
goric_sesam_2c <- goric(fit_Sesame, hypotheses = list(H2 = H2), comparison = "complement")
#summary(goric_sesam_2c)
goric_sesam_2c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
model
                                             loglik.weights penalty.weights
                                                                               goric.weights
                 loglik
                         penalty
                                      goric
           H2
               -916.540
                                                      0.765
                                                                        0.855
                                                                                       0.950
                           3.095
                                  1839.270
1
               -917.718
                           4.870
                                  1845.176
                                                      0.235
                                                                        0.145
                                                                                       0.050
  complement
```

The order-restricted hypothesis 'H2' has 19.17 times more support than its complement.

From the output, it is concluded that H1 is about 19 times more likely than its complement, showing quite some support for H2.

Calculate GORICA values and weights for H2 and its complement

```
set.seed(123)
gorica_sesam_2c <- goric(fit_Sesame, hypotheses = list(H2 = H2), type = "gorica", comparison = "complem
#summary(gorica_sesam_2c)
gorica_sesam_2c</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion approximation:

Results:

```
        model
        loglik
        penalty
        gorica
        loglik.weights
        penalty.weights
        gorica.weights

        1
        H2
        -5.125
        2.095
        14.438
        0.762
        0.855
        0.950

        2
        complement
        -6.288
        3.870
        20.317
        0.238
        0.145
        0.050
```

The order-restricted hypothesis 'H2' has 18.90 times more support than its complement.

Notes on weights

Here, the example of Lucas is used again; thus, using the following hypotheses of interest:

```
H1 <- 'group5 = group3 > group1 > group2; group3 > group4 > group2'

# Note: H1 is not full row-rank;

# for more details, see below and/or the goric tutorial.

H2 <- 'group3 > group1 > group4 = group5 > group2'
```

Note 1: GORIC weights versus GORICA weights

The GORICA weights (asymptotically) equal the GORIC weights. The differences are minor and often not notable with 2 decimals. Because of these minor differences, the relative weights (i.e., ratio of weights) can differ. Those differences in relative weights can even be large (as is in the Berzonsky et al Example), when dividing a very large number by a very small number with minor differences in these values.

Note 2: complement in case of two hypotheses of interest

One cannot compare the support of hypotheses when comparing them to their complements. Instead, one should evaluate them simultaneously in one set (like done above):

```
# Calculate goric for H1 and its complement
set.seed(123) # Set seed value
output_c_H1 <- goric(lm_fit_Lucas, hypotheses = list(H1), comparison = "complement")
#summary(output_c_H1)
output_c_H1</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
        model
        loglik
        penalty
        goric
        loglik.weights
        penalty.weights
        goric.weights

        1
        H1
        -278.051
        3.458
        563.019
        0.499
        0.903
        0.903

        2
        complement
        -278.048
        5.691
        567.479
        0.501
        0.097
        0.097
```

The order-restricted hypothesis 'H1' has 9.30 times more support than its complement.

```
# Calculate goric for H2 and its complement
set.seed(123) # Set seed value
output_c_H2 <- goric(lm_fit_Lucas, hypotheses = list(H2 = H2), comparison = "complement")
#summary(output_c_H2)
output_c_H2</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
        model
        loglik
        penalty
        goric
        loglik.weights
        penalty.weights
        goric.weights

        1
        H2
        -281.761
        3.136
        569.794
        0.024
        0.938
        0.270

        2
        complement
        -278.048
        5.853
        567.803
        0.976
        0.062
        0.730
```

The order-restricted hypothesis 'H2' has 0.37 times more support than its complement.

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	H1	-278.051	3.458	563.019	0.493	0.407	0.899
2	Н2	-281.761	3.136	569.794	0.012	0.561	0.030
3	unconstrained	-278.048	6.000	568.097	0.495	0.032	0.071

Ratio GORIC-weights:

	vs. H1	vs. H2	vs.	unconstrained
H1	1.000	29.593		12.668
H2	0.034	1.000		0.428
unconstrained	0.079	2.336		1.000

```
output_H1H2$ratio.gw[1,2]
[1] 29.59315
# The latter is not equal to:
output_c_H1$ratio.gw[1,2] / output_c_H2$ratio.gw[1,2]
[1] 25.16131
Extra
# Notably, you could derive the support from H1 vs H2 from their support versus
# that of Hu:
set.seed(123) # Set seed value
output_u_H1 <- goric(lm_fit_Lucas, hypotheses = list(H1))</pre>
#summary(output_u_H1)
output_u_H1
restriktor (0.5-80): generalized order-restricted information criterion:
Results:
                                       goric loglik.weights penalty.weights goric.weights
           model
                    loglik penalty
              H1 -278.051
                              3.458 563.019
                                                       0.499
                                                                        0.927
                                                                                        0.927
1
2 unconstrained -278.048
                              6.000 568.097
                                                       0.501
                                                                        0.073
                                                                                        0.073
Advise: Are you certain you wish to assess the order-restricted hypothesis in comparison to the unconst
The order-restricted hypothesis 'H1' has 0.927 / 0.073 > 1 times more support than the unconstrained.
set.seed(123) # Set seed value
output_u_H2 <- goric(lm_fit_Lucas, hypotheses = list(H2 = H2))</pre>
#summary(output_u_H2)
output_u_H2
restriktor (0.5-80): generalized order-restricted information criterion:
Results:
                                       goric loglik.weights penalty.weights goric.weights
           model
                    loglik penalty
              H2 -281.761
                              3.136 569.794
                                                       0.024
                                                                        0.946
                                                                                       0.300
1
                              6.000 568.097
                                                       0.976
                                                                        0.054
                                                                                        0.700
2 unconstrained -278.048
Advise: Are you certain you wish to assess the order-restricted hypothesis in comparison to the unconst
The order-restricted hypothesis 'H2' has 0.300 / 0.700 < 1 times more support than the unconstrained.
output_u_H1$ratio.gw[1,2] / output_u_H2$ratio.gw[1,2]
[1] 29.59099
# which (approximately) equals:
output_H1H2$ratio.gw[1,2]
```

[1] 29.59315

Note 3: weights when using complement not per se higher

In the Lucas example:

- The order-restricted hypothesis H_1 has 12.7 times more support than H_u (unconstrained).
- The order-restricted hypothesis H_1 has 9.3 times more support than its complement.

Now, the complement does not render a higher weight than if H_u was used.

If H_m is not in agreement with the data, the complement does not always render a higher weight. This is actually a good thing: Against H_u , H_m might obtain too much support then.

Some more explanation

This is because for this particular example the mean of group 3 and 5 are close (and they are compared in Hypothesis H_1). Notably, it will hold for all of the following three hypothesis:

```
H1_gr <- 'group5 > group3 > group1 > group2; group3 > group4 > group2'
# not full row-rank
H1_sm <- 'group5 < group3 > group1 > group2; group3 > group4 > group2'
# not full row-rank
H1 <- 'group5 = group3 > group1 > group2; group3 > group4 > group2'
# not full row-rank
```

Because the means of groups 3&5 are similar (and the other restrictions are in agreement with the data), the log likelihood values (LL's) of H_1 , H_u , and H_c (i.e., the complement of H_1) will be close - since the restricted/bounded solution (which is in agreement with H_1) is near the (unconstrained) maximum likelihood estimate (mle). Since the penalty for the complement is (always) lower than for H_u , it will receive more support than H_u - thus H_1 receives less. Notably, when H_1 is very specific (as in Berzonsky et al Example below), the penalty of H_u and H_c are almost the same.

When the means of groups 3 and 5 differ more, then evaluating against the complement does render a higher weight (than if H_u were used): Let us increase the mean of group 5 with 0.5 points (and change the name of the data & the analysis):

```
Lucas2 <- Lucas
Lucas2$Influence[Lucas$group == 5] <- Lucas$Influence[Lucas$group == 5] + 0.5

#describeBy(Lucas$Influence, Lucas$group, mat = TRUE)

#describeBy(Lucas2$Influence, Lucas$group, mat = TRUE)

lm_fit_Lucas2 <- lm(Influence ~ group-1, data = Lucas2)

# Now, from the three hypotheses H1_gr, H1_sm, and H1, # the first (H_gr) is

# correct. Hence, that one is used to illustrate the case where # a correct

# hypothesis obtains more support when it is evaluated against its complement:

H1_gr <- 'group5 > group3 > group1 > group2; group3 > group4 > group2'

# not full row-rank

set.seed(123) # Set seed value

output_u_gr <- goric(lm_fit_Lucas2, hypotheses = list(H1_gr = H1_gr))

#summary(output_u_gr)

output_u_gr
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
loglik.weights penalty.weights goric.weights
        model
                 loglik
                         penalty
                                    goric
        H1_gr
               -278.048
                           3.683
                                  563.464
                                                     0.500
                                                                      0.910
                                                                                     0.910
unconstrained -278.048
                           6.000 568.097
                                                     0.500
                                                                      0.090
                                                                                     0.090
```

Note: Hypotheses 'H1_gr' and 'unconstrained' have equal likelihood values (i.e., the hypotheses overlap

Advise: Are you certain you wish to assess the order-restricted hypothesis in comparison to the unconst

```
# Notably, this is the maximum support H1 can
# receive versus Hu (because the log likelihoods (LLs) are the same).
set.seed(123) # Set seed value
output_c_gr <- goric(lm_fit_Lucas2, hypotheses = list(H1_gr = H1_gr), comparison = "complement")
#summary(output_c_gr)
output_c_gr</pre>
```

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
model
                                           loglik.weights penalty.weights goric.weights
                 loglik
                         penalty
                                    goric
               -278.048
                           3.683
                                  563.464
                                                     0.706
                                                                      0.902
                                                                                      0.957
        H1_gr
                           5.900
                                                     0.294
                                                                      0.098
                                                                                      0.043
2
               -278.923
                                 569.646
  complement
```

The order-restricted hypothesis 'H1_gr' has 22.00 times more support than its complement.

Now, when the means of groups 3 and 5 differ more, then evaluating against the complement does render more support than if H_u were used: 22 versus 10.

Extra: Two methods to calculate the penalty

There are two methods that can be used in calculating the penalty. The default method is often much faster (if the number of parameters is not too high) and needs less input specification. It can, however, not deal with hypotheses that are not of full row-rank (like H_1 above). In that case, **restriktor** uses automatically the other (bootstrap) method.

To use this bootstrap method use 'mix weights = "boot" ':

restriktor (0.5-80): generalized order-restricted information criterion:

Results:

```
goric loglik.weights penalty.weights goric.weights
           model
                    loglik penalty
1
              H1
                  -278.051
                              3.457
                                      563.016
                                                        0.493
                                                                          0.406
                                                                                         0.899
2
              H2
                  -281.761
                                                        0.012
                                                                          0.562
                                                                                         0.030
                              3.131
                                      569.784
3 unconstrained
                 -278.048
                              6.000
                                     568.097
                                                        0.495
                                                                          0.032
                                                                                         0.071
```

Ratio GORIC-weights:

	vs. H1	vs. H2	vs. unconstrained
H1	1.000	29.483	12.685
H2	0.034	1.000	0.430

unconstrained 0.079 2.324 1.000

This, of course, renders the same results as above (if there is a difference, it is in the second decimal of the penalty).

Note on not full row-rank

If the restriction matrix is not of full row-rank, this means one of the following:

a) There is at least one redundant restriction.

Then, either a.1) leave the redundant one out or a.2) use another (more time-consuming) way of obtaining the level probabilities for the penalty term (the goric function does this by default): Bootstrapping, as discussed above.

b) There is at least one range restriction (e.g., -2 < group1 < 2).

Such a restriction can be evaluated but there is a sensitivity (of a scaling factor in the covariance matrix, like with a prior in a Bayes factor) which currently cannot be checked for.

c) There is at least one conflicting restriction (e.g., 2 < group1 < -2).

Such a restriction can evidently never hold and is thus impossible to evaluate. To prevent this type of error delete the one that is incorrect and run goric() again.