# Tutorial for updating hypotheses using the GORIC(A)

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This is a tutorial for updating hypotheses using the GORIC(A). It uses the goric() function of the restriktor package to calculate the GORIC and the GORICA.

Updating hypotheses, consists of the following three steps:

- 1. Conduct (exploratory) model selection to the first study.
- 2. Specify the (confirmatory/informative) hypotheses for the second/replication study using the results from Step 1, that is, update your hypotheses.
- 3. Conduct model selection to the second study.

#### Two examples will be used:

- 1. The first is based on Monin and Holubar, a replication study of Monin. Here, the study of Monin is used to update the exploratory hypotheses evaluated in Monin to theory-based, order-restricted hypotheses to be evaluated in Holubar. By examining these, one can check whether Holubar replicated the findings of Monin or not. Both these studies use an ANOVA model.
- 2. The second example is based on Altinisik et al.. Here, theory-based hypotheses are based on theories and on findings of previous studies. In this example, the GORICA is applied to the results of a logistic regression model.

At the end, there is code to automatically create all possible exploratory hypotheses, which can be helpful when there are a lot of parameters.

# Example 1: ANOVA

In this example, the study of Monin will be used to render order-constrained hypotheses for the Holubar study, which is a (direct) replication study of Monin. For this, an exploratory analysis will be performed on the Monin data and inspect the sample means.

# Step 1: Model selection 1st study

### 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 <code>getwd()</code> shows you your current working directory. You can change the working directory to the one you prefer using the function <code>setwd()</code> 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\_Monin.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
Monin <- read.table("data/Data_Monin.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
Monin$group <- factor(Monin$group)</pre>
```

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

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

```
attract group
1 1.19560213 1
2 0.05852551 1
3 -1.10092218 1
4 1.78095312 1
5 1.86726943 1
6 1.12279975 1
```

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

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

```
item group1 vars n mean
                                sd median trimmed
                                                                             skew kurtosis
                                                    mad
                                                            min
                                                                  max range
                                                                                               se
                                            1.869 0.868 -1.101 5.050 6.151 0.201
                                                                                      0.135 0.317
X11
                   1 19 1.88 1.38
                                    1.781
       1
              1
X12
       2
                   1 19 2.54 1.95
                                    2.711
                                            2.507 1.931 -1.099 6.746 7.845 0.024
                                                                                     -0.514 0.447
X13
       3
                   1 29 0.02 2.38
                                   0.438
                                          -0.008 2.897 -3.911 4.916 8.828 0.028
                                                                                     -1.0340.442
```

### Preparation for GORIC(A)

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

```
lm_fit_Monin <- lm(attract ~ group - 1, data = Monin)</pre>
```

Note that:

- 1. y ~ group 1 instructs the function lm (linear model) to regress y on 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. On the other hand, if the intercept is not dropped, 'y ~ group' would estimate an intercept, representing the mean of the reference group, and the mean differences between the other (here, two) groups and the reference group.
- 3. The results are collected in, what is called, an R-object, named lm\_fit\_Monin.

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

```
names(coef(lm_fit_Monin))
```

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

**Set of (exploratory) hypotheses** On the Monin data set, an exploratory analysis will be performed, which means evaluating all combinations with equalities (and no restrictions). To specify hypotheses, remember the following:

- 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").
  - 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-group3).

• 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').

All possible exploratory hypotheses are denoted by:

```
H00 <- "group1 = group2 = group3"

H01 <- "group1 = group2"

H02 <- "group1 = group3"

H03 <- "group2 = group3"
```

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.

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

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

### Model selection using GORIC(A)

```
set.seed(123) # Set seed value
goric(lm_fit_Monin, hypotheses = list(H00 = H00, H01 = H01, H02 = H02, H03 = H03))
```

## Calculate GORIC values and weights

restriktor (0.5-50): generalized order-restricted information criterion:

#### Results:

	model	loglik	penalty	goric	loglik.weights	penalty.weights	goric.weights
1	НОО	-149.907	2.000	303.815	0.000	0.447	0.000
2	HO1	-141.191	3.000	288.383	0.369	0.164	0.610
3	H02	-145.404	3.000	296.809	0.005	0.164	0.009
4	Н03	-148.907	3.000	303.815	0.000	0.164	0.000
5	unconstrained	-140.665	4.000	289.330	0.625	0.060	0.380

Calculate GORICA values and weights In case you want to use the GORICA values and weights instead, use: type = "gorica". Thus:

```
set.seed(123) # Set seed value
goric(lm_fit_Monin, hypotheses = list(H00 = H00, H01 = H01, H02 = H02, H03 = H03),
     type = "gorica")
```

#### Step 2: Update hypotheses

It can be seen that  $H_01$  ( $\mu_1 = \mu_2, \mu_3$ ) receives the most support. From the sample means (see descript), it is concluded that  $\mu_1$  and  $\mu_2$  are both larger than  $\mu_3$ . Therefore, the following hypothesis will be evaluated in the Holubar data:

$$H_1: \mu_1 = \mu_2 > \mu_3.$$

Since  $H_u$  obtained some support as well, one could specify (using the sample means of the Monin data) the following competing hypothesis:

$$H_2: \mu_2 > \mu_1 > \mu_3.$$

# Step 3: Model selection 2nd study

## Data preparation

First, read in the Holubar dataset (the replication study of Monin), 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
                                                                                 -0.7780.268
                  1 27 0.02 1.88 0.169
X12
      2
             2
                                          0.111 1.951 -4.961 2.921 7.883 -0.517
                                                                                 -0.104 0.362
X13
                  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: R-object** 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))</pre>
```

```
[1] "gr1" "gr2" "gr3"
```

**Set of hypotheses** Based on the results and sample means of Monin, the following two competing hypotheses were created:

```
H1 <- "gr1 = gr2 > gr3"
H2 <- "gr2 > gr1 > gr3"
```

# Model selection using GORIC(A)

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

#### Calculate GORIC values and weights

restriktor (0.5-50): generalized order-restricted information criterion:

#### Results:

```
goric loglik.weights penalty.weights goric.weights
          model
                   loglik penalty
                 -144.981
                             2.500
                                    294.962
                                                      0.111
                                                                      0.510
                                                                                     0.303
1
             H1
2
             H2 -144.981
                             2.803 295.569
                                                      0.111
                                                                      0.376
                                                                                     0.224
3 unconstrained -143.038
                             4.000 294.076
                                                      0.777
                                                                      0.114
                                                                                     0.473
```

Ratio GORIC-weights:

H1 H2 unconstrained	vs. H1 1.000 0.738 1.557	1.000	vs. unconstrained 0.642 0.474 1.000
Ratio loglik-w	eights:		
	vs. H1	vs. H2	vs. unconstrained
H1	1.000	1.000	0.143
H2	1.000	1.000	0.143
unconstrained	6.978	6.978	1.000
Ratio penalty-	weights:		
	vs. H1	vs. H2	vs. unconstrained
H1	1.000	1.355	4.482
H2	0.738	1.000	3.309
unconstrained	0.223	0.302	1.000

order-restricted hypotheses:

```
H1:
gr1 = gr2 > gr3
H2:
gr2 > gr1 > gr3
```

Since the support for  $H_1$  and  $H_2$  is lower than for  $H_u$ , it can be concluded that both are weak hypotheses. Hence, the study of Holubar did not replicate the findings of Monin.

**Alternative set** In case you are only interested in the 'main hypothesis'  $H_1$  found in Monin, you could also evaluate this against its complement:

```
set.seed(123) # Set seed value
goric(lm_fit_Holubar, hypotheses = list(H1), comparison = "complement")
```

restriktor (0.5-50): generalized order-restricted information criterion:

#### Results:

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

        1
        H1
        -144.981
        2.500
        294.962
        0.125
        0.731
        0.280

        2
        complement
        -143.038
        3.500
        293.076
        0.875
        0.269
        0.720
```

The order-restricted hypothesis 'H1' has 0.39 times more support than its complement.

Since  $H_1$  has only 0.39 (lower than 1) times more support than its complement, it is a weak hypothesis. Hence, the study of Holubar did not replicate the findings of Monin.

Calculate GORICA values and weights Notably, in case you want to use the GORICA, you can use the following commands:

```
set.seed(123) # Set seed value
goric(lm_fit_Holubar, hypotheses = list(H1, H2), type = "gorica")
```

When you want to calculate the GORICA for  $H_1$  and its complement, use:

```
set.seed(123) # Set seed value
goric(lm_fit_Holubar, hypotheses = list(H1), type = "gorica", comparison = "complement")
```

# Example 2: Logistic regression example

In this example, the order-constrained hypotheses for a replication data set are based on the results of the original study and based on theories from two previous studies. More details can be found in Altinisik et al. (2021). Since the data cannot be shared, the estimates of the parameters of interest and their covariance matrix are used, which can serve as input for the GORICA.

#### Load libraries

Start with loading 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 package, if you have not done this already:
if (!require("restriktor")) install.packages("restriktor")

## Then, load the packages:
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
```

#### Load data

The estimates of structural parameters (i.e., the parameters of interest, that is, the parameters used in the hypotheses) are

```
est <- c(0.4765484, 0.2759022, 0.8669835, -0.443003, 0.8066336, -0.246261)
names(est) <- c("RS_Gr1", "RSES_Gr1", "RSES_Gr2", "RSES_Gr2", "RSES_Gr3")
```

Notably, the parameters need to be labelled such that those can be used in specifying the hypotheses.

Their covariance matrix is:

Notably, the covariance matrix does not need to be labelled.

## Set of (updated) hypotheses

As discussed in Altinisik et al. (2021), the set of hypotheses is based on theories which are updated by findings from previous studies. Note that the same names/labels as the ones above are used.

```
H1 <- "RS_Gr1 + RSES_Gr1 = 0; RS_Gr1 > 0;

RS_Gr2 + RSES_Gr2 = 0; RS_Gr2 > 0;

RSES_Gr3 = 0; RS_Gr3 > 0"

H2 <- "RS_Gr1 + RSES_Gr1 = 0; RS_Gr1 > 0;
```

```
RSES_Gr2 = 0; RS_Gr2 = 0;
RSES_Gr3 = 0; RS_Gr3 > 0"
```

## Model selection using GORICA

Next, the GORICA values and weights are calculated:

```
set.seed(123)
output_gorica_Altinisik <- goric(est, hypotheses = list(H1, H2), VCOV = VCOV, type = "gorica")
summary(output_gorica_Altinisik)</pre>
```

restriktor (0.5-50): generalized order-restricted information criterion approximation:

#### Results:

	model	loglik	penalty	gorica	loglik.weights	penalty.weights	gorica.weights
1	H1	-1.373	1.500	5.746	0.202	0.376	0.760
2	Н2	-3.168	1.000	8.335	0.034	0.620	0.208
3	unconstrained	-0.045	6.000	12.089	0.764	0.004	0.032

#### Ratio GORICA-weights:

	vs. Hl	vs. H2	vs. unconstrained
H1	1.000	3.650	23.847
H2	0.274	1.000	6.534
unconstrained	0.042	0.153	1.000
unconstrained	0.042	0.153	

Ratio loglik-weights:

vs. H1	vs. H2	vs. unconstrained
1.000	6.018	0.265
0.166	1.000	0.044
3.775	22.715	1.000
	1.000	vs. H1 vs. H2 1.000 6.018 0.166 1.000 3.775 22.715

Ratio penalty-weights:

	vs. H1	vs. H2	vs.	unconstrained
H1	1.000	0.607		90.017
H2	1.649	1.000		148.413
${\tt unconstrained}$	0.011	0.007		1.000

order-restricted hypotheses:

```
H1:

RS_Gr1 + RSES_Gr1 = 0; RS_Gr1 > 0;

RS_Gr2 + RSES_Gr2 = 0; RS_Gr2 > 0;

RSES_Gr3 = 0; RS_Gr3 > 0

H2:

RS_Gr1 + RSES_Gr1 = 0; RS_Gr1 > 0;

RSES_Gr2 = 0; RS_Gr2 = 0;

RSES_Gr3 = 0; RS_Gr3 > 0
```

It can be seen that both  $H_1$  and  $H_2$  are not weak hypotheses and that  $H_1$  is the preferred one: It is 3.65 times more supported than  $H_2$ . Hence, the support for these theories is also found in this replication and,

based on this replication study, it is concluded that  $H_1$  is more supported.

# Extra: Generating all exploratory (=) restrictions

```
# make columns per number of variables in hypothesis
h <- paste(names(coef(lm_fit_Monin)), " = 0")</pre>
col <- NULL
for (i in 1:length(h)) {
    col <- c(col, combn(h, i, FUN = function(x) {</pre>
        return(paste(x, collapse = "; "))
}
hypotheses <- data.frame(col)</pre>
\# make dataframe without the missings and split in two columns
if (!require("tidyr")) install.packages("tidyr")
Loading required package: tidyr
library(tidyr)
hypotheses <- gather(hypotheses, na.rm = TRUE, factor_key = TRUE)
# all hypothesis in text, by R
hypotheses$value
[1] "group1 = 0"
                                             "group2 = 0"
                                                                                       "group3 = 0"
# number of hypotheses
nrhypos <- sum(!is.na(hypotheses$value))</pre>
# create all hypotheses HO1 to HOnrhypos
for (i in 1:nrhypos) {
    assign(paste("HO", i, sep = ""), hypotheses$value[i])
# Now all the exploratory (=) hypotheses are specified
# In case you want to see them all, paste the following result in the R
# console:
cat(paste0("H0", 1:nrhypos, ";"))
H01; H02; H03; H04; H05; H06; H07;
# and past result in R console and you see them.
# paste result from following into goric function
cat(paste0("HO", 1:nrhypos, ","))
H01, H02, H03, H04, H05, H06, H07,
```