

Informative hypotheses evaluation

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ANOVA Example: Comparisons of 3 Means

Palmer and Gough (2007) examined the difference in the attribution of importance of defective education as an explanation for criminal behavior between three types of “offenders”:

(1) non-offenders, (2) property offenders, and (3) person offenders.

ANOVA

$$y_j = \mu_1 d_{1j} + \mu_2 d_{2j} + \mu_3 d_{3j} + \epsilon_j,$$

where μ_i ($i = 1, 2, 3$) is the mean in group i ,
 d_{ij} is a dummy variable denoting group membership,
 ϵ_j is an error term and $\epsilon_j \sim N(0, \sigma^2)$.

$(n_i = 20, 20, 31)$

Hypotheses of interest

Palmer and Gough (2007) expect that

- non-offenders (1) attribute more importance to defective education for explaining crime than the other two offenders;
i.e., $\mu_1 > \mu_2$ and $\mu_1 > \mu_3$,
- property offenders (2) attribute more importance to defective education than person offenders (3);
i.e., $\mu_2 > \mu_3$.

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

Note that ‘<’ denotes “smaller than” and ‘>’ denotes “larger than”.

Conclusions w.r.t. hypothesis of interest $H_1 : \mu_1 > \mu_2 > \mu_3$

With post-hoc test (e.g., Bonferroni) and on basis of the sample means, we could say a bit more.

But, what if

- $\mu_1 \neq \mu_2, \mu_1 \neq \mu_3, \mu_2 = \mu_3,$
- $\bar{y}_1 > \bar{y}_2,$

which implies $\mu_1 > \mu_2 = \mu_3$.

But also not interested in this.

Furthermore, inconsistent results are possible (like in this data example):

$$\mu_1 \neq \mu_3, \mu_1 = \mu_2, \text{ and } \mu_2 = \mu_3,$$

which is logically impossible, since latter two imply $\mu_1 = \mu_3$.

Using planned contrasts would overcome this, but still = restrictions then.

NHST on Palmer & Gough (2007) Data

```
PandG_data <- read.table("Data_PalmerAndGough.txt",  
                          header=TRUE)  
PandG_data$group <- factor(PandG_data$group)  
pairwise.t.test(PandG_data$Importance, PandG_data$group,  
                p.adj = 'bonferroni')  
  
##  
## Pairwise comparisons using t tests with pooled SD  
##  
## data: PandG_data$Importance and PandG_data$group  
##  
##      1      2  
## 2 0.191 -  
## 3 0.011 1.000  
##  
## P value adjustment method: bonferroni
```

Exploration and Confirmation

Exploration (like in post hoc tests and possibly AIC)

Test all possible pairs/subsets of means whether significant different (“,”) or not (“=”).

For example, when $k = 3$:

$$H_{0E} : \mu_1 = \mu_2 = \mu_3$$

$$H_{1E} : \mu_1 = \mu_2, \mu_3$$

$$H_{2E} : \mu_1, \mu_2 = \mu_3$$

$$H_{3E} : \mu_1 = \mu_3, \mu_2$$

$$H_a : \mu_1, \mu_2, \mu_3$$

When $k = 5$, there are even 52 hypotheses.

Confirmation

Limited set: Compare only prespecified hypotheses including order restrictions ($<$, $>$, but also $=$).

GORIC on Palmer & Gough (2007) Data

$$H_0 : \mu_1 = \mu_2 = \mu_3,$$

$$H_1 : \mu_1 > \mu_2 > \mu_3,$$

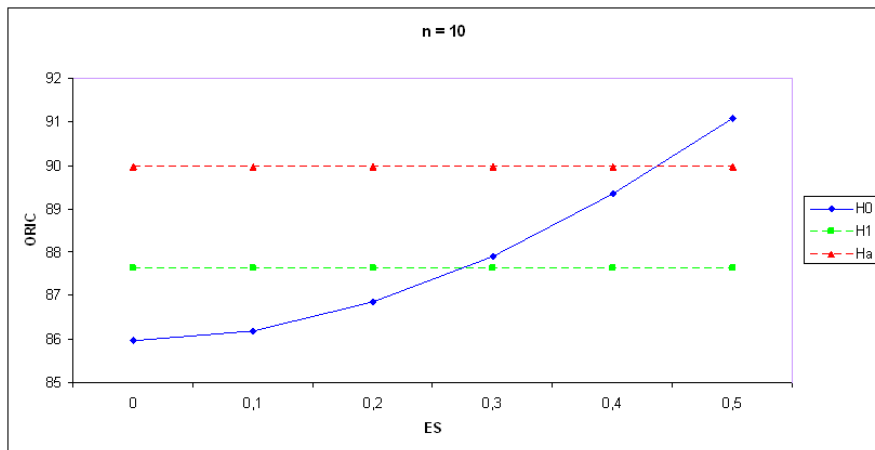
$$H_u : \mu_1, \mu_2, \mu_3.$$

GORIC

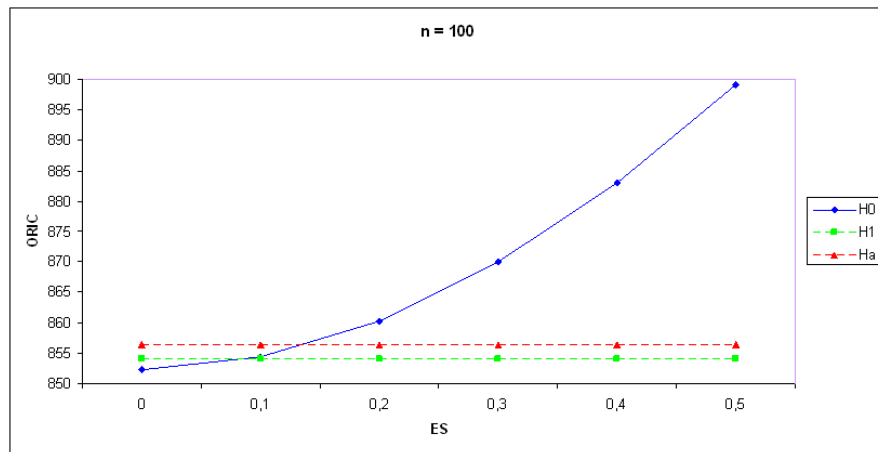
Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.02
H_1	-191.89	2.81	389.41	0.75
H_u	-191.89	4.00	391.79	0.23

Confirmatory methods have more “power” than their exploratory counterparts (cf. one-sided testing).

Confirmation more power: 1 data set. GORIC values for 3 groups, effect size ES , and $n = 10$ observations per group



Confirmation more power: 1 data set. GORIC values for 3 groups, effect size ES , and $n = 100$ observations per group



Confirmatory methods - some of my references

- Kuiper, R. M., and Hoijtink, H. (2010). Comparisons of Means Using Exploratory and Confirmatory Approaches. *Psychological Methods*, 15(1), 69–86.
- Kuiper, R. M., Klugkist, I., and Hoijtink, H. (2010). A Fortran 90 Program for Confirmatory Analysis of Variance. *Journal of Statistical Software*, 34(8), 1–31.
- Kuiper, R.M., Hoijtink, H. and Silvapulle, M.J. (2011). An Akaike type information criterion for model selection under inequality constraints. *Biometrika*, 98, 495-501. (GORIC)
- Kuiper, R.M., Nederhof, T., and Klugkist, I. (2015). Properties of hypothesis testing techniques and (Bayesian) model selection for exploration-based and theory-based (order-restricted) hypotheses. *British Journal of Mathematical and Statistical Psychology*, 68(2), 220 – 245.
- Altınışık, Y., Van Lissa, C. J., Hoijtink, H., Oldehinkel, A. J., and Kuiper, R. M. (2021). Evaluation of inequality constrained hypotheses using a generalization of the AIC. *Psychological Methods*, 26(5), 599-621. (GORICA)

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Generalized Order-Restricted Information Criterion

GORIC

$$\text{'IC'} = -2 \text{ fit} + 2 \text{ complexity}$$

Fit = Maximized order-restricted log likelihood

Maximized log likelihood based on parameters in agreement with H_m .

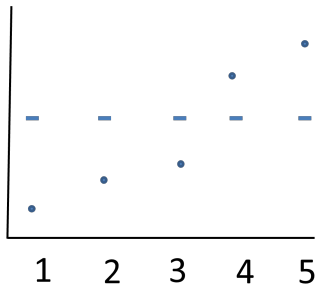
Complexity = Penalty

Represents: Expected number of distinct parameters.

Here, expected number of distinct mean values plus 1 (because of the unknown variance term).

Details: Function of level probabilities.

Intermezzo: Balance Fit and Complexity

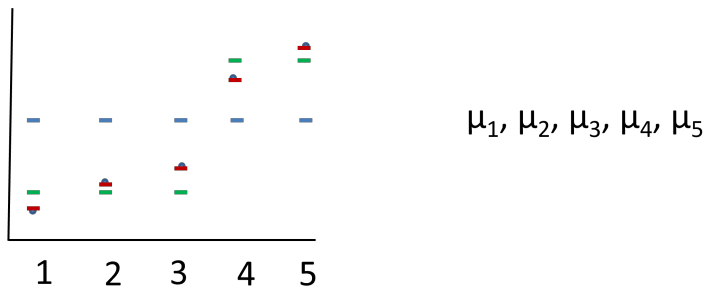


$$\mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5 = \mu$$

parameters: 1 mean parameter (i.e., μ)

Fit: Bad fit

Intermezzo: Balance Fit and Complexity

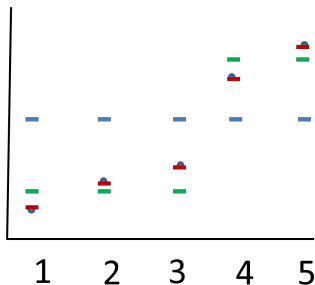


parameters: 5 mean parameter (complex)
Fit: Better than with two parameters, even best possible fit.

Thus: Best fit (= highest likelihood), but also most complex (= highest penalty).

In this example: Two means may be best trade-off between fit & complexity.

Intermezzo: Balance Fit and Complexity

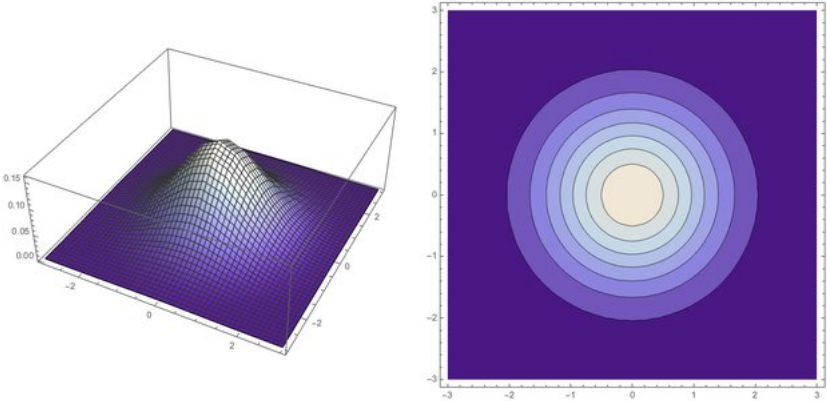


Now, we also incorporate order-restrictions (e.g., $\mu_1 < \mu_2$).

Then, helpful to look at likelihood (not scatter plot).

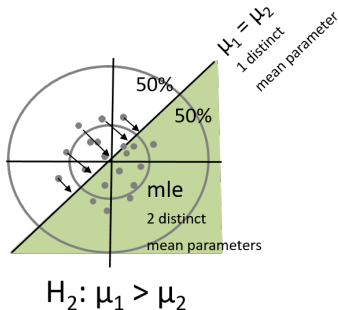
In presentation, I will use contour plots (in case of two means, not five).

Intermezzo: Contour plot



Idea complexity

penalty (PT)



Note: sampling from null distribution: $H_0: \mu_1 = \mu_2 = 0$.

complexity $H_2 = PT_2 = 1 + 0.5 \times 1 + 0.5 \times 2 = 2.5$.

PT = number of expected distinct parameters (under H_0).

GORIC on Palmer & Gough (2007) Data

$$H_0 : \mu_1 = \mu_2 = \mu_3,$$

$$H_1 : \mu_1 > \mu_2 > \mu_3,$$

$$H_2 : \mu_1 > \mu_2 < \mu_3,$$

$$H_3 : \mu_1 < \mu_2 < \mu_3,$$

$$H_u : \mu_1, \mu_2, \mu_3.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.01
H_1	-191.89	2.81	389.41	0.56
H_2	-192.34	3.19	391.05	0.25
H_3	-196.36	2.81	398.34	0.01
H_u	-191.89	4.00	391.79	0.17

GORIC: Lowest value is best

GORIC is like AIC expected distance from the truth (KL-distance). Hence, smallest value is best.

GORIC on Palmer & Gough (2007) Data

$$H_0 : \mu_1 = \mu_2 = \mu_3,$$

$$H_1 : \mu_1 > \mu_2 > \mu_3,$$

$$H_u : \mu_1, \mu_2, \mu_3.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.02
H_1	-191.89	2.81	389.41	0.75
H_u	-191.89	4.00	391.79	0.23

Hands-on/Demo (1): GORIC

Start Rstudio and let's practice.

- Go to <https://github.com/rebeccakuiper/Tutorials>:
 1. Click on green button called Code.
 2. Download zip (last option in list).
 3. Unzip it on your machine (that folder is now your working directory).
- Optional: Open 'Tutorial_GORIC_restriktor_ANOVA.html'.
- Start R (optional: make project).
Optional: open 'PalmerAndGough_and_Lucas.R'.
- Install packages and load them.
- Read and inspect data. Use Data_PalmerAndGough.txt.
- Run model (lm()).
- Specify hypotheses (make up your own).
Note: Use names used in the model, or overwrite those.
- Run goric().
- Inspect and interpret output.
Note: 'GORIC weights' will be explained later on.

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Interpretation: GORIC weights

GORIC values

GORIC values cannot be interpreted, only compared:
Smallest is best.

GORIC weights (w_m) and ratios ($w_m/w_{m'}$)

GORIC weight quantifies how much H_m is more supported than others in set.

Ratio of GORIC weights quantifies relative support of H_m vs $H_{m'}$.

The bigger, the better.

Reference:

Kuiper, R.M., Hoijtink, H. and Silvapulle, M.J. (2012). Generalization of the order restricted information criterion for multivariate normal linear models. *Journal of Statistical Planning and Inference*, 142, 2454-2463.

Illustration of the GORIC weights (w_m)

$$H_0 : \mu_1 = \mu_2 = \mu_3,$$

$$H_1 : \mu_1 > \mu_2 > \mu_3,$$

$$H_u : \mu_1, \mu_2, \mu_3.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.02
H_1	-191.89	2.81	389.41	0.75
H_u	-191.89	4.00	391.79	0.23

Note on GORIC weights vs BF and PMPs

ratio GORIC weights ($w_m/w_{m'}$) \sim Bayes factor ($BF_{mm'}$).

GORIC weight (w_m) \sim posterior model probability (PMP).

1 - w_m = conditional error probability.

Like PMP, w_m depends on set of hypotheses.

Example

$$w_m = .8$$

$$w_{m'} = .2$$

H_m is $.8/.2 = 4$ times more supported than $H_{m'}$.

There is an error probability of $(1 - .8) * 100\% = 20\%$ that H_m is true.

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R code: GORIC

Palmer & Gough

$$H_0 : \mu_1 = \mu_2 = \mu_3$$

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

$$H_u : \quad \mu_1, \mu_2, \mu_3$$

```
# GORIC (using goric function in restriktor package)
library(restriktor)
set.seed(123) # Set seed value
goric.PandG <- goric(fit.PandG,
                     hypotheses = list(H0, H1))
```


GORIC on Palmer & Gough Data

$$H_0 : \mu_1 = \mu_2 = \mu_3$$

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

$$H_u : \mu_1, \mu_2, \mu_3$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.02
H_1	-191.89	2.81	389.41	0.75
H_u	-191.89	4.00	391.79	0.23

Another illustration: one-way ANOVA

Lucas: 5 groups

Lucas (2003) investigated difference between female and male leadership w.r.t. influence of the leader.

Five experimental groups:

1. a randomly selected male leader
2. a randomly selected female leader
3. male leader selected via task
4. female leader selected via task
5. female leader selected via task + institutionalized female leadership via movie

(Two informative) hypotheses of interest

- $H_0 : \mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5$ (include only when of interest),
- $H_1 : \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$,
- $H_2 : \mu_3 > \mu_1 > \mu_4 = \mu_5 > \mu_2$,
- $H_u : \mu_1, \mu_2, \mu_3, \mu_4, \mu_5$ (includes as failsafe).

Descriptive statistics of Lucas' Data

Group	Mean Influence	SD	n
1 (male, random)	2.33	1.86	30
2 (female, random)	1.33	1.15	30
3 (male, selected)	3.20	1.79	30
4 (female, selected)	2.23	1.45	30
5 (female, selected+)	3.23	1.50	30

R code: Data and fit object

Lucas

```
# Data
```

```
lucas_data <- read.table("Data_Lucas.txt", header=TRUE)
```

```
lucas_data$group <- factor(lucas_data$group)
```

```
# fit object (needed as input for goric function)
```

```
fit.lucas <- lm(Influence ~ group - 1, data = lucas_data)
```

R code: Hypotheses

Lucas

$$H_0: \mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5,$$

$$H_1: \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$$

$$H_2: \mu_3 > \mu_1 > \mu_4 = \mu_5 > \mu_2,$$

$$H_u : \quad \mu_1, \mu_2, \mu_3, \mu_4, \mu_5.$$

Hypotheses

```
H0 <- 'group1 = group2 = group3 = group4 = group5'
```

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

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


GORIC on Lucas' Data

$$H_0 : \mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5,$$

$$H_1 : \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$$

$$H_2 : \mu_3 > \mu_1 > \mu_4 = \mu_5 > \mu_2,$$

$$H_u : \mu_1, \mu_2, \mu_3, \mu_4, \mu_5.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-292.27	2.00	588.54	0.00
H_1	-278.05	3.20	562.49	0.92
H_2	-281.76	3.14	569.79	0.02
H_u	-278.05	6.00	568.10	0.06

GORIC on Lucas' Data: GORIC weights (w_m)

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-292.27	2.00	588.54	0.00
H_1	-278.05	3.20	562.49	0.92
H_2	-281.76	3.14	569.79	0.02
H_u	-278.05	6.00	568.10	0.06

H_1 is $.921/.024 \approx 38.6$ (∞) times more supported than H_2 (H_0 , resp.). Thus, there is quite some evidence that H_1 is the best of this set.

Note:

H_1 is $.921/.056 \approx 16.5 > 1$ times more supported than H_ν .

Hence, H_1 not weak and thus comparable to the other informative hypotheses (as explained next).

Hands-on/Demo (2): GORIC weights

Start Rstudio and let's practice.

- Optional: Open 'Tutorial_GORIC_restriktor_ANOVA.html'.
- If needed: Start R again (optional: make project) and then also load packages again.
Optional: open 'PalmerAndGough_and_Lucas.R'.
- Read and inspect data.
Use Data_Lucas.txt and/or Data_PalmerAndGough.txt.
- Run model (lm()).
- Specify hypotheses (make up your own). Note: Use names used in the model, or overwrite those.
- Run goric().
- Inspect and interpret output: Focus on GORIC weights.

Include “unconstrained” hypothesis

If set of hypotheses does not contain a reasonable/good one:
 Select the best of set of weak hypotheses.
 E.g.: $w_0 = .2$ and $w_1 = .8$.

Prevent choosing a weak hypothesis

Include unconstrained hypothesis H_u (or H_a):

$$\begin{aligned}
 H_0 : & \quad \mu_1 = \mu_2 = \mu_3, \\
 H_1 : & \quad \mu_1 > \mu_2 > \mu_3, \\
 H_u : & \quad \mu_1, \mu_2, \mu_3, \\
 & \quad \text{(i.e., no restrictions).}
 \end{aligned}$$

H_u highest fit but also most complex, thus failsafe.

E.g.: $w_0 = .02$, $w_1 = .08$, and $w_u = .90$.

What if another hypothesis is true?

What if another informative hypothesis is true?

The failsafe hypothesis will be preferred over the informative hypotheses in the set, if the sample size is large enough.

E.g.: $w_0 = .02$, $w_1 = .08$, and $w_u = .90$.

Use of H_u : Palmer & Gough (2007) Data

$$H_0 : \mu_1 = \mu_2 = \mu_3,$$

$$H_1 : \mu_1 > \mu_2 > \mu_3,$$

$$H_u : \mu_1, \mu_2, \mu_3.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_0	-196.36	2.00	396.71	0.02
H_1	-191.89	2.81	389.41	0.75
H_u	-191.89	4.00	391.79	0.23

If at least one informative hypothesis not weak ($w_1 > w_u$ or $w_1/w_u > 1$), then compare informative hypotheses.

Hence: H_u is only a failsafe not another hypothesis of interest.

H_1 vs H_u : Palmer & Gough (2007) Data

What if only one informative hypothesis:

$$H_1 : \quad \mu_1 > \mu_2 > \mu_3,$$

$$H_u : \quad \mu_1, \mu_2, \mu_3.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_1	-191.89	2.81	389.41	0.77
H_u	-191.89	4.00	391.79	0.23

H_1 is $.77/.23 \approx 3.3$ times more supported than H_u .

BUT: H_u includes H_1 .

So, support for H_u contains support for H_1 .

H_1 vs H_u : Palmer & Gough (2007) Data

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H_1	-191.89	2.81	389.41	0.77
H_u	-191.89	4.00	391.79	0.23

Unconstrained is always true, so will always receive support.
Support for H_u contains support for H_1 (if any).

Now: Fit H_1 and H_u the same;
thus, GORIC weights are based on only complexity.

Consequently, $.77/.23 \approx 3.3$ is an upper bound.

H_1 vs H_u : Upper bound

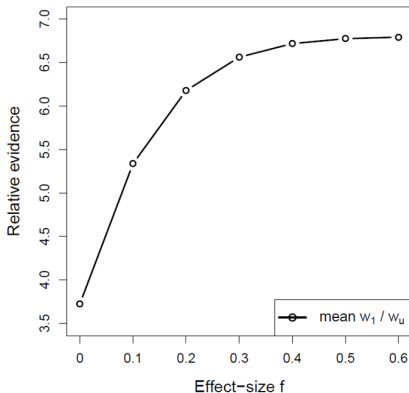
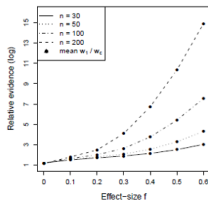


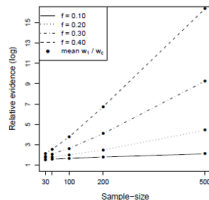
Figure: IC weights can have an upper bound, when informative hypothesis has maximum fit (i.e., is fully in agreement with the data).

Alternative safeguard: Complement of H_m using GORIC

vs complement

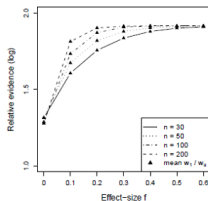


(a)

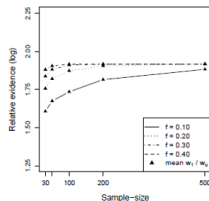


(b)

vs unconstrained



(c)



(d)

Palmer & Gough (2007): H_1 vs H_c

$$H_1 : \quad \mu_1 > \mu_2 > \mu_3,$$

$$H_c : \quad \text{not } H_1.$$

GORIC

Model	Fit	Complexity	GORIC	GORIC weights
H1	-191.89	2.81	389.41	0.79
complement	-192.34	3.69	392.05	0.21

H_1 is $.79/.21 \approx 3.8$ times more supported than its complement, that is, any other hypothesis.

Hands-on/Demo (3): GORIC - H_m vs complement

Start Rstudio and let's practice.

- Optional: Open 'Tutorial_GORIC_restriktor_ANOVA.html'.
- If needed: Start R again (optional: make project) and then also load packages again.
- Read and inspect data.
Use Data_Lucas.txt and/or Data_PalmerAndGough.txt.
- Run model (`lm()`).
- Specify hypotheses (make up your own). E.g., for Lucas:

$$H_1 : \quad \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$$

Note: Use names used in the model, or overwrite those.

- Run `goric()`; now, add:
`comparison = "complement"`

- Inspect and interpret output.

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GORICA

Similarities with GORIC

- Form: $GORICA_m = -2 \text{ fit} + 2 \text{ complexity}$.
- Broad type of restrictions.

Differences compared to GORIC

- Uses asymptotic expression of the likelihood (is a normal):
can therefore be easily applied to all types of models.
Disadvantage: might work less well in case of small samples.
- Does not need data set; mle's and their covariance matrix suffice.
- Can leave out nuisance parameters (i.e., not part of hypotheses).

Note

In case of normal linear models and/or not too small samples:
GORICA weights = GORIC weights.

R code: GORICA

Palmer & Gough

$$H_1 : \quad \mu_1 > \mu_2 > \mu_3,$$

$$H_c : \quad \text{not } H_1.$$

GORICA: type = "gorica"

```
H1 <- 'group1 > group2 > group3'  
# GORICA (using goric function in restriktor package)  
library(restriktor)  
set.seed(123) # Set seed value  
gorica.lucas_C <- goric(fit.PandG,  
  hypotheses = list(H1),  
  comparison = 'complement',  
  type = "gorica")
```


Hands-on/Demo (4): GORICA - H_m vs complement

Start Rstudio and let's practice.

- Optional: Open 'Tutorial_GORIC_restriktor_ANOVA.html'.
- If needed: Start R again (optional: make project) and then also load packages again.
- Read and inspect data.
Use Data_Lucas.txt and/or Data_PalmerAndGough.txt.
- Run model (lm()).
- Specify hypotheses (make up your own). E.g., for Lucas:

$$H_1 : \quad \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$$

Note: Use names used in the model, or overwrite those.

- Run `goric()`; now, add:
`type = "gorica"`

- Inspect and interpret output.

GORIC in JASP: Lucas

- Open JASP and 'Data_Lucas_JASP.txt'. Check measurement levels.
- Go to ANOVA and denote variables of interest.
- Go to Order Restricted Hypotheses tab.
- Specify hypotheses. For example,

$$H_1 : \quad \mu_5 = \mu_3 > \{\mu_1, \mu_4\} > \mu_2$$

$$\begin{aligned} \text{JASP :} \quad & \text{group5} = \text{group3} > \text{group1} > \text{group2} \\ & \text{group3} > \text{group4} > \text{group2} \end{aligned}$$

- Press Ctrl+Enter. Inspect and interpret output.

GORIC and GORICA in JASP

	GORIC R	GORIC JASP	GORICA R	GORICA JASP
ANOVA	✓	✓	✓	
RM-ANOVA			✓	✓
Regression	✓	Not yet	✓	
SEM		Future?	✓	Future?
Other model		Future?	✓	Future?

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GORICA

GORIC(A) in JASP

End & Extra

The End

Thanks for listening!

Are there any questions?

Websites

www.uu.nl/staff/RMKuiper/Software
www.uu.nl/staff/RMKuiper/Websites%20%2F%20Shiny%20apps
informative-hypotheses.sites.uu.nl/software/goric/

E-mail

r.m.kuiper@uu.nl

Promo: Publish in special issue

Applying GORIC(A)?

If you collected data or will collect,
if you have one or more a-priori hypotheses,
then you can apply the GORIC(A).

FYI: Special issue

Journal (mdpi): Mathematics (ISSN 2227-7390)

Special Issue:

Evaluation of Theory-Driven Hypotheses: No Hypothesis, No G(L)ORIC

Guest Editor: Rebecca M. Kuiper

https://www.mdpi.com/journal/mathematics/special_issues/97C26430R3

Contact: r.m.kuiper@uu.nl

Please contact me (r.m.kuiper@uu.nl), if you want to explore the possibilities (and to obtain discount).

What's next

Depending on time and wishes:

- Some notes
- Demo in R
- Demo in JASP
- Evidence synthesis / Support aggregation

We end with:

- Lab
- Discussion

Notes

Note on using complement

Complement not always higher weight

In case H_m is almost true, but not true:

1. H_m does not have maximum fit (but much lower penalty).
2. Support for H_m is less when evaluating it against its complement (than H_u).

This is of course a good thing, since H_m is not true.

Explanation:

Penalty of H_c is smaller than that of H_u .

Against H_u , you choose H_m 'sooner' because of low penalty for H_m .

Note on hypotheses

1. Only include hypotheses with sound theoretical and/or empirical basis.
Often a null hypothesis is not of interest.
2. Keep the number of hypotheses included as small as possible.
3. This is a subjective endeavor, aim for inter-peer / inter-subjective agreement.

Note on possibilities multiple studies

- Update GORIC(A) and GORIC(A) weights:
More data collected, (re-)calculate the GORIC(A) weights.
- Update hypotheses:
First data set (or a part of it) generates one or more hypotheses.
Other data set (or part) used to determine evidence / support.
See this html tutorial and/or this R script tutorial.
- Aggregate evidence for hypotheses:
Aggregate the support for theories (diverse designs allowed).
Bear in mind: Meta-analysis aggregates parameter estimates or effect sizes which need to be comparable (often same designs required).
See this html tutorial and/or this R script tutorial.

Note on BF as an IC

The fit in GORICA refers to the maximum log likelihood.

The fit in BF refers to the maximum likelihood.

Notably, BF can be written as an IC (and vice versa):

$$\begin{aligned} -2 \log BF_{1u} &= -2 \log \frac{f_1}{c_1} \\ &= -2 \log f_1 + 2 \log c_1 \\ &= -2 \log \text{fit } H_1 + 2 \log \text{complexity } H_1. \end{aligned}$$

Note: complexity value in BF depends on prior.

Note on GORIC weights vs BF and PMPs

$$w_m/w_{m'} = \text{relative weight} \sim \text{Bayes factor } (BF_{mm'}).$$

$w_m \sim$ posterior model probability (PMP).

$1 - w_m$ = conditional error probability.

Note: w_m depends on set of hypotheses.

Note on conditional error probability

using PMPS and GORIC(A) weights

H_m	weights
H_1 : Sex Match	.04
H_2 : Gender Role Match	.81
H_3 : Sex Mismatch	.01
H_4 : Gender Role Mismatch	.00
H_u :	.14

The conditional error probability for preferred hypotheses H_2 is $1 - .81 = .19$.

Note on conditional error probability

using PMPS and GORIC(A) weights

What if we compare many hypotheses?

H_m	weights
H_1 : Sex Match	.013
H_2 : Gender Role Match	.270
H_3 : Sex Mismatch	.003
H_4 : Gender Role Mismatch	.000
H_5 : Lets try this one too	.180
...	...
H_{12} : Don't miss something	.040
H_u :	.047

The conditional error probability for H_2 becomes $1 - .27 = .77$ (was .19).

namely, included additional hypotheses may obtain some support.
Hence, the conditional error probabilities becomes larger.

Note on conditional error probability

using PMPS and GORIC(A) weights

Hypotheses that overlap can also share support (like H_u does).

My advise:

Only use error probabilities if one hypothesis versus its complement.

(Or when you are sure that there is no overlap in hypotheses)

H_m	weights
H_2 : Gender Role Match	.82
H_c : complement H_2	.18

Extra material

- GORICA on SEM, using R scripts from https://github.com/rebeccakuiper/GORICA_in_SEM.
This is material that belongs to the 'GORICA on SEM' article on <https://www.tandfonline.com/doi/full/10.1080/10705511.2020.1836967>.
- GORICA on cross-lagged panel model (CLPM), using R scripts from https://github.com/rebeccakuiper/GORICA_in_CLPM.

Note: On github site, go to Code (green button) and download zip.

Simulation study

Properties of Methods: Simulation study

1. All methods (e.g., (G)ORIC and BMS) programmed in Fortran.
2. Generate data sets with know structure (conditions).
3. Apply methods to all data sets (within one condition).
4. Summarize results (per condition):
proportion of (correct) chosen hypotheses.

Reference:

Kuiper, R.M., Nederhof, T., and Klugkist, I. (2015). Properties of hypothesis testing techniques and (Bayesian) model selection for exploration-based and theory-based (order-restricted) hypotheses. *British Journal of Mathematical and Statistical Psychology*, 68(2), 220 – 245.

Population Values in Simulated Data Sets when $k = 3$

Values of Population Means (μ_i) and Standard Deviations (σ_i) Used to Simulate 1000 Data Sets

$k = 3$	Case Number ('ES')	μ_1	μ_2	μ_3	ES (Cohen)
	0	0.00	0.00	0.00	
	1	-0.10	0.00	0.10	.1
	2	-0.20	0.00	0.20	.2
	3	-0.30	0.00	0.30	.3
	4	-0.40	0.00	0.40	.4
	5	-0.50	0.00	0.50	.5
		σ_1	σ_2	σ_3	
	0 - 5	1.00	1.00	1.00	

NB in Case 0 $H_{30C} : \mu_1 = \mu_2 = \mu_3$ is true
 in Cases 1 to 5 $H_{31C} : \mu_1 < \mu_2 < \mu_3$ is true.

Results Exploration vs Confirmatory ($k = 3$ and $n = 50$)

$H_{30E} : \mu_1 = \mu_2 = \mu_3$	$H_{30C} : \mu_1 = \mu_2 = \mu_3$
$H_{31E} : \mu_1 = \mu_2, \mu_3$	$H_{31C} : \mu_1 < \mu_2 < \mu_3$
$H_{32E} : \mu_1, \mu_2 = \mu_3$	$H_{32C} : \mu_1 = \mu_2 < \mu_3$
$H_{33E} : \mu_1 = \mu_3, \mu_2$	$H_{33C} : \mu_1 < \mu_2 > \mu_3$
$H_{34E} : \mu_1, \mu_2, \mu_3$	$H_{34C} : \mu_1, \mu_2, \mu_3$

'ES'	Method	H_{30E}	H_{31E}	H_{32E}	H_{33E}	H_{34E}
0	PCIC: AIC	0.658	0.113	0.115	0.112	0.002
2	PCIC: AIC	0.201	0.356	0.358	0.028	0.057
5	PCIC: AIC	0.000	0.135	0.136	0.000	0.729

'ES'	Method	H_{30C}	H_{31C}	H_{32C}	H_{33C}	H_{34C}
0	ORIC	0.662	0.065	0.119	0.120	0.034
2	ORIC	0.148	0.506	0.311	0.029	0.006
5	ORIC	0.000	0.955	0.044	0.001	0.000

Results Confirmation ($k = 3$ and $n = 50$)

$H_{30C} : \mu_1 = \mu_2 = \mu_3$

$H_{31C} : \mu_1 < \mu_2 < \mu_3$

$H_{32C} : \mu_1 = \mu_2 < \mu_3$

$H_{33C} : \mu_1 < \mu_2 > \mu_3$

$H_{34C} : \mu_1, \mu_2, \mu_3$

'ES'	Method	'Prior'	H_{30C}	H_{31C}	H_{32C}	H_{33C}	H_{34C}
0	ORIC		0.662	0.065	0.119	0.120	0.034
0	BMS	3	0.865	0.012	0.075	0.036	0.012
0	BMS	2	0.774	0.022	0.124	0.058	0.022
0	BMS	1	0.656	0.047	0.147	0.120	0.033
2	ORIC		0.148	0.506	0.311	0.029	0.006
2	BMS	3	0.299	0.299	0.371	0.031	0.001
2	BMS	2	0.252	0.328	0.381	0.040	0.000
2	BMS	1	0.125	0.494	0.321	0.063	0.000
5	ORIC		0.000	0.955	0.044	0.001	0.000
5	BMS	3	0.000	0.887	0.112	0.001	0.000
5	BMS	2	0.001	0.890	0.107	0.002	0.000
5	BMS	1	0.000	0.928	0.070	0.002	0.000

Results Confirmation ($k = 3$ and $n = 50$)

$$H_{30C} : \mu_1 = \mu_2 = \mu_3$$
$$H_{31C} : \mu_1 < \mu_2 < \mu_3$$
$$H_{32C} : \mu_1 = \mu_2 < \mu_3$$
$$H_{33C} : \mu_1 < \mu_2 > \mu_3$$
$$H_{34C} : \mu_1, \mu_2, \mu_3$$

1 – Type I error (here):

not preferring the true null hypothesis,
given the set of hypotheses (!).

'ES'	Method	'Prior'	H_{30C}	H_{31C}	H_{32C}	H_{33C}	H_{34C}
0	ORIC		0.662	0.065	0.119	0.120	0.034
0	BMS	3	0.865	0.012	0.075	0.036	0.012
0	BMS	2	0.774	0.022	0.124	0.058	0.022
0	BMS	1	0.656	0.047	0.147	0.120	0.033
2	ORIC		0.148	0.506	0.311	0.029	0.006
2	BMS	3	0.299	0.299	0.371	0.031	0.001
2	BMS	2	0.252	0.328	0.381	0.040	0.000
2	BMS	1	0.125	0.494	0.321	0.063	0.000
5	ORIC		0.000	0.955	0.044	0.001	0.000
5	BMS	3	0.000	0.887	0.112	0.001	0.000
5	BMS	2	0.001	0.890	0.107	0.002	0.000
5	BMS	1	0.000	0.928	0.070	0.002	0.000

Results Confirmation ($k = 3$ and $n = 50$)

$$H_{30C} : \mu_1 = \mu_2 = \mu_3$$

$$H_{31C} : \mu_1 < \mu_2 < \mu_3$$

$$H_{32C} : \mu_1 = \mu_2 < \mu_3$$

$$H_{33C} : \mu_1 < \mu_2 > \mu_3$$

$$H_{34C} : \mu_1, \mu_2, \mu_3$$

1 – Type II error (here):
not preferring the true hypothesis,
given the set of hypotheses (!).

'ES'	Method	'Prior'	H_{30C}	H_{31C}	H_{32C}	H_{33C}	H_{34C}
0	ORIC		0.662	0.065	0.119	0.120	0.034
0	BMS	3	0.865	0.012	0.075	0.036	0.012
0	BMS	2	0.774	0.022	0.124	0.058	0.022
0	BMS	1	0.656	0.047	0.147	0.120	0.033
2	ORIC		0.148	0.506	0.311	0.029	0.006
2	BMS	3	0.299	0.299	0.371	0.031	0.001
2	BMS	2	0.252	0.328	0.381	0.040	0.000
2	BMS	1	0.125	0.494	0.321	0.063	0.000
5	ORIC		0.000	0.955	0.044	0.001	0.000
5	BMS	3	0.000	0.887	0.112	0.001	0.000
5	BMS	2	0.001	0.890	0.107	0.002	0.000
5	BMS	1	0.000	0.928	0.070	0.002	0.000

Extra: Prior choice for BMS

Prior

$$P(\boldsymbol{\mu}, \sigma^2) = P(\mu_1) \times \dots \times P(\mu_k) \times P(\sigma^2),$$

where $P(\mu_i) = P(\mu) \forall i = 1, \dots, k$.

$P(\mu)$ is a data-based normal distribution, with parameters β_0 and τ_0^2 , and $P(\sigma^2)$ an inverse chi-squared distribution (see Klugkist et al., 2005)

β_0 and τ_0^2 data-based hyperparameters

For each μ_i ($i = 1, \dots, k$) a credibility interval is computed:

estimated group mean \pm **PV** \times *the group standard error*.

The credibility interval for μ_i is represented by " $[LB_i, UB_i]$ ".

Lowest lower bound: LB_{min} .

Highest upper bound: UB_{max} .

Then, $\beta_0 = \frac{LB_{min} + UB_{max}}{2}$ and $\tau_0^2 = [\frac{UB_{max} - LB_{min}}{2}]^2$.

Three types of prior vagueness: BMS PV 1, BMS PV 2 and BMS PV 3