Examples of focused model comparison: skew-normal models

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Abstract

The following example (from Claeskens and Hjort (2008)) illustrates how focused model comparison can be performed using the **fic** package in a situation where:

- a novel class of models is defined and fitted by custom R functions
- a simple model is extended in two different directions to define the models being compared

Keywords: models.

1. Skew-normal models

An outcome y_i and a covariate x_i are observed for individuals i = 1, ..., n. Four different models are compared: two normal models with a constant variance, one without (1) and one with (2) a linear regression term, and two "skew-normal" models without (3) and with (4) the linear regression term. The skew-normal model is defined by an error term $\sigma \epsilon_i$, where the ϵ_i are independently distributed with a density $f(u|\lambda) = \lambda \Phi(u)^{\lambda-1} \phi(u)$. All four models are nested in the "wide" model (4).

(1)
$$y_i \sim N(\beta_0, \sigma^2)$$

(2)
$$y_i \sim N(\beta_0 + \beta_1 x_i, \sigma^2)$$

(3)
$$y_i = \beta_0 + \sigma \epsilon_i$$

(4)
$$y_i = \beta_0 + \beta_1 x_i + \sigma \epsilon_i$$

2. Fitting the models in R

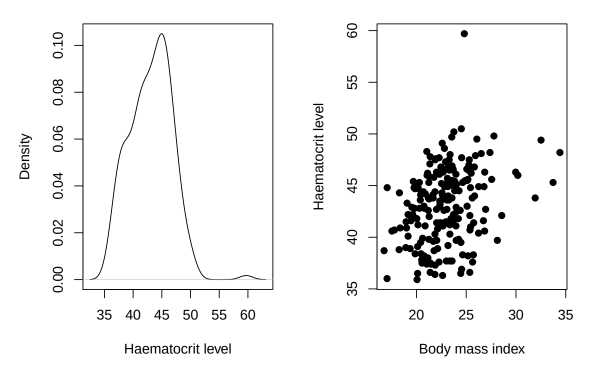
To implement this class of models in R, firstly we define the log density function of the general skew-normal model with mean, scale and skewness parameters μ , σ , λ indicated by arguments mean, sigma and lambda. This defines the distribution of y_i in model (3) with mean $\mu_i = \beta_0$, and in (4) with $\mu_i = \beta_0 + \beta_1 x_i$. Models (1) and (2) are defined by setting $\lambda = 1$ in models (3) and (4) respectively.

```
ldsnorm <- function(x, mean, sd, lambda){
    log(lambda) + (lambda-1)*pnorm(x, mean, sd, log=TRUE) +
        dnorm(x, mean, sd, log=TRUE)
}</pre>
```

The models are fitted to data from the Australian Institute of Sports (Cook and Weisberg 1994), available as **ais** from the **sn** package (Azzalini 2018). The outcome y_i is haematocrit level Hc, and the covariate is body mass index BMI. Observe the skewed distribution of the outcome and a mild association between the variables.

```
if (!require("sn"))
    stop("The `sn` package should be installed to run code in this vignette")
data(ais)
par(mfrow=c(1,2))
plot(density(ais$Hc), xlab="Haematocrit level")
plot(ais$BMI, ais$Hc, pch=19,
    xlab="Body mass index", ylab="Haematocrit level")
```

density.default(x = ais\$Hc)



The following defines the minus log likelihood for these data as a function of four parameters β_0, β_1, σ and λ .

```
mloglik <- function(b0, b1, sd, lambda){
    -sum(ldsnorm(ais$Hc, b0 + b1*ais$BMI, sd, lambda))
}</pre>
```

Then to obtain the maximum likelihood estimates under models 1 to 4, mloglik is rewritten as a function of a single vector par, containing either 2, 3 or 4 parameters to be minimised over, depending on the model. Note here that models 1 and 3 have β_2 fixed at 0, and models 1 and 2 have λ fixed at 1. The positive parameters σ and λ will be estimated by unconstrained maximisation on the log scale. These functions can be passed to the nlm function for optimisation.

```
fn1 <- function(par) mloglik(par[1], 0, exp(par[2]), 1)
fn2 <- function(par) mloglik(par[1], par[2], exp(par[3]), 1)
fn3 <- function(par) mloglik(par[1], 0, exp(par[2]), exp(par[3]))
fn4 <- function(par) mloglik(par[1], par[2], exp(par[3]), exp(par[4]))</pre>
```

nlm also requires plausible initial values for the parameters. A vector of these (ini) is obtained as follows by fitting model (2) with lm and extracting the coefficients with coef (for β_0 and β_1) and the residual standard deviation for $\log(\sigma)$. An initial value of 0 is used for $\log(\lambda)$.

The appropriate objective function (fn1-fn4) is then optimised for each model, starting from the given initial values².

```
opt1 <- nlm(fn1, ini[c("beta0","logsigma")], hessian=TRUE)
opt2 <- nlm(fn2, ini[c("beta0","beta1","logsigma")], hessian=TRUE)
opt3 <- nlm(fn3, ini[c("beta0","logsigma","loglambda")], hessian=TRUE)
opt4 <- nlm(fn4, ini, hessian=TRUE)</pre>
```

Finally, the information required by the fic function (the estimates and covariance matrices) is extracted from the nlm results and arranged into a list, for each of the four models.

```
mod1 <- list(est=opt1$estimate, vcov=solve(opt1$hessian) )
mod2 <- list(est=opt2$estimate, vcov=solve(opt2$hessian) )
mod3 <- list(est=opt3$estimate, vcov=solve(opt3$hessian) )
mod4 <- list(est=opt4$estimate, vcov=solve(opt4$hessian) )</pre>
```

¹Note these are the exact maximum likelihood estimates for model 2. The added value of nlm in fitting models 1 and 2, compared to simply using lm, is to conveniently provide the covariance matrix at the maximum likelihood estimates, which is required for focused model comparison

²A warning message of NA/Inf replaced by maximum positive value can be ignored and is the result of nlm trying out extreme and implausible values on the way to finding the maximum likelihood.

3. Focused model comparison

We now perform a focused comparison of the four models. Two alternative focuses are investigated: the mean and median outcome at a covariate value of interest. Expressions for the mean and median of the skew normal, in the parameterisation used here, are given by Claeskens and Hjort (2008) and implemented in the following R functions:

```
mean_snorm <- function(mu, sigma, lambda){
    f <- function(u){u*exp(ldsnorm(u, 0, 1, lambda))}
    mu + sigma * integrate(f, -Inf, Inf)$value
}
median_snorm <- function(mu, sigma, lambda){
    mu + sigma * qnorm(0.5^(1/lambda))
}</pre>
```

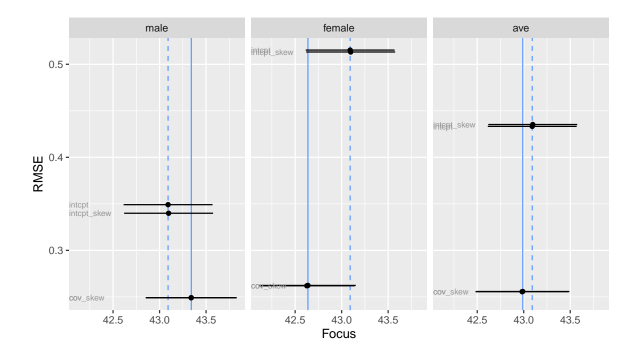
As described in the main fic package vignette, the focus function supplied to fic should have arguments defined by a vector par of parameters of the biggest model (in this case the four-parameter model 4), and a matrix of covariate values X, and return the corresponding focus quantity. In this example, the two focus functions are

The inds matrix, required by fic, is now constructed. Recall this indicates which parameters (columns) are included in each of the models (rows) being compared. The narrow model is in the first row, and the wide model in the last. The rows are given names to describe the models, so the output is easier to read.

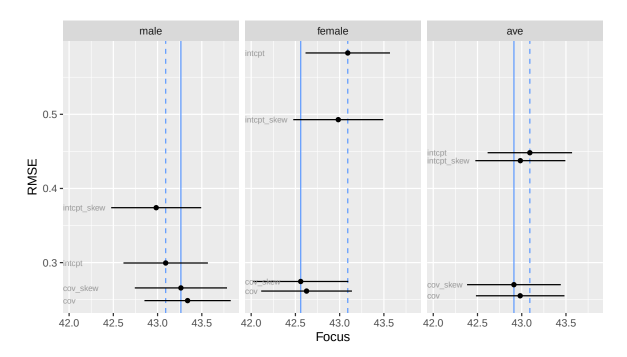
The functions **fns** required to extract the estimates and covariance matrix from the fitted model objects are then defined.

Finally fic is called to compare the four models for focuses defined by the mean and median for average men and women, with covariate values defined by med.bmi. The sub argument is supplied to ensure the focus estimates are returned too – note that fic can only automatically fit the submodels for standard R model classes such as glm.

```
vcov=function(x)x$vcov,
       nobs=function(x)nrow(ais))
med.bmi <- rbind(male=23.56, female=21.82)</pre>
fmean <- fic(mod4, inds=inds, fns=fns, focus=focus1, X=med.bmi, FIC=TRUE,</pre>
        sub=list(mod1, mod2, mod3, mod4))
fmean
##
     vals
            mods rmse rmse.adj
                                se
                                   FIC focus
                           bias
## 1
     male
           ## 4
     male
             cov 0.249 0.249 0.000 0.249 1.05 43.3
     ## 7
## 10
     male cov_skew 0.249 0.249 0.000 0.249 1.05 43.3
## 2 female
           intcpt 0.515  0.515  0.454  0.244  41.62  43.1
## 5 female
             ## 11 female
## 3
     ave
          0.256 0.000 0.256 1.30 43.0
## 6
             cov 0.256
     ave
## 9
     ## 12
        cov_skew 0.256
                    0.256 0.000 0.256 1.32 43.0
     ave
fmed <- fic(mod4, inds=inds, fns=fns, focus=focus2, X=med.bmi, FIC=TRUE,</pre>
       sub=list(mod1, mod2, mod3, mod4))
fmed
            mods rmse rmse.adj
##
     vals
                           bias
                                se
                                  FIC focus
## 1
     male
           intcpt 0.299 0.299 -0.174 0.244 6.14 43.1
## 4
     male
            cov 0.242 0.249 0.000 0.249 2.12
     ## 7
     male cov_skew 0.266 0.266 0.000 0.266 4.60 43.3
## 10
           ## 2 female
## 5 female
             cov 0.256 0.262 0.000 0.262 4.55 42.6
## 11 female
          cov_skew 0.275
                    0.275 0.000 0.275 6.50 42.6
## 3
     ave
          intcpt 0.448   0.448   0.376   0.244   30.29   43.1
## 6
             cov 0.249
                    0.255 0.000 0.255 2.24 43.0
     ave
## 9
     ## 12 ave cov_skew 0.270 0.270 0.000 0.270 4.46 42.9
```



ggplot_fic(fmed)



For both the mean and the median, including the covariate is judged to be essential. Including the skewness is pointless for estimating the mean, as discussed by Claeskens and Hjort (2008), the skewness term provides no extra information. The utility of including the skewness is also doubtful for estimating the median too, given the covariate is includes, the focus estimates and RMSE are very similar whether or not the skewness is also included.

References

Azzalini A (2018). The R package sn: The Skew-Normal and Related Distributions such as the Skew-t (version 1.5-3). Università di Padova, Italia. URL http://azzalini.stat.unipd.it/SN.

Claeskens G, Hjort N (2008). *Model selection and model averaging*. Cambridge University Press.

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