Different models for different purposes: focused model comparison in R

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Abstract

Typical methods of model comparison pick one "best" model, no matter what the estimates from the model are used for. "Focused" model comparison, by contrast, considers that different models may be better for different purposes. Different models may be preferred for estimating different "focus" quantities, functions of the basic parameters.

In the "focused information criterion" of Claeskens and Hjort (2006), data are assumed to be generated by a "wide" model, in which all models we would consider are nested. Fitting the wide model to the observed data, however, may give estimates that are not sufficiently precise. Therefore we might accept some bias in the estimate in return for greater precision. The optimal submodel for a particular focus is the one which minimises the mean squared error of the estimate of that focus from the submodel, assuming that the wide model is true. [maybe other losses if can get bootstrap to work]

The fic package calculates this error, and related quantities, straightforwardly for any class of models fitted by maximum likelihood. There are shortcuts for commonly-used model classes such as GLMs and parametric survival models. Cox regression models are also supported [todo]

Keywords: FIC, model comparison, AIC, BIC.

1. Introduction: principles for model comparison

To compare a set of statistical models fitted to the same data by maximum likelihood, it is common to rank them according to some "criterion". For example, Akaike's information criterion (AIC, Akaike (1973)) takes the form

$$-2\log\ell(\hat{\theta};\mathbf{x}) + 2p$$

where $\ell(\hat{\theta}; \mathbf{x})$ is the maximised likelihood for the model fitted to the dataset \mathbf{x} , the likelihood is maximised at parameters $\hat{\theta}$, and p is the number of parameters.

The Bayesian information criterion (BIC, Schwarz (1978)) is

$$-2\log\ell(\hat{\theta}; \mathbf{x}) + p\log(n)$$

These two criteria are based on very different principles. Thus they often rank models differently. The AIC is designed to choose the model with the best predictive ability, thus it tends to favour bigger models as the sample size increases. BIC is an approximation to Bayesian model comparison by Bayes factors, and selects the model with the highest posterior probability under an implicit weak prior (with an amount of information equivalent to one

observation, see Kass and Wasserman (1995)). If there is a "true" model, the BIC will select it "consistently" as the sample size increases. In many situations there may not be a true model, and collecting more data will uncover more complexity in the process generating the data, in which case AIC may be more suitable. See e.g. Burnham and Anderson (2003), Claeskens and Hjort (2008) for more theory behind these criteria.

However both of these methods select one "best fitting" model for a given dataset. But that might not always be appropriate. Different models may be better for different purposes. This is the idea behind "focused" model comparison.

[describe what package does, structure of paper]

2. Focused model comparison: principles and formulae

Suppose the range of models we are willing to use is bounded by

- ullet a wide model, in which all models we would use are nested, with parameters $(m{ heta}, \gamma)$
- a narrow model, the smallest model we are willing to use, defined by setting $\gamma = \gamma_0$ in the wide model.

[list of examples. covariate selection obvious, but lots more too]

Suppose also that the purpose of the model is to estimate some *focus* quantity, which could be any function of the basic parameters

$$\mu = g(\boldsymbol{\theta}, \boldsymbol{\gamma})$$

In focused model comparison, we prefer models which give better estimates of μ . A typical way to define "better" is by the *mean square error*. The mean square error of the estimate $\hat{\mu}_S$ under a submodel S of the wide model, compared to the true value μ , is

$$E\left\{(\hat{\mu}_S - \mu)^2\right\}$$

This expectation is calculated under the assumption that the data are generated from the wide model. While we believe the wide model is the most realistic, we also accept that there may not be enough data to give sufficiently precise estimates of μ . Therefore we are willing to accept some bias in this estimate, in return for a smaller variance, by selecting a smaller model than the wide model. The submodel S with the lowest mean square error is the one which makes the optimal trade-off between bias and variance.

The mean squared error MSE_S under model S can be decomposed as a sum of the squared bias B_S^2 and the variance V_S .

$$MSE_S = E\{(\hat{\mu}_S - \mu)^2\} = \{E(\hat{\mu}_S) - \mu\}^2 + E\{(\hat{\mu}_S - E(\hat{\mu}_S))^2\}$$

= $B_S^2 + V_S$ (1)

Estimators for these quantities are constructed by Claeskens and Hjort (2003) under an asymptotic framework in which the data are assumed to be n independent observations generated

from the wide model, but reparameterised so that $\gamma = \gamma_0 + \delta/\sqrt{n}$. Thus as the sample size increases, we aim to detect more subtle departures from the narrow model.

[any more basic assumptions?]

An obvious estimator for the bias B_S is $\hat{B_S} = \hat{\mu_S} - \hat{\mu_W}$, where $\hat{\mu_W}$ is the estimate of the focus quantity under the wide model, which is assumed to be unbiased. However, Claeskens and Hjort (2003) derive a more accurate estimator for the *squared* bias as

$$\widehat{B_S^2} = (\hat{\psi}_W - \hat{\psi}_S)^2 / n$$

where

- $\hat{\psi}_W = \hat{\omega}^T \hat{\delta}$ and $\hat{\psi}_S = \hat{\omega}^T G_S \hat{\delta}$ are estimates of $\omega^T \delta$ under the wide model and submodel respectively, where $\omega^T \delta$ is the bias of the estimate of $\sqrt{n}\mu$ under the narrow model N, that is, the asymptotic mean of $\sqrt{n}(\hat{\mu}_N \mu)$. Thus ω acts as a linear transformation from the biases of the basic parameters γ to the biases of the focus parameter μ . [why not let all the ns cancel, and explain everything in terms of biases on the focus scale?]
- ω is estimated as $\hat{\omega} = J_{10}J_{00}^{-1}\frac{d\mu}{d\theta} \frac{d\mu}{d\gamma}$ using Taylor approximation arguments, where J is the information (inverse covariance) matrix under the wide model divided by n, [but the ns cancel, so could we work in terms of the info under the wide model?] and subscripts 0 and 1 select the rows and columns forming the submatrices of J that correspond to parameters θ and γ respectively. The partial derivatives of the focus μ are evaluated at the estimates from the wide model.
- $G_S = \pi^T Q_S \pi Q^{-1}$ is an estimate of the transformation that maps the wide model estimate of δ to the submodel S estimate, where $Q_S = (\pi Q^{-1} \pi^T)^{-1}$, $Q^{-1} = J_{11}$ and π is the projection matrix consisting of 0s and 1s which maps a vector of the same length as (θ, γ) to a subvector containing the elements corresponding to submodel S. [again don't the ns in the Q_S cancel]
- $\hat{\boldsymbol{\delta}} = \hat{\boldsymbol{\gamma}} \sqrt{n}$, where $\hat{\boldsymbol{\gamma}}$ is the estimate of $\boldsymbol{\gamma}$ under the wide model.

The estimator for the variance of $\hat{\mu}_S$ under the wide model, derived by Claeskens and Hjort (2003), is

$$\hat{V} = (\hat{\tau_0}^2 + \hat{\omega}^T Q_S^0 \hat{\omega})/n$$

where $\hat{\tau_0}^2/n$ estimates the variance of the narrow model focus estimate (using "delta method" principles, $\hat{\tau}_0^2 = \frac{d\mu}{d\theta}^T J_{00}^{-1} \frac{d\mu}{d\theta}$), and the additional term $(\hat{\omega}^T Q_S^0 \hat{\omega})/n$ is the increase in variance we accept by using a wider but still misspecified model S, with $Q_S^0 = \pi^T Q_S \pi$.

[again if we define J as info instead of info/n, then we wouldn't need to supply n] Thus we compare models on the basis of the root mean square error, estimated by

$$\sqrt{\widehat{MSE}_S} = \sqrt{\widehat{B}_S^2 + \widehat{V}_S} \tag{2}$$

2.1. Bias-corrected MSE

Claeskens and Hjort (2003) derive a further correction for the bias estimator which is necessary when the above estimate is negative. The adjusted squared bias estimator is

$$\widehat{B^{*2}} = \max \left\{ 0, \quad \hat{\omega}^T (I - G_S) (\hat{\boldsymbol{\delta}} \hat{\boldsymbol{\delta}}^T - Q) (I - G_S)^T \hat{\omega} / n \right\}$$
 (3)

The corresponding estimate of the bias B is $sign(\hat{\psi}_W - \hat{\psi}_S)\sqrt{\widehat{B^{*2}}}$, and the bias-corrected root MSE is $\sqrt{\widehat{MSE}} = \sqrt{\widehat{B^{*2}} + \hat{V}}$.

2.2. Ingredients needed for the MSE calculation

Therefore in order to calculate MSE_S for a submodel S, we just need to know

- the estimates $\hat{\boldsymbol{\theta}}_W$ and $\hat{\boldsymbol{\gamma}}_W$ under the wide model,
- \bullet the information matrix J or covariance matrix of these estimates,
- the focus function $\mu(\boldsymbol{\theta}, \boldsymbol{\gamma})$ and its derivatives, evaluated at $\hat{\boldsymbol{\theta}}_W, \hat{\boldsymbol{\gamma}}_W,$
- the definition of which parameters are included in submodel S and which are included in the narrow model N.
- [doubtful we also need to know n]

Claeskens and Hjort (2003) define the "focused information criterion" (FIC), which has a slightly simpler form due to excluding terms common to all submodels S, and is related to the MSE as

$$FIC_S = MSE_S - \hat{\tau}_0^2 + \hat{\omega}^T Q \hat{\omega} \tag{4}$$

Models with lower FIC give better estimates of the focus quantity. However we prefer to use the (root) MSE as the model comparison statistic, due to its direct interpretation as the error of the focus estimate.

2.3. Average MSE over a range of focuses

Often we want a model that performs well in a range of situations. In covariate selection problems, for example, we might want to estimate a focus quantity accurately for a defined range of covariate values. We might simply define the "averaged MSE"

$$AMSE = \int MSE(u)dW(u)du$$

as a weighted average of the mean squared errors (1) for focuses defined by different covariate values u, weighted by their prevalence W(u). However Claeskens and Hjort (2008) derived an alternative formula, so that if bias correction analogous to (3) is required, it only needs to be performed once.

$$AMSE = max(IS, 0) + IIS \tag{5}$$

where

$$IS = Tr((I - G_S)(\hat{\boldsymbol{\delta}}\hat{\boldsymbol{\delta}}^T - Q)(I - G_S)^T A), \quad IIS = Tr(Q_S^0 A)$$

$$A = J_{10}J_{00}^{-1}B_{00}J_{00}^{-1}J_{01} - J_{10}J_{00}^{-1}B_{01} - B_{10}J_{00}^{-1}J_{01} + B_{11}$$

$$B = \int \begin{pmatrix} d\mu(u)/d\boldsymbol{\theta} \\ d\mu(u)/d\boldsymbol{\gamma} \end{pmatrix} \begin{pmatrix} d\mu(u)/d\boldsymbol{\theta} \\ d\mu(u)/d\boldsymbol{\gamma} \end{pmatrix}^T dW(u) = \begin{pmatrix} B_{00} & B_{01} \\ B_{10} & B_{11} \end{pmatrix}$$

This is equivalent to AIC [where we average over all covariates in the data ?] [What about equivalence to AIC where the focus is the log-likelihood?]

3. Example: covariate selection in logistic regression

This example is used by Claeskens and Hjort (2008) (Example 6.1) to illustrate the focused information criterion. The dataset was originally presented by Hosmer and Lemeshow (1989). Data are taken from n = 189 women with newborn babies, and the binary outcome is whether the baby is born with a weight less than 2500g. We build a logistic regression model to predict the outcome, but are uncertain about what covariates should be included.

The data are provided as an object birthwt in the fic package. This is the same as birthwt in MASS (Venables and Ripley 2002) with the addition of a few extra columns defining interactions and transformations as in Claeskens and Hjort (2008).

The following covariates are always included (coefficient vector $\boldsymbol{\theta}$)

• x_1 Weight of mother in kg, lwtkg

The following covariates will be selected from (coefficient vector γ)

- z_1 age, in years, age
- z₂ indicator for smoking, smoke
- z₃ history of hypertension, ht
- z_4 uterine irritability, ui
- interaction $z_5 = z_1 z_2$ between smoking and age, smokeage
- interaction $z_6 = z_2 z_4$ between smoking and uterine irritability, smokeui

Firstly the wide model, that includes all the above covariates, is defined and fitted.

The focus function is then defined. This should be an R function, mapping the parameters par of the wide model to the quantity of interest. The focus can optionally have an second argument. If supplied, this must be called X, and can be used to supply covariate values at which the focus function should be evaluated. Here we take the probability of low birth weight as the focus, for two covariate categories:

- 1. smokers with average or typical values of the other covariates. These values are given in the order supplied when specifying the model (for smokers: intercept, lwtkg=58.24, age=22.95, smoke=1, ht=0, ui=0, smokeage=22.95, smokeui=0).
- 2. non-smokers with average values of the other covariates

```
focus <- function(par, X)plogis(X %*% par)
vals.smoke <- c(1, 58.24, 22.95, 1, 0, 0, 22.95, 0)
vals.nonsmoke <- c(1, 59.50, 23.43, 0, 0, 0, 0, 0)
X <- rbind("Smokers"=vals.smoke, "Non-smokers"=vals.nonsmoke)</pre>
```

We can illustrate these functions by calculating the probability of low birth weight, given the parameters of the fitted wide model, for each group. This is about twice as high for smokers.

```
focus(coef(wide.glm), X=X)

## [,1]
## Smokers 0.345
## Non-smokers 0.168
```

The fic function can then be used to calculate the mean square error of the focus for one or more given submodels. For illustration we will compare two models, both including maternal weight, one including age and smoking, but the other including age, smoking and hypertension.

```
\label{eq:mod1.glm} $$ \leftarrow \mbox{glm}(\mbox{low $^{\circ}$ lwtkg + age + smoke, data=birthwt, family=binomial)} $$ mod2.glm $<-\mbox{glm}(\mbox{low $^{\circ}$ lwtkg + age + smoke + ht, data=birthwt, family=binomial)} $$
```

We supply the following arguments to the fic function.

- wide: the fitted wide model. All the model fit statistics are computed using the estimates and covariance matrix from this model. fic will automatically recognise that this is a GLM fitted by the glm function in R, and extract the relevant information.
- inds: indicators for which parameters are included in the submodel, that is, which elements of (θ, γ) are fixed to γ_0 . This should have number of rows equal to the number of submodels to be assessed, and number of columns equal to $dim(\theta) + dim(\gamma)$, the total number of parameters in the wide model, 8 in the case of wide.glm, which includes the intercept and the coefficients of seven covariates. It contains 1s in the positions where the parameter is included in the submodel, and 0s in positions where the parameter is excluded. This should always be 1 in the positions defining the narrow model, as specified in inds0 below. If just one submodel is to be assessed, inds can also be supplied as a vector of length $dim(\theta) + dim(\gamma)$.

Note that inds indexes parameters rather than linear model terms, that is, in covariate selection problems where a variable is a factor with more than two levels, inds should contain separate entries for the coefficient of each factor level relative to the baseline level, not just one entry indicating the presence of the factor as a whole. [TODO can we construct these automatically?]

• inds0 vector of indicators for which parameters are included in the narrow model, in the same format as inds. This can be omitted, in which case the narrow model is assumed to be given by the first row of inds. In this case, just the first two parameters are included, the intercept and the coefficient of lwtkg.

```
inds <- rbind(mod1 = c(1,1,1,1,0,0,0,0),

mod2 = c(1,1,1,1,1,0,0,0))

inds0 <- c(1,1,0,0,0,0,0,0,0)
```

- focus the focus function.
- sub a list of the fitted submodels to be assessed. This is optional, and is only needed so that the model comparison statistics can be presented alongside the estimates of the focus under the submodels. Note, if just one submodel is to be assessed, this should still be enclosed in a list, e.g. list(mod1.glm).

The main fic function then returns an object containing the model fit statistics and the estimate of the focus quantity for each model.

```
sub <- list(mod1.glm, mod2.glm)</pre>
fic1 <- fic(wide=wide.glm, inds=inds, inds0=inds0, focus=focus, X=X, sub=sub)
fic1
##
            vals mods
                        FIC
                               rmse rmse.adj
                                                bias bias.adj
## 1
         Smokers mod1 1.187 0.0723
                                      0.0723 0.0548
                                                       0.0459
## 4
         Smokers mod2 0.783 0.0556
                                      0.0572 0.0237
                                                       0.0000
## 2 Non-smokers mod1 1.305 0.0804
                                      0.0804 0.0765
                                                       0.0731
## 5 Non-smokers mod2 0.755 0.0596
                                      0.0596 0.0525
                                                       0.0484
## 3
             ave mod1 1.246 0.0844
                                      0.0764 0.0657
                                                       0.0610
## 6
             ave mod2 0.769 0.0678
                                      0.0576 0.0381
                                                       0.0329
         se focus
##
## 1 0.0558 0.398
## 4 0.0572 0.366
## 2 0.0334 0.243
## 5 0.0348 0.215
## 3 0.0460 0.320
## 6 0.0473 0.291
```

The object returned by fic is a matrix containing one row for each combination of focus covariate values indicated in the column vals and submodels indicated in the column mods. The focus estimate is returned in the final column focus, while the remaining columns contain the following model comparison statistics:

• FIC The FIC as originally defined by ?? (equation 4),

- rmse The root mean square error of the submodel focus estimate, calculated assuming the wide model is true (equation 2),
- rmse.adj The bias-adjusted root mean square error (Section 2.1),
- bias The estimated bias $\sqrt{\widehat{B}^2}$ (which may be undefined if \widehat{B}^2 is negative),
- bias.adj The adjusted bias estimate (Section 2.1),
- se The standard error $\sqrt{\hat{V}}$ of the submodel focus estimate, calculated assuming the wide model is true.

As well as the specific covariate categories, fic calculates model comparison statistics which are averaged over the categories, indicated by a value of ave in the column vals. TODO EXPLAIN WEIGHTS

Recall that mod2 contains one more covariate than mod1. For each of the two focuses, and the average, the unadjusted and adjusted bias estimates are lower due to the inclusion of this covariate, while the standard error se is higher. Given the lower rmse and rmse.adj under mod2, the reduction in bias is deemed to be worth the increase in uncertainty.

3.1. Comparing a wide range of models

In covariate selection problems, we may want to examine a broad range of models. The function all_inds (a wrapper around expand.grid) creates a matrix of indicators that defines all submodels spanned by a given wide model (here wide.glm and a narrow model (here defined by inds0). This function works for all classes of model objects x for which the terms(x) function is understood, which includes standard R regression models such as lm and glm. [can we use terms.formula more generally, e.g. in flexsurvreg?] Factors are handled naturally.

```
combs <- all_inds(wide.glm, inds0)</pre>
```

The resulting matrix can be used as the inds argument to fic to calculate focused model comparison statistics for all submodels in this example, again for a focus defined by the probability of low birth weight at covariate values defined by X. However before calling fic again, we redefine combs to exclude models with interactions but not both corresponding main effects.

We can actually fit the submodels in a loop as follows, by extracting the design matrix XZ of the wide model and removing the first column (the intercept). At each iteration, the

design matrix of the submodel is defined by extracting the columns of XZ indexed by the corresponding row of combs. The submodel is fitted by placing this submatrix XZi on the right hand side of the model formula passed to glm (with -1 appended to instruct glm not to include a second intercept term, since an intercept is already included in XZi). The list of fitted models sub can then be passed to the fic function, so that so that the focus estimate is displayed alongside the model comparison statistics.

```
nmod <- nrow(combs)
sub <- vector(nmod, mode="list")
XZ <- model.matrix(wide.glm)
for (i in 1:nmod){
    XZi <- XZ[,which(combs[i,]==1)]
    sub[[i]] <- glm(low ~ XZi - 1, data=birthwt, family=binomial)
}
ficres <- fic(wide=wide.glm, inds=combs, inds0=inds0, focus=focus, X=X, sub=sub)</pre>
```

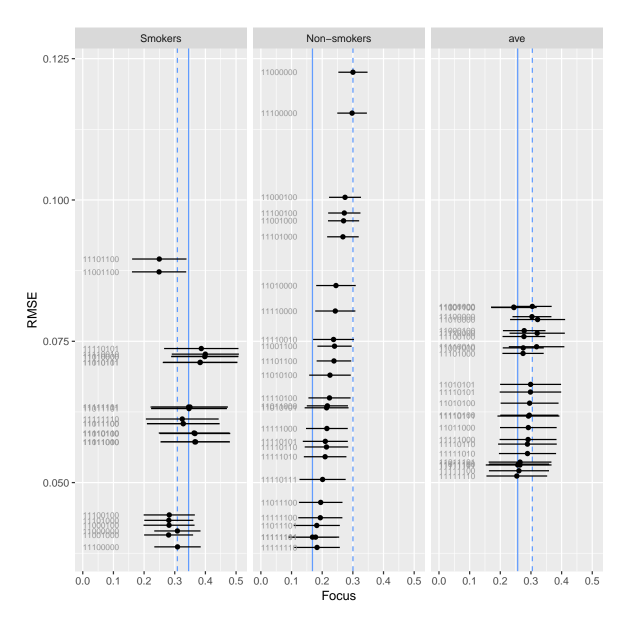
Notice that some of the rmse elements of ficres are NaN, since the first squared bias estimator $\widehat{B^2}$ is negative. The alternative estimate $\sqrt{\widehat{B^{*2}}}$, rmse.adj, can simply be used in these cases. [this raises the question: why we don't just use the adjusted one all the time. Is the unadjusted one better in any circumstances?]

A comparison of many models can be illustrated by a scatterplot of the focus estimate against the RMSE of each submodel. The default plot method for fic objects accomplishes this using base R graphics: try

```
plot(ficres)
```

Alternatively a graph can be plotted using ggplot2 if this package is installed. This is illustrated here.

```
ggplot_fic(ficres)
```



There is one panel for each of the two covariate categories (smokers and non-smokers) defining the focus (probability of low birth weight) and an average over the two categories. The solid blue line is the focus estimate under the wide model, and the dashed blue line is the focus estimate under the narrow model. An informal illustration of the uncertainty around the estimate of the focus quantity from each submodel is given by the estimate $\pm 1.96 \times \sqrt{\hat{V}}$.

Each submodel is labelled faintly using the row names of the matrix supplied as the inds argument to fic. In this case, these names were automatically constructed by the function all_inds and contain a string of binary 0/1 indicators for the inclusion of eight parameters. For smokers, the narrow model (labelled 11000000) and similar smaller models give estimates of the probability of low birth weight with the lowest MSE, while by contrast, for non-smokers, the wide model (labelled 11111111) and similar larger models give the most accurate estimates of the focus quantity. Note that in this dataset, there are 115 non-smokers and 74 smokers, thus more data enables bigger models to be identified for non-smokers. [discuss average?]

4. Other classes of models

Illustrate new concepts. Could be in separate vignettes.

Linear Illustrates alternative focuses: expected outcome at given covariate values, quantile for given covariate values. Polynomial order selection as well as covariate selection. Use a well known dataset, e.g. mtcars?

Multi-state Illustrates focuses that are complicated functions of the model parameters: package should facilitate this

Survival Increasingly flexible parametric models. Challenge to express models as nested within each other. Important health economic application: restricted mean survival.

Skew-normal Novel class of models, user-written model fitting function. Store est, vcov, nobs in result list. Narrow model parameters could be in the middle.

4.1. Calling "fic" for an unfamiliar class of models

Above, the fic function recognised the fitted model objects as GLMs, that is, objects of class "glm" returned by the glm() function in base R. But the package can be used to calculate focused model comparison statistics for any class of models, not just the special classes it recognises. To do this, it needs to know where three things are stored inside the fitted model objects:

- 1. coef: the vector of maximum likelihood estimates $(\hat{\boldsymbol{\theta}}, \hat{\boldsymbol{\gamma}})$,
- 2. nobs: the number of observations n contributing to the model fit,
- 3. vcov: the covariance matrix of the maximum likelihood estimates, $(nJ)^{-1}$.

Given a fitted model object called mod, the fic() function assumes by default that coef(mod), nobs(mod) and vcov(mod) respectively return these pieces of information. If one or more of these assumptions is not true, the defaults can be changed by supplying the argument fns to fic(), which should be a named list of three components. Each component should be a function with one argument (the fitted model) which extracts the required information from the fitted model and returns it. For example, the first component of the list below is a function which, when applied to a glm object, returns the maximum likelihood estimates of the regression coefficients. (TODO better explained with a more obscure class?)

5. Focused covariate selection in Cox proportional hazards regression

In a Cox regression model, time-to-event outcomes t_i are observed on individuals i, potentially with right-censoring. At time t, individual i is assumed to have a hazard $h_i(t)$ which is proportional to their covariate values. We wish to select between models that have different sets of covariates. In the most general "wide" model, $h_i(t) = h_0(t) \exp(\boldsymbol{\theta}^T x_i + \boldsymbol{\gamma}^T z_i)$. The baseline hazard $h_0(t)$ is left unspecified, while $\boldsymbol{\theta}$ and $\boldsymbol{\gamma}$ are estimated by maximum partial likelihood.

We compare submodels of this wide model, which include different subsets of covariates, according to how accurately they estimate some focus quantity $\mu = \mu(\theta, \gamma, H_0()|\mathbf{x}, t)$, where $H_0()$ is the cumulative baseline hazard function. Typical focus quantities might depend on depend on time t as well as covariate values \mathbf{x} (REF EARLIER), e.g. the probability that a person with covariates \mathbf{x} will survive t years.

Again the mean square error $MSE_S = B_S^2 + V_S$ of μ of the focus quantity under submodel S is estimated as $\widehat{B_S^2} + \widehat{V_S}$, using similar formulae to REF (CITE book, and/or paper?)

•
$$\widehat{B}_S^2 = (\hat{\psi}_W - \hat{\psi}_S)^2 / n$$
, where $\hat{\psi}_W = (\hat{\omega} - \hat{\kappa})^T \hat{\delta}$ and $\hat{\psi}_S = (\hat{\omega} - \hat{\kappa})^T G_S \hat{\delta}$.

•
$$\hat{V} = \{\hat{\tau}_0^2 + (\hat{\omega} - \hat{\kappa})^T Q_S^0(\hat{\omega} - \hat{\kappa})\}/n$$

$$\bullet \hat{\omega} = J_{10}J_{00}^{-1}\frac{d\mu}{d\theta} - \frac{d\mu}{d\gamma}$$

• $\hat{\kappa}(t) = (J_{10}J_{00}^{-1}F_0(t) - F_1(t))\frac{d\mu}{dH_0}$ [intuitive explanations of $\hat{\omega}$ and $\hat{\kappa}$, note the formulae are like before except with $\hat{\omega} + \hat{\kappa}$ instead of $\hat{\omega}$]

Requires some new quantities

$$F(t) = \int_0^t G_n^{(1)} / G_n^{(0)} dH_0(u) = \begin{pmatrix} F_0(t) \\ F_1(t) \end{pmatrix}$$

where $F_0(t)$ and $F_1(t)$ have p and q components, and

•
$$G_n^{(0)} = \frac{1}{n} \sum_{i=1}^n Y_i(u) \exp(x_i^T \theta + z_i^T \gamma)$$

•
$$G_n^{(1)} = \frac{1}{n} \sum_{i=1}^n Y_i(u) \exp(x_i^T \boldsymbol{\theta} + z_i^T \boldsymbol{\gamma}) \begin{pmatrix} x_i \\ z_i \end{pmatrix}$$

•
$$Y_i(t) = I(t_i \ge t)$$
 at-risk indicator

other quantities e.g. $Q_S^0, G_S, \hat{\delta}, \tau_0^2, J_{10}, J_{00}$ are defined as before, using the partial likelihood where necessary instead of the likelihood. The asymptotic theory underlying these formulae is discussed by CITE

hazard or cumulative hazard

6. Bootstrap

Would like to

- illustrate alternative losses to the mean square error
- use resampling to illustrate FIC principles

7. Discussion

Post-selection inference, model averaging High dimensional regression

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