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<td>Author(s)</td>
<td>Krnjajic, Milovan</td>
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<td>Publication Date</td>
<td>2011</td>
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<td>Item record</td>
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Bayesian model specification: some problems related to model choice and calibration

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Abstract

In the development of Bayesian model specification for inference and prediction we focus on the conditional distributions \( p(\theta|B) \) and \( p(D|\theta, B) \), with data \( D \) and background assumptions \( B \), and consider calibration (an assessment of how often we get the right answers) as an important integral step of the model development. We compare several predictive model-choice criteria and present related calibration results. In particular, we have implemented a simulation study to compare predictive model-choice criteria \( LS_{CV} \), a log-score based on cross-validation, \( LS_{FS} \), a full-sample log score, with deviance information criterion, \( DIC \). We show that for several classes of models \( DIC \) and \( LS_{CV} \) are (strongly) negatively correlated; that \( LS_{FS} \) has better small-sample model discrimination performance than either \( DIC \), or \( LS_{CV} \); we further demonstrate that when validating the model-choice results, a standard use of posterior predictive tail-area for hypothesis testing can be poorly calibrated and present a method for its proper calibration.

Keywords: log-score, deviance information criterion, posterior predictive tail areas, hypothesis testing.

Introduction

Bayesian approach to modeling comprises inference, prediction and decision-making and considers three main objects: \( \theta \), a model parameter vector; \( D \), an information (data) source about \( \theta \); and \( B \), a set of propositions summarizing background assumptions about of \( \theta \) and \( D \), for example, that \( \theta > 0 \) if \( \theta \) represents the mean remission time for a specified set of patients with a given disease; or that the data set arose as the result of a randomized controlled trial with the specified design). From the results of Cox (1946) and Ramsey (1926) each of these three basic Bayesian statistical activities is governed conceptually by a single equation and requires a series of specification tasks:

- (inference) \( p(\theta|D, B) = c p(\theta|B) p(D|\theta, B) \), where \( c > 0 \), and \( p(\theta|D, B) \) posterior distribution, quantifies the information about \( \theta \), both internal and external to \( D \);
- (prediction) \( p(D^*|D, B) = \int_\Theta p(D^*|\theta, B) p(\theta|D, B) d\theta \), where \( D^* \) is future data;
- (decision) The optimal action is given by \( a^* = \arg\max_{a \in A} E_{|\theta|D, B} U(a, \theta) \).

In problems of realistic complexity it is uncertain how to specify \( p(D|\theta, B) \). In our view, a leading principle governing this specification should be calibration, which consists of checking how often one obtains the right answers. For example, a statement such as “\( p(a < \theta < b|D, B) = 0.9 \)” should be verifiably correct about 90% of the time. To address the uncertainty in specifying a model, \( p(D|\theta, B) \), we search for an ensemble, \( M \), of such specifications in a well calibrated manner, carefully avoiding a double use of data (to specify priors on model space and again to update this prior when carrying out inference and prediction).
we present calibration results related to the following basic questions in Bayesian model specification, “Q1: Is model $M_j$ better than $M_k$?” and “Q2: Is model $M_j$ good enough?” These questions are not complete without a clear reference to the purpose of the models. However, once the purpose is made explicit, the inferential task transforms into a decision problem, best solved by maximizing expectation of utility (MEU) specific for the model’s purpose.

A standard way to answer question $Q_1$ is to use Bayes factors and related criteria, for example (for reasons of space we comment on this very briefly). A well known problem with Bayes factors is a possibly extreme sensitivity to the way diffuse priors are specified on the model parameters, (see e.g. Bernardo and Smith (1994)). The consequence of this instability is that the evidence in favor of one model over the other may be made arbitrarily large, based on a range of plausible parameter values, even regardless of the data set. Motivated in part by this well known problem, we focus on stable model-choice criteria based on the posterior predictive distribution (of the future data, $D^*$, given the observed sample, $D$), which has a sound basis as a utility for model comparison and is entirely stable relative to the specification of diffuse priors: $p(D^*|D, M_j, B) = E_{(y_j|D, M_j, B)}p(D^*|y_j, M_j, B)$.

We argue that the quality of model prediction is also a solid basis for a useful generic utility in model comparison and hence we focus on working with posterior predictive distributions. In order to compare a predictive distribution with the actual data point, $y^*$, we use two log-score criteria, $LS_{CV}$, based on cross validation and defined as $n LS_{CV}(M_j|y, B) = \sum_{i=1}^{n} \log p(y_i|y_{-i}, M_j, B)$, and $LS_{FS}$, the full-sample log-score defined as $n LS_{FS}(M_j|y, B) = \sum_{i=1}^{n} \log p(y_i|y, M_j, B)$, and which uses all data in the sample only once (see, for example, Gelfand and Dey (1994) and Laud and Ibrahim (1995)). Considering how to address the question $Q_1$, we contrast deviance information criterion, $DIC$ with the log-score rules, $LS_{CV}$ and $LS_{FS}$.

The plan of the paper is as follows: In Sections 1 and 2 we present aspects of some answers to question $Q_1$ whereas Section 3 addresses a calibration issue arising from $Q_2$. Specifically, in Section 1, we consider how to obtain answers to $Q_1$, explore similarities and differences between $DIC$ and $LS_{CV}$ in Gaussian and Poisson models, and show results on the small-sample performance of $DIC$, $LS_{CV}$ and $LS_{FS}$ in discriminating between nested models. In Section 3 we show that the posterior predictive tail areas (Gelman et al. (1996)), a standard method for answering “could model $M_j$ have generated the data?” (a question related to $Q_2$) can be poorly calibrated, and we document an approach to calibrating the answer.

1 $LS_{CV}$ and $DIC$

In order to show the relationship between $LS_{CV}$ and $DIC$ as model-comparison criteria let us consider a simple parametric model, $M_0$, for continuous outcomes, where: $(y_i|\mu, B) \overset{IID}{\sim} N(\mu, s^2)$ and $(\mu|B) \sim N(a, b^2)$.

With $(s^2, a, b^2)$ known and a diffuse prior on $\mu$ (large $b^2$), the posterior for $\mu$ is: $p(\mu|y, B) \overset{\sim}{=} N(\bar{y}, s^2/n)$, where $\bar{y}$ is the sample mean of $y = (y_1, \ldots, y_n)$. The predictive distribution for the next observation is then $p(y_{n+1}|y, B) \overset{\sim}{=} N(\bar{y}, s^2/(1 + 1/n))$. Similarly, $p(y_i|y_{-i}, B) \overset{\sim}{=} N(\bar{y}_{-i}, s^2_i)$, where $\bar{y}_{-i}$ is the sample mean with observation $i$ omitted and $s^2_i = s^2(1 + 1/(n - 1))$, so that $LS_{CV}(M_0|y, B) = c_1 - c_2 \sum_{i=1}^{n} (y_i - \bar{y}_{-i})^2$ for some constants $c_1$ and $c_2$ with $c_2 > 0$.

With a bit of algebra it can be shown that $LS_{CV}(M_0|y, B) = c_1 - c_2 \sum_{i=1}^{n} (y_i - \bar{y})^2$, $(c_2 > 0)$, meaning that for $M_0$ with a diffuse prior the $LS_{CV}$ is almost perfectly negatively correlated with the sample variance.
In model $M_0$ the deviance is $D(\mu) = -2 \ln(\mu|y, B) = c_0 + c_3 \sum_{i=1}^{n} (y_i - \mu)^2$ for some $c_3 > 0$. Given a parametric model $p(y|\theta)$, Spiegelhalter et al. (2002) define the deviance information criterion (DIC) as: $DIC(M|y, B) = D(\theta) + 2\hat{p}_D$, where $p_D$ is the effective number of model parameters, and $\theta$ is the posterior mean of $\theta$, so that models with low DIC values are to be preferred over those with higher values. When $p_D$ is difficult to read directly from the model (e.g., in complex hierarchical settings with random effects), it can be estimated from standard MCMC output as $\hat{p}_D = D(\bar{\theta}) - D(\theta)$, where $D(\bar{\theta})$ is the posterior mean of the deviance and $D(\theta)$ is the deviance evaluated at the posterior mean of $\theta$.

Model $M_0$ has just one parameter ($p_D = 1$), a diffuse prior for which implies $\bar{\theta} \cong \bar{y}$, so that we get $DIC(M_0|y, B) = c_0 + c_3 \sum_{j=1}^{n} (y_j - \bar{y})^2 + 2$ concluding that

$$-DIC(M_0|y, B) = c_1 + c_2 LS_{CV}(M_0|y, B)$$

for $c_2 > 0$. In other words, in this simple setting, choosing a model by maximizing $LS_{CV}$ and by minimizing $DIC$ are approximately equivalent behaviors. This argument readily generalizes to any situation in which the predictive distribution is approximately Gaussian.

As a second example of the relationship between $LS_{CV}$ and $DIC$ we consider two models for count data a fixed-effects Poisson (FEP), model $M_1$ where $(y_i|\lambda, B) \sim \text{Poisson}(\lambda)$ and $(\lambda|B) \sim p(\lambda|B)$, and random-effects Poisson (REP), model $M_2$:

$$(y_i|\lambda, B) \overset{\text{indep}}{\sim} \text{Poisson}(\lambda_i)$$

$$\log(\lambda_i) = \beta_0 + e_i$$

$$(\beta_0, \sigma^2|B) \sim p(\beta_0, \sigma^2|B)$$

$$(e_i|\sigma^2, B) \overset{\text{indep}}{\sim} N(0, \sigma^2).$$

where $i = 1, ..., n$. $M_1$ is of course a special case of $M_2$ with $(\sigma^2 = 0, \lambda = e^{\beta_0})$; the likelihood in $M_2$ is a Lognormal mixture of Poissons.

We conducted a partial-factorial simulation study with factors \{n = 18, 32, 42, 56, 100\}, \{\beta_0 = 0.0, 1.0, 2.0\}, and \{\sigma^2 = 0.0, 0.5, 1.0, 1.5, 2.0\} in which \{(data-generating mechanism, assumed model)\} = \{(M_1, M_1), (M_1, M_2), (M_2, M_1), (M_2, M_2)\}; in each cell of this grid we used 100 simulation replications. Here we summarize only a small part of the results of this simulation (see Krnjajić (2005) for additional details).

When both the data-generating model and the assumed model were $M_1$ (the fixed-effects Poisson), $LS_{CV}$ and $DIC$ are almost perfectly negatively correlated (graph not shown); By contrast, the Figure 1 shows that when the data-generating and assumed models were $M_2$ (the random-effects Poisson), $LS_{CV}$ and $DIC$ are less strongly negatively correlated, although the correlation increases with $n$ (graph not shown).

## 2 Model-comparison criteria and small data samples

In addition to $LS_{CV}$, which requires $n$ model fitting exercises, our interest was drawn to another version of the log-score idea in which no cross-validation is employed. Instead, in the one-sample situation, for instance, it suffices to compute only a single predictive distribution $p(\cdot|y, M_j)$ for future data, for each model $M_j$ under consideration and based on the entire data set $y$. Thus, we define the full-sample log score $n LS_{FS}(M_j|y, B) = \sum_{i=1}^{n} \log p(y_i|y, M_j, B)$ (cf. Laud and Ibrahim (1995)). Remark. This appears to use the data twice, but (a) all $LS_{FS}$ is actually doing is evaluating the posterior predictive distribution for the next data value at the observed data, and (b) when $n$ is even moderate in size, any effect this may induce is
small. The calculation of $L_{FS}$, as opposed to Bayes factors, is entirely stable and does not have any difficulties related to the way diffuse priors may be specified.

Here we examine three model-choice rules: \{maximize $L_{CV}$, maximize $L_{FS}$, minimize $DIC$\}, and consider two models $M_1$ and $M_2$ to choose between. Our objective is to find out how accurately do these rules discriminate between $M_1$ and $M_2$?

As an extension of the previous simulation study, we generated data from the random-effects Poisson model $M_2$ (equation (2)) and computed $L_{CV}$, $L_{FS}$, and $DIC$ for models $M_1$ (the fixed-effects Poisson, FEP) and $M_2$ (the random-effects Poisson, REP) in the full-factorial grid $\{n = 32, 42, 56, 100\}$, $\{\beta_0 = 0, 0.0, 1.0\}$, $\{\sigma^2 = 0.1, 0.25, 0.5, 1.0, 1.5, 2.0\}$, with 1000 simulation replications in each cell, and we monitored the percentages of correct choice for each model specification method (in this simulation $M_2$ is always correct).

Table 1 gives examples of the results of this simulation, using $L_{CV}$ for illustration. Even with a sample size of only 32, $L_{CV}$ makes the right model choice more than 90% of the time when $\sigma^2 > 0.5$ for $\beta_0 = 1$ and when $\sigma^2 > 1.0$ for $\beta_0 = 0$ (these are parameter ranges that lead to large enough amounts of extra-Poisson variability that random-effects
Table 1: Percentages of correct model choice and mean absolute difference in $L_{SCV}$ between $M_1$ and $M_2$ when the right model is $M_2$, for $n = 32$.

<table>
<thead>
<tr>
<th>$\sigma^2$</th>
<th>$\beta_0$</th>
<th>% Correct Decision</th>
<th>Mean Absolute Difference in $L_{SCV}$</th>
</tr>
</thead>
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<tr>
<td></td>
<td></td>
<td></td>
<td>$\sigma^2$</td>
</tr>
<tr>
<td>0.10</td>
<td>31</td>
<td>47</td>
<td>0.10</td>
</tr>
<tr>
<td>0.25</td>
<td>49</td>
<td>85</td>
<td>0.25</td>
</tr>
<tr>
<td>0.50</td>
<td>76</td>
<td>95</td>
<td>0.50</td>
</tr>
<tr>
<td>1.00</td>
<td>97</td>
<td>100</td>
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</tr>
<tr>
<td>1.50</td>
<td>98</td>
<td>100</td>
<td>1.50</td>
</tr>
<tr>
<td>2.00</td>
<td>100</td>
<td>100</td>
<td>2.00</td>
</tr>
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</table>

models would be contemplated). The right part of the table shows that even rather small differences in $L_{SCV}$ can separate correct and incorrect model choice, which begs the question “When a difference on the log score scale is big enough?” (we return to this point in Section 3). Based on model discrimination results for $L_{SCV}$, $L_{FS}$, and $DIC$ we created a series of performance graphs (not shown) and observed (expectedly) that increasing $\sigma^2$ makes it easier for all three methods to conclude that random effects model (the nesting model) is needed to accommodate the Poisson over-dispersion. Interestingly, in this simulation environment $L_{FS}$ was more accurate, with small samples of data, at identifying the correct model than $L_{SCV}$ or $DIC$; for this reason, we focus on $L_{FS}$ in what follows.

3 Calibration of posterior predictive tail areas

Section 2 shows that full-sample log scores can stably and reliably decide between two models by choosing one with higher $L_{FS}$ (or $L_{SCV}$) value. However, this still leaves open model specification question $Q_2$: Is $M_1$ good enough?

In our view, a full judgment of adequacy requires real-world input about the purpose of the model, so it does not seem possible to propose generic methodology to answer $Q_2$. Instead, the somewhat related question “$Q'_2$: Could model $M_j$ have generated the data?” can be answered in a general way by simulating from $M_j$ many times, developing a distribution of (e.g.) $L_{FS}$ values, and seeing how unusual the actual data set’s log score is in this distribution.

This is related to the posterior predictive model-checking method of Gelman et al. (1996). However, this kind of simulation needs to be done carefully (Draper (1996)), or the result will be poor calibration; indeed, Berger (2000) and Robins et al. (2000) have demonstrated that the procedure in (Gelman et al. (1996)) may be (sharply) conservative. Using a modification of an idea suggested by Robins et al., we have developed a method for accurately calibrating the log score scale.

The inputs to our procedure are: a data set and a model (which may be parametric or non-parametric). For simplicity, consider a one-sample data set, $D$, of counts and suppose the goal is to quantify whether this data set could have come from the model $(y_i | \lambda, B) \text{ IID Poisson}(\lambda)$, and $(\lambda | B) \sim$ diffuse (call it model (*)). Now, consider the following procedure:

**Step 1:** Calculate $L_{FS}$ for data set $D$ and call it the actual log score (ALS). Obtain
the posterior for \( \lambda \) given \( y \) based on data set \( D \); call this the actual posterior. \[ \text{Step 2:} \]

\[
\text{for ( i in 1:m1 )} \{ \\
\text{Let \( \text{lambda}[\ i\ ] \) be a draw from the actual posterior.} \\
\text{Sample \( n \) data points from model (*) above, using \( \text{lambda} = \text{lambda}[\ i\ ] \).} \\
\text{Compute the full-sample log-score, \( \text{LS.FS}[\ i\ ] \), for this data set.} \\
\}
\]

The output of this loop is a vector of log scores; call this \( V.LS \). Locate the ALS in the distribution of \( \text{LS}_FS \) values by computing the percentage of \( \text{LS}_FS \) values in \( V.LS \) that are no greater than ALS; call this percentage the unadjusted actual tail area (suppose, e.g., that this comes out 0.22).

So far this is similar to Gelman et al. with \( \text{LS}_FS \) as the discrepancy function. We know from our own simulations (summarized below) and the literature such as Berger (2000), Robins et al. (2000)) that this tail area (a \( P \)-value for a composite null hypothesis, e.g., Poisson(\( \lambda \)) with \( \lambda \) unspecified) is conservative, i.e., with the 0.22 example above an adjusted version of it that is well calibrated would be smaller (and might be much smaller, e.g., 0.02). We have modified and implemented one of the ways suggested by Robins et al. for improving calibration, and we have shown that it does indeed work even in rather small-sample situations, although implementing the basic idea can be computationally intensive. \[ \text{Step 3:} \]

\[
\text{for ( j in 1:m2 )} \{ \\
\text{Let \( \text{lambda}* \) be a draw from the actual posterior.} \\
\text{Generate a data set of size \( n \) from the model (*) above,} \\
\text{using \( \text{lambda} = \text{lambda}* \); call this the simulated data set.} \\
\text{Repeat Steps 1 and 2 above on this simulated data set.} \\
\}
\]

The result will be a vector of unadjusted tail areas; call this \( V.P \). Compute the percentage of tail areas in \( V.P \) that are no greater than the unadjusted actual tail area; this is the adjusted actual tail area.

The claim is that the 3–step procedure above is well-calibrated: if the sampling part of model (*) really did generate the observed data, the distribution of adjusted actual tail areas would be approximately uniform, since \( X \sim F_X \) implies \( F_X(X) \sim U(0,1) \). This claim of calibration can be verified by building a further loop around steps 1–3 as follows:

Choose a \( \lambda \) value of interest; call it \( \text{lambda.sim} \).

\[
\text{for ( k in 1:m3 )} \{ \\
\text{Generate a data set of size \( n \) from the model (*) above,} \\
\text{using \( \text{lambda} = \text{lambda.sim} \); call this the validation data set.} \\
\text{Repeat Steps 1-3 on the validation data set.} \\
\}
\]

The result here is a vector, \( V.TA \), of adjusted tail areas. We have verified (via simulation, performed on a cluster of 100 Linux-based CPUs) in several settings that the distribution of values in \( V.TA \) is (very) close to \( U(0,1) \) indeed.

Figure 2 summarizes a set of histograms of the uncalibrated actual tail areas from one-sample Poisson model, indicating that in many cases the tail areas (\( p \)-values) are far from the target (uniform) distribution.
Consider, for example, the case \((n = 100, \lambda = 0.14)\) in the fourth row and first column of the Figure 2: if the uncalibrated tail area came out 0.35 in this situation, it would be natural to conclude that the data could very well have come from the Poisson model, but this part of Figure 2 demonstrates clearly that in fact an uncalibrated tail area of 0.35 with \((n = 100, \lambda = 0.14)\) is highly unusual under the Poisson model. Our procedure solves the calibration problem by asking “How often would one get 0.35 or less for an uncalibrated tail area in this situation?”, and it is evident from Figure 2 that the answer is not very often (in fact, only about 0.035 of the time, i.e., in this case the calibrated version of the uncalibrated Gelman et al. tail area is 10 times smaller).

Figure 2 shows also that the calibration of the unadjusted approach improves in the one-sample Poisson setting, for increasing \(\lambda\) (even for small \(n\)), but in case of the Gaussian model with both \(\mu\) and \(\sigma^2\) unknown, the unadjusted approach remains poorly calibrated across the entire subset \([-1 \leq \mu \leq +1] \times [0.1 \leq \sigma^2 \leq 10]\) of parameter space we examined, and things actually seem to get worse as \(n\) increases (not shown). However, the adjusted results, for the

\[
\begin{align*}
\text{null Poisson model: uncalibrated p-values} \\
\text{Figure 2: Poisson model: uncalibrated tail-area values.}
\end{align*}
\]
Gaussian model, are nearly perfectly calibrated, having distributions close to $U(0,1)$ for all examined parameter values and sample sizes, (again, not shown). Note that for the reason of limited space here we could show only a small fraction of results and graphs.

**Conclusions**

We have argued that calibration (checking how often one obtains the right answer) is an important principle that that arises naturally in good Bayesian modeling; the question “$Q_1$: Is model $M_j$ better than $M_k$?” is central to the process of well-calibrated Bayesian model specification; and it is not well formed unless the purpose of the model is considered. Once the purpose of the model is explicitly stated, the task of Bayesian model specification turns into a decision problem of maximizing expected utility (MEU), with a purpose-specific utility function (which may be computationally intensive).

$LS_{FS}$ appears as a useful improvement upon $DIC$, with three advantages: $LS_{FS}$ may well have better small-sample model discrimination behavior (as in the simulation of Section 3.1); $LS_{FS}$ is insensitive to model parameterization; and $LS_{FS}$ can be used both in Bayesian nonparametric and parametric settings; To decide when to stop looking for a better model, the question “$Q'_2$: Could model $M_j$ have generated the data?” can be answered in a well-calibrated manner, using $LS_{FS}$ as a model choice criterion, as shown in the last section.

**References**


