# Bayesian Model Specification

#### 4: Dealing With Model Uncertainty

#### David Draper

Department of Applied Mathematics and Statistics University of California, Santa Cruz draper@ucsc.edu

SHORT COURSE (DAY 4) UNIVERSITY OF READING (UK)

26 Nov 2015

users.soe.ucsc.edu/~draper/Reading-2015-Day-4.html

© 2015 David Draper (all rights reserved)

## Getting From the Context and Design to the Model

Definition. In model specification, optimal = {conditioning only on propositions rendered true by the context of the problem and the design of the data-gathering process, while at the same time ensuring that the set of conditioning propositions includes all relevant problem context}.

This seems hard to achieve; for example, in the IHGA case study, visualizing the data set before it arrives, it would look like the table shell presented back on page 2 of Part 1 of the Lecture Notes:

	Num	ber of	Hospit	alizations			
Group	0	1		k	n	Mean	SD
					$n_C = 287$		
Treatment	$n_{T0}$	$n_{T1}$		$n_{Tk}$	$n_T = 285$	$ar{y}_{\mathcal{T}}$	ST

The problem context and design make this table shell something You can condition on, and the lack of previous trials with IHGA (this was the first time it was implemented anywhere) implies that You can also condition on a diffuse choice for  $p(\theta|\mathcal{B})$  (with 572 observations, it won't matter much how this diffuseness is specified), but context and design don't seem to have anything to say about the predictive (sampling) distribution  $p(D|\theta|\mathcal{B})$ .

# Model Uncertainty

In **problems** of **realistic complexity** You'll generally **notice** that (a) You're **uncertain** about  $\theta$  but (b) You're also **uncertain** about how to **quantify Your uncertainty about**  $\theta$ , i.e., **You** have **model uncertainty**.

Cox's Theorem says that You can draw logically-consistent inferences about an unknown  $\theta$ , given data D and background information  $\mathcal{B}$ , by specifying  $M = \{p(\theta|M\mathcal{B}), p(D|\theta M\mathcal{B})\}$ , but item (b) in the previous paragraph implies that there will typically be more than one such plausible M; what should You do about this?

It would be nice to be able to solve the inference problem by using Bayes's Theorem to compute  $p(\theta|D\,\mathcal{M}_{\mathit{all}}\,\mathcal{B})$ , where  $\mathcal{M}_{\mathit{all}}$  is the set of all possible models, but this is not feasible: just as Kolmogorov had to resort to  $\sigma$ -fields because the set of all subsets of an  $\Omega$  with uncountably many elements is too big to meaningfully assign probabilities to all of the subsets, with a finite data set D,  $\mathcal{M}_{\mathit{all}}$  is too big for D to permit meaningful plausibility assessment of all the models in  $\mathcal{M}_{\mathit{all}}$ .

Having adopted the Calibration Principle, it makes sense to talk about an underlying data-generating model  $M_{DG}$ , which is unknown to You (more on this below).

#### An Ensemble $\mathcal{M}$ of Models

Not being able to compute  $p(\theta|D\mathcal{M}_{all}\mathcal{B})$ , in practice the best You can do is to compute  $p(\theta|D\mathcal{M}\mathcal{B})$ , where  $\mathcal{M}$  is an ensemble of models (finite or countably or uncountably infinite) chosen "well" by You, where "well" can and should be brought into focus by the Calibration Principle (and some of the other Principles to be introduced later): evidently what You want, among other things, is for  $\mathcal{M}$  to contain one or more models that are identical (or at least close) to  $M_{DG}$ .

Suppose **initially**, for the sake of **discussion**, that You've **identified** such an **ensemble** (I'll present some **ideas** for how to do this later) and that it turns out to be **finite**:  $\mathcal{M} = (M_1, \dots, M_k)$  for  $2 \le k < \infty$ ; **what next?** 

Are You **supposed** to try to **choose** one of these **models** (the **model selection problem**) and **discard** the rest, or **combine** them in some way (if so, **how?**), or **what?** 

Solving the model uncertainty problem. People used to "solve" the problem of what to do about model uncertainty by ignoring it: it was common, at least through the mid-1990s, to

# **Dealing With Model Uncertainty**

- (a) use the data D to conduct a search among possible models, settling on a single (apparently) "best" model  $M^*$  arising from the search, and then
  - (b) draw **inferences** about  $\theta$  **pretending** that  $M^*$  "="  $M_{DG}$ .

This of course can lead to quite bad calibration, almost always in the direction of pretending You know more than You actually do, so that, e.g., Your nominal 90% posterior predictive intervals for data values not used in the modeling process would typically include substantially fewer than 90% of the actual observations.

The  $M^*$  approach "solves" the problem of how to specify  $\mathcal M$  by setting  $\mathcal M = \{M^*\}$ ; I'll continue to postpone for the moment how You might do a better job of arriving at  $\mathcal M$ .

Having chosen  $\mathcal M$  in some way, how can You assess Your uncertainty across the models in  $\mathcal M$ , and appropriately propagate this through to Your uncertainty about  $\theta$ , in a well-calibrated way?

I'm aware of three approaches to improved assessment and propagation of model uncertainty: BMA, BNP, CCV.

#### BMA, BNP

ullet Bayesian model averaging (BMA): If interest focuses on something that has the same meaning across all the models in  $\mathcal{M}$  — for example, a set of future data values  $D^*$  to be predicted — calculation reveals (e.g., Leamer, 1978) that

$$p(D^*|D\mathcal{M}\mathcal{B}) = \int_{\mathcal{M}} p(D^*|D\mathcal{M}\mathcal{B}) p(M|D\mathcal{M}\mathcal{B}) dM, \qquad (1)$$

which is **eminently reasonable**: equation (1) tells You to form a **weighted average** of Your **conditional predictive distributions**  $p(D^*|DMB)$ , given particular **models**  $M \in \mathcal{M}$ , **weighted** by those models' **posterior probabilities** p(M|DMB).

This approach typically provides (substantially) better calibration than that obtained by the  $M^*$  method.

Bayesian nonparametric (BNP) modeling: The BMA integral in

 (1) can be thought of as an approximation to the (unattainable?)
 ideal of averaging over all worthwhile models; a better approximation to this ideal can often be achieved with Bayesian nonparametric modeling, which dates back to de Finetti (1937).

## Exchangeability

Continuing the Kaiser example on page 14 (Part 1), suppose You also observe (for each of the n=112 randomly-sampled patients from the population  $\mathbb P$  of N=8,561 heart-attack patients) a real-valued conceptually-continuous quality-of-care score  $y_i$ , and (following de Finetti) You're thinking about Your predictive distribution  $p(y_1 \ldots y_n | \mathcal B)$  for these scores before any data have arrived.

de Finetti pointed out that, if You have no covariate information about the patients, Your predictive distribution  $p(y_1 \ldots y_n | \mathcal{B})$  should remain the same under arbitrary permutation of the order in which the patients are listed, and he coined the term exchangeability to describe this state of uncertainty.

He (and later **Diaconis/Freedman**) went on to **prove** that, if Your judgment of **exchangeability** extends from  $(y_1 \ldots y_n)$  to  $(y_1 \ldots y_N)$  (as it certainly **should** here, given the **random sampling**) and  $N \gg n$  (as is **true** here), then all **logically-internally-consistent predictive distributions** can **approximately** be expressed **hierarchically** as follows:

# Bayesian Nonparametric (BNP) Modeling

letting F stand for the **empirical CDF** of the **population values**  $(y_1 \ldots y_N)$ , the **hierarchical model** is (for  $i = 1, \ldots, n$ )

$$\left\{ \begin{array}{ccc} (F|\mathcal{B}) & \sim & p(F|\mathcal{B}) \\ (y_i|F|\mathcal{B}) & \stackrel{\text{IID}}{\sim} & F \end{array} \right\}.$$

This requires placing a scientifically-appropriate prior distribution  $p(F|\mathcal{B})$  on the set  $\mathcal{F}$  of all CDFs on  $\Re$ , which de Finetti didn't know how to do in 1937; thanks to work by Freedman, Ferguson, Lavine, Escobar/West, and others, two methods for doing this sensibly — Pólya trees and Dirichlet-process (DP) priors — are now in routine use: this — placing distributions on function spaces — is Bayesian nonparametric (BNP) modeling.

IHGA Example, Revisited: Once again visualizing the IHGA data set before it arrives, here's the table shell one more time:

		Num	ber of	Hospita	alizations			
	Group	0	1		k	n	Mean	SD
						$n_C = 287$		
Tr	eatment	$n_{T0}$	$n_{T1}$		$n_{Tk}$	$n_T = 285$	$ar{y}_{\mathcal{T}}$	ST

## BNP Case Study

Letting (as before)  $\mu_C$  and  $\mu_T$  be the mean hospitalization rates (per two years) in the population  $\mathcal P$  (of all elderly non-institutionalized people in Denmark in the early 1980s) under the  $\mathcal C$  and  $\mathcal T$  conditions, respectively, the inferential quantity of main interest is still  $\theta = \frac{\mu_T - \mu_C}{\mu_C}$  (or this could be redefined without loss as  $\theta = \frac{\mu_T}{\mu_C}$ ); how can You draw valid and accurate inferences about  $\theta$  while coping with Your uncertainty about the population  $\mathcal C$  and  $\mathcal T$  CDFs — call them  $\mathcal F_C$  and  $\mathcal F_T$ , respectively — of numbers of hospitalizations per person (per two years)?

One approach: Bayesian qualitative-quantitative inference (Draper 2013): exchangeability implies a multinomial sampling distribution on the qualitative outcome variable with category labels  $0, 1, \ldots$ , and this permits optimal model specification here (this approach treats the hospitalization outcome categorically but permits quantitative inference about  $\theta$ ).

Another approach: Bayesian nonparametric modeling — it turns out that DP priors put all their mass on discrete distributions, so one BNP model for this data set would involve placing parallel DPs priors on  $F_C$  and  $F_T$ ; see KKD (2008) for details on the results.

## BNP Case Study (continued)

To serve as the **basis** of the  $M^*$  (cheating) approach (in which You look at the data for inspiration on which models to fit), here's a table of the actual data values:

	Number of Hospitalizations										
Group	0	1	2	3	4	5	6	7	n	Mean	SD
Control	138	77	46	12	8	4	0	2	287	0.944	1.24
Treatment	147	83	37	13	3	1	1	0	285	0.768	1.01

Evidently (description) IHGA lowered the mean hospitalization rate (for these elderly Danish people, at least) by (0.944-0.768)=0.176, which is a  $\left\{100\left(\frac{0.768-0.944}{0.944}\right) \doteq\right\}$  19% reduction from the control level, a difference that's large in clinical terms, but (inference) how strong is the evidence for a positive effect in  $\mathcal{P}=\{\text{all people similar}\ \text{to those}\ \text{in the experiment}\}$ ?

It's **natural** to think **initially** of **parallel Poisson**( $\lambda_C$ ) and Poisson( $\lambda_T$ ) modeling ( $M_1$ ), but there's **substantial over-dispersion**: the C and T **variance-to-mean ratios** are  $\frac{1.24^2}{0.944} \doteq 1.63$  and  $\frac{1.01^2}{0.768} \doteq 1.33$ .

# Bayesian Parametric Modeling

Unfortunately we have **no covariates** to help **explain** the **extra-Poisson variability**, and there's **little information external** to the **data set** about the **treatment effect**; this latter **state of knowledge** is expressed in **prior distributions** on **parameters** by making them **diffuse** (i.e., ensuring they have **large variability** to express **substantial uncertainty**).

In this **situation** You could fit **parallel Negative Binomial models**  $(M_2)$ , but a **parametric choice** that more readily **generalizes** is obtained by letting  $(x_i, y_i) = (C/T \text{ status, outcome})$  — so that  $x_i = 1$  if **Treatment**, 0 if **Control** and  $y_i = \text{the number of hospitalizations}$  — for person  $i = 1, \ldots, n$  and considering the **random-effects Poisson regression model**  $(M_3)$ :

$$(y_i|\lambda_i \ M_3 \ \mathcal{B}) \stackrel{\text{indep}}{\sim} \operatorname{Poisson}(\lambda_i) \ \log(\lambda_i) = \gamma_0 + \gamma_1 x_i + \epsilon_i \ (\epsilon_i|\sigma_\epsilon^2 \ M_3 \ \mathcal{B}) \stackrel{\text{IID}}{\sim} N(0,\sigma_\epsilon^2) \ (\gamma_0 \ \gamma_1 \ \sigma_\epsilon^2|M_3 \ \mathcal{B}) \sim \text{diffuse}.$$

In this model the unknown of main policy interest is

## **BNP** Example

 $\theta=rac{ ext{population } ar{ au}}{ ext{population } ar{ au}}=e^{\gamma_1};$  the **other parameters** can be collected in a **vector**  $\eta=(\gamma_0,\sigma_\epsilon^2);$  and the **random effects**  $\epsilon_i$  can be thought of as **proxying** for the **combined main effect**  $\sum_{j=2}^J \gamma_j (x_{ij}-ar{x}_j)$  of all the **unobserved relevant covariates (age, baseline health status**, ...).

The first line of (2) makes good scientific sense (the  $y_i$  are counts of relatively rare events), but the Gaussian assumption for the random effects is conventional and not driven by the science; a potentially better model ( $M_4$ ) is obtained by putting a prior distribution on the CDF of the  $\epsilon_i$  that's centered at the  $N(0,\sigma_\epsilon^2)$  distribution but that expresses substantial prior uncertainty about the Gaussian assumption:

# Dirichlet-Process Mixture Modeling

Many Bayesian prior distributions  $p(\theta|M_j|\mathcal{B})$  have two user-friendly inputs: a quantity  $\theta_0$  that acts like a prior estimate of the unknown  $\theta$ , and a number  $n_0$  that behaves like a prior sample size (i.e., a measure of how tightly the prior is concentrated around  $\theta_0$ ); DP priors are no exception to this pattern.

In equation (3),  $DP(\alpha, F_0)$  is a **Dirichlet-process prior** on F with **prior estimate**  $F_0 = N(0, \sigma_\epsilon^2)$  and a **quantity**  $(\alpha)$  that behaves something like a **prior sample size**; this is referred to as **Dirichlet-process mixture modeling**, because (3) is a **mixture model** — each **person** in the study has her/his **own**  $\lambda$ , drawn from  $F_C$  (control) or  $F_T$  (treatment) — in which **uncertainty** about  $F_C$  and  $F_T$  is **quantified** via a **DP**.

NB Bayesian model averaging (BMA) with a finite set of models can be regarded as a crude approximation to what Bayesian nonparametric (BNP) modeling is trying to do, namely average over Your uncertainty in model space to provide an honest representation of Your overall uncertainty that doesn't condition on things You don't know are true.

#### Cross-Validation

• Calibration cross-validation (CCV): The way the IHGA example unfolded looks a lot like the  $M^*$  approach I condemned previously: I used the entire data set to suggest which models to consider.

This has the (strong) potential to underestimate uncertainty;
Bayesians (like everybody else) need to be able to look at the data to suggest alternative models, but all of us need to do so in a way that's well-calibrated.

Cross-validation — partitioning the data (e.g., exchangeably) into subsets used for different tasks (modeling, validation, ...) can help.

- The  $M^*$  approach is an example of what might be called **1CV** (one-fold cross-validation): You use the entire data set D both to model and to see how good the model is (this is clearly inadequate).
- 2CV (two-fold cross-validation) is frequently used: You (a) partition the data into modeling (M) and validation (V) subsets, (b) use M to explore a variety of models until You've found a "good" one  $M^*$ , and (c) see how well  $M^*$  validates in V (a useful Bayesian way to do this is to use the data in M

# Calibration Cross-Validation (CCV)

- to construct **posterior predictive distributions** for **all of the data values** in V and see how the **latter compare** with the **former**).
- **2CV** is a **lot better** than **1CV**, but what do You do (as frequently happens) if  $M^*$  doesn't validate well in V?
- CCV (calibration cross-validation): going out one more term in the Taylor series (so to speak),
  - (a) partition the data into modeling (M), validation (V) and calibration (C) subsets,
  - (b) use M to explore a variety of models until You've found one or more plausible candidates  $\mathcal{M} = \{M_1, \dots, M_m\}$ ,
    - (c) see how well the models in  $\mathcal{M}$  validate in V,
  - (d) if **none of** them do, **iterate (b) and (c)** until You do get **good** validation. and
- (e) **fit** the **best model** in  $\mathcal{M}$  (or, better, **use BMA**) on the **data** in M + V, and report both (i) **inferential conclusions** based on **this fit** and (ii) the **quality of predictive calibration** of **Your model/ensemble**) in C.

# CCV (continued)

#### The **goal** with this **method** is both

- (1) a good answer, to the main scientific question, that has paid a reasonable price for model uncertainty (the inferential answer is based only on M + V, making Your uncertainty bands wider) and
- (2) an indication of how well calibrated {the iterative fitting process yielding the answer in (1)} is in C (a good proxy for future data).

You can use **decision theory** (Draper and Southwood, 2013) to decide **how much data** to put in each of M, V and C: the **more important calibration** is to You, the **more data** You want to put in C, but **only up to a point**, because getting a **good answer** to the **scientific question** is also **important** to You.

This is **related** to the **machine-learning** practice (e.g., **Hastie, Tibshirani, Friedman** [HTF] 2009) of **Train/Validation/Test** partitioning, with one **improvement** (**decision theory** provides an **optimal way** to choose the **data subset sizes**); I **don't agree** with HTF that this can **only be done with large data sets**: it's even **more important** to do it with **small** and **medium-size data sets** (You just need to work with **multiple** (**M**, **V**, **C**) **partitions** and **average**).

#### Modeling Algorithm

CCV provides a way to pay the right price for hunting around in the data for good models, motivating the following modeling algorithm:

- (a) Start at a model  $M_0$  (how choose?); set the current model  $M_{\text{current}} \leftarrow M_0$  and the current model ensemble  $\mathcal{M}_{\text{current}} \leftarrow \{M_0\}$ .
- (b) If  $\textit{M}_{current}$  is good enough to stop (how decide?), return  $\mathcal{M}_{current}$ ; else
- (c) Generate a new candidate model  $M_{\text{new}}$  (how choose?) and set  $\mathcal{M}_{\text{current}} \leftarrow \mathcal{M}_{\text{current}} \cup M_{\text{new}}$ .
- (d) If  $M_{\text{new}}$  is better than  $M_{\text{current}}$  (how decide?), set  $M_{\text{current}} \leftarrow M_{\text{new}}$ .
- (e) Go to (b).

For human analysts the choice in (a) is not hard, although it might not be easy to automate in full generality; for humans the choice in (c) demands creativity, and as a profession, at present, we have no principled way to automate it; here I want to focus on the questions in (b) and (d):

 $Q_1$ : Is  $M_1$  better than  $M_2$ ?  $Q_2$ : Is  $M_1$  good enough?

## The Modeling-As-Decision Principle

These questions **sound fundamental** but **are not**: better **for what purpose**? Good enough **for what purpose**? This **implies** (see, e.g., Bernardo and Smith, 1995; Draper, 1996; Key et al., 1999) a

Modeling-As-Decision Principle: Making clear the purpose to which the modeling will be put transforms model specification into a decision problem, which should be solved by maximizing expected utility with a utility function tailored to the specific problem under study.

Some examples of this may be found (e.g., Draper and Fouskakis, 2008: variable selection in generalized linear models under cost constraints), but this is hard work; there's a powerful desire for generic model-comparison methods whose utility structure may provide a decent approximation to problem-specific utility elicitation.

Two such methods are Bayes factors and log scores.

• Bayes factors. It looks natural to compare models on the basis of their posterior probabilities; from Bayes's Theorem in odds form,

#### **Bayes Factors**

$$\frac{p(M_2|DB)}{p(M_1|DB)} = \left[\frac{p(M_2|B)}{p(M_1|B)}\right] \cdot \left[\frac{p(D|M_2B)}{p(D|M_1B)}\right]; \tag{4}$$

the **first term** on the right is just the **prior odds** in favor of  $M_2$  over  $M_1$ , and the **second term** on the right is called the **Bayes factor**, so in words equation (4) says

$$\begin{pmatrix} \mathbf{posterior} \\ \mathbf{odds} \\ \text{for } M_2 \\ \text{over } M_1 \end{pmatrix} = \begin{pmatrix} \mathbf{prior odds} \\ \text{for } M_2 \\ \text{over } M_1 \end{pmatrix} \cdot \begin{pmatrix} \mathbf{Bayes factor} \\ \text{for } M_2 \\ \text{over } M_1 \end{pmatrix}. \quad (5)$$

(Bayes factors seem to have first been considered by Turing and Good  $(\sim 1941)$ , as part of the effort to break the German Enigma codes.)

**Odds** o are related to **probabilities** p via  $o = \frac{p}{1-p}$  and  $p = \frac{o}{1+o}$ ; these are **monotone increasing transformations**, so the **decision rules** {choose  $M_2$  over  $M_1$  if the **posterior odds** for  $M_2$  are greater} and {choose  $M_2$  over  $M_1$  if  $p(M_2|DB) > p(M_1|DB)$ } are **equivalent**.

## Decision-Theoretic Basis for Bayes Factors

This approach does have a **decision-theoretic basis**, but it's rather **odd**: if You pretend that the **only possible data-generating mechanisms** are  $\mathcal{M} = \{M_1, \ldots, M_m\}$  for finite m, and You pretend that one of the models in  $\mathcal{M}$  must be the **true data-generating mechanism**  $M_{DG}$ , and You pretend that the **utility function** 

$$U(M, M_{DG}) = \left\{ \begin{array}{ll} 1 & \text{if } M = M_{DG} \\ 0 & \text{otherwise} \end{array} \right\}$$
 (6)

reflects Your real-world values, then it's decision-theoretically optimal to choose the model in  $\mathcal{M}$  with the highest posterior probability (i.e., that choice maximizes expected utility).

If it's scientifically appropriate to take the prior model probabilities  $p(M_j|\mathcal{B})$  to be equal, this rule reduces to choosing the model with the highest Bayes factor in favor of it; this can be found by (a) computing the Bayes factor in favor of  $M_2$  over  $M_1$ ,

$$BF(M_2 \text{ over } M_1|D\mathcal{B}) = \frac{p(D|M_2\mathcal{B})}{p(D|M_1\mathcal{B})},\tag{7}$$

#### Parametric Model Comparison

favoring  $M_2$  if  $BF(M_2 \text{ over } M_1|D\mathcal{B}) > 1$ , i.e., if  $p(D|M_2\mathcal{B}) > p(D|M_1\mathcal{B})$ , and calling the **better model**  $M^*$ ; (b) **computing the Bayes factor** in favor of  $M^*$  over  $M_3$ , calling the **better model**  $M^*$ ; and so on up through  $M_m$ .

Notice that there's something else a bit funny about this:  $p(D|M_j|\mathcal{B})$  is the prior (not posterior) predictive distribution for the data set D under model  $M_j$ , so the Bayes factor rule tells You to choose the model that does the best job of predicting the data before any data arrives.

Let's look at the **general problem** of **parametric model comparison**, in which model  $M_j$  has **its own parameter vector**  $\gamma_j$  (of length  $k_j$ ), where  $\gamma_i = (\theta, \eta_i)$ , and is **specified** by

$$M_{j}: \left\{ \begin{array}{c} (\gamma_{j}|M_{j}\mathcal{B}) \sim p(\gamma_{j}|M_{j}\mathcal{B}) \\ (D|\gamma_{j}M_{j}\mathcal{B}) \sim p(D|\gamma_{j}M_{j}\mathcal{B}) \end{array} \right\}. \tag{8}$$

Here the quantity  $p(D|M_i|\mathcal{B})$  that **defines the Bayes factor** is

## Integrated Likelihoods

$$p(D|M_j \mathcal{B}) = \int p(D|\gamma_j M_j \mathcal{B}) p(\gamma_j | M_j \mathcal{B}) d\gamma_j; \qquad (9)$$

this is called an **integrated likelihood** (or **marginal likelihood**) because it tells You to take a **weighted average** of the **sampling distribution/likelihood**  $p(D|\gamma_j M_j \mathcal{B})$ , but  $\boxed{\text{NB}}$  weighted by the  $\boxed{\text{prior}}$  for  $\gamma_j$  in model  $M_j$ ; as noted above, this may seem **surprising**, but it's **correct**, and it can lead to **trouble**, as follows.

The first trouble is **technical**: the **integral** in (9) can be **difficult to compute**, and may not even be easy to **approximate**.

The second thing to **notice** is that (9) can be **rewritten** as

$$p(D|M_j \mathcal{B}) = E_{(\gamma_j|M_j \mathcal{B})} p(D|\gamma_j M_j \mathcal{B}). \tag{10}$$

In other words the **integrated likelihood** is the **expectation** of the **sampling distribution** over the **prior** for  $\gamma_j$  in model  $M_j$  (evaluated at the **observed data set** D).

A few additional words about prior distributions on parameters:

#### Instability of Bayes Factors

A distribution (density) for a real-valued parameter  $\theta$  that summarizes the information

 $\{\theta \text{ is highly likely to be near } \theta_0\}$ 

will have **most of its mass** concentrated **near**  $\theta_0$ , whereas the **information** 

 $\{$ **not much is known** about  $\theta\}$ 

would correspond to a **density** that's rather **flat** (or **diffuse**) across a broad range of  $\theta$  values; thus when the **scientific context** offers **little information** about  $\gamma_j$  **external** to the data set D, this translates into a **diffuse prior** on  $\gamma_j$ , and this spells **trouble** for **Bayes factors**:

$$p(D|M_j \mathcal{B}) = E_{(\gamma_i|M_j \mathcal{B})} p(D|\gamma_j M_j \mathcal{B}).$$

You can see that if the **available information** implies that  $p(\gamma_j|M_j\,\mathcal{B})$  should be **diffuse**, the **expectation** defining the **integrated likelihood** can be **highly unstable** with respect to **small details** in how the **diffuseness is specified**.

**Example:** Integer-valued data set  $D = (y_1 \dots y_n); \ \bar{y} = \frac{1}{n} \sum_{i=1}^n y_i;$ 

# Instability of Bayes Factors (continued)

 $M_1 = \mathbf{Geometric}(\theta_1)$  likelihood with a  $\mathbf{Beta}(\alpha_1, \beta_1)$  prior on  $\theta_1$ ;

 $M_2 = \mathbf{Poisson}(\theta_2)$  likelihood with a  $\mathbf{Gamma}(\alpha_2, \beta_2)$  prior on  $\theta_2$ .

The **Bayes factor** in favor of  $M_1$  over  $M_2$  turns out to be

$$\frac{\Gamma(\alpha_1 + \beta_1)\Gamma(n + \alpha_1)\Gamma(n\bar{y} + \beta_1)\Gamma(\alpha_2)(n + \beta_2)^{n\bar{y} + \alpha_2}\left(\prod_{i=1}^n y_i!\right)}{\Gamma(\alpha_1)\Gamma(\beta_1)\Gamma(n + n\bar{y} + \alpha_1 + \beta_1)\Gamma(n\bar{y} + \alpha_2)\beta_2^{\alpha_2}}.$$
 (11)

With standard diffuse priors — take  $(\alpha_1, \beta_1) = (1, 1)$  and  $(\alpha_2, \beta_2) = (\epsilon, \epsilon)$  for some  $\epsilon > 0$  — the **Bayes factor** reduces to

$$\frac{\Gamma(n+1)\Gamma(n\bar{y}+1)\Gamma(\epsilon)(n+\epsilon)^{n\bar{y}+\epsilon}\left(\prod_{i=1}^{n}y_{i}!\right)}{\Gamma(n+n\bar{y}+2)\Gamma(n\bar{y}+\epsilon)\epsilon^{\epsilon}}.$$
 (12)

This goes to  $+\infty$  as  $\epsilon \downarrow 0$ , i.e., You can make the evidence in **favor** of the **Geometric model** over the **Poisson** as **large** as You want, **no matter what the data says**, as a function of a quantity near 0 that **scientifically** You have **no basis** to specify.

If instead You **fix and bound**  $(\alpha_2, \beta_2)$  away from 0 and let  $(\alpha_1, \beta_1) \downarrow 0$ , You can **completely reverse** this and make the evidence in **favor** of the **Poisson model** over the **Geometric** as **large** as You want (for **any** y).

## Approximating Integrated Likelihoods

The bottom line is that, when scientific context suggests diffuse priors on the parameter vectors in the models being compared, the integrated likelihood values that are at the heart of Bayes factors can be hideously sensitive to small arbitrary details in how the diffuseness is specified.

This has been well-known for quite awhile now, and it's given rise to an amazing amount of fumbling around, as people who like Bayes factors have tried to find a way to fix the problem: at this point the list of attempts includes {partial, intrinsic, fractional} Bayes factors, well-calibrated priors, conventional priors, intrinsic priors, expected posterior priors, ... (e.g., Pericchi 2004), and all of them exhibit a level of ad-hockery that's otherwise absent from the Bayesian paradigm.

Approximating integrated likelihoods. The goal is

$$p(D|M_j B) = \int p(D|\gamma_j M_j B) p(\gamma_j | M_j B) d\gamma_j; \qquad (13)$$

maybe there's an **analytic approximation** to this that will suggest how to **avoid trouble**.

#### Laplace Approximation

Laplace (1785) already faced this problem 225 years ago, and he offered a solution that's often useful, which people now call a Laplace approximation in his honor (it's an example of what's also known in the applied mathematics literature as a saddle-point approximation).

Noticing that the **integrand**  $P^*(\gamma_j) \equiv p(D|\gamma_j M_j \mathcal{B}) p(\gamma_j|M_j \mathcal{B})$  in  $p(D|M_j \mathcal{B})$  is an **un-normalized version** of the **posterior distribution**  $p(\gamma_j|D|M_j \mathcal{B})$ , and appealing to a **Bayesian version** of the **Central Limit Theorem** — which says that **with a lot of data**, such a **posterior distribution** should be **close to Gaussian**, **centered** at the **posterior mode**  $\hat{\gamma}_j$  — You can see that (with a **large sample size** n)  $\log P^*(\gamma_j)$  should be **close to quadratic** around that mode; the **Laplace idea** is to take a **Taylor expansion** of  $\log P^*(\gamma_j)$  around  $\hat{\gamma}_j$  and **retain** only the terms out to **second order**; the result is

$$\log p(D|M_j \mathcal{B}) = \log p(D|\hat{\gamma}_j M_j \mathcal{B}) + \log p(\hat{\gamma}_j | M_j \mathcal{B}) + \frac{k_j}{2} \log 2\pi - \frac{1}{2} \log |\hat{I}_j| + O\left(\frac{1}{n}\right); \quad (14)$$

here  $\hat{\gamma}_j$  is the maximum likelihood estimate of the parameter vector  $\gamma_j$  under model  $M_j$  and  $\hat{l}_j$  is the observed information matrix under  $M_j$ .

Notice that the **prior** on  $\gamma_j$  in model  $M_j$  enters into this **approximation** through  $\log p(\hat{\gamma}_j|M_j\mathcal{B})$ , and this is a term that **won't go away with more data**: as n increases this term is O(1).

Using a less precise Taylor expansion, Schwarz (1978) obtained a different approximation that's the basis of what has come to be known as the Bayesian information criterion (BIC):

$$\log p(y|M_j \mathcal{B}) = \log p(y|\hat{\gamma}_j M_j \mathcal{B}) - \frac{k_j}{2} \log n + O(1). \tag{15}$$

People often work with a multiple of this for model comparison:

$$BIC(M_j|DB) = -2\log p(D|\hat{\gamma}_j M_j B) + k_j \log n$$
 (16)

(the -2 multiplier comes from deviance considerations); multiplying by -2 induces a search (with this approach) for models with small BIC.

This model-comparison method makes an explicit trade-off between model complexity (which goes up with  $k_j$  at a log n rate) — and model lack of fit (through the  $-2 \log p(D|\hat{\gamma}_i M_i \mathcal{B})$  term).

#### BIC and the Unit-Information Prior

BIC is called an information criterion because it resembles AIC (Akaike, 1974). which was derived using information-theoretic reasoning:

$$AIC(M_j|D\mathcal{B}) = -2\log p(D|\hat{\gamma}_j M_j \mathcal{B}) + 2k_j. \tag{17}$$

AIC penalizes model complexity at a linear rate in  $k_j$  and so can have different behavior than BIC, especially with moderate to large n (BIC tends to choose simpler models; more on this later).

It's possible to work out what **implied prior BIC** is using, from the point of view of the **Laplace approximation**; the result is

$$(\gamma_j|M_j\,\mathcal{B}) \sim N_{k_j}(\hat{\gamma}_j,n\hat{l}_j^{-1}). \tag{18}$$

In the literature this is called a unit-information prior, because in large samples it corresponds to the prior being equivalent to 1 new observation yielding the same sufficient statistics as the observed data.

This **prior** is **data-determined**, but this **effect** is **close to negligible** even with only **moderate** *n*.

#### Bayes Factors; Log Scores

The BIC approximation to Bayes factors has the extremely desirable property that it's free of the hideous instability of integrated likelihoods with respect to tiny details, in how diffuse priors are specified, that do not arise directly from the science of the problem; in my view, if You're going to use Bayes factors to choose among models, You're well advised to use a method like BIC that protects You from Yourself in mis-specifying those tiny details.

I said back on page 18 that there are two generic utility-based model-comparison methods: Bayes factors and log scores.

• Log scores are based on the

Prediction Principle: Good models make good predictions, and bad models make bad predictions; that's one scientifically important way You know a model is good or bad.

This suggests developing a **generic utility structure** based on **predictive accuracy**: consider first a **setting** in which  $D = y = (y_1 \dots y_n)$  for real-valued  $y_i$  and the **models** to be **compared** are (as before)

## Log Scores

$$M_{j:} \left\{ \begin{array}{c} (\gamma_{j} | M_{j} \mathcal{B}) \sim p(\gamma_{j} | M_{j} \mathcal{B}) \\ (y | \gamma_{j} M_{j} \mathcal{B}) \sim p(y | \gamma_{j} M_{j} \mathcal{B}) \end{array} \right\}. \tag{19}$$

When comparing a (future) data value  $y^*$  with the predictive distribution  $p(\cdot|y|M_j|\mathcal{B})$  for it under  $M_j$ , it's been shown that (under reasonable optimality criteria) all optimal scores measuring the discrepancy between  $y^*$  and  $p(\cdot|y|M_j|\mathcal{B})$  are linear functions of  $\log p(y^*|y|M_j|\mathcal{B})$  (the log of the height of the predictive distribution at the observed value  $y^*$ ).

Using this **fact**, perhaps the most **natural-looking** form for a **composite measure** of **predictive accuracy** of  $M_j$  is a **cross-validated** version of the resulting **log score**,

$$LS_{CV}(M_j|y|\mathcal{B}) = \frac{1}{n} \sum_{i=1}^{n} \log p(y_i|y_{-i}|M_j|\mathcal{B}), \qquad (20)$$

in which  $y_{-i}$  is the y **vector** with observation i **omitted**.

Somewhat **surprisingly**, Draper and Krnjajić (2010) have shown that a **full-sample log score** that **omits** the **leave-one-out idea**,

# Full-Sample Log Score

$$LS_{FS}(M_j|y|\mathcal{B}) = \frac{1}{n} \sum_{i=1}^n \log p(y_i|y|M_j|\mathcal{B}), \qquad (21)$$

made operational with the rule {favor  $M_2$  over  $M_1$  if  $LS_{FS}(M_2|y|\mathcal{B}) > LS_{FS}(M_1|y|\mathcal{B})$ }, can have better small-sample model discrimination ability than  $LS_{CV}$  (in addition to being faster to approximate in a stable way).

If, in the spirit of **calibration**, You're prepared to **think about** an **underlying data-generating model**  $M_{DG}$ ,  $LS_{FS}$  also has a **nice interpretation** as an **approximation** to the **Kullback-Leibler divergence** between  $M_{DG}$  and  $p(\cdot|y\ M_j\ \mathcal{B})$ , in which  $M_{DG}$  is **approximated** by the **empirical CDF**:

$$KL[p(\cdot|y|M_j|\mathcal{B})||M_{DG}] = E_{M_{DG}}\log M_{DG} - E_{M_{DG}}\log p(\cdot|y|M_j|\mathcal{B})$$

$$\stackrel{\cdot}{=} E_{M_{DG}}\log M_{DG} - LS_{FS}(M_j|y|\mathcal{B}); \qquad (22)$$

the first term on the right side of (22) is constant in  $p(\cdot|y|M_j|\mathcal{B})$ , so minimizing  $KL[p(\cdot|y|M_j|\mathcal{B}||M_{DG})]$  is approximately the same as maximizing  $LS_{FS}$ .

# Bayes Factors/BIC Versus Log Scores

What follows is a **sketch** of **recent results** (Draper, 2013) based on **simulation experiments** with **realistic sample sizes**; in my view **standard asymptotic calculations** — **choosing between** the **models** in  $\mathcal{M} = \{M_1, M_2\}$  as  $n \to \infty$  with  $\mathcal{M}$  **remaining fixed** — are **essentially irrelevant** in **calibration studies**, for **two reasons**:

- (1) With increasing n, You'll want  $\mathcal{M}$  to grow to satisfy Your desire to do a better job of capturing real-world complexities, and
- (2) **Data** usually **accumulate over time**, and with **increasing** *n* it **becomes more likely** that the **real-world process** You're modeling is **not stationary**.
- Versions of Bayes factors that behave sensibly with diffuse priors on the model parameters (e.g., intrinsic Bayes factors: Berger and Pericchi, 1996, and more recent cousins) tend to have model discrimination performance similar to that of BIC in calibration (repeated-sampling with known MDG) environments; I'll show results for BIC here

**Example:** Consider assessing the performance of a drug, for lowering

# Clinical Trial to Quantify Improvement

systolic blood pressure (SBP) in hypertensive patients, in a phase–II clinical trial, and suppose that a Gaussian sampling distribution for the outcome variable is reasonable (possibly after transformation).

Two **frequent designs** in **settings** of this type have as their goals **quantifying improvement** and **establishing bio-equivalence**.

(quantifying improvement) Here You want to estimate the mean decline in blood pressure under this drug, and it would be natural to choose a repeated-measures (pre-post) experiment, in which SBP values are obtained for each patient, both before and after taking the drug for a sufficiently long period of time for its effect to become apparent.

Let  $\theta$  stand for the **mean difference**  $(SBP_{before} - SBP_{after})$  in the **population** of **patients** to which it's **appropriate** to **generalize** from the **patients** in Your **trial**, and let  $D = y = (y_1 \dots y_n)$ . where  $y_i$  is the **observed difference**  $(SBP_{before} - SBP_{after})$  for **patient** i  $(i = 1, \dots, n)$ .

The real-world purpose of this experiment is to decide whether to take the drug forward to phase III; under the weight of 20th-century

#### Decision, Not Inference

inertia (in which decision-making was strongly — and incorrectly — subordinated to inference), Your first impulse might be to treat this as an inferential problem about  $\theta$ , but it's not; it's a decision problem that involves  $\theta$ .

This is an example of the

Decision-Versus-Inference Principle: We should all get out of the habit of using inferential methods to make decisions: their implicit utility structure is often far from optimal.

The action space here is  $\mathcal{A}=(a_1,a_2)=$  (don't take the drug forward to phase III, do take it forward), and a sensible utility function  $U(a_j,\theta)$  should be continuous and monotonically increasing in  $\theta$  over a broad range of positive  $\theta$  values (the bigger the SBP decline for hypertensive patients who start at (say) 160 mmHg, the better, up to a drop of about 40 mmHg, beyond which the drug starts inducing fainting spells).

However, to facilitate a comparison between BIC and log scores, here I'll compare two models  $M_1$  and  $M_2$  that dichotomize the  $\theta$  range,

#### Models For Quantifying Improvement

but not at 0: despite a century of textbook claims to the contrary, there's nothing special about  $\theta=0$  in this setting, and in fact You know scientifically that  $\theta$  is not exactly 0 (because the outcome variable in this experiment is conceptually continuous).

What matters here is whether  $\theta > \Delta$ , where  $\Delta$  is a practical significance improvement threshold below which the drug is not worth advancing into phase III (for example, any drug that did not lower SBP for severely hypertensive patients — those whose pre-drug values average 160 mmHg or more — by at least 15 mmHg would not deserve further attention).

With little information about  $\theta$  external to this experimental data set, what counts in this situation is the comparison of the following two models:

$$M_1$$
:  $\left\{ \begin{array}{cc} (\theta | \mathcal{B}) & \sim & \text{diffuse for } \theta \leq \Delta \\ (y_i | \theta | \mathcal{B}) & \stackrel{\text{IID}}{\sim} & N(\theta, \sigma^2) \end{array} \right\}$  and (23)

$$M_2$$
:  $\left\{ \begin{array}{cc} (\theta | \mathcal{B}) & \sim & \text{diffuse for } \theta > \Delta \\ (y_i | \theta | \mathcal{B}) & \stackrel{\text{IID}}{\sim} & N(\theta, \sigma^2) \end{array} \right\},$  (24)

# Quantifying Improvement: Model Comparison Methods

in which for simplicity I'll take  $\sigma^2$  to be known (the results are similar with  $\sigma^2$  learned from the data).

This gives rise to **three model-selection methods** that can be **compared calibratively**:

- Full-sample log scores: choose  $M_2$  if  $LS_{FS}(M_2|yB) > LS_{FS}(M_1|yB)$ .
  - Posterior probability: let

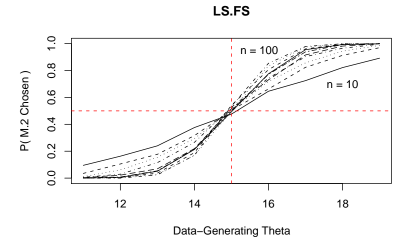
$$M^* = \{(\theta|\mathcal{B}) \sim \text{diffuse on } \Re, (y_i|\theta|\mathcal{B}) \stackrel{\text{IID}}{\sim} \mathcal{N}(\theta, \sigma^2)\} \text{ and choose } M_2 \text{ if } p(\theta > \Delta|y|M^*|\mathcal{B}) > 0.5.$$

• BIC: choose  $M_2$  if  $BIC(M_2|y|\mathcal{B}) < BIC(M_1|y|\mathcal{B})$ .

Simulation experiment details, based on the SBP drug trial:  $\Delta=15$ ;  $\sigma=10;\ n=10,20,\ldots,100;$  data-generating  $\theta_{DG}=11,12,\ldots,19;$   $\alpha=0.05;\ 1,000$  simulation replications; Monte-Carlo approximations of the predictive ordinates in  $LS_{FS}$  based on 10,000 posterior draws.

The **figures** below give **Monte-Carlo estimates** of the **probability that**  $M_2$  **is chosen**.

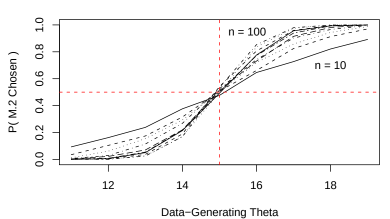
# LS<sub>FS</sub> Results: Quantifying Improvement



This **exhibits all** the **monotonicities** that it **should**, and **correctly yields 0.5** for all n with  $\theta_{DG} = 15$ .

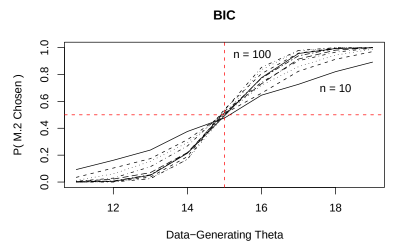
# Posterior Probability Results: Quantifying Improvement

#### **Posterior Probability**



Even though the  $LS_{FS}$  and posterior-probability methods are quite different, their information-processing in discriminating between  $M_1$  and  $M_2$  is identical to within  $\pm\,0.003$  (well within simulation noise with 1,000 replications).

### BIC Results: Quantifying Improvement



Here BIC and the posterior-probability approach are algebraically identical, making the model-discrimination performance of all three approaches the same in this problem.

## Establishing Bio-Equivalence

• (establishing bio-equivalence) In this case there's a previous hypertension drug B (call the new drug A) and You're wondering if the mean effects of the two drugs are close enough to regard them as bio-equivalent.

A good design here would again have a repeated-measures character, in which each patient's SBP is measured four times: before and after taking drug A, and before and after taking drug B (allowing enough time to elapse between taking the two drugs for the effects of the first drug to disappear).

Let  $\theta$  stand for the **mean difference** 

$$[(SBP_{before,A} - SBP_{after,A}) - (SBP_{before,B} - SBP_{after,B})]$$
 (25)

in the **population** of **patients** to which it's **appropriate** to **generalize** from the **patients in Your trial**, and let  $y_i$  be the **corresponding difference** for patient i (i = 1, ..., n).

Again in this setting there's nothing special about  $\theta = 0$ , and as before You know scientifically that  $\theta$  is not exactly 0;

# Bio-Equivalence Modeling

what matters here is whether  $|\theta| \le \lambda$ , where  $\lambda > 0$  is a practical significance bio-equivalence threshold (e.g., 5 mmHg).

Assuming as before a Gaussian sampling story and little information about  $\theta$  external to this experimental data set, what counts here is a comparison of

$$M_3$$
:  $\left\{ \begin{array}{cc} (\theta | \mathcal{B}) & \sim & \text{diffuse for } |\theta| \leq \lambda \\ (y_i | \theta | \mathcal{B}) & \sim & N(\theta, \sigma^2) \end{array} \right\}$  and (26)

$$M_4$$
:  $\left\{ \begin{array}{cc} (\theta | \mathcal{B}) & \sim & \text{diffuse for } |\theta| > \lambda \\ (y_i | \theta | \mathcal{B}) & \sim & N(\theta, \sigma^2) \end{array} \right\},$  (27)

in which  $\sigma^2$  is again taken for **simplicity** to be **known**.

A natural alternative to BIC and  $LS_{FS}$  here is again based on posterior probabilities: as before, let  $M^* = \{(\theta|\mathcal{B}) \sim \text{diffuse on } \Re, (y_i|\theta|\mathcal{B}) \stackrel{\text{IID}}{\sim} \mathcal{N}(\theta,\sigma^2)\}$ , but this time favor  $M_4$  over  $M_3$  if  $p(|\theta| > \lambda|y|M^*|\mathcal{B}) > 0.5$ .

As before, a careful real-world choice between  $M_3$  and  $M_4$  in this case would be based on a utility function that quantified the

## Bio-Equivalence Model Comparison

#### costs and benefits of

{claiming the two drugs were bio-equivalent when they were, concluding that they were bio-equivalent when they were not, deciding that they were not bio-equivalent when they were, judging that they were not bio-equivalent when they were not},

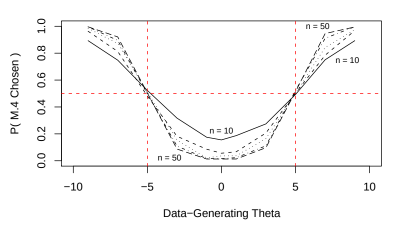
but here I'll again simply compare the calibrative performance of  $LS_{FS}$ , posterior probabilities, and BIC.

Simulation experiment details, based on the SBP drug trial:  $\lambda=5$ ;  $\sigma=10;\ n=10,20,\ldots,100;$  data-generating  $\theta_{DG}=\{-9,-7,-5,-3,-1,0,1,3,5,7,9\};\ \alpha=0.05;\ \textbf{1,000}$  simulation replications, M=10,000 Monte-Carlo draws for  $LS_{FS}$ .

NB It has previously been established that when making the (unrealistic) sharp-null comparison  $\theta=0$  versus  $\theta\neq 0$  in the context of  $(y_i|\theta|\mathcal{B})^{\text{IID}} \sim N(\theta,\sigma^2)$ , as  $n\to\infty$  LSFS selects the  $\theta\neq 0$  model with probability  $\to 1$  even when  $\theta_{DG}=0$ ; this "inconsistency of log scores at the null model" has been used by some people as a reason to dismiss log scores as a model-comparison method.

### *LS<sub>FS</sub>* Results: Bio-Equivalence

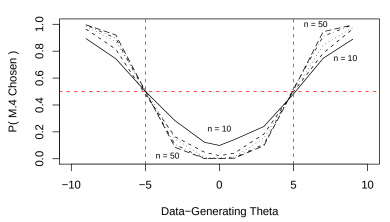




In this more realistic setting, comparing  $|\theta| \leq \lambda$  versus  $|\theta| > \lambda$  with  $\lambda > 0$ ,  $LS_{FS}$  has the correct large-sample behavior, both when  $|\theta_{DG}| \leq \lambda$  and when  $|\theta_{DG}| > \lambda$ .

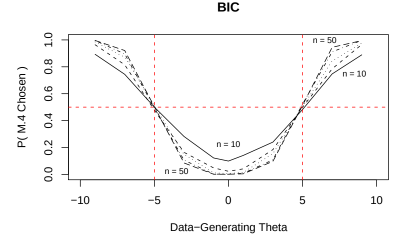
## Posterior Probability Results: Bio-Equivalence

#### **Posterior Probability**



The qualitative behavior of the  $LS_{FS}$  and posterior-probability methods is identical, although there are some numerical differences (highlighted later).

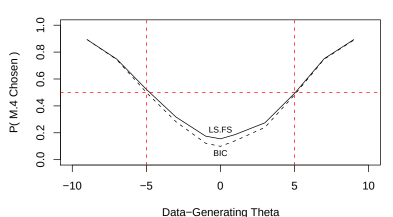
## BIC Results: Bio-Equivalence



In the quantifying-improvement case, the BIC and posterior-probability methods were algebraically identical; here they nearly coincide (differences of  $\pm\,0.001$  with 1,000 simulation repetitions).

## LS<sub>FS</sub> Versus BIC Results: Bio-Equivalence

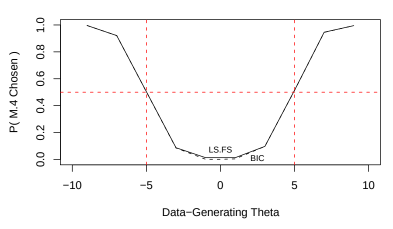
#### LS.FS Versus BIC (n = 10)



If You call **choosing**  $M_4$ :  $|\theta| > \lambda$  when  $|\theta_{DG}| \le \lambda$  a **false-positive** error and **choosing**  $M_3$ :  $|\theta| \le \lambda$  when  $|\theta_{DG}| > \lambda$  a **false-negative** mistake, with n=10 there's a **trade-off**:  $LS_{FS}$  has more **false positives** and BIC has more **false negatives**.

## LS<sub>FS</sub> Versus BIC Results: Bio-Equivalence





By the time You reach n = 50 in this problem,  $LS_{FS}$  and BIC are essentially equivalent.

### The Decision-Versus-Inference Principle, Revisited

In the context of the quantifying-improvement example, the real-world purpose of the experiment was to decide whether or not to take the drug forward to phase III.

Suppose that You tried to solve this decision problem with a popular inferential tool: frequentist hypothesis-testing of  $H_0$ :  $\theta \leq \Delta$  versus  $H_A$ :  $\theta > \Delta$  at significance level  $\alpha$ .

Decision-theoretically this is already wrong; as noted back on page 34, the utility function should actually be continuous in  $\theta$  rather than artificially dichotomizing  $\Theta$  into  $(-\infty, \Delta]$  and  $(\Delta, \infty)$ .

Even if You temporarily buy into this incorrect dichotomization, to solve the problem properly You'd have to quantify the real-world consequences of each of the cells in this table specifying  $U(a,\theta)$  (here  $u_{ij} \geq 0$ ):

<b>Action</b>
$a_1$ (stop)
a <sub>2</sub> (phase III)

Truth			
$\theta \leq \Delta$	$\theta > \Delta$		
<i>u</i> <sub>11</sub>	$-u_{12}$		
$-u_{21}$	u <sub>22</sub>		

# Decision-Theory (Not Inference) For Decision Problems

	Truth	
<u>Action</u>	$\theta \leq \Delta$	$\theta > \Delta$
$a_1$ (stop)	<i>u</i> <sub>11</sub>	$-u_{12}$
a <sub>2</sub> (phase III)	$-u_{21}$	u <sub>22</sub>

- $u_{11}$  is the gain from correctly not taking the drug forward to phase III (this is clearly 0);
- $u_{12}$  is the loss from incorrectly failing to take the drug forward to phase III;
- $u_{21}$  is the loss from incorrectly taking the drug forward to phase III;
- $u_{22}$  is the gain from correctly taking the drug forward to phase III.

The optimal Bayesian decision turns out to be: choose  $a_2$  (go forward to phase III) iff

$$P(\theta > \Delta | y \mathcal{B}) \ge \frac{u_{21}}{u_{12} + u_{21} + u_{22}} = u^*.$$
 (28)

The frequentist (hypothesis-testing) inferential approach is equivalent to this only if

# Optimal Decision-Making in Phase-II Trials

$$\alpha = 1 - u^* = \frac{u_{12} + u_{22}}{u_{12} + u_{21} + u_{22}}.$$
 (29)

The implicit trade-off between false positives and false negatives in BIC and  $LS_{FS}$  — and the built-in trade-off in level- $\alpha$  hypothesis-testing for any given  $\alpha$  — may be close to optimal or not, according to the real-world values of  $\{u_{12}, u_{21}, u_{22}\}$ .

In phase-II clinical trials or micro-array experiments, when You're screening many drugs or genes for those that may lead to an effective treatment and — from the drug company's point of view — a false-negative error (of failing to move forward with a drug or gene that's actually worth further investigation) can be much more costly than a false-positive mistake, this corresponds to  $u_{12}\gg u_{21}$  and leads in the hypothesis-testing approach in phase-II trials to a willingness to use (much) larger  $\alpha$  values than the conventional 0.01 or 0.05, something that good frequentist biostatisticians have long known intuitively.

(In work I've done with a Swiss pharmaceutical company, this approach led to  $\alpha$  values on the order of 0.45, which is close to the implicit trade-off in BIC and  $LS_{FS}$ .)

# For People Who Like to Test Sharp-Null Hypotheses

An extreme example of the false-positive/false-negative differences between  $LS_{FS}$  and BIC in this setting may be obtained, albeit unwisely, by letting  $\lambda \downarrow 0$ .

This is unwise here (and is often unwise) because it amounts, in frequentist language, to testing the sharp-null hypothesis  $H_0$ :  $\theta=0$  against the alternative  $H_A$ :  $\theta\neq0$ .

It's necessary to distinguish between problems in which there is or is not a structural singleton in the (continuous) set  $\Theta$  of possible values of  $\theta$ : settings where it's scientifically important to distinguish between  $\theta=\theta_0$  and  $\theta\neq\theta_0$  — an example would be discriminating between {these two genes are on different chromosomes (the strength  $\theta$  of their genetic linkage is  $\theta_0=0$ )} and {these two genes are on the same chromosome  $(\theta>0)$ }.

Sharp-null testing without structural singletons is always unwise because

(a) You already know from scientific context, when the outcome variable is continuous, that  $H_0$  is false, and (relatedly)

## Testing Sharp-Null Hypotheses (continued)

(b) it's silly from a measurement point of view: with a (conditionally) IID  $N(\theta,\sigma^2)$  sample of size n, your measuring instrument  $\bar{y}$  is only accurate to resolution  $\frac{\sigma}{\sqrt{n}}>0$ ; claiming to be able to discriminate between  $\theta=0$  and  $\theta\neq 0$  — with realistic values of n — is like someone with a scale that's only accurate to the nearest ounce telling You that Your wedding ring has 1 gram (0.035 ounce) less gold in it than the jeweler claims it does.

Nevertheless, for people who like to test sharp-null hypotheses, here are some results: here I'm comparing the models (i = 1, ..., n)

$$M_5$$
:  $\left\{ \begin{array}{cc} (\sigma^2 | \mathcal{B}) & \sim & \text{diffuse on (0, large)} \\ (y_i | \sigma^2 \mathcal{B}) & \stackrel{\text{IID}}{\sim} & \mathcal{N}(0, \sigma^2) \end{array} \right\}$  and (30)

$$M_6$$
:  $\left\{ \begin{array}{ccc} (\theta \, \sigma^2 | \mathcal{B}) & \sim & \text{diffuse on } (-\text{large}, \text{large}) \times (0, \text{large}) \\ (y_i | \theta \, \sigma^2 \, \mathcal{B}) & \stackrel{\text{IID}}{\sim} & N(\theta, \sigma^2) \end{array} \right\}, \quad (31)$ 

In this case a natural Bayesian competitor to BIC and  $LS_{FS}$  would be to construct the central  $100(1-\alpha)\%$  posterior interval for  $\theta$  under  $M_6$  and choose  $M_6$  if this interval doesn't contain  $\mathbf{0}$ .

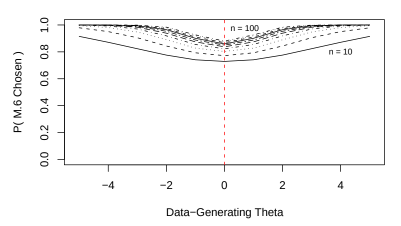
### Testing Sharp-Null Hypotheses (continued)

```
Simulation experiment details: data-generating \sigma_{DG}=10; n=10,20,\ldots,100; data-generating \theta_{DG}=\{0,1,\ldots,5\}; 1,000 simulation replications, M= 100,000 Monte-Carlo draws for LS_{FS}; the figures below give Monte-Carlo estimates of the probability that M_6 is chosen.
```

As before, let's call **choosing**  $M_6$ :  $\theta \neq 0$  when  $\theta_{DG} = 0$  a **false-positive** error and **choosing**  $M_5$ :  $\theta = 0$  when  $\theta_{DG} \neq 0$  a **false-negative** mistake.

# LS<sub>FS</sub> Results: Sharp-Null Testing

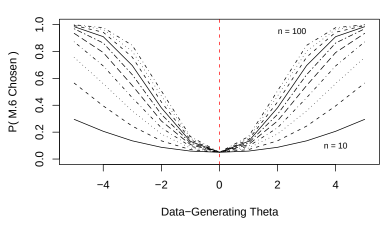




In the limit as  $\lambda \downarrow 0$ , the  $LS_{FS}$  approach makes hardly any false-negative errors but quite a lot of false-positive mistakes.

# Interval ( $\alpha = 0.05$ ) Results: Sharp-Null Testing

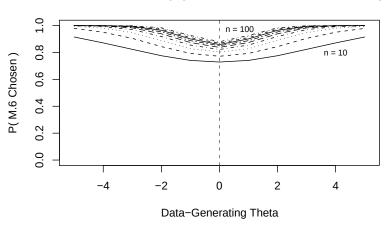
#### Posterior Interval (alpha = 0.05)



The behavior of the posterior interval approach is of course quite different: it makes many false-negative errors because its rate of false-positive mistakes is fixed at 0.05.

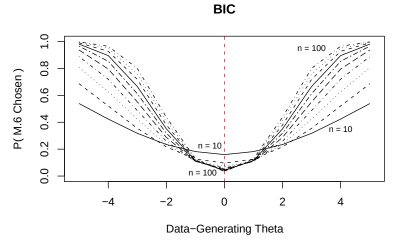
### Interval ( $\alpha$ Modified to $LS_{FS}$ Behavior) Results

#### Posterior Interval (alpha Modified to LS.FS Behavior)



When the **interval method** is **modified** so that  $\alpha$  **matches** the  $LS_{FS}$  behavior at  $\theta_{DG}=0$  (letting  $\alpha$  vary with n), the **two approaches** have **identical model-discrimination ability**.

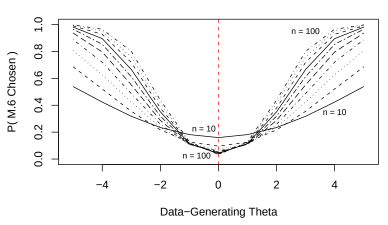
## BIC Results: Sharp-Null Testing



BIC's behavior is quite different from that of  $LS_{FS}$  and fixed- $\alpha$  posterior intervals: its false-positive rate decreases as n grows, but it suffers a high false-negative rate to achieve this goal.

### Interval ( $\alpha$ Modified to BIC Behavior) Results

#### Posterior Interval (alpha Modified to BIC Behavior)



When the **interval method** is **modified** so that  $\alpha$  **matches** the **BIC** behavior at  $\theta_{DG} = 0$  (again letting  $\alpha$  vary with n), the **two approaches** have **identical model-discrimination ability**.

### LS<sub>FS</sub> Versus BIC: Geometric Versus Poisson

As another model-comparison example, suppose You have an integer-valued data set  $D = y = (y_1 \dots y_n)$  and You wish to compare

 $M_7 = \mathbf{Geometric}(\theta_1)$  sampling distribution with a  $\mathbf{Beta}(\alpha_1, \beta_1)$  prior on  $\theta_1$ , and

 $M_8 = \mathbf{Poisson}(\theta_2)$  sampling distribution with a  $\mathbf{Gamma}(\alpha_2, \beta_2)$  prior on  $\theta_2$ .

*LS<sub>FS</sub>* and **BIC** both have **closed-form expressions** in this **situation**: with  $s = \sum_{i=1}^{n} y_i$  and  $\hat{\theta}_1 = \frac{\alpha_1 + n}{\alpha_1 + \beta_1 + s + n}$ ,

$$LS_{FS}(M_{7}|y\,\mathcal{B}) = \log \Gamma(\alpha_{1} + n + \beta_{1} + s) + \log \Gamma(\alpha_{1} + n + 1) - \log \Gamma(\alpha_{1} + n) - \log \Gamma(\beta_{1} + s)$$
(32)  
 
$$+ \frac{1}{n} \sum_{i=1}^{n} [\log \Gamma(\beta_{1} + s + y_{i}) - \log \Gamma(\alpha_{1} + n + \beta_{1} + s + y_{i} + 1)],$$

$$BIC(M_{7}|y\,\mathcal{B}) = -2[n \log \hat{\theta}_{1} + s \log(1 - \hat{\theta}_{1})] + \log n,$$
(33)

## Geometric Versus Poisson (continued)

$$LS_{FS}(M_8|y\,\mathcal{B}) = (\alpha_2 + s)\log(\beta_2 + n) - \log\Gamma(\alpha_2 + s) - (\alpha_2 + s)\log(\beta_2 + n + 1)$$

$$+ \frac{1}{n} \sum_{i=1}^{n} [\log\Gamma(\alpha_2 + s + y_i) - y_i \log(\beta_2 + n + 1) - \log\Gamma(y_i + 1)], \text{ and}$$

$$BIC(M_8|y\,\mathcal{B}) = -2[s\log\hat{\theta}_2 - n\,\hat{\theta}_2 - \sum_{i=1}^{n} \log(y_i!)] + \log n,$$

$$\text{where } \hat{\theta}_2 = \frac{\alpha_2 + s}{\beta_3 + n}.$$
(34)

Simulation details:  $n = \{10, 20, 40, 80\}$ ,  $\alpha_1 = \beta_1 = \alpha_2 = \beta_2 = 0.01$ , 1,000 simulation replications; it turns out that with  $(\theta_1)_{DG} = 0.5$  (Geometric) and  $(\theta_2)_{DG} = 1.0$  (Poisson), both data-generating distributions are monotonically decreasing and not easy to tell apart by eye.

Let's call **choosing**  $M_8$  (Poisson) when  $M_{DG} =$  **Geometric** a **false-Poisson** error and **choosing**  $M_7$  (Geometric) when  $M_{DG} =$  **Poisson** a **false-Geometric** mistake.

### Geometric Versus Poisson (continued)

The **table below** records the **Monte-Carlo probability** that the **Poisson model** was **chosen**.

M.DG = Poisson	M.DG = Geometric
n LS.FS BIC	n LS.FS BIC
10 0.8967 0.8661	10 0.4857 0.4341
20 0.9185 0.8906	20 0.3152 0.2671
40 0.9515 0.9363	40 0.1537 0.1314
80 0.9846 0.9813	80 0.0464 0.0407

Both methods make more false-Poisson errors than false-Geometric mistakes; the results reveal once again that neither BIC nor  $LS_{FS}$  uniformly dominates — each has a different pattern of false-Poisson and false-Geometric errors ( $LS_{FS}$  correctly identifies the Poisson more often than BIC does, but as a result BIC gets the Geometric right more often than  $LS_{FS}$ ).

### Properties of LS<sub>FS</sub>

• Log scores are entirely free from the diffuse-prior problems bedeviling Bayes factors:

$$LS_{FS}(M_{j}|y \mathcal{B}) = \frac{1}{n} \sum_{i=1}^{n} \log p(y_{i}|y M_{j} \mathcal{B}),$$
in which
$$p(y_{i}|y M_{j} \mathcal{B}) = \int p(y_{i}|\gamma_{j} M_{j} \mathcal{B}) p(\gamma_{j}|y M_{j} \mathcal{B}) d\gamma_{j}$$

$$= E_{(\gamma_{i}|y M_{i} \mathcal{B})} p(y_{i}|\gamma_{j} M_{i} \mathcal{B});$$
(36)

this **expectation** is over the **posterior** (not the prior) distribution for the parameter vector  $\gamma_j$  in model  $M_j$ , and is therefore completely stable with respect to small variations in how prior diffuseness (if scientifically called for) is specified, even with only moderate n.

 Following the Modeling-As-Decision Principle, the decision-theoretic justification for Bayes factors involves not only the Bayes factors themselves but also the prior model probabilities, which can be hard to specify in a scientifically-meaningful way: under the Bayes-factor (possibly unrealistic) 0/1 utility structure,

## Properties of $LS_{FS}$ (continued)

You're supposed to **choose the model** with the **highest posterior probability**, not the one with the **biggest Bayes factor**.

By contrast, **specification** of **prior model probabilities** doesn't arise with **log scores**, which have a **direct decision-theoretic justification** based on the **Prediction Principle**.

- It may seem that log scores have no penalty for unnecessary model complexity, but this is not true: for example, if one of Your models carries around a lot of unnecessary parameters, this will needlessly inflate its predictive variances, making the heights of its predictive densities go down, thereby lowering its log score.
- It may also seem that the behavioral rule based on posterior Bayes factors (Aitkin 1991) is the same as the rule based on  $LS_{FS}$ , which favors model  $M_j$  over  $M_{j'}$  if

$$n LS_{FS}(M_j|y,\mathcal{B}) > n LS_{FS}(M_{j'}|y,\mathcal{B}). \tag{37}$$

But this is **not true either**: for example, in the **common situation** in which the **data set** D consists of **observations**  $y_i$  that are **conditionally IID** from  $p(y_i|\eta_i, M_i, \mathcal{B})$  under  $M_i$ ,

### Summary

$$nLS_{FS}(M_j|y,\mathcal{B}) = \log \prod_{i=1}^n \left[ \int p(y_i|\eta_j, M_j, \mathcal{B}) \, p(\eta_j|y, M_j, \mathcal{B}) \, d\eta_j \right], \quad (38)$$

and this is not the same as

$$\log \int \left[ \prod_{i=1}^{n} p(y_i | \eta_j, M_j, \mathcal{B}) \right] p(\eta_j | y, M_j, \mathcal{B}) d\eta_j = \bar{L}_j^{PBF}$$
 (39)

because the **product** and **integral operators do not commute**.

- Some take-away messages:
- In the bio-equivalence example, even when You (unwisely) let  $\lambda \downarrow 0$ , thereby testing a sharp-null hypothesis, the asymptotic behavior of log scores is irrelevant; what counts is the behavior of log scores and Bayes factors with Your sample size and the models being compared, and for any given n it's not possible to say that the false-positive/false-negative trade-off built into Bayes factors is universally better for all applied problems than the false-positive/false-negative trade-off built into log scores,

or vice versa — You have to think it through in each problem.

For instance, the tendency of log scores to choose the "bigger" model in a nested-model comparison is exactly the right qualitative behavior in the following two examples (and many more such examples exist):

- Variable selection in searching through many compounds or genes to find successful treatments: here a false-positive mistake (taking an ineffective compound or gene forward to the next level of investigation) costs the drug company C, but a false-negative error (failing to move forward with a successful treatment, in a highly-competitive market) costs C with C
  - In a two-arm clinical-trial setting, consider the random-effects

    Poisson regression model

$$(y_{i}|\lambda_{i},\mathcal{B}) \stackrel{\text{indep}}{\sim} \text{Poisson}(\lambda_{i})$$

$$\log \lambda_{i} = \beta_{0} + \beta_{1}x_{i} + e_{i}$$

$$(e_{i}|\sigma_{e}^{2},\mathcal{B}) \stackrel{\text{IID}}{\sim} N(0,\sigma_{e}^{2}), \quad (\beta_{0},\beta_{1},\sigma_{e}^{2}) \sim \text{diffuse},$$

$$(40)$$

where the  $y_i$  are counts of a relatively rare event and  $x_i$  is 1 for the treatment group and 0 for control; You would consider fitting this model instead of its fixed-effects counterpart, obtained by setting  $\sigma_e^2 = 0$ , to describe unexplainable heterogeneity (Poisson over-dispersion).

In this setting, Bayes factors will make the mistake of {telling You that  $\sigma_e^2=0$  when it's not} more often than log scores, and log scores will make the error of {telling You that  $\sigma_e^2>0$  when it's actually 0} more often than Bayes factors, but the former mistake is much worse than the latter, because You will underpropagate uncertainty about the fixed effect  $\beta_1$ , which is the whole point of the investigation.

All through this discussion it's vital to keep in mind that

the **gold standard** for **false-positive/false-negative behavior** is provided **neither by Bayes factors nor by log scores** but instead by **Bayesian decision theory in Your problem**.

Asymptotic conclusions are often misleading: while it's true that

**Old Theorem:** 
$$P_{\theta_{DG}=0}(LS_{FS} \text{ chooses } \theta=0) \to 0 \text{ as } n \to \infty,$$

it's also true that

and the **second theorem** would seem to **call the relevance of the first theorem into question**.

• As a profession, we need to strengthen the progression

$$\textbf{Principles} \rightarrow \textbf{Axioms} \rightarrow \textbf{Theorems}$$

in optimal model specification; the Calibration Principle, the Modeling-As-Decision Principle, the Prediction Principle and the Decision-Versus-Inference Principle seem helpful in moving toward this goal.

# Is M<sub>1</sub> Good Enough?

What about  $Q_2$ : Is  $M_1$  good enough?

As discussed previously, by the Modeling-As-Decision Principle a full judgment of adequacy requires real-world input ("To what purpose will the model be put?"), so it's not possible to propose generic methodology to answer  $Q_2$  (apart from maximizing expected utility, with a utility function that's appropriately tailored to the problem at hand), but the somewhat related question

 $\overline{Q_{2'}}$ : Could the data have arisen from model  $M_j$ ?

can be answered in a general way by simulating from  $M_j$  many times, developing a distribution of (e.g.)  $LS_{FS}$  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), which **produces** a *P*-value.

However, **this sort of thing** needs to be **done carefully** (Draper 1996), or the result will be **poor calibration**; indeed, Bayarri and Berger (2000) and Robins et al. (2000) have **demonstrated** that the

# Is $M_1$ Good Enough? (continued)

**Gelman et al. procedure** may be **(sharply) conservative**: You may get  $P = \mathbf{0.4}$  from Gelman et al. (indicating that **Your model is fine**) when a **well-calibrated** version of **their idea** would have  $P = \mathbf{0.04}$  (indicating that it's **not fine**).

Using a modification of an idea suggested by Robins et al., Draper and Krnjajić (2010) have developed a simulation-based method for accurately calibrating the log-score scale (I'd be happy to send You the paper).

How should You **judge how unusual** the **actual data set's log score** is in **the simulation distribution**?

In all of Bayesian inference, prediction and decision-making, except for calibration concerns, there's no need for *P*-values, but — since this is a calibrative question — it's no surprise that tail areas (or something else equally ad-hoc, such as the ratio of the attained height to the maximum height of the simulation distribution) arise.

I don't see how to avoid this ad-hockery except by directly answering  $Q_2$  with decision theory (instead of answering  $Q_{2'}$  with a tail area).

### Summary

- I've offered an axiomatization of inferential, predictive and decision-theoretic statistics based on information, not belief, and RT Cox's (1946) notion of probability as a measure of the weight of evidence in favor of the truth of a true-false proposition whose truth status is uncertain for You.
  - Cox's Theorem lays out a progression from

 $\textbf{Principles} \rightarrow \textbf{Axioms} \rightarrow \textbf{Theorem}$ 

to prove that Bayesian reasoning is justified under natural logical consistency assumptions; for me this secures the foundations of applied probability.

- But Cox's Theorem does not go far enough for statistical work in science, in two ways related to model specification:
  - Nothing in its consequences requires You to pay attention to how often You get the right answer, which is a basic scientific concern, and

- it doesn't offer any advice on how to specify the required ingredients: with  $\theta$  as the unknown of principal interest,  $\mathcal B$  as Your relevant background assumptions and judgments, and an information source (data set) D relevant to decreasing Your uncertainty about  $\theta$ , the ingredients are
  - \*  $\{p(\theta|\mathcal{B}), p(D|\theta|\mathcal{B})\}$  for inference and prediction, and
- \* in addition  $\{\mathcal{A}, U(a,\theta)\}$  for decision, where  $\mathcal{A}$  is Your set of available actions and  $U(a,\theta)$  is Your utility function (mapping from actions a and unknown  $\theta$  to real-valued consequences).
- To secure the foundations of statistics, work is needed laying out the logical progression

 $\textbf{Principles} \rightarrow \textbf{Axioms} \rightarrow \textbf{Theorems}$ 

for model specification; progress in this area is part of the Theory of Applied Statistics.

 A Calibration Principle helps address the first of the two deficiencies above:

Calibration Principle: In model specification, You should pay attention to how often You get the right answer, by creating situations in which You know what the right answer is and seeing how often Your methods recover known truth.

Interest in calibration can be seen to be natural in Bayesian work by thinking decision-theoretically, with a utility function that rewards both quality of scientific conclusions and good calibration of the modeling process yielding those conclusions.

- In problems of **realistic complexity** You'll generally notice that (a) You're **uncertain** about  $\theta$  but (b) You're also **uncertain** about how to **quantify Your uncertainty about**  $\theta$ , i.e., **You** have **model uncertainty**.
- This acknowledgment of Your model uncertainty implies a willingness by You to consider two or more models in an ensemble  $\mathcal{M} = \{M_1, M_2, \dots\}$ , which gives rise immediately to two questions:
  - $Q_1$ : Is  $M_1$  better than  $M_2$ ?  $Q_2$ : Is  $M_1$  good enough?

• These questions sound fundamental but are not: better for what purpose? Good enough for what purpose? To address the second of the two deficiencies above (lack of guidance from Cox's Theorem on model specification), this implies a

Modeling-As-Decision Principle: Making clear the purpose to which the modeling will be put transforms model specification into a decision problem, solvable by maximizing expected utility with a utility function tailored to the specific problem under study.

This solves the model-specification problem but is hard work; there's a powerful desire for generic model-comparison methods whose utility structure may provide a decent approximation to problem-specific utility elicitation.

Two such methods are **Bayes factors** (whose **utility justification** is **less than compelling**) and **log scores**, which are based on the

**Prediction Principle:** Good models make good predictions, and bad models make bad predictions; that's one scientifically important way You know a model is good or bad.

- I'm aware of three approaches to improved assessment and propagation of model uncertainty: Bayesian model averaging (BMA), Bayesian nonparametric (BNP) modeling, and calibration (3-fold) cross-validation (CCV).
- CCV provides a way to pay the right price for hunting around in the data for good models, motivating the following modeling algorithm:
  - (a) Start at a model  $M_0$  (how choose?); set the current model  $M_{\text{current}} \leftarrow M_0$  and the current model ensemble  $\mathcal{M}_{\text{current}} \leftarrow \{M_0\}$ .
  - (b) If  $M_{current}$  is good enough to stop (how decide?), return  $\mathcal{M}_{current}$ ; else
  - (c) Generate a new candidate model  $M_{\text{new}}$  (how choose?) and set  $\mathcal{M}_{\text{current}} \leftarrow \mathcal{M}_{\text{current}} \cup M_{\text{new}}$ .
  - (d) If  $M_{\text{new}}$  is better than  $M_{\text{current}}$  (how decide?), set  $M_{\text{current}} \leftarrow M_{\text{new}}$ .
  - (e) Go to (b).
  - For the **choice** in **(a)**, there's usually a **default off-the-shelf initial model** based on the **structure** of the **data set** *D* and the **scientific context**.

- In manual model search the choice in (c) is typically based on the results of a variety of diagnostics, with the new model suggested by deficiencies revealed in this way; at present, we have no better way to automate this choice in many cases than choosing  $M_{new}$  at random (I offer no new ideas on this topic today).
- In comparing  $M_1$  with  $M_2$  (the choice in (d)), consider a calibrative scenario in which the the data-generating model  $M_{DG}$  is one or the other of  $\mathcal{M} = \{M_1, M_2\}$  (apart from parameter estimation), and call {choosing  $M_2$  when  $M_{DG} = M_1$ } a false positive and {choosing  $M_1$  when  $M_{DG} = M_2$ } a false negative; then
- The **right way** to do this, following the **Modeling-As-Decision Principle**, is to build a **utility function** by **quantifying** the **real-world consequences** of

```
{choosing M_1 when M_{DG} = M_1, choosing M_1 when M_{DG} = M_2, choosing M_2 when M_{DG} = M_1, choosing M_2 when M_{DG} = M_2}
```

and maximize expected utility.

- If instead You contemplate using Bayes factors/BIC or log scores, it is not the case that one of these two methods uniformly dominates the other in calibrative performance; in some settings they behave the same, in others (for Your sample size) they will have a different balance of false positives and false negatives; it's a good idea to investigate this before settling on one method or the other.
  - See Draper and Krnjajić (2013) for a **method** for **answering the question**  $Q_{2'}$ : Could the data have arisen from model  $M_j$ ? in a well-calibrated way.
  - CCV provides an approach to finding a good ensemble  $\mathcal{M}$  of models, and gives You a decent opportunity both to arrive at good answers to Your main scientific questions and to evaluate the calibration of the iterative modeling process that led You to Your answers.
- Decision-Versus-Inference Principle: We should all get out of the habit of using inferential methods to make decisions: their implicit utility structure is often far from optimal.

#### Another Unsolved Foundational Problem

 One more unsolved foundational problem: how can good decisions be arrived at when "You" is a collective of individuals, all with their own utility functions that imply partial cooperation and partial competition?

have agreed to band together in some sense (i.e., politics, at the level of family or nation or ...).

An instance of this: Defining and funding good quality of health care — the actors in the drama include

{patient, doctor, hospital, state and local regulatory bodies, federal regulatory system};

all are in partial agreement and partial disagreement on how (and how many) resources should be allocated to the problem of addressing this patient's immediate health needs.

(But that's for **another day**, as is the topic of **Bayesian computing** with **large data sets**.)

#### Cromwell's Rule, Part 1: Inference and Prediction

The following two facts are easy consequences of the definition of conditional probability: for any two propositions A and B and any background information  $\mathcal{B}$ :

- Cromwell's Rule, Part 1(a) If P(A|B) = 0 then P(A|BB) = 0 for all B for which P(A|BB) is defined (i.e., for which P(B|B) > 0).
- Cromwell's Rule, Part 1(b) If P(A|B) = 1 then P(A|BB) = 1 for all B for which P(A|BB) is defined (i.e., for which P(B|B) > 0).

To see the implications of these facts, let A be a proposition about something unknown to You, such as  $(\theta < 0)$ , and let B be a proposition about Your data set D, such as  $(y_1 = 3, y_2 = -0.4, \dots, y_n = 6.9)$ .

Then Part 1(a) of Cromwell's Rule says that any proposition about the unknown  $\theta$  to which You give prior probability 0 must have posterior probability 0, no matter how the data set D comes out, and Part 1(b) of Cromwell's Rule says the same thing with 0 replaced by 1.

Bayes's Theorem is supposed to be a piece of machinery that permits You to learn, about unknowns from new data, in an optimal way; Cromwell's Rule Part 1 says that if You dogmatically place prior probability 0 or 1 on something, no learning is possible when new data values arrive.

This is obviously a way to break the Bayes's Theorem learning machine, so the practical consequence of Cromwell's Rule Part 1 is captured in the following piece of advice:

You should try hard never to put prior probability 0 or 1 on anything that might later have posterior probability between 0 and 1, depending on how new information comes out.

This has direct consequences for Bayesian model specification: for example, consider the NB10 data set from Day 1, for which (by exchangeability) Your basic sampling model for the data values  $y = (y_1, \ldots, y_n)$  before You see the data is  $(y_i|FB) \stackrel{\text{IID}}{\sim} F$  for some (unknown) CDF F.

If, before You see the data, You put all Your modeling eggs in the Gaussian basket, so that You replace  $(y_i|F\mathcal{B}) \stackrel{\text{IID}}{\sim} F$  with  $(y_i|\mu\,\sigma\,\mathcal{B}) \stackrel{\text{IID}}{\sim} \mathcal{N}(\mu,\sigma^2)$ , this means that You've placed all of your prior probability on the Gaussian family, thereby implicitly giving prior probability 0 to all non-Gaussian behavior (such as multimodality, skewness and/or heavy tails).

Now what do You do when the data set arrives and — as in the case of the NB10 data — demonstrates much heavier tails than the Gaussian family can accommodate?

Strictly speaking, by Cromwell's Rule Part 1(a), all such non-Gaussian behavior must have posterior probability 0, and yet the data set clearly makes You wish that You hadn't been so dogmatic in Your "prior on model space."

Going back and changing Your prior on model space based on how the data set came out is a clear violation of the dichotomization  $\{\text{information internal to } D, \text{ information external to } D\}$ , which was part of the axiomatization in the Day 3 (Part 1) Lecture Notes:

#### Cromwell's Rule, Part 2: Decision-Making

in effect, when You do this You're using the data twice — once in specifying the prior on model space, and again in updating that prior with the data set D — and the typical consequence will be understatement of Your actual uncertainty.

I see only two ways out of this dilemma:

- Bayesian nonparametric methods, which when used properly give positive prior probability to all possible CDFs F), and
- Calibration cross-validation, which (1) allows You to "cheat" by looking at the data and changing Your prior on model space but (2) forces You to pay an appropriate price for having done so.

Cromwell's Rule, Part 2: Part 1 of Cromwell's Rule is about

inference and prediction; Part 2 concerns decision-making, and it also has a part (a) and a part b), which give advice on how to specify Your action space  $\mathcal{A}$  and Your utility function  $U(a,\theta)$  (respectively).

• Cromwell's Rule, Part 2(a): In enumerating the possible actions  $\{a_1,a_2,\dots\}$  while specifying Your action space  $\mathcal A$  in the problem  $\mathbb P$  on which You're working, You should try hard not to omit any action  $a_i$  that might turn out to be optimal if it's included in  $\mathcal A$ .

The point is that Bayesian decision theory only optimizes over the possible actions You remember to put in  $\mathcal{A}$ : if You forget a feasible action  $a_i$  that (unknown to You) is better than all of the actions in Your current  $\mathcal{A}$ , maximization of expected utility cannot protect You from this omission.

Example: HIV screening with *ELISA* and Western Blot (Day 1, Lecture Notes Part 1). In that case study it was tempting to think that the only two possible actions were  $a_1 = \{\text{test the blood sample with ELISA}\}$  and  $a_2 = \{\text{test the blood sample with Western Blot}\}$ , but a third action —  $a_3 = \{\text{test half of the blood sample with ELISA}\}$ ; if negative, declare HIV negative; if positive, test the other half of the blood sample with Western Blot} — turned out to dominate the others.

Example: People at eBay are constantly running randomized controlled trials on the eBay web experience, looking for variations on things like (i) search algorithms and (ii) {presentation of items for sale to the users} that may create a better marketplace.

Having run an experiment in which the control group gets the current best eBay website and the treatment group gets {the current best eBay website plus a particular intervention I}, thereby obtaining a data set D, the unknown  $\theta$  is {what the future would be like, as far as important outcome variables (such as user satisfaction) are concerned, if intervention I is or is not implemented}, and it appears that there are only two possible actions:  $a_1 = \{\text{implement } I\}$  and  $a_2 = \{\text{don't implement } I\}$ .

However, as with the HIV case study, there's a third possible action that has an adaptive flavor: perhaps there's still too much uncertainty, on the basis of D, to make a good choice between  $a_1$  and  $a_2$ , so (if You're not in a tremendous hurry to choose) why not include  $a_3 = \{\text{get more data before deciding}\}$  in Your  $\mathcal{A}$ ?

Bayesian sequential experimental design and analysis has this adaptive character — get some data, see if the optimal choice is clearcut yet, if so make it, if not get more data — and have been shown to yield results with good false-positive and false-negative rates that involve collecting (far) less data than approaches with sample sizes that are fixed at the design stage.

- Cromwell's Rule, Part 2(a): In specifying Your utility function for the problem ℙ on which You're working, You should try hard to ensure that
- (1) Your vector of unknowns  $\theta$  contains all relevant unknowns, and
  - (2) Your utility function  $U(a, \theta)$  captures <u>all relevant</u> costs and benefits to be balanced against each other.

#### The **point** is that

• any relevant unknown that You mistakenly omit from  $\theta$  has no opportunity to influence Your decision; the result will often be decisions that don't hedge sufficiently against uncertainty,

because **omitting** a **relevant unknown** is **tantamount** to **pretending that it's known**; and

ullet any relevant cost or benefit that You mistakenly omit from  $U(a, \theta)$  has no opportunity to influence the optimal trade-off between costs and benefits, and (dramatically) sub-optimal decisions can result when this happens.

In enumerating the relevant costs and benefits, You'll have to fight against the following three basic human tendencies:

(1) Things that are easy to measure tend to get measured; things that are hard to measure tend to get ignored.

With respect to a particular cost or benefit, the classifications  $\mathcal{C}_1 = \{\text{important, not important}\}$  and  $\mathcal{C}_2 = \{\text{hard to quantify, easy to quantify}\}$  have nothing to do with each other; including or omitting costs and benefits solely on the basis of  $\mathcal{C}_2$  provides no assurance that the included costs and benefits are correct with respect to  $\mathcal{C}_1$ .

- People with optimistic world views tend to exaggerate benefits and downplay costs; pessimists tend to make the opposite mistakes; not many people get this right unless they're on the lookout for it.
- in Your utility function, You'll typically find it relatively easy to choose the scale on which to quantify the costs (e.g., in monetary terms), but it may then be quite difficult to quantify the benefits on the same scale.

Example: The new drug Olysio (simeprevir, approved for use in the U.S. in Nov 2013) has been shown to be quite effective at reducing the viral load of Hepatitis C patients, permitting their livers to begin to heal, as long as the drug is taken for at least three months.

The drug company that markets *Olysio*, Janssen, has set the wholesale price of a 12-week supply of this drug at \$66,360 (about £41,000).

Q: Should the NHS decide to approve *Olysio* for routine treatment of Hepatitis C in the UK?

The cost term in the NHS's utility function clearly comes out in £; to trade this off against the health benefits (e.g., lengthened life span, better quality of life), the NHS has to be prepared to measure those benefits in monetary terms, leading to unpleasant questions such as "What's the monetary value of human life?"

Example: Draper (1995) examines an attempt by economists in 1980 to predict future oil prices over the horizon 1981–2020; many companies and governments routinely make investment decisions based on such predictions.

The OPEC oil embargo of 1973–74 created a large spike in oil prices, because demand stayed constant and supply dramatically dropped; this was the first time anything like that had ever occurred.

If You had undertaken the 1980 predictive exercise mentioned above prior to 1973, You would have had no straightforward way to include {Will another OPEC oil embargo occur, and if so when?} as part of the unknowns in Your  $\theta$  vector, but You would have no excuse for omitting this unknown after 1973–74, and indeed this omission could lead to a dramatic understatement of Your future uncertainty about the price of oil, causing You to make decisions that fail to hedge sufficiently against the totality of Your uncertainty.

Example: You're about to make a long drive by car, and You're wondering about the optimal driving speed: the faster You drive the quicker You get to Your destination (good), but undesirable outomes increase in probability as You speed up (bad).

The action space clearly consists of possible speeds (that was easy), but what about  $\theta$  and  $U(a, \theta)$ ?

As a first pass, You might include in  $\theta$  only the unknown {will You get a speeding ticket?}, in which case Your utility function would have only two terms, which could be combined additively:

a benefit (quantified in monetary terms) based on how short the journey is, and a cost (also expressed in money) based on what happens if You get a ticket (You have to pay a fine, and Your insurance costs may rise).

I've found that this formulation typically leads to a recommendation to drive quite rapidly.

However, a number of additional relevant unknowns have been omitted from this first-pass specification of A and  $U(a, \theta)$ :

{Will You get into an accident?} If so, {How serious is the accident?} {How badly is Your car damaged?} {Does the accident injure or kill You?} If other people are involved in the accident, {How badly is their vehicle damaged?} {Does the accident injure or kill any of them?}

Increasingly unfavorable answers to all of these questions will all result in additional cost terms in Your utility function, with the result that Your optimal driving speed will monotonically decrease as You increase the realism of Your utility specification.