

Bayesian Statistics for Genetics

Lecture 6: Model Selection and Model Averaging

July, 2023

Motivation

Diabetes example:

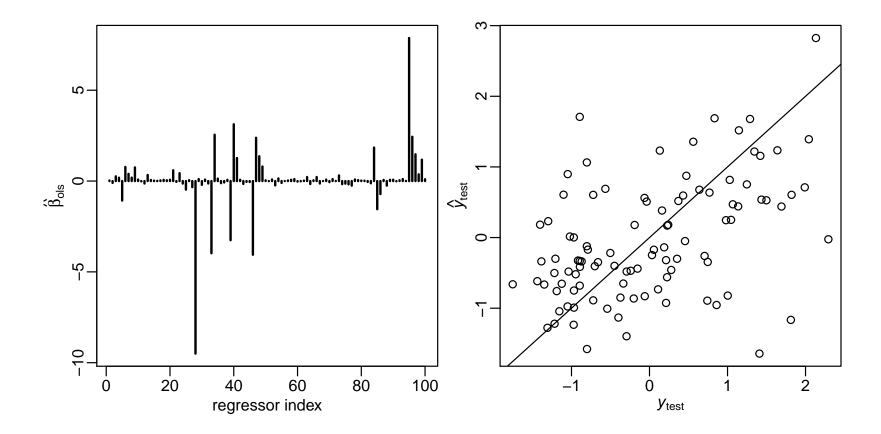
- n=342 subjects
- $y_i = \text{diabetes progression}$
- $x_i = explanatory variables.$

Each x_i includes

- 13 subject specific measurements $(x_{age}, x_{sex}, \ldots)$;
- 78 = $\binom{13}{2}$ interaction terms $(x_{age} \cdot x_{sex}, ...)$;
- 9 quadratic terms (x_{sex} and three genetic variables are binary)

100 explanatory variables total!

OLS regression



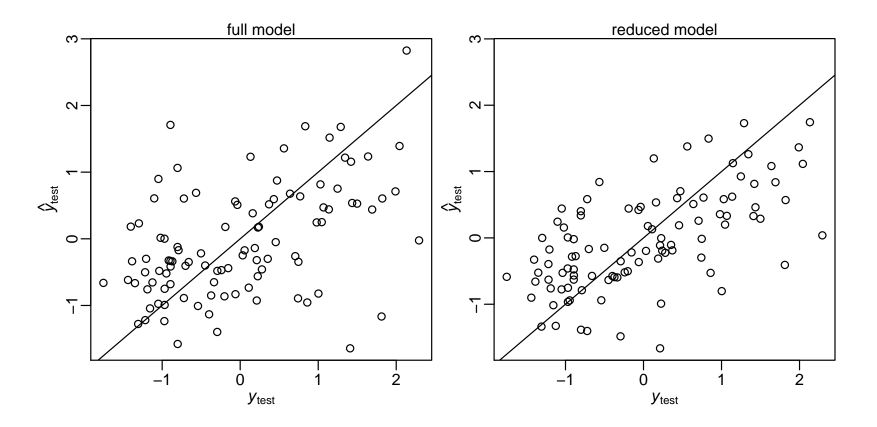
$$\frac{1}{100} \sum (y_{\text{test},i} - \hat{y}_{\text{test},i})^2 = 0.93, \text{ while } \frac{1}{100} \sum (y_{\text{test},i} - 0)^2 = \sum y_{\text{test},i}^2 = 1.01$$

Backwards elimination

1. Obtain the estimator $\hat{\beta}_{ols} = (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T y$ and its *t*-statistics.

- 2. If there are any regressors j such that $|t_j| < t_{cutoff}$, (a) find the regressor j_{min} having the smallest value of $|t_j|$ and remove column j_{min} from **X**.
 - (b) return to step 1.
- 3. If $|t_j| > t_{cutoff}$ for all variables j remaining in the model, then stop.

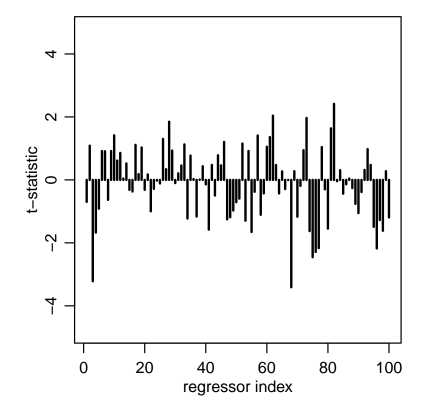
Backwards elimination



 $\frac{1}{100}\sum(y_{\text{test},i} - \hat{y}_{\text{test},i}^{bel})^2 = 0.64, \ \dots \ \text{annoyingly!}$

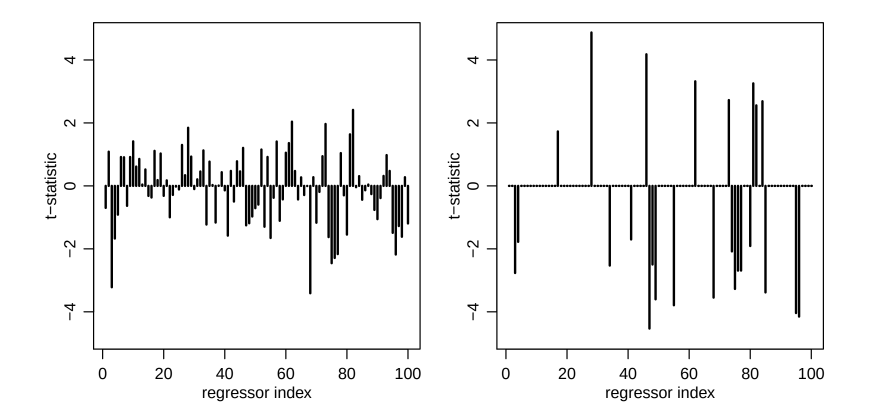
Spurious associations

Now try modeling permuted $y_{\pi(i)} = \boldsymbol{\beta}^T \boldsymbol{x}_i + \epsilon_i$ (and backwards-select)



Spurious associations

Now try modeling permuted $y_{\pi(i)} = \boldsymbol{\beta}^T \boldsymbol{x}_i + \epsilon_i$ (and backwards-select)



Spurious associations

```
sum(abs(t.bslperm)>2 )
## [1] 21
sum(abs(t.bslperm)>3 )
## [1] 12
sum(abs(t.bslperm)>4 )
## [1] 5
```

- 21 regressors have *t*-stats > 2 ($p \approx 0.05$)
- 12 regressors have t-stats > 3 ($p \approx 0.003$)
- 5 regressors have t-stats > 4 ($p \approx 0.00006$)

Often want some way to pick a sparse model – but this approach is not smart if we want to say how reliable our pick is.

We have prior belief that $\beta_j \approx 0$ for many j's; a model allowing this specifies $\beta_j = z_j \times b_j$, where $z_j \in \{0, 1\}$ and $b_j \in \mathbb{R}$.

$$y_i = z_1 b_1 x_{i,1} + \dots + z_p b_p x_{i,p} + \epsilon_i.$$

For example, in Session 4's FTO experiment,

$$\mathbb{E}[Y|x, b, z = (1, 0, 1, 0)] = b_1 x_1 + b_3 x_3$$

= $b_1 + b_3 \times age$
$$\mathbb{E}[Y|x, b, z = (1, 1, 0, 0)] = b_1 x_1 + b_2 x_2$$

= $b_1 + b_2 \times group$
$$\mathbb{E}[Y|x, b, z = (1, 1, 1, 0)] = b_1 x_1 + b_2 x_2 + b_3 x_3$$

= $b_1 + b_2 \times group + b_3 \times age.$

Can think of each value of $z = (z_1, \ldots, z_p)$ representing a different model.

But easier to implement thinking of z_j as unknown components in one (big) model – written informally as;

$$z_{j} \stackrel{i.i.d.}{\sim} Bern(0.5)$$

$$b_{j} \sim p(b_{j})$$

$$\epsilon_{i} \stackrel{i.i.d.}{\sim} N(0, \sigma^{2})$$

$$\sigma^{2} \sim p(\sigma^{2})$$

$$y_{i} = z_{1}b_{1}x_{i,1} + \dots + z_{p}b_{p}x_{i,p} + \epsilon_{i}$$

Each of the 2^p possible values of of z has a posterior probability. (In the prior we treat them as a 'coin toss', equally likely to be 'in' or 'out'.)

Bayesian model comparison

The posterior probability of the submodels is obtained from

$$p(\boldsymbol{z}|\boldsymbol{y}, \boldsymbol{X}) = \frac{p(\boldsymbol{z})p(\boldsymbol{y}|\boldsymbol{X}, \boldsymbol{z})}{p(\boldsymbol{y}|\boldsymbol{X})}$$

To compare submodels a and b, usually consider the odds of each, and how they compare:

$$\frac{p(\boldsymbol{z}_a | \boldsymbol{y}, \boldsymbol{X})}{p(\boldsymbol{z}_b | \boldsymbol{y}, \boldsymbol{X})} = \frac{p(\boldsymbol{z}_a)}{p(\boldsymbol{z}_b)} \times \frac{p(\boldsymbol{y} | \boldsymbol{X}, \boldsymbol{z}_a)}{p(\boldsymbol{y} | \boldsymbol{X}, \boldsymbol{z}_b)}$$
sosterior odds = prior odds × "Bayes factor"

Importantly, the Bayes Factor (BF) does not depend on the prior for z – so the 'coin toss' prior is not crucial for this approach.

Parsimony

The formula for p(y|x,z) is complex, but

$$\frac{p(\boldsymbol{y}|\mathbf{X}, \boldsymbol{z}_{a})}{p(\boldsymbol{y}|\mathbf{X}, \boldsymbol{z}_{b})} = (1+n)^{(p_{z_{b}} - p_{z_{a}})/2} \left(\frac{s_{z_{a}}^{2}}{s_{z_{b}}^{2}}\right)^{1/2} \times \left(\frac{s_{z_{b}}^{2} + SSR_{g}^{z_{b}}}{s_{z_{a}}^{2} + SSR_{g}^{z_{a}}}\right)^{(n+1)/2}$$

where SSR_g denotes a form of sum of squared residuals.

So a model z_a is penalized if;

- it is too complex (number of covariates p_A is large)
- it doesn't fit well (SSR $_q^a$ is large)

$\mathbb{E}[Y_i|\boldsymbol{\beta}, \boldsymbol{x}_i] = \beta_1 x_{i,1} + \beta_2 x_{i,2} + \beta_3 x_{i,3} + \beta_4 x_{i,4} \\ = \beta_1 + \beta_2 \times \operatorname{grp}_i + \beta_3 \times \operatorname{age}_i + \beta_4 \times \operatorname{grp}_i \times \operatorname{age}_i.$

effect of group \Leftrightarrow one of more of β_2, β_4 not zero

z	model	$\log p(m{y} m{X},m{z})$	$p(oldsymbol{z} oldsymbol{y},oldsymbol{X})$
(1,0,0,0)	β_1	-71.82	0
(1,1,0,0)	$\beta_1 + \beta_2 \times \operatorname{grp}_i$	-70.04	0
(1,0,1,0)	$\beta_1 + \beta_3 \times age_i$	-67.04	0
(1,1,1,0)	$\beta_1 + \beta_2 \times \operatorname{grp}_i + \beta_3 \times \operatorname{age}_i$	-61.19	0.63
(1,1,1,1)	$\beta_1 + \beta_2 \times \operatorname{grp}_i + \beta_3 \times \operatorname{age}_i + \beta_4 \times \operatorname{grp}_i \times \operatorname{age}_i$	-61.72	0.37

 $\mathbb{P}[\beta_2 \text{ or } \beta_4 \neq 0] = 0.60$ $\mathbb{P}[\beta_2 \text{ or } \beta_4 \neq 0 | \boldsymbol{y}, \boldsymbol{X}] \approx 1$

Using the conjugate g-prior is a little artificial here;

- Each sub-model has a prior that corresponds to one observation's information, but those observations are not the same.
- It's strange to support the model with all $\beta_j = 0$, i.e. where $\mathbb{E}[Y_i|x_i]$ is exactly zero for everyone

So we'll instead use a general-purpose Gibbs sampler for the same model, but with $z_1 = 1$ (forcing an intercept) and

$$\begin{array}{rcl} z_{j} & \stackrel{i.i.d.}{\sim} & Bern(0.5) \\ b_{j} & \sim & N(0,10), \text{ for } j = 2,3,4 \\ \epsilon_{i} & \stackrel{i.i.d.}{\sim} & N(0,\sigma^{2}) \\ 1/\sigma^{2} & \sim & \Gamma(0.5,1.839) \ \dots \ \text{as in Lec } 4 \\ y_{i} & = & z_{1}b_{1}x_{i,1} + \dots + z_{p}b_{p}x_{i,p} + \epsilon_{i} \end{array}$$

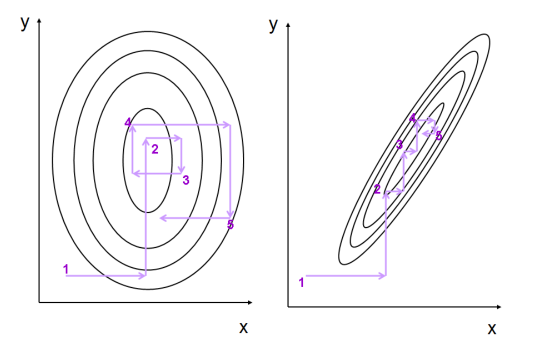
Goal: A Monte Carlo approximation to p(x, y, z)

Given $\{x^{(s)}, y^{(s)}, z^{(s)}\},\$

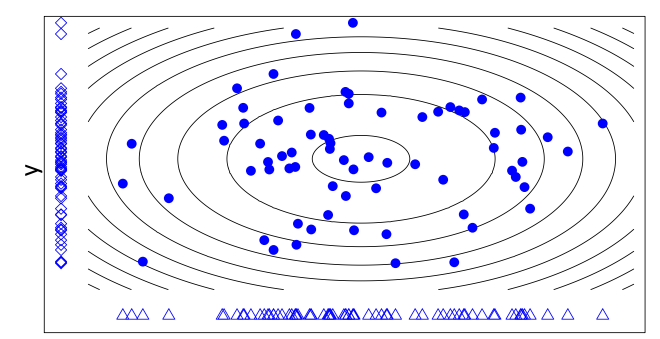
- 1. simulate $x^{(s+1)} \sim p(x|y^{(s)}, z^{(s)})$,
- 2. simulate $y^{(s+1)} \sim p(y|x^{(s+1)}, z^{(s)})$,
- 3. simulate $z^{(s+1)} \sim p(z|x^{s+1}), y^{(s+1)})$.

This generates $\{x^{(s+1)}, y^{(s+1)}, z^{(s+1)}\}$ – and then 'go round' again, many times. Repeated many times, this generates $\{x^{(1)}, y^{(1)}, z^{(1)}\}, \dots, \{x^{(S)}, y^{(S)}, z^{(S)}\}$

For a couple of two-dimensional examples;



Output from a short sampler;



Sample (points) approximate distribution (contours)

Repeated many times, this gives $\{x^{(1)}, y^{(1)}, z^{(1)}\}, \dots, \{x^{(S)}, y^{(S)}, z^{(S)}\}$

The distribution of this sequence approximates p(x, y, z):

$$\frac{1}{S} \sum x^{(s)} \approx \mathbb{E}x = \int x \ p(x, y, z) \ dx \ dy \ dz$$
$$\frac{\#(x^{(s)} \in A)}{S} \approx \Pr(x \in A) = \int \int \int_A p(x, y, z) \ dx \ dy \ dz$$
$$\frac{\#(\{x^{(s)}, y^{(s)}, z^{(s)}\} \in B)}{S} \approx \int \int_B \int p(x, y, z) \ dx \ dy \ dz$$

By necessity, the sequence will frequently visit regions where p(x, y, z) is large.

Gibbs sampling: for model selection

Goal Approximate $p(z_1, \ldots, z_p | \boldsymbol{y}, \boldsymbol{X})$.

Gibbs sampler: Given $z^{(s)} = (z_1^{(s)}, \dots, z_p^{(s)}),$ $z_1^{(s+1)} \sim p(z_1 | z_2^{(s)}, \dots, z_p^{(s)}, y, X)$ $z_2^{(s+1)} \sim p(z_2 | z_1^{(s+1)}, z_3^{(s)}, \dots, z_p^{(s)}, y, X)$: $z_p^{(s+1)} \sim p(z_p | z_1^{(s+1)}, \dots, z_{p-1}^{(s+1)}, y, X)$

This generates $z^{(s+1)}$ from $z^{(s)}$.

Repeating this generates $z^{(1)}, \ldots, z^{(S)}$ with which to approximate $p(\boldsymbol{z}|\boldsymbol{y}, \boldsymbol{X})$.

Stan can't handle discrete parameters (yet) so we'll use JAGS – Just Another Gibbs Sampler. JAGS writes and executes MCMC code, given a model and data.

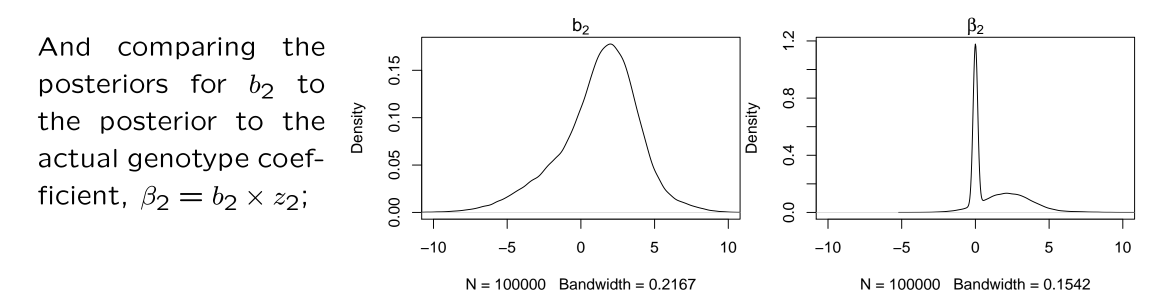
```
library("rjags")
# first, write the model as to a text file
cat(file="linearprog2.txt", "model{
   for(j in 1:p){
      b[j]~dnorm(0, 0.1) }
   z[1] <- 1 # fix the intercept to be in the model
   for(j in 2:p){
      z[j] ~ dbern(0.5) }
inv.sigma2 ~ dgamma( 0.5, 1.839 )
   sigma <- sqrt(1/inv.sigma2)</pre>
   for(i in 1:n){
mu[i] <- x[i,1]*b[1]*z[1] + x[i,2]*b[2]*z[2] + x[i,3]*b[3]*z[3] + x[i,4]*b[4]*z[4]</pre>
y[i] ~ dnorm(mu[i], inv.sigma2) }
}")
# compile code based on model and data, then run chain
jags1 <- jags.model("linearprog2.txt", data=list(y=y,x=X, n=nrow(X), p=ncol(X)) )</pre>
update(jags1, 50000) # initial iteraions
```

> jags1.out <- coda.samples(jags1, c("b","inv.sigma2", "z"), n.iter=100000)[[1]]</pre> And some of > summary(jags1.out) the output; Tterations = 50001:150000Number of chains = 1Sample size per chain = 1e+051. Empirical mean and standard deviation for each variable & std err of the mean: SD Naive SE Time-series SE Mean b[1] 0.7593 1.26609 0.0040037 0.0184052 b[2] 1.2431 2.71152 0.0085746 0.0300475 b[3] 2.6202 0.39962 0.0012637 0.0057575 b[4] 2.1791 0.62138 0.0019650 0.0091990 inv.sigma2 0.2676 0.09069 0.0002868 0.0004338 z[1] 1.0000 0.00000 0.000000 0.000000 z[2] 0.5604 0.49634 0.0015696 0.0058886 z[3] 1.0000 0.00000 0.000000 0.000000 z[4] 0.9928 0.08431 0.0002666 0.0015052

The coefficient of genotype is $\neq 0$ with 56% posterior support; the interaction term being $\neq 0$ has 99% support. The chain never moved from supporting age term $\neq 0$, so it has (approximately) 100% support.

All 100,000 steps in the chain are stored, so we can assess posterior for other terms – for example the support for each set of included/excluded variables;

> table(apply(jags1.out[,c("z[1]","z[2]","z[3]","z[4]")], 1, paste, collapse=""))/100000 1011 1110 1111 0.43851 0.00693 0.55456



Using MCMC, we have to start the 'chain' somewhere – but this arbitrary choice shouldn't affect analysis, if we run the chains for long enough.

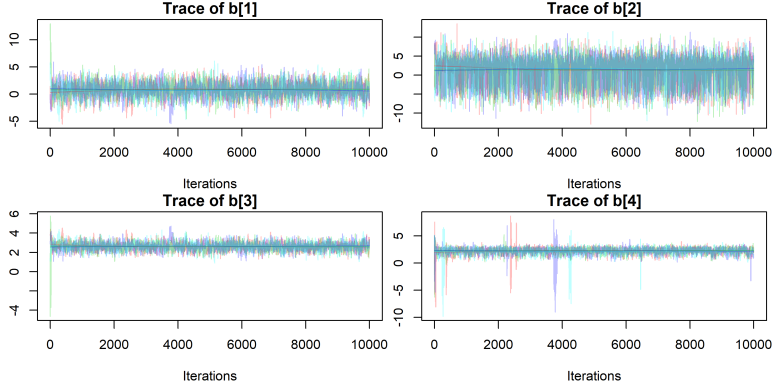
- After running long enough, the chains from any two starting points should converge to cover the posterior in the same way
- Less formally, after running long enough, chains forget where they started
- It's pragmatic (but not perfect) to run chains from a few different starting points, and check they give similar answers

JAGS makes this fairly painless – here for 4 short chains;

```
set.seed(4)
inits1 <- list( b=rnorm(4,0,1),inv.sigma2=0.5,z=c(NA,0,1,0))
inits2 <- list( b=rnorm(4,0,1),inv.sigma2=0.5,z=c(NA,0,0,0))
inits3 <- list( b=rnorm(4,0,1),inv.sigma2=0.5,z=c(NA,1,1,0))
inits4 <- list( b=rnorm(4,0,1),inv.sigma2=0.5,z=c(NA,1,1,1))
jags2 <- jags.model("linearprog2.txt", data=list(y=y,x=X, n=nrow(X), p=ncol(X)),
inits=list( inits1, inits2, inits3, inits4), n.chains=4 )
jags2.out <- coda.samples(jags2, c("b"), n.iter=10000)</pre>
```

An informal way to check for convergence is to look for differences in each chain's traceplot; (no issues seen here)

plot(jags2.out, trace=TRUE, density=FALSE, auto.layout=FALSE, col=adjustcolor(2:5, alpha.f=0.25), lty=1)



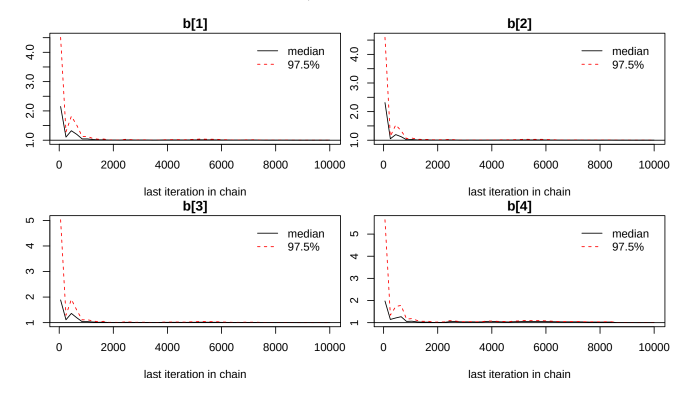
To more formally check convergence of the chains for individual parameters, the *Gelman-Rubin diagnostic* compares within-chain variance (W) to between-chain variance (B), using tools from mixed models. For a converged chain their ratio R = W/B should be $\approx 1...$

> gelman.diag(jags2.out)					
Potential scale reduction factors:					
	Point	est.	Upper	C.I.	
b[1]		1		1.00	
b[2]		1		1.00	
b[3]		1		1.00	
b[4]		1		1.01	

Similar ideas provide the effective sample size, i.e. roughly how big a simple random sample from the posterior is represented by the (auto-correlated) chain

```
> effectiveSize(jags2.out)
        b[1]        b[2]        b[3]        b[4]
1860.972 3057.044 1898.274 1586.170 # each from 40,000 iterations
```

gelman.plot(jags2.out) shows how W/B evolves over iterations;



Ideally, don't start using the chain output until it looks like it converged -& even then, use as long a chain as you can manage. Thin it, if memory is an issue.

Stochastic search: High dimensional regression

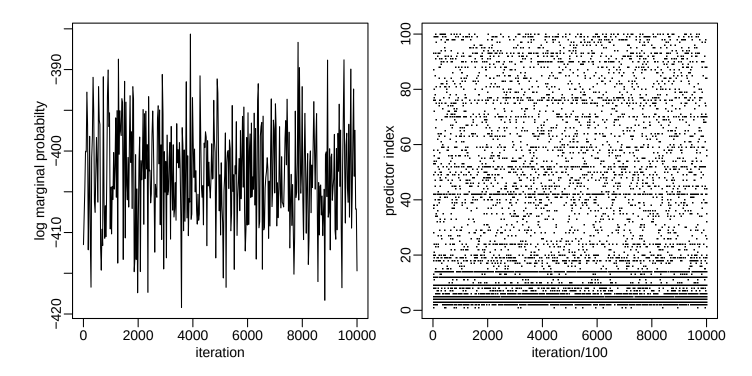
Back to the diabetes example, with p=100, meaning there are $2^{100} \approx 10^{30}$ models to consider.

That's a huge number! We can't compute $\mathbb{P}[z|y,X]$ for each z. Instead, we hope to;

- search for models z with high posterior probability;
- approximate $\beta_j = z_j \times b_j$ for each j;
- build a predictive model for y.

We can view Gibbs Sampling here as a way to explore possible models – not to fully cover the whole parameter space. It will tend to sample models that better support the data.

Diabetes example



Marginal inference

What is the estimate of β ? Recall

$$\boldsymbol{\beta} = (\beta_1, \ldots, \beta_p) = (b_1 z_1, \ldots, b_p, z_p)$$

Our Monte Carlo samples are

$$\beta^{(1)} = (0 -.299 \quad 0 \ .427 \ \cdots \ .845)$$

$$\beta^{(2)} = (0 -.235 \ .834 \ .374 \ \cdots \ 0)$$

$$\vdots \qquad \vdots$$

$$\beta^{(S)} = (0 -.315 \quad 0 \ .536 \ \cdots \ 0)$$

A posterior mean for β is obtained in the usual way:

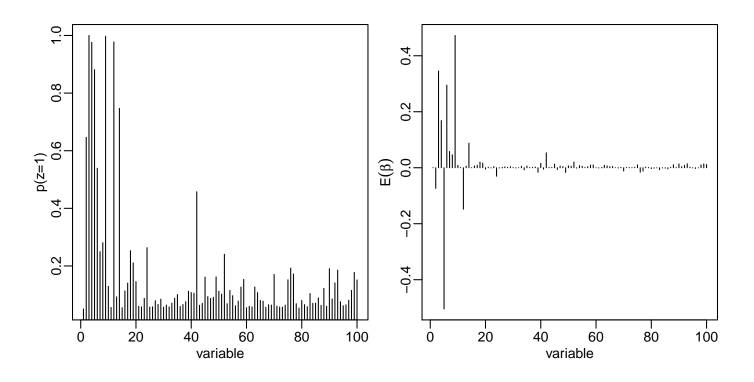
$$\widehat{\boldsymbol{eta}}^{\mathsf{bayes}} = rac{1}{S} \sum \boldsymbol{eta}^{(s)} pprox \mathbb{E} \boldsymbol{eta} | \boldsymbol{y}, \mathsf{X}$$

Out of sample predictions can be made with $\hat{\beta}_{\text{bayes}}$:

$$\widehat{y}_{\text{test},i}^{bayes} = \widehat{\boldsymbol{\beta}}_{\text{bayes}}^T \boldsymbol{x}_{\text{test},i}$$

Marginal inference

How does it do?



Out of sample prediction error:

$$\frac{1}{S}\sum(y_{\text{test},i} - \hat{y}_{\text{test},i}^{bayes})^2 = 0.485$$

A standard (if ill-defined!) question is "what variables matter most?"

Here, we can order covariates by either their posterior probability of being in the model, or the p-value from non-Bayesian backwards selection. The answers (below) are not identical, but some variables appear high on the list both ways.

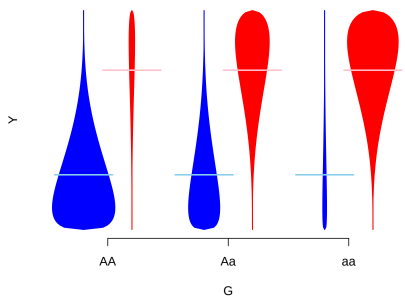
```
colnames(X)[ order(z.pmean,decreasing=TRUE)[1:10] ]
## [1] "bmi" "ltg" "g2" "map" "tc" "sex.age" "sex"
## [8] "ldl" "ltg.age" "tch"
```

```
colnames(X)[ order(b.pmean,decreasing=TRUE)[1:10] ]
## [1] "ltg" "bmi" "ldl" "map" "sex.age" "hdl" "ltg.age"
## [8] "tch" "glu.bmi" "map.sex"
```

Model-averaging in this way gives an honest statement of uncertainty. But;

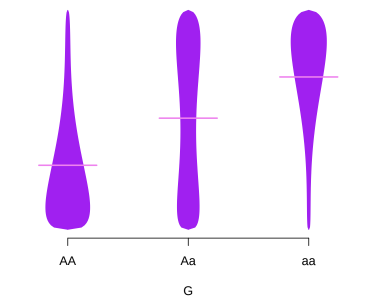
- Not all variables are in the model for the same reason may want to 'force' some covariates into the model
- When selecting a single, parsimonious model, may want to maximize its ability to predict – not its probability of being true

'Confounding' means not being able to distinguish between a signal of interest, and some other cause. Here's a genetic 'signal';



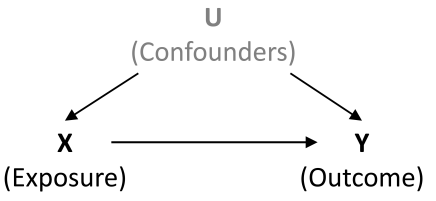
...which can be explained by ancestry, i.e. is confounded by ancestry

≻



However, analysis that adjusts for ancestry would be of interest – even if models without it are better-supported.

Directed Acyclic Graphs (DAGs) are a general language for confounding;



Arrows indicate causal relationships; confounding means 'backdoor paths' exist; these can be removed by adjustment for confounders. In genetic association work, typically ancestry is the only plausible confounder - expression and methylation work is more complex.

Bayesian Adjustment for Confounding (BAC, Wang et al 2012) specifies a model with

- 1. Dependence of outcome on the exposure and the set of confounders
- 2. Dependence of exposure on the set of confounders
- 3. Dependence between these models, making variable inclusion in (1) more likely if it is included in (2)

So BAC fits two set of z indicators, and links them. Modeling exposures is unusual – doing it well takes careful work.

The method is implemented in BEAU, a stand-alone R package, using approximate calculations for the posterior.

Prediction

Understanding causes (and confounding) is often very important – but ability to predict can matter too;

- Remaining lifetime
- Drug response
- Telling 'good' genotyping from 'bad'

To pick a model here, it's reasonable to ask how well it would predict *in similarly-collected data*. This choice may not be the same as asking what the causes are, e.g. TV ownership rates predict child mortality but are not a cause.

Cross-validation

A natural way to assess how well a fitted model predicts is to fit it, and predict!



SSR is a common measure of predictive accuracy

Cross-validation

- *SSR* (squared error loss) is not the only option need to consider the *loss* (*utility*) of particular predictions
- For categorical outcomes, could also weight misclassification rates (e.g. P(1|0) and P(0|1)) some mistakes may be worse than others
- Trickier still for dependent outcomes
- 10-fold cross-validation is typical
- Fitting multiple models with Gibbs sampling, and cross-validating each can be too slow

Approximate prediction measures

The standard 'score' is log *posterior predictive density*

$$\log p_{\text{ppost}}(y) = \log \int p(y|\theta) p(\theta|y)_{\text{obs}} d\theta).$$

Expected out-of-sample accuracy (over new datasets \tilde{y}) is defined as

$$elpd = E(\log p_{ppost}(\tilde{y})) = \int \log p_{ppost}(\tilde{y})q(\tilde{y})d\tilde{y}$$

for true density $q(\tilde{y})$. A natural way to estimate this is through the 'in sample accuracy',

$$lpd = \log \int p(y_{\text{obs}}|\theta)p(\theta|y)_{\text{obs}}d\theta,$$

but its double-use of the posterior leads to bias – worse with more parameters.

Approximate prediction measures

- Akaike's Information Criterion (AIC) approximates lpd by $\log p(y_{obs}|\hat{\theta}_{MLE})$ so is not Bayesian, and adds bias-correction k, the number of parameters
- Deviance Information Criterion (DIC) approximates lpd by $\log p(y_{obs}|E(\theta|y_{obs}))$ and adds the *effective number of parameters*,

 $p_D = 2\left(\log p(y_{\text{obs}} | \mathbb{E}[\theta | y_{\text{obs}}]) - \mathbb{E}_{\theta}[\log p(y_{\text{obs}} | \theta)]\right)$

For either, in large samples – and under some conditions – choosing the model with the lowest value is equivalent to doing cross-validation.

Note: several other versions are available; AIC, DIC2, WAIC...

DIC examples

- Shriner and Yi 2009 use DIC in the context of multiple QTL Mapping to select how many QTLs there are, and their locations
- Yu et al, 2012 use DIC studying gene×environment interactions, with a model that 'clusters' nearby* variants, so they have similar interaction effects. DIC is used to choose how many clusters
- * ...using the Potts model