Lecture 22 Model Comparison and Ensembling



Decision Theory

Predictions (or actions based on predictions) are described by a utility or loss function, whose values can be computed given the observed data.

Utility = -Loss = -Energy





Point Predictions: squared loss

Sometimes we want to make point predictions. In this case *a* is a single number.

squared error loss:
$$l(a, y^*) = (a - y^*)^2$$

The optimal point prediction that minimizes the expected loss (negative) expected utility):

$$ar{l}\left(a
ight)=\int\!dy^{*}\left(a-y^{*}
ight)^{2}p(y^{*}|D,M)$$





is the posterior predictive mean:

$$\hat{a}=E_p[y^*].$$

The expected loss then becomes:

$$ar{l}(\hat{a}) = \int dy^* \, (\hat{a} - y^*)^2 \, p(y^* | D, M) = \int dy^* \, (E_p[y^*] - y^*)^2 \, p(y^* | D, M)$$

Squared loss \implies we dont care about skewness or kurtosis

This action \hat{a} is called the **Bayes Action**.



$p(y^*|D,M) = Var_p[y^*]$

Process

First define the distribution-averaged utility:

$$ar{u}(a) = \int d\omega \, u(a,\omega) \, p(\omega|D)$$

We then find the *a* that maximizes this utility:

$$\hat{a} = rg\max_{a} ar{u}(a)$$

This action is called the **bayes action**.



The resulting maximized expected utility is given by:

$$ar{u}(\hat{a},p)=ar{u}(\hat{a})=\int d\omega\, u(\hat{a},\omega)\, p(\omega$$

sometimes referred to as the entropy function, and an associate **divergence** can be defined:

$$d(a,p)=ar{u}(p,p)-ar{u}(a,p)$$

Then one can think of minimizing d(a, p) with respect to a to get \hat{a} , so that this discrepancy can be thought of as a loss function.







Log score: probabilistic prediction

Here we want to find a distribution *a*.

The utility is defined as:

$$u(a, y^*) = \log a(y^*),$$

The expected utility then is

$$ar{u}(a) = \int dy^* \log(a(y^*)) \, p(y^*|D, I)$$





The *a* that maximizes this utility is the predictive itself (in the bayesian context, the posterior predictive)!

$$\hat{a}(y^*) = p(y^*|D,M)$$

Maximized utility:
$$ar{u}(a) = \int dy^* \log(p(y^*|D,M))$$

This is just the negative entropy of the predictive distribution, and the associated divergence is our old friend the KL-divergence.

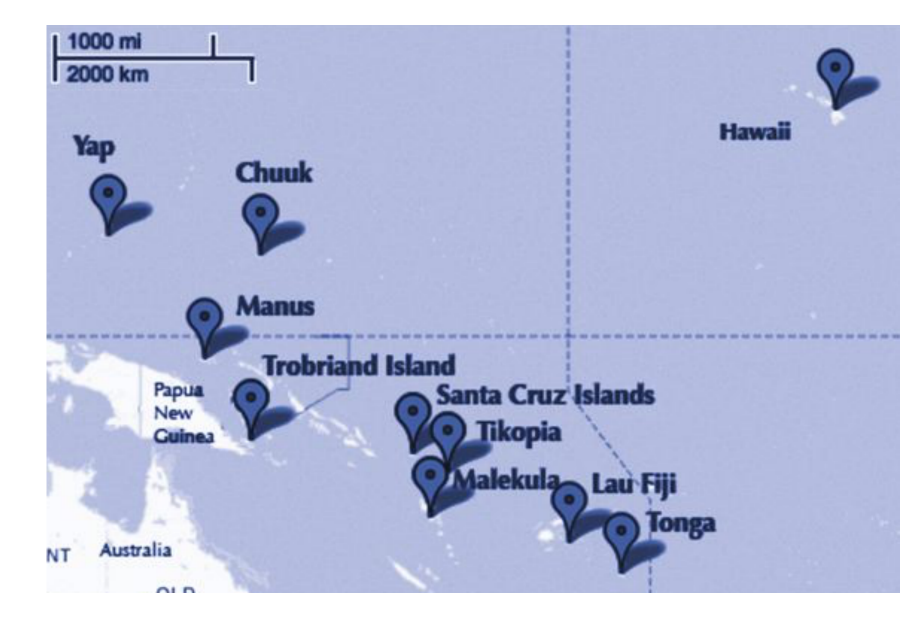


$p(y^*|D, M).$

Back to Poisson GLMs

From Mcelreath:

The island societies of Oceania provide a natural experiment in technological evolution. Different historical island populations possessed tool kits of different size. These kits include fish hooks, axes, boats, hand plows, and many other types of tools. A number of theories predict that larger populations will both develop and sustain more complex tool kits. So the natural variation in population size induced by natural variation in island size in Oceania provides a natural experiment to test these ideas. It's also suggested that contact rates among populations effectively increase population size, as it's relevant to technological evolution. So variation in contact rates among Oceanic societies is also relevant. (McElreath 313)



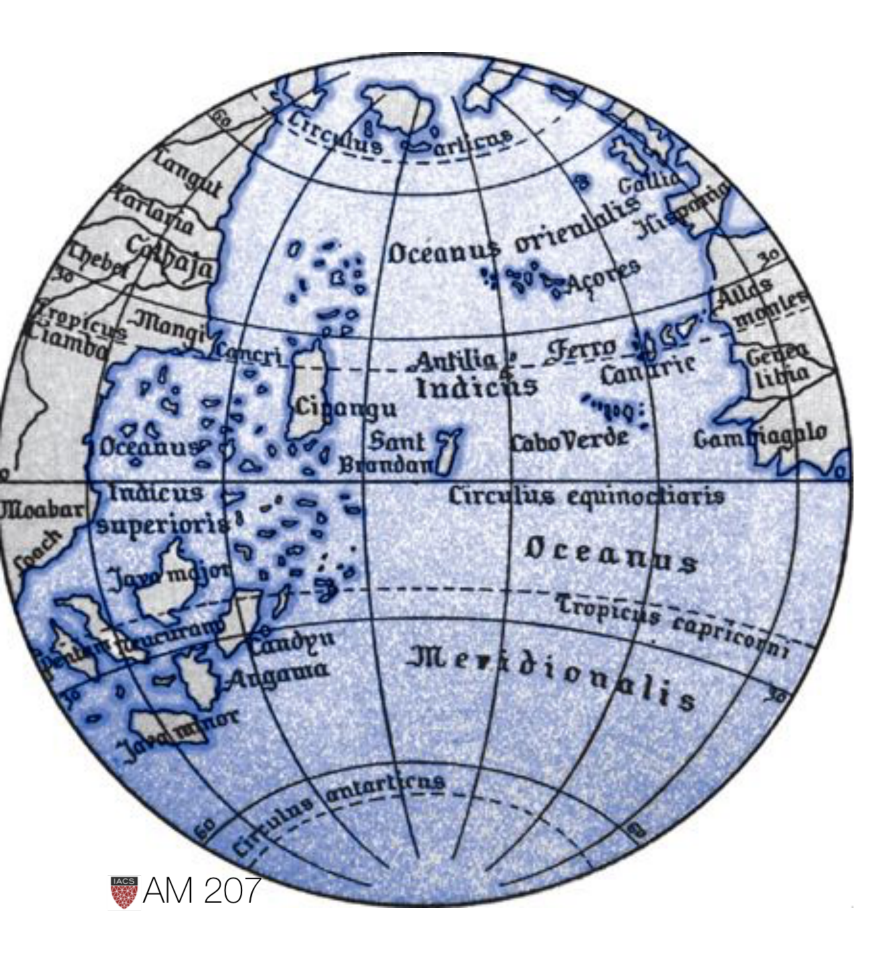


Were the contacts really needed?

Let us compare models:

m2c_onlyic: loglam = alpha m2c_onlyc: loglam = alpha + betac*df.clevel m2c_onlyp: loglam = alpha + betap*df.logpop_c m2c_nopc: loglam = alpha + betap*df.logpop_c + betac*df.clevel m1c: loglam = alpha + betap*df.logpop_c + betac*df.clevel + betapc*df.clevel*df.logpop_c





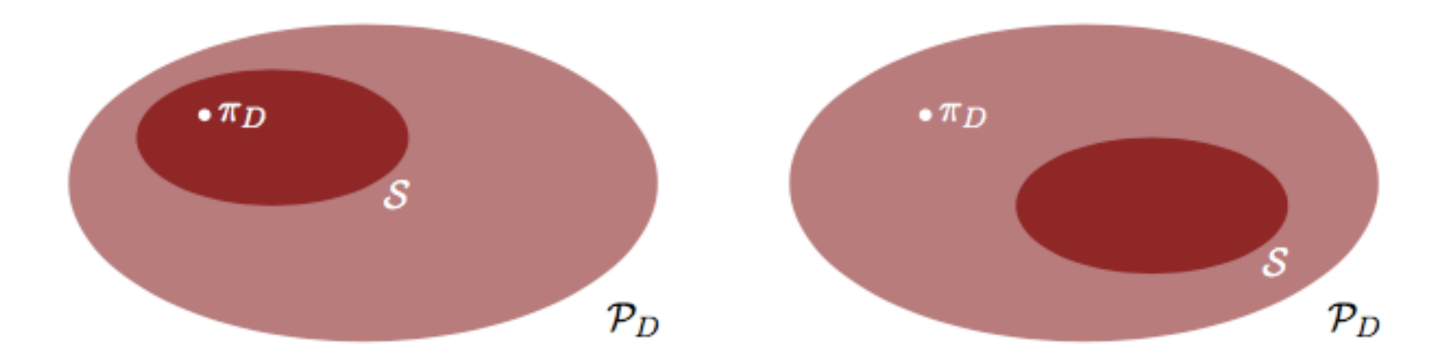
SMALL World vs BIG World

- "hyperparameters".

• *Small World* answers the question: given a model class (i.e. a Hypothesis space, whats the best model in it). It involves parameters. Its model checking.

• BIG World compares model spaces. Its model comparison with or without

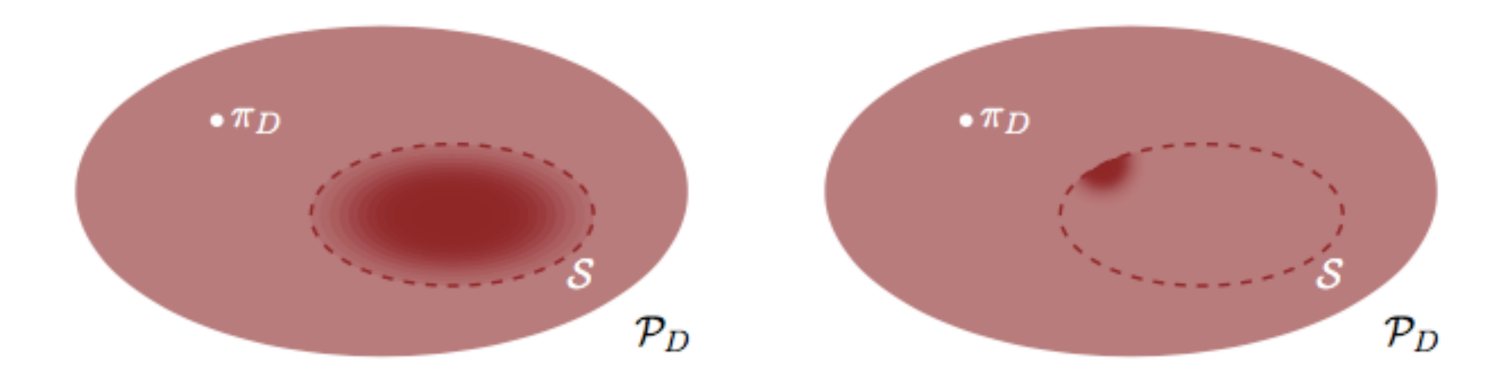
Bayesian Inference works in the small world



(some times includes the true generating process p_{tb} or p or $\pi_{\mathcal{D}}$)



Inference in the small world

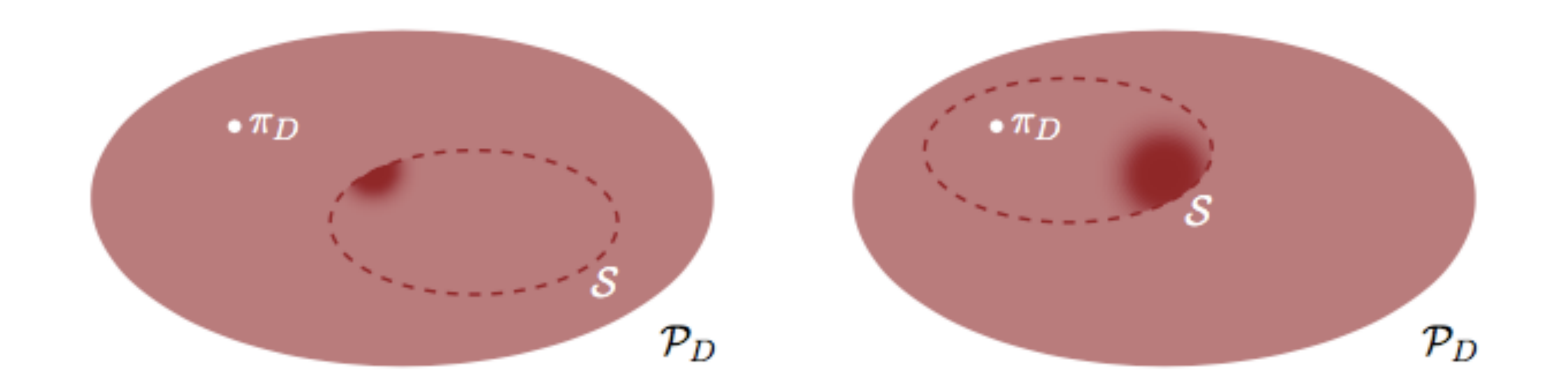


we go from prior to posterior





Bias and Variance



Overfitting can occur even if the small world includes the true data generating process p_{tb} .



How to compare against p_{tb}

In model comparison scenario we might use the "true" distribution:

$$ar{u}_t(\hat{a}) = \int dy^* u(\hat{a},y^*) p_{tb}(y^*)$$

Notice that we use $u(\hat{a}, y^*)$. The \hat{a} has already been found by optimizing over our posterior predictive.

But we dont know p_{tb} . Does this remind you of something...?



True-belief distribution

- the "p" we used in KL-divergence formulae eons ago
- that is, the known or empirically estimated true distribution, or
- model M_{tb} that has undergone posterior predictive checks and is very expressive, a model we can use as a reference model.
- often non-parametric or found via bayesian model averaging.
- if the true generating process is outside the hypothesis set of the models you are using, true belief model never = true. This is called misfit or bias.



Model comparison

The key idea in model comparison is that we will sort our average utilities in some order. The exact values are not important, and may be computed with respect to some true distribution or true-belief distribution M_{tb} .

Utility is maximized with respect to some model $M_k \in \mathcal{H}$ whereas the average of the utility is computed with respect to either the true belief distribution.



$$ar{u}(M_k, \hat{a}_k) = \int dy^* u(\hat{a}_k, y^*) p(y^*|D,$$

where a_k is the optimal prediction under the model M_k . Now we compare the actions, that is, we want:

$$\hat{M} = rg\max_k ar{u}(M_k, \hat{a}_k)$$

No calibration, but calculating the standard error of the difference can be used to see if the difference is significant.



M_{tb})

The MSE or R^2 in regression problems...

For the squared loss the first step gives us $\hat{a}_k = E_{p(y^*|D,M_k)}[y^*]$. Then:

$$ar{l}\left(\hat{a_k}
ight) = \int dy^* \, (\hat{a}_k - y^*)^2 \, p(y^*|D, M) \ = \int dy^* \, (E_{p_k}[y^*] - y^*)^2 \, p(y^*|D, M_{tb}) = Var_{p_{tb}}[y^*] + 0$$

We have bias if M_{tb} is not in our Hypothesis set \mathcal{H} .



 $M_{tb})$

 $(E_{p_{th}}[y^*] - E_{p_k}[y^*])^2$

So far...

- Decisions are made on some validation or test set
- ensures that we dont use data twice
- while we are not picking hyper-parameters in the bayesian scenario
- we are still maximizing utility/minimizing risk
- and furthermore choosing a "best" model according to this utility



Information criteria

- we dont want to go out-of-sample
- use information criteria to decide between models

$$ext{KL:} \ D_{KL}(p,q) = E_p[log(p/q)] = \sum_i p_i log(rac{p_i}{q_i}) \ or$$

Use **law or large numbers** to replace the true distribution by empirical estimate



 $r \int dPlog(\frac{p}{a})$

Deviance

$$D_{KL}(p,q) = E_p[log(p/q)] = rac{1}{N}\sum_i (log(p_i))$$

Deviance
$$D(q) = -\frac{N}{2}E_p[log(q)] = -2\sum_i log(q_i)$$

then
$$D_{KL}(p,q)-D_{KL}(p,r)=rac{2}{N}(D(q)-D(r))$$



$_i) - log(q_i))$

),

Deviance of a predictive

$$D(q) = -rac{N}{2} E_p[log(q)]$$

We want to estimate the "true-belief" average of a predictive:

$$E_p[log(pred(y^*))]$$

where $pred(y^*)$ is the predictive for points y^* on the test set or future data.



Do it pointwise instead

Call the expected log predictive density at a "new" point:

$$elpd_i = E_p[log(pred(y_i^*)]$$

Then the "expected log pointwise predictive density" is

$$elppd = \sum_i E_p[log(pred(y_i^*))] = \sum_i$$



 $elpd_i$

What predictive distribution *pred* do we use? We start from the frequentist scenario of using the likelihood at the MLE for the AIC, then move to using the likelihood at the posterior mean (a sort of plug in approximation) for the DIC, and finally to the fully Bayesian WAIC.

Specifically, in the first two cases, we are writing the predictive distribution conditioned on a point estimate from the posterior:

$$elpd_i = E_p[log(pred(y^*_i \mid \hat{ heta}))]$$



The game we will play in these first two cases is:

(1) Conditional on fixed θ , the full predictive splits into a product per point so the writing of elppd as a sum over pointwise elpd is exact

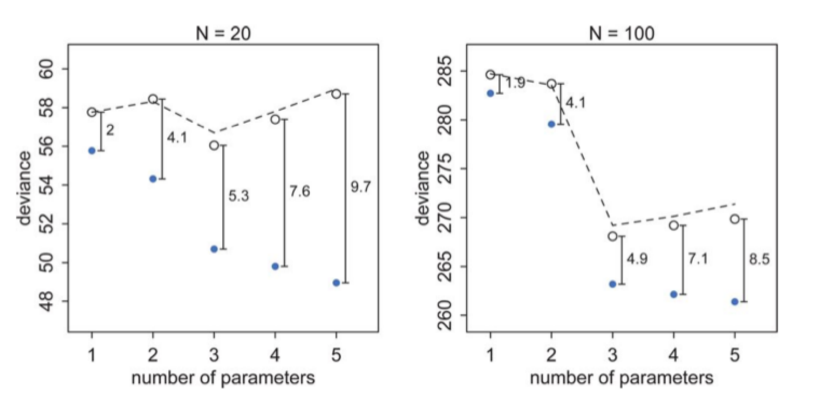
- (2) However we dont know p_{tb} (or just p), so we use the empirical distribution on the training set
- (3) this underestimates the test set deviance as we learnt in the case of the AIC, so we must apply a correction factor.



Akaike Information Criterion, or AIC:

$$D_{train} = -$$

- multivariate gaussian posterior
- flat priors
- data >> parameters





AIC

$= D_{train} + 2p$

$-2*log(p(y| heta_{mle})$

DIC

Uses the posterior distribution, calculable from MCMC, and assumes multivariate gaussian posterior distribution.

$$D_{train} = -2 * log(p(y| heta_{postmean}), DIC = D_{train})$$

$$p_{DIC} = 2*(log(p(y| heta_{postmean}) - E_{post}[log(p(y| heta)])$$

alternative fomulation for p_D , guaranteed to be positive, is

$$p_{DIC} = 2 * Var_{post} [log(p(y| heta_{postmean}))]$$



$+2p_{DIC}$ where

(by monte carlo)

 $_{n}))]$

Bayesian deviance

 $D(q) = -\frac{N}{2}E_p[log(pp(y))]$ posterior predictive for points y^* on the test set or future data

replace joint pp over new points y by product of marginals:

$$elpd_i = E_p[log(pp(y_i^*)]]$$

$$elppd = \sum_i E_p[log(pp(y^*_i))] = \sum_i el$$



 lpd_i

Game is to REPLACE

$$elppd = \sum_i E_p[log(pp(y_i^*))]$$
 where y_i^* are

by the computed "log pointwise predictive density" (lppd) insample

$$lppd = log \left(\prod_{j} pp(y_{j})
ight) = \sum_{j} log \langle p(y_{j}| heta)
angle_{post} = \sum_{j} log$$



new points

 $\log\left(rac{1}{S}\sum_{s\sim nost}p(y_j| heta_s)
ight)$

- As we know now, is that the *lppd* of observed data y is an overestimate of the *elppd* for future data.
- Hence the plan is to like to start with the *llpd* and then apply some sort of bias correction to get a reasonable estimate of elppd.
- This gives us the WAIC (Widely Applicable Information Criterion or Watanable-Akaike Information Criterion)



WAIC

$WAIC = lppd + 2p_W$

where

$$p_W = 2\sum_i (log(E_{post}[p(y_i| heta)] - E_{post}[log(x_i| heta)])$$

Once again this can be estimated by $\sum_{i} Var_{post}[log(p(y_i|\theta))]$





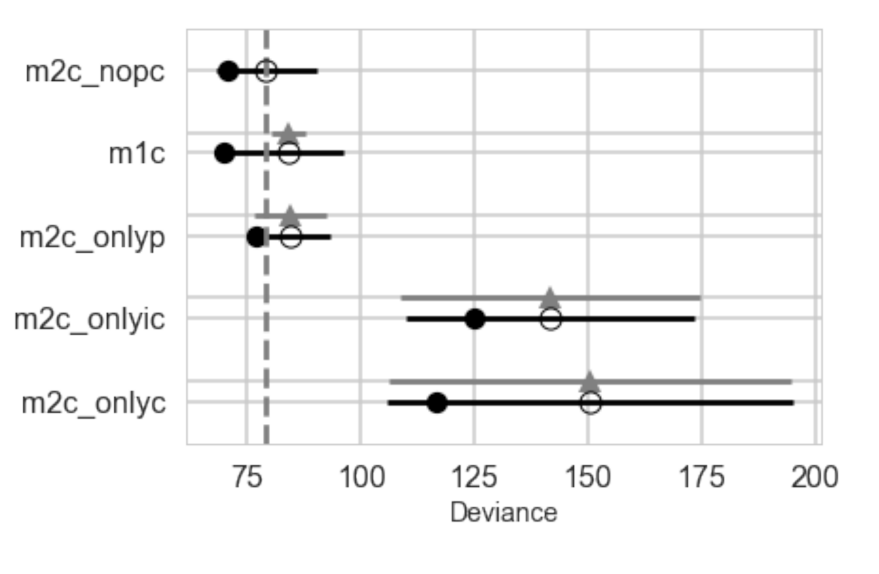
Oceanic tools

Lets use the WAIC to compare models

m2c_onlyic: loglam = alpha m2c_onlyc: loglam = alpha + betac*df.clevel m2c_onlyp: loglam = alpha + betap*df.logpop_c m2c_nopc: loglam = alpha + betap*df.logpop_c + betac*df.clevel m1c: loglam = alpha + betap*df.logpop_c + betac*df.clevel + betapc*df.clevel*df.logpop_c



Centered



- and the lowest WAIC.
- ranked model.

$$w_i = rac{exp(-rac{1}{2}dWAIC_i)}{\sum_j exp(-rac{1}{2}dWAIC_j)}$$

read each weight as an estimated probability that each model will perform best on future data.



dWAIC is the difference between each WAIC

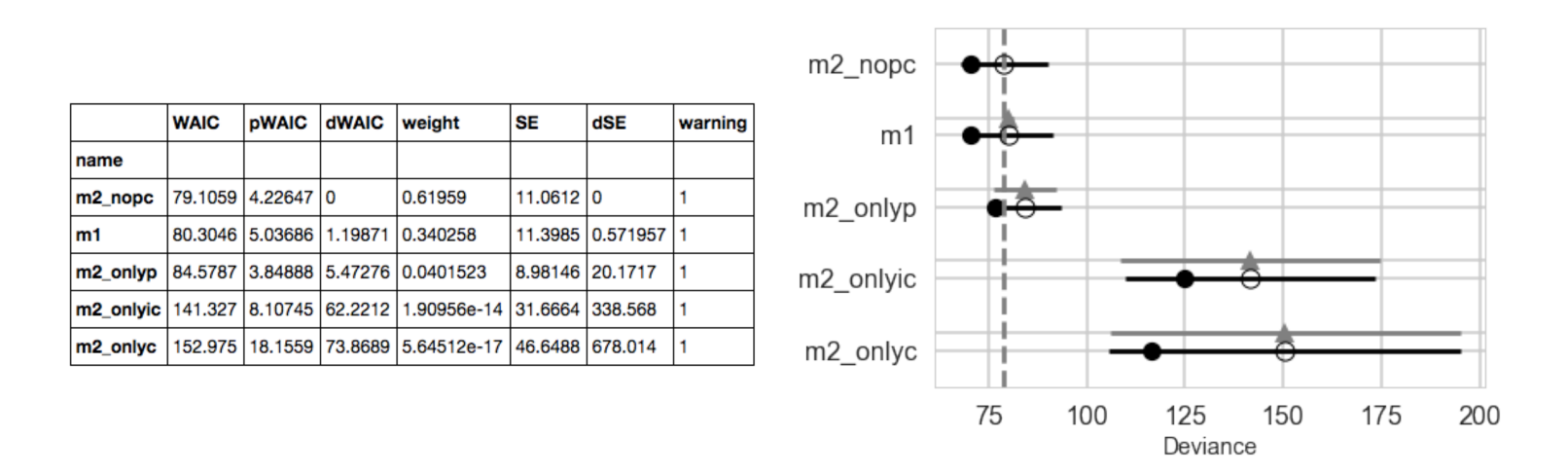
• SE is the standard error of the WAIC estimate.

• dSE is the standard error of the difference in WAIC between each model and the topFrom McElreath, here is how to read this table:

- 1. WAIC is obviously WAIC for each model. Smaller WAIC indicates better estimated out-of-sample deviance.
- 2. pWAIC is the estimated effective number of parameters. This provides a clue as to how flexible each model is in fitting the sample.
- 3. dWAIC is the difference between each WAIC and the lowest WAIC. Since only relative deviance matters, this column shows the differences in relative fashion.
- 4. weight is the AKAIKE WEIGHT for each model. These values are transformed information criterion values. I'll explain them below.
- 5. SE is the standard error of the WAIC estimate. WAIC is an estimate, and provided the sample size N is large enough, its uncertainty will be well approximated by its standard error. So this SE value isn't necessarily very precise, but it does provide a check against overconfidence in differences between WAIC values.
- 6. dSE is the standard error of the difference in WAIC between each model and the top-ranked model. So it is missing for the top model.



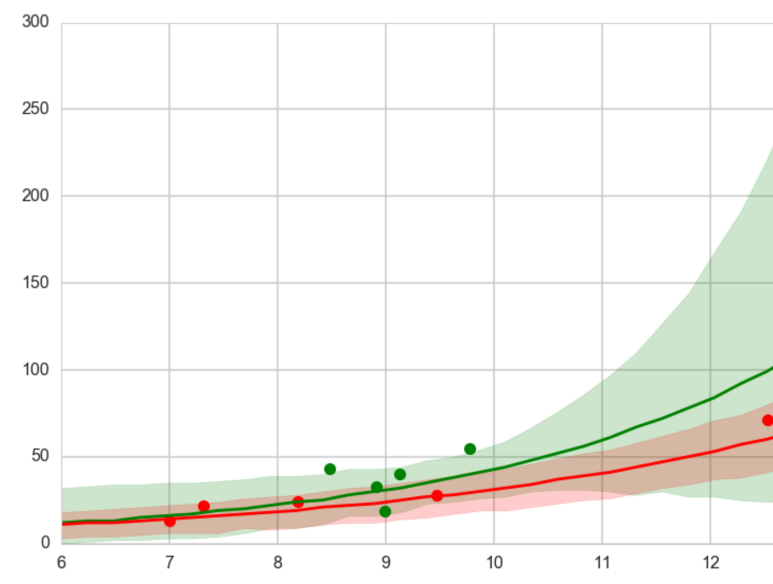
Uncentered



interaction is overfit. centering decorrelates

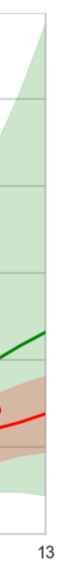


Counterfactual Posterior predictive









Bayes Theorem in model space

 $p(M_k|D) \propto p(D|M_k)p(M_k)$

where:

$$p(D|M_k) = \int d heta_k \, p(y| heta_k, M_k) p(heta_k)$$

is the marginal likelihood under each model. Can use these "Bayes" Factors" to compare but high sensitivity to prior.



$|M_k)$

Bayesian Model Averaging

$$p_{BMA}(y^*|x^*,D) = \sum_k p(y^*|x^*,D,M_k)p_k$$

where the averaging is with respect to weights $w_k = p(M_k|D)$, the posterior probabilities of the models M_k .

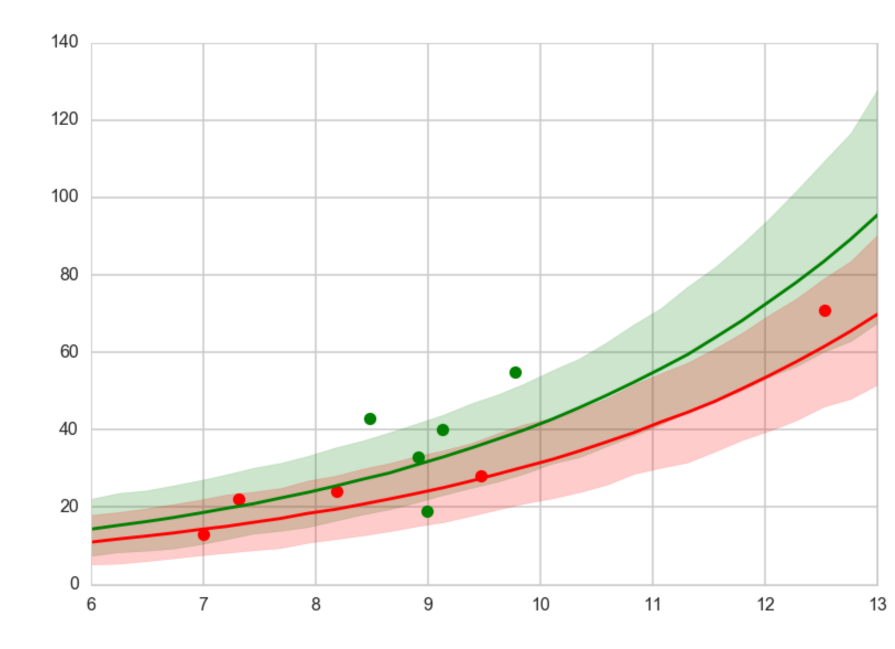
We will use the "Akaike" weights from the WAIC. This is called pseudo-BMA



 $p(M_k|D)$

Ensembling

- use WAIC based akaike weights for top
 3
- regularizes down the green band at high population by giving more weight to the no-interaction model.

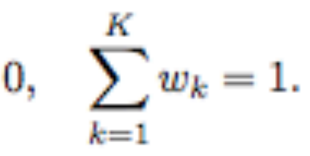




- BMA is appropriate in the M-closed case, which is when the true generating process is one of the models
- what we will use here is to estimate weights by the WAIC, following McElreath (pseudo-BMA)
- But see Yao et. al. which claims log-score stacking is better. Implemented in pymc3

$$\max_{w} \frac{1}{n} \sum_{i=1}^{n} \log \sum_{k=1}^{K} w_k p(y_i | y_{-i}, M_k), \quad s.t. \quad w_k \ge 0$$





Pseudo BMA vs stacking

		WAIC	pWAIC	dWAIC	weight	SE	dSE	warning		WAIC	pWAIC	dWAIC	weight	SE	dSE	warning
nar	ne								name							
m	2c_nopc	79.06	4.24	0	0.87	11.06	0	1	m2c_nopc	79.06	4.24	0	0.76	11.06	0	1
	m1c	84.09	7.05	5.04	0.07	12.19	3.77	1	m1c	84.09	7.05	5.04	0	12.19	3.77	1
m2	2c_onlyp	84.43	3.75	5.37	0.06	8.94	7.93	1	m2c_onlyp	84.43	3.75	5.37	0.24	8.94	7.93	1
m2	c_onlyic	141.65	8.38	62.6	0	31.7	32.84	1	m2c_onlyic	141.65	8.38	62.6	0	31.7	32.84	1
m	2c_onlyc	150.44	16.94	71.38	0	44.67	44.44	1	m2c_onlyc	150.44	16.94	71.38	0	44.67	44.44	1



... it is tempting to use information criteria to compare models with different likelihood functions. Is a Gaussian or binomial better? Can't we just let WAIC sort it out? Unfortunately, WAIC (or any other information) criterion) cannot sort it out. The problem is that deviance is part normalizing constant. The constant affects the absolute magnitude of the deviance, but it doesn't affect fit to data.

– McElreath



How to handle non-nested models?

- cross-validation
- less data to fit so biased models
- we are not talking here about cross-validation to do hyperparameter optimization
- specifically we will use Leave-One-Out-Cross-Validation (LOOCV) with importance sampling



- The idea here is that you fit a model on N-1 data points, and use the Nth point as a validation point. Clearly this can be done in N ways.
- the N-point and N-1 point posteriors are likely to be quite similar, and one can sample one from the other by using importance sampling.

$$E_f[h] = rac{\sum_s w_s h_s}{\sum_s w_s}$$
 where $w_s = f_s$ /



 $/g_s$.

Fit the full posterior once. Then we have

$$w_s = rac{p(heta_s|y_{-i})}{p(heta_s|y)} \propto rac{1}{p(y_i| heta_s,y_{-i})}$$

- the importance sampling weights can be unstable out in the tails.
- importance weights have a long right tail, pymc (pm.loo) fits a generalized pareto to the tail (largest 20% importance ratios) for each held out data point i (a MLE fit). This smooths out any large variations.



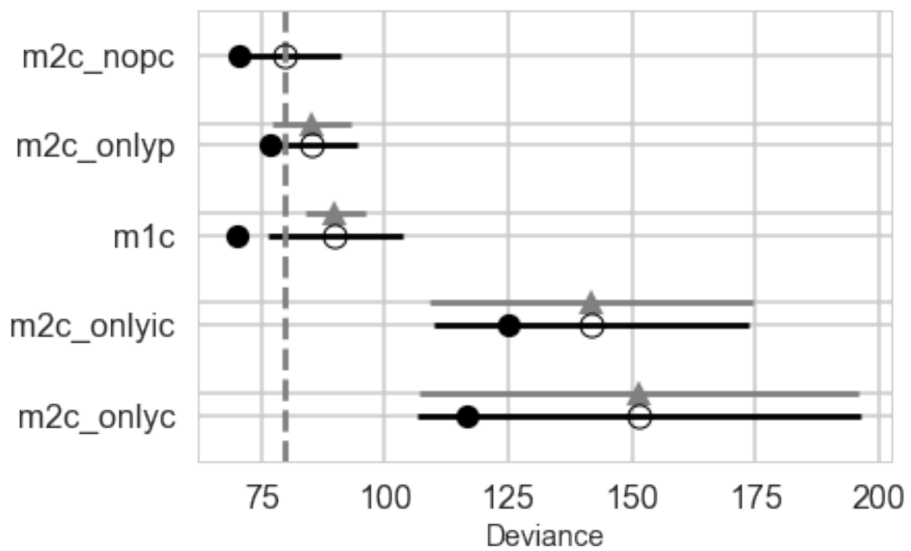
$$elpd_{loo} = \sum_i log(p(y_i|y_{-i}))$$

$$=\sum_i log\left(rac{\sum_s w_{is} p(y_i| heta_s)}{\sum_s w_{is}}
ight)$$

over the training sample.



Oceanic tools LOOCV





What should you use?

- 1. LOOCV and WAIC are fine. The former can be used for models not having the same likelihood, the latter can be used with models having the same likelihood.
- 2. WAIC is fast and computationally less intensive, so for same-likelihood models (especially nested models where you are really performing feature selection), it is the first line of attack
- 3. One does not always have to do model selection. Sometimes just do posterior predictive checks to see how the predictions are, and you might deem it fine.
- 4. For hierarchical models, WAIC is best for predictive performance within an existing cluster or group. Cross validation is best for new observations from new groups

