Week 4: Multilevel Models

PLSC 40502 - Statistical Models

Review

Previously

Bayesian inference

- Parameters of interest are random variables (randomness captures "beliefs")
- Contrast with frequentist approach: parameter is fixed, data is random
- \circ Bayesian target of inference is the **posterior** distribution $p(\theta|\mathbf{Y})$
- **Bernstein-von Mises theorem** in large samples, bayesian posterior distributions approximate MLE sampling distributions.

Posterior inference

- Some posteriors have an analytically tractable form -- typically we get this through the use of conjugate priors
- Most posterior distributions are not analytically tractable -- need to do inference numerically via sampling
- Markov Chain-Monte Carlo Construct a chain of samples that converge to a stationary distribution which happens to be the target posterior.
- o Metropolis-Hastings An algorithm to generate a chain that approximates a target posterior
- Gibbs Sampling Special case of MH where samples are from alternating conditional distributions

This week

MCMC via Stan

- Stan is a C++ library for specifying statistical models and conducting inference via Hamiltonian
 MCMC
- R has bindings to Stan via the rstan package
- Extremely flexible any model that you can write can be translated to machine code an estimated using an algorithm that converges much more quickly than conventional MH

Model diagnostics

- How do we know if our model is good?
- Posterior predictive checks
- Cross-validation, Leave-one-out CV

Multilevel models

- What happens when we impose further structure on parameters (e.g. random slopes and intercepts)
- Choices of hyperpriors governs the extent of partial pooling

Gibbs Sampling Continued

- When we discussed **Gibbs Sampling** we illustrated the significant gains we can obtain over naive Metropolis-Hastings
 - Faster convergence, less correlation between samples
- These gains come from **theory** being able to derive the form of the conditional distribution of θ_k given the other parameters θ_{-k} as well as the data **Y**.
- Deriving these conditionals often relies on inspecting a density that is proportional to the conditional density and recognizing its form.
 - To do this, it's useful to remember the connection between **conditional** densities and **joint** densities as well as how **joint** densities factor.

- Consider our **normal regression** example, we want to obtain samples from the joint density $f(\beta, \sigma^2 | \mathbf{Y})$
 - \circ To do so, we need the conditionals $f(\beta|\mathbf{Y},\sigma^2)$ and $f(\sigma^2|\mathbf{Y},\beta)$.
 - Can we derive them (or at least their distributional forms?)
- Remember the definition of a conditional expectation: joint divided by marginal

$$f(eta|\mathbf{Y},\sigma^2) = rac{f(eta,\sigma^2|\mathbf{Y})}{f(\sigma^2|\mathbf{Y})}$$

• Since we treat σ^2 here as a constant, we can write

$$f(eta|\mathbf{Y},\sigma^2) \propto f(eta,\sigma^2|\mathbf{Y}) \equiv f(\mathbf{Y}|eta,\sigma^2)f(eta,\sigma^2)$$

where the last part follows from the definition of the posterior

Next, remember how distributions factor

$$f(eta,\sigma^2)=f(eta|\sigma^2)f(\sigma^2)$$

- If β and σ^2 are marginally independent, then the joint distribution factors into the **marginals** \circ But even if they're not independent, we can always factor into a product of conditionals
- So we have

$$f(eta|\mathbf{Y},\sigma^2) \propto f(\mathbf{Y}|eta,\sigma^2) f(eta|\sigma^2) f(\sigma^2)$$

• Again, σ^2 is a constant here, so that last density drops as a multiplicative constant

$$f(eta|\mathbf{Y},\sigma^2) \propto f(\mathbf{Y}|eta,\sigma^2) f(eta|\sigma^2)$$

• Remember, our special case of the normal model from last week assumed that β and σ^2 are marginally independent

$$f(eta|\mathbf{Y},\sigma^2) \propto f(\mathbf{Y}|eta,\sigma^2)f(eta)$$

And independence of observations gives

$$f(eta|\mathbf{Y},\sigma^2) \propto igg[\prod_{i=1}^N f(Y_i|eta,\sigma^2)igg]f(eta)$$

• The key is finding densities that are given by our model description - now all of these factors are known

$$f(eta|\mathbf{Y},\sigma^2) \propto igg[\prod_{i=1}^N \underbrace{f(Y_i|eta,\sigma^2)}_{ ext{Normal}(X_i'eta,\sigma^2)}igg] imes \underbrace{f(eta)}_{ ext{Normal}(b_0,B_0^{-1})}$$

- One useful property from probability theory is that products of two normal PDFs are proportional to the normal PDF
 - So $\beta | \mathbf{Y}, \sigma^2$ is normal!

- From normal regression theory, we can derive the mean and variance of that posterior distribution
- Variance-Covariance matrix

$$V = \sigma^2 (B_0 + \mathbf{X}'\mathbf{X})^{-1}$$

And Mean

$$\mu = (B_0 + \mathbf{X}'\mathbf{X})^{-1}(B_0b_0 + \mathbf{X}'\mathbf{Y})$$

• Note again how the posterior distribution combines the data and the prior with the influence of the prior diminishing as the number of observations grows.

Revisiting: Predicting Elections

```
# Load the data
elections <- read csv("data/us-house-wide.csv")</pre>
# Aggregate the house data to counties
elections county <- elections %>% group by(fipscode) %>% summarize(state=state[1], county=count
                                                                     total.votes = sum(total.vote)
                                                                     dem = sum(dem))
# Merge in 2015 Presidential
pres 2016 <- read csv("data/clinton 2016 vote.csv")</pre>
elections county <- elections county %>% left join(pres 2016 %>% dplyr::select(county fips, car
                                                    by=c(fipscode="county fips"))
# Generate vote shares
elections county$dem2018 <- elections county$dem/elections county$total.votes
elections county$dem2016 <- elections county$candidatevotes/elections county$totalvotes
# Drop missing
elections county <- elections county %>% filter(!is.na(dem2018)&!is.na(dem2016))
```

Gibbs and Metropolis

• Set up the regression

```
X_mat <- model.matrix(dem2018 ~ dem2016, data=elections_county)
Y <- elections_county$dem2018
K <- ncol(X_mat) # Number of beta parameters</pre>
```

Set up a diffuse prior

```
beta_0 <- rep(0, K)
B_0 <- diag(rep(1/9, K))
B_inv_0 <- solve(B_0)
c_0 = 0.001
d_0 = 0.001</pre>
```

• Set up the MCMC

```
M <- 40000 # Number of MCMC samples
burnin <- 5000
beta_mcmc <- matrix(nrow = M, ncol=K) # Vector to store our samples
beta_mcmc[1,] <- c(0,1) # Pick a starting value
sigma_mcmc <- rep(NA, M)
sigma_mcmc[1] <- 1</pre>
```

Gibbs and Metropolis

• Write some functions to evaluate the likelihood and priors

```
log_lik_norm <- function(b, sigma, Y, X){
  linpred <- X%*%b
  sum(dnorm(Y, mean=linpred, sd=sigma, log=T))
}</pre>
```

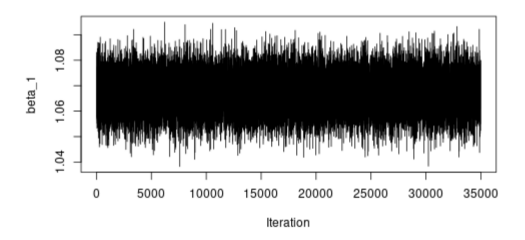
Gibbs and Metropolis

```
set.seed(60637)
for (i in 1:(M-1)){ # For i in 1:(M-1)
 ## Beta
   ## Gibbs Step!
    var b <- solve(t(X mat)%*%X mat + B 0)</pre>
    mean b <- var b%*%(B 0%*%beta 0 + t(X mat)%*%Y)
    beta mcmc[i+1,] <- mvtnorm::rmvnorm(1, mean = mean b, sigma=(sigma mcmc[i]^2)*var b)
 ## Sigma
    ## Step 1 - Proposal
    sigma log <- rnorm(1, mean = log(sigma mcmc[i]), sd=.01)</pre>
    sigma star <- exp(sigma log)</pre>
    ## Step 2 - Accept/Reject
    lik star sigma <- log lik norm(beta mcmc[i+1,], sigma star, Y, X mat)</pre>
    lik current sigma <- log lik norm(beta mcmc[i+1,], sigma mcmc[i], Y, X mat)</pre>
    # The trick here is independence - otherwise if \sigma^2 appeared in the prior for \beta, v
    prior star sigma <- log(MCMCpack::dinvgamma(sigma star^2, shape = c 0/2, scale = d 0/2))
    prior current sigma \leftarrow log(MCMCpack::dinvgamma(sigma mcmc[i]^2, shape = c ^0/2, scale = d
    ## Accept/reject
    ar_sigma <- exp(lik_star_sigma + prior_star_sigma - lik_current sigma - prior current sigma
    accept sigma <- rbinom(1,1,min(1,ar sigma))</pre>
    sigma \ mcmc[i+1] <- sigma \ star*accept \ sigma + sigma \ mcmc[i]*(1-accept \ sigma)
beta mcmc use <- beta mcmc[burnin:M,] # Toss our burn-in period</pre>
```

Convergence

• β_1

```
plot(y=beta_mcmc_use[,2], x=1:length(beta_mcmc_use[,2]), xlab="Iteration", ylab="beta_1", type=
```

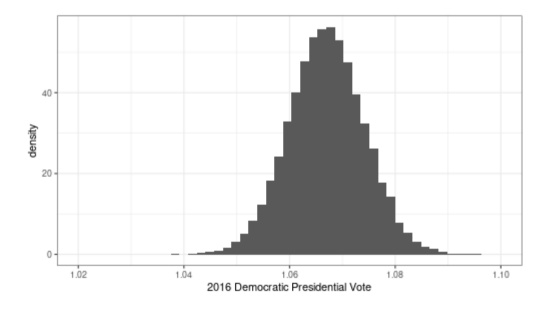


• Much better than before with the naive "random walk" proposal distribution.

Results

 eta_1

```
beta_mcmc_out <- as.data.frame(beta_mcmc_use)
colnames(beta_mcmc_out) <- c("Intercept", "dem2016")
beta_mcmc_out %>% ggplot(aes(x=dem2016)) + theme_bw() + xlim(1.02, 1.10) + xlab("2016 Democrat")
```



Summaries

dem2016

• Posterior means and 95% credible intervals

dem2016 1.067

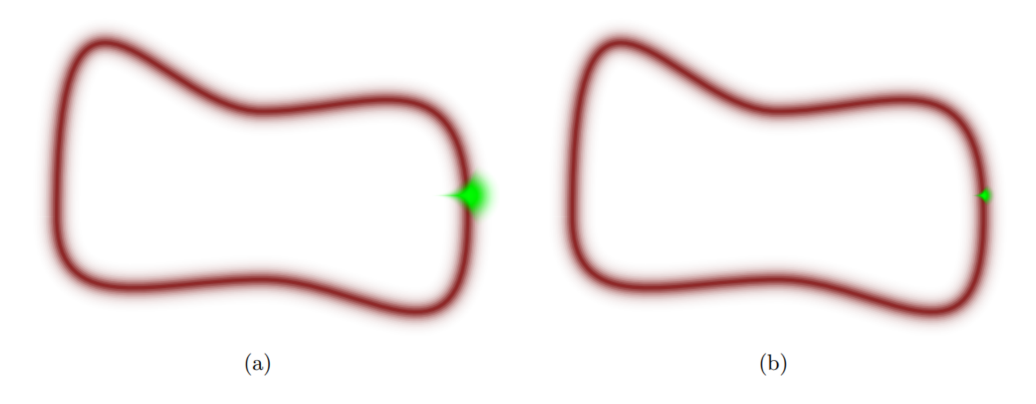
1.0810

1.0530

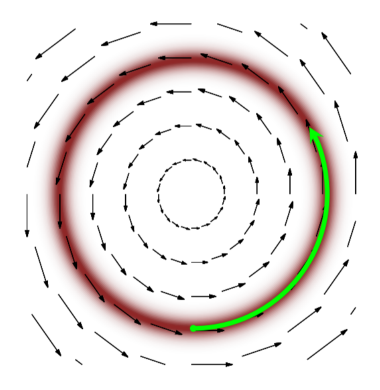
Intro to Stan/HMC

- Hamiltonian Monte Carlo is a method that improves upon the conventional Metropolis-Hastings algorithm by improving the **proposals** such that they are accepted with very high probability
 - The specific method implemented in Stan is the "No U-Turn Sampler" (Hoffman and Gelman, 2011)
- The method draws on a theoretical core from differential geometry, but it's not strictly necessary to get the intuition for it.
 - Betancourt (2014) "A Conceptual Introduction to Hamiltonian Monte Carlo" is a good overview for applied researchers (familiarity with statistics but not physics)
- MCMC (ideally) works by 3 general phases
 - 1. From the starting values, reach the "typical set" of parameter values that occupy most of the posterior density mass
 - 2. The MCMC traverses the "typical set" of values once.
 - 3. Repeated traversals improve the quality of the MCMC approximation.
- Problems arise when Markov Chains
 - 1. Don't traverse high density parts of the posterior
 - 2. Get *stuck* in parts of the density.
- How do we get proposals that are likely to get accepted?

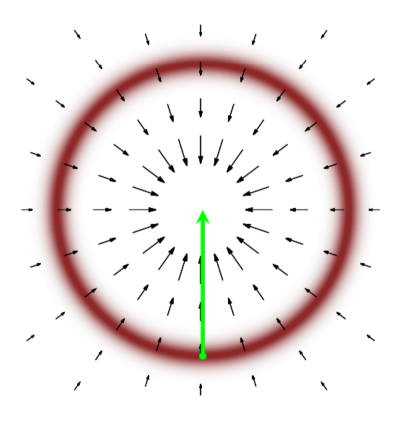
- ullet Typical distributions for Q are random walks
 - In high dimensions, these tend to be **bad** proposals.
- One solution is to choose a low variance for the random walk
 - This leads to high autocorrelation in the proposals



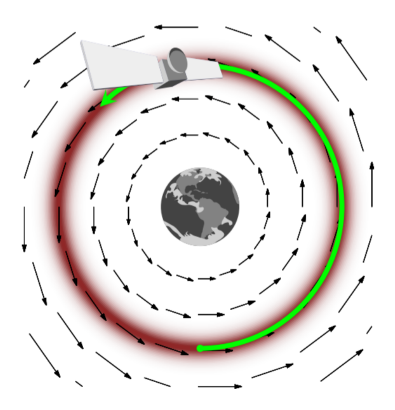
• What if we could come up with a "direction" to point our proposal (or sequence of proposals) that stayed in the "typical set"



• The gradient of the posterior gets us close, but we need to augment it to get the desired field.



- ullet Analogizing to a physical system, HMC introduces a "momentum" variable p for each parameter q
 - \circ H(p,q) is the "Hamiltonian" for each point -- a sum of "kinetic" and a "potential" energies -- the potential energy is the log density.
 - Using Hamilton's equations of motion, we obtain a method for generating updates that "conserve" energy



Central intuitions

- Hamiltonian Monte Carlo is designed to generate proposals with high acceptance probability
- These proposals require knowing the **gradient** of the unnormalized posterior with respect to the parameters
- Analytical computation of the gradient is tedious
 - Numeric computation can be unstable.
 - Many modern techniques use automatic differentiation
- **Intuition**: Decompose the unnormalized posterior into a series of elementary arithmetic operationns and functions where derivatives are trivially known
 - Use the chain rule to construct the gradient from the component functions + their derivatives.

Overview of Stan

- Stan is a C++ library that implements a version of Hamiltonian Monte Carlo
 - Latest iteration of the "model-to-MCMC" libraries like JAGS and BUGS (which are gibbs samplers)
- Key Features
 - 1. Language agnostic has bindings in R, Python, Julia, STATA, etc...
 - 2. Fast run-time
 - 3. High-level language for specifying models to be converted to C++ code.
- Complications
 - Running Stan requires you to have the ability to compile C++ code locally
 - Stan also requires learning a slightly new syntax

Stan resources

- Installation Quick Start Guide
- Stan User Guide
- Stan Reference Manual
- Stan Functions Reference
- A good model building tutorial for social scientists by Jim Savage

Installing Rstan

- We're working with R, so we'll be using the Rstan interface to Stan
- To Install
 - Install the latest development version (2.34) directly from mc-stan.org
 - Currently, the version on CRAN uses slightly dated syntax

```
# run the next line if you already have rstan installed
# remove.packages(c("StanHeaders", "rstan"))
install.packages("StanHeaders", repos = c("https://mc-stan.org/r-packages/", getOption("repos"))
install.packages("rstan", repos = c("https://mc-stan.org/r-packages/", getOption("repos")))
```

Installing Rstan

- In order to run Stan, your computer needs to be able to compile C++ code. This requires some auxiliary installs that are dependent on your operating system
 - Windows: On Windows, you need to install Rtools. See instructions: https://github.com/stan-dev/rstan/wiki/Configuring-C---Toolchain-for-Windows
 - Mac: Mac has a nice installer package for its C++ toolchain https://github.com/rmacoslib/r-macos-rtools#how-do-i-use-the-installer
 - Linux: You probably already have it, but you need g++ or clang++ either tends to be installed by default or is available from your repositories.

Writing a Stan program

- Stan programs define a particular statistical model through a sequence of blocks that identify the data (modeled and unmodeled), parameters, and statistical model.
 - These are then translated to C++ and compiled. Because Stan uses C++, variables need to be declared differently from a language like R.
 - C++ is **statically typed** which means that the *type* of variable needs to be fixed at initialization.
- Primary variable types and their syntax
 - int: Integer and real: continuous values
 - vector and matrix for real-valued column vectors and matrices
 - o **array** allows you to generate an array structure for any arbitrary type (like an integer) (note pre-2.26 Stan used a different syntax for this)
- If variables are constrained, these constraints can be added via <upper = .> and <lower = .> statements during variable declaration.

Minor notes

- Stan programs in R can be saved as separate files (with a .stan suffix) or in a length 1 character vector (directly typed into R).
- Lines end with semi-colons;
- // denotes a comment

- data: Defines the modeled (e.g. Y) and unmodeled data (e.g. X) that are used in the model.
 - \circ Includes any important constants (like sample size N) or number of parameters K that you don't hard-code.
 - When compiling and estimating the model, you'll pass in R objects for each entry defined in data through a list
- Example data block (for a logistic regression):

```
data{
  int N; // number of observations
  int K; // number of covariates
  matrix[N, K] X; //matrix of covariates
  array[N] int<lower=0, upper=1> y; // binary outcome
}
```

- parameters: Defines the parameters of interest whose posterior density we're trying to sample from
- Example parameters block (for a logistic regression):

```
parameters{
  vector[K] beta; // beta coefficients
}
```

- model: Defines the statistical distributions of the observed data + the parameters
 - Likelihood and prior distributions
- Example model block (for a logistic regression):

```
model{
  beta ~ normal(0, 10); // normal prior
  y ~ bernoulli_logit(X * beta); //bernoulli outcome (logit link)
}
```

• We can use for loops as well (can be useful for more complicated DGPs)

```
model{
    for (k in 1:K){
        beta[k] ~ normal(0, 10)
    }
    for (n in 1:N){
        y[n] ~ bernoulli_logit(X[n] * beta)
    }
}
```

- Other useful (but not required) blocks
 - **functions**: Contains user-defined functions that can be used within the remainder of the model definition (or made available to R)
 - transformed data, transformed parameters: Contains deterministic transformations of either the data or the parameters.
 - **generated quantities**: Contains functions that generate or simulate quantities from the model after estimation (e.g. predictions, posterior means, model summary statistics)

Application: Elections

• Let's implement the "Normal Regression" model in Stan

```
model structure <- "
data {
 int N; // number of observations
  int K; // number of covariates
 matrix[N, K] X; //covariate matrix
  vector[N] v; //outcome vector
parameters {
  vector[K] beta; //regression coefficients
  real<lower = 0> sigma; // standard deviation
model {
  beta ~ normal(0, 9); // multivariate normal prior
  sigma \sim inv gamma(0.001/2, 0.001/2); // inverse gamma
  y ~ normal(X * beta, sigma); // * is matrix multiplication if terms are matrices
generated quantities {
  array[N] real y rep = normal rng(X * beta, sigma);
 vector[N] log lik;
  for (n in 1:N) log lik[n] = normal lpdf(y[n] | X[n, ] * beta, sigma);
```

Application: Elections

• Load the relevant packages

library(rstan)

• Pass the actual parameters as a list

```
data_source <- list(N = nrow(X_mat), K = ncol(X_mat), X=X_mat, y=Y)
```

Application: Elections

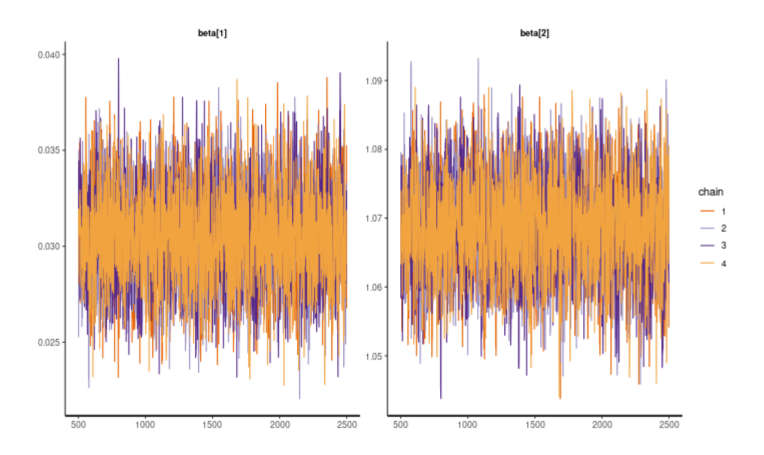
• Run Stan!

```
model_fit <- stan(
    model_code = model_structure,  # Stan code
    data = data_source,  # named list of data
    chains = 4,  # number of Markov chains
    warmup = 500,  # number of warmup iterations per chain
    iter = 2500,  # total number of iterations per chain
    cores = 4,  # number of cores (could use one per chain - by default uses however
    refresh = 0,  # no progress shown
    seed = 60637
    )
</pre>
```

Convergence

• The traceplot function lets us plot trace plots of any parameters or generated quantities. Let's do it for β

```
traceplot(model_fit, pars = c("beta"))
```



Summary

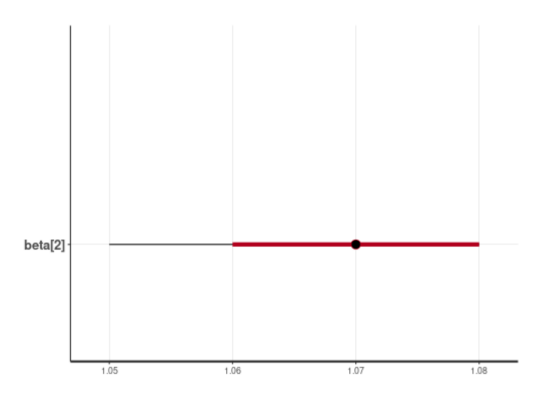
Summarize the fit

```
print(model fit, pars = c("beta", "sigma"))
## Inference for Stan model: anon model.
## 4 chains, each with iter=2500; warmup=500; thin=1;
## post-warmup draws per chain=2000, total post-warmup draws=8000.
##
          mean se mean sd 2.5% 25% 50% 75% 97.5% n eff Rhat
## beta[1] 0.03
                     0 0.00 0.03 0.03 0.03 0.03 0.04 4150
## beta[2] 1.07
                    0 0.01 1.05 1.06 1.07 1.07 1.08 3928
## sigma 0.06
                     0 0.00 0.06 0.06 0.06 0.06 0.06 3681
##
## Samples were drawn using NUTS(diag_e) at Thu Jan 25 10:28:51 2024.
## For each parameter, n eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor on split chains (at
## convergence, Rhat=1).
```

Summary

• Plot the results

```
stan_plot(model_fit, pars = c("beta[2]"))
```

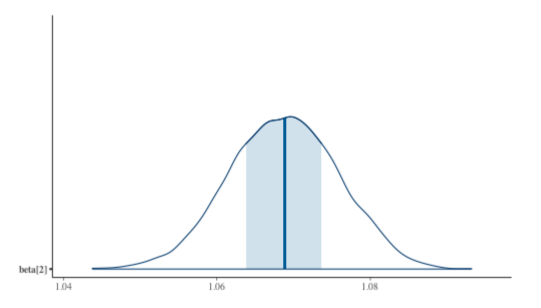


Summary

• There's cooler-looking stuff in bayesplot

```
library(bayesplot)
```

```
coef_plot <- mcmc_areas(model_fit, pars = c("beta[2]"))
coef_plot</pre>
```



Model diagnostics

Posterior predictive distribution

• A very useful quantity to obtain is the **predictive** distribution of y given the model.

$$f(ilde{y}|\mathbf{Y}) = \int f(ilde{y}, heta|\mathbf{Y})d heta = \int f(ilde{y}|\mathbf{Y}, heta)f(heta|\mathbf{Y})d heta$$

- Suppose our prediction target is the in-sample distribution of the covariates
 - We can get the posterior predictive distribution directly from our MCMC algorithm by $drawing^*$ a value of y for each value of θ that we sample.
- (e.g.) for the logit

```
generated quantities {
  y_rep ~ bernoulli_logit_rng(X * beta); //generate coin flips for each sample
}
```

- It is common to use the posterior predictive distribution as a model diagnostic ("posterior predictive checking")
 - 1. Check in-sample fit: Does the model actually reproduce the distribution of the data?
 - 2. Check against simulated data: Does the model recover the simulated distributions?

• Pull the y_rep from the model (in the form of a list)

```
y_ppc <- rstan::extract(model_fit)$y_rep</pre>
```

• Calculate the upper and lower credible intervals

```
y_ppc_ci <- t(apply(y_ppc, 2, function(x) quantile(x, c(.025, .975))))</pre>
```

What's the share that cover the truth?

```
cover_95 <- y_ppc_ci[,1]<Y&Y<y_ppc_ci[,2]
mean(cover_95)</pre>
```

[1] 0.958

What's the root mean squared deviation from the truth?

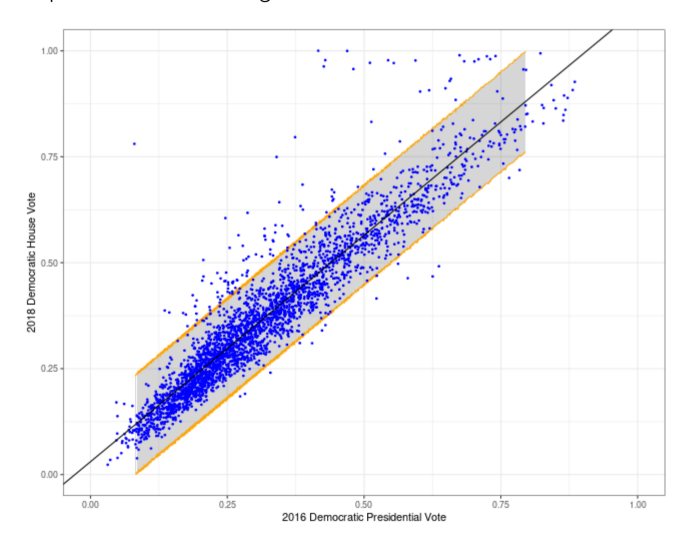
```
sqd_error <- apply((Y - t(y_ppc))^2, 2, mean)
print(sqrt(mean(sqd_error)))</pre>
```

[1] 0.0847

• Make a dataframe for the predictive "ribbon" plot

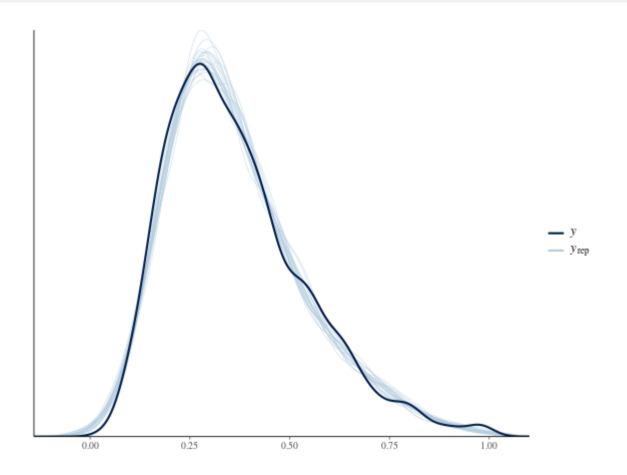
```
elections_county$lowerppd <- y_ppc_ci[,1]
elections_county$upperppd <- y_ppc_ci[,2]</pre>
```

• Plot the posterior predictive intervals against the data



• Or plot the empirical density vs. the posterior predicted curves

```
bayesplot::ppc_dens_overlay(y = elections_county$dem2018, yrep = y_ppc[1:25,])
```



Model Diagnostics

- There are two general approaches to evaluating model quality and comparing the predictive power of two or more statistical models given the same data
 - Information criteria
 - Cross-validation
- Both apporaches actually share common goals
 - Reward quality of predictions
 - **Penalize** model complexity
- We want to avoid models that overfit to the observed data
 - Information criteria address this with an explicit penalty term for the number of parameters
 - Cross-validation methods do this by evaluating fit on a "held-out" part of the data that's not used to estimate the model.

Information Criteria

• Frequentist/MLE Inference - the Akaike Information Criterion

$$\mathrm{AIC} = 2k - 2\ell(\hat{ heta})$$

where k is the number of parameters and $\ell(\hat{\theta})$ is the value of the maximized log-likelihood.

- Pick the model with the lowest AIC
 - Bayesian Inference the Deviance Information Criterion

$$ext{DIC} = 2p_{DIC} - 2\log p(y|\hat{ heta}_{ ext{PM}})$$

where $\hat{ heta}_{\mathrm{PM}}$ is the posterior mean and the "effective number of parameters" is

$$p_{DIC} = (\log p(y|\hat{ heta}_{ ext{PM}}) - E_{ ext{post}}[\log p(y| heta)])$$

Log-predictive density

- In general, a very common objective function for evaluating the quality of model "fit" in a Bayesian setting is the **log-predictive density** or $\log p(y|\theta)$
 - Think of it as a more general version of criteria like "mean-squared error"
- For a new "out-of-sample" \tilde{y}_i , the posterior predictive distribution is

$$\log p_{ ext{post}}({ ilde{y}}_i) = \log \int p({ ilde{y}}_i| heta) imes p_{ ext{post}}(heta) d heta$$

• We want to pick a model that has a high **expected log pointwise-predictive density** -- for a "new" size n sample of observation drawn from a common distribution $f(\tilde{y}_i)$,

$$ext{elppd} = \sum_{i=1}^n E_f[\log p_{ ext{post}}({ ilde{y}}_i)] = \sum_{i=1}^n \int \log p_{ ext{post}}({ ilde{y}}_i) f({ ilde{y}}_i) d{ ilde{y}}_i$$

Our goal is to try to estimate this from the data that we have.

Estimating elppd

ullet One approach would be to use the "in-sample" fit criterion using in-sample observations y_i

$$ext{lppd} = \sum_{i=1}^n \log p_{ ext{post}}(y_i)$$

- But here we know that it's going to be biased for the "out-of-sample" predictive error due to over-fitting.
- An improvement would be to correct the "in-sample" fit using a measure of the model complexity
 - This is the intuition behind AIC, DIC, (and other information criteria like WAIC)
- An alternative would be to use **cross-validation**:
 - \circ Split the sample into K "folds" (typical choice is 10)
 - \circ For each fold k, estimate the model on the K-1 other folds
 - \circ Using that model, calculate the log-predictive density for the observations in the kth fold
 - \circ The kth fold is the "held-out" (or "test" set), the K-1 remaining folds are the "training" set
- When K = N, this is often known as "leave-one out" cross-validation

Leave-one-out CV

- **loo** implements a leave-one-out cross-validation method that does not require re-fitting the model by using importance-sampling (Vehtari, Gelman and Gabry, 2017)
 - And interestingly, it's asymptotically equivalent to the WAIC ("Watanabe-Akaike (or widely applicable) information criterion")
- In order to use loo, we need the log-likelihood evaluated at each observation. This comes by default in canned rstanarm models but for custom models we need to specify to return it in the generated quantities block
- For the typical normal regression:

```
vector[N] log_lik;
for (n in 1:N) log_lik[n] = normal_lpdf(y[n] | X[n, ] * beta, sigma);
```

• But this will depend on the *actual* log-likelihood of the model

Leave-one-out CV

-8538.8 230.79

looic

• Once specified, we can obtain an estimate of the expected log predictive accuracy

```
library(loo)
## This is loo version 2.6.0
## - Online documentation and vignettes at mc-stan.org/loo
## - As of v2.0.0 loo defaults to 1 core but we recommend using as many as possible. Use the 'cores' are
##
## Attaching package: 'loo'
## The following object is masked from 'package:rstan':
##
       loo
##
 loo reg <- loo(model fit, pars="log lik")</pre>
 loo reg$estimates
##
           Estimate
                         SE
## elpd loo 4269.4 115.40
## p loo 10.5 2.94
```

Multilevel Models

Hierarchical Models

- In our simple normal regression from before, we assumed both a common mean and intercept for the data.
 - The underlying model is the same regardless of state.
- We also assumed that observations were completely exchangeable
 - $\circ Y_i \sim \operatorname{Normal}(X_i'\beta, \sigma^2)$ implies no residual correlation in the outcomes across **all** counties.
- This may be a poor modeling choice if states are very heterogeneous and information about the state in which a county is in would provide additional predictive power beyond knowing the 2016 presidential vote.
 - Additionally, exchangeability of observations may be violated -- unobserved factors may lead observations within a particular state to be correlated (e.g. if all counties in a region are exposed to some common shock).

Hierarchical Models

- **Hierarchical** or **Multilevel** linear models try to address this by incorporating additional structure on the regression parameters.
 - \circ Suppose observations belong to a group $j \in \{1,2,3,\ldots,J\}$ (e.g. states, time periods, schools, etc...)
 - We observe Y_{ij} as the outcome for unit i in grouping j.
- We can incorporate group membership into the data-generating process.
- Instead of a **common prior** on $\beta \sim \mathrm{Normal}(b_0, B_0^{-1})$, we may do something like:

$$eta_j \sim ext{Normal}(\mu_eta, \Sigma_eta)$$

$$Y_{ij} \sim ext{Normal}(X_{ij}'eta_j, \sigma^2)$$

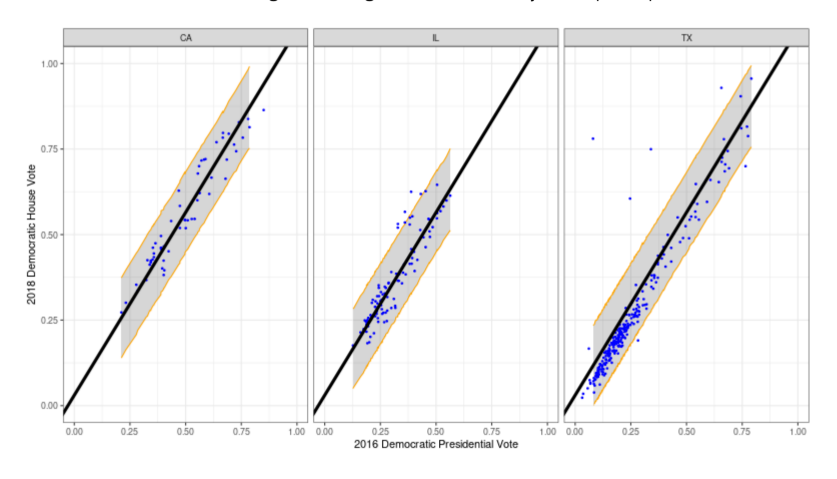
- This model assumes that each group has it's own set of group-level coefficients β_j that are drawn from a common distribution centered at β .
- The choice of prior on β then dictates the degree of **pooling** across groupings.
 - \circ Flat prior: **no pooling** β_i are estimated entirely separately
 - \circ Zero variance prior: complete pooling (all β_i are equal to β)

Hierarchical Models

- There are lots of choices about how we want to incorporate hierarchy into the model the coefficients across groupings
 - What levels of aggregation and how many?
 - Which parameters should be allowed to vary (all of the coefficients or just the intercepts)?
 - Where does the structure enter into the model (the means? the variances?)

Fully pooled model

- Our regression from before is an example of a fully pooled regression.
 - Within individual states though, the regression line may be a poor predictor



- The simplest fix is to allow the "intercept" to shift across units.
- Assume:

$$Y_{ij} \sim ext{Normal}(X_{ij}'eta + lpha_j, \sigma^2) \ lpha_j \sim ext{Normal}(0, \sigma_lpha^2)$$

And keep the same priors as before on the β and σ_j^2 parameters. Here, we now omit the intercept from the betas.

- α_j can be interpreted as the group-specific "shift" in the intercept from the "grand" intercept β_0
 - An equivalent parameterization would be to remove the intercept from β and write $\alpha_j \sim Normal(\mu_{\alpha}, \Sigma_{\alpha})$

• Let's implement this in Stan - first, our data block

```
data{
  int N; // number of observations
  int J; // Number of groups
  int K; // number of covariates
  array[N] int J_i; // group membership indicator
  matrix[N, K] X; //matrix of covariates
  vector[N] y; //outcome
}
```

• Next, our parameters.

```
parameters{
  vector[K] beta; // beta coefficients
  vector[J] alpha; // random intercepts
  real<lower = 0> sigma; // variance of outcome
  real<lower=0> sigma_a; // variance of intercepts
}
```

• Finally, our model

```
model{
  beta ~ normal(0, 9); // normal prior on coefficients
  alpha ~ normal(0, sigma_a); // normal distribution on random intercepts
  sigma ~ inv_gamma(0.001/2, 0.001/2);
  sigma_a ~ inv_gamma(0.001/2, 0.001/2);
  y ~ normal(alpha[J_i] + X*beta, sigma);
}
```

Put it into Stan!

```
model_var_intercept <- stan(
  model_code = varying_intercepts_model, # Stan code
  data = varying_intercepts_data, # named list of data
  chains = 4, # number of Markov chains
  warmup = 500, # number of warmup iterations per chain
  iter = 2500, # total number of iterations per chain
  cores = 4, # number of cores (this is much slower if = 1)
  refresh = 0, # no progress shown
  seed = 60637 # random seed
)</pre>
```

Summarize the fit.

```
print(model var intercept, pars = c("beta", "sigma", "alpha[1]"))
## Inference for Stan model: anon model.
## 4 chains, each with iter=2500; warmup=500; thin=1;
## post-warmup draws per chain=2000, total post-warmup draws=8000.
##
##
           mean se mean sd 2.5% 25% 50% 75% 97.5% n eff Rhat
                      0 0.01 0.03 0.04 0.04 0.04 0.05 610 1.01
## beta[1] 0.04
## beta[2] 1.07 0 0.01 1.05 1.06 1.06 1.07 1.08 9100 1.00
## sigma 0.05 0 0.00 0.05 0.05 0.05 0.05 0.05 10327 1.00
## alpha[1] 0.03
                      0 0.01 0.01 0.02 0.03 0.03 0.04 1076 1.01
##
## Samples were drawn using NUTS(diag_e) at Thu Jan 25 11:34:30 2024.
## For each parameter, n eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor on split chains (at
## convergence, Rhat=1).
```

• Pull the y_rep from the model (in the form of a list)

```
y_ppc_varint <- rstan::extract(model_var_intercept)$y_rep</pre>
```

• Calculate the upper and lower credible intervals

```
y_ppc_ci_varint <- t(apply(y_ppc_varint, 2, function(x) quantile(x, c(.025, .975))))</pre>
```

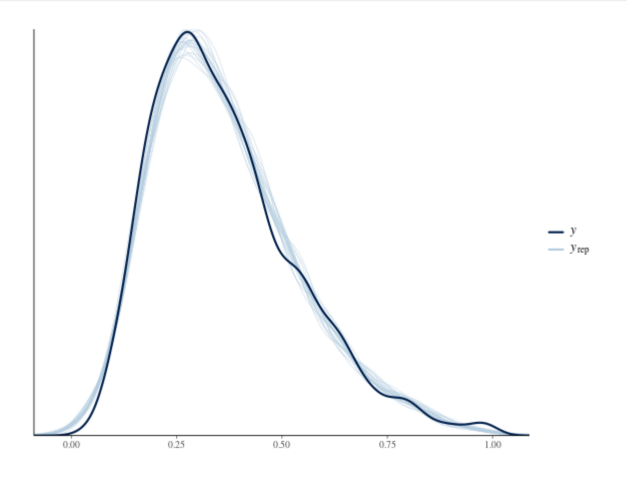
What's the share that cover the truth?

```
cover_95_varint <- y_ppc_ci_varint[,1]<Y&Y<y_ppc_ci_varint[,2]
mean(cover_95_varint)</pre>
```

[1] 0.962

• Empirical density vs. predicted

```
bayesplot::ppc_dens_overlay(y = elections_county$dem2018, yrep = y_ppc_varint[1:25,])
```



What's the root mean squared deviation from the truth?

```
sqd_error_varint <- apply((Y - t(y_ppc_varint))^2, 2, mean)
print(sqrt(mean(sqd_error_varint)))</pre>
```

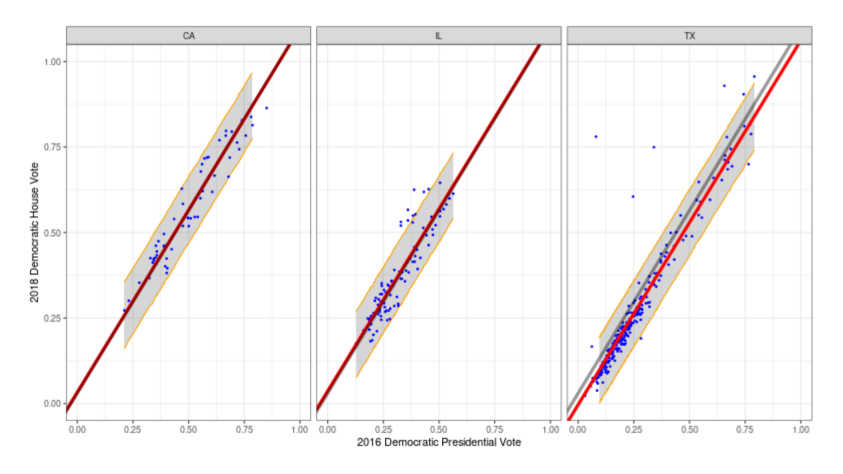
[1] 0.0693

• Make a dataframe for the predictive "ribbon" plot

```
elections_county$lowerppdVI <- y_ppc_ci_varint[,1]
elections_county$upperppdVI <- y_ppc_ci_varint[,2]</pre>
```

Outcome plots

• Compare the two regression lines

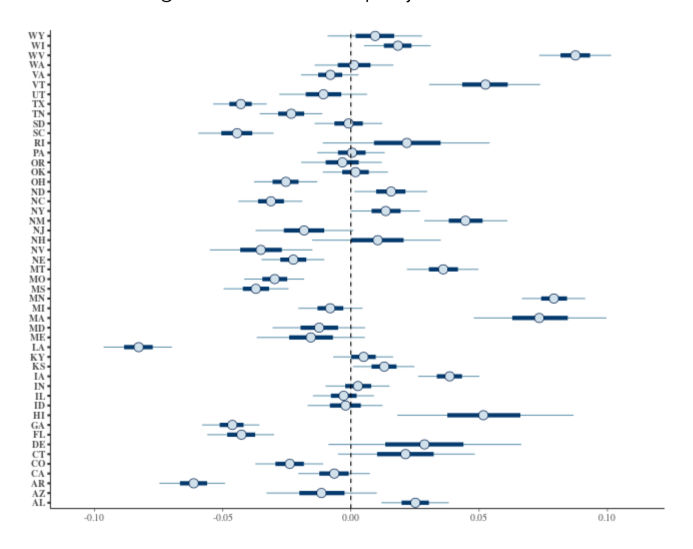


Leave-one-out statistic

• Calculate the leave-one-out statistic

```
library(loo)
 loo_varint <- loo(model_var_intercept, pars="log_lik")</pre>
 loo varint$estimates
##
           Estimate SE
## elpd loo 4855.2 162.13
## p_loo 60.9 9.79
## looic -9710.4 324.26
 loo compare(loo varint, loo reg)
## elpd diff se diff
## model1 0.0 0.0
## model2 -585.8
                     62.9
```

• Plot the deviations from the "grand mean" intercept by state



• What if we allow all of the parameters to vary by group?

$$eta_j \sim ext{Normal}(\mu_eta, \Sigma_eta)$$

$$Y_{ij} \sim ext{Normal}(X_{ij}'eta_j, \sigma^2)$$

- This is a more general and flexible hierarchical model
 - But introduces a slight complication -- we now need a prior distribution on an entire **matrix** Σ_{β} (the variances and covariances of the parameters).
- The conjugate prior is the inverse-Wishart, the multivariate extension of the inverse-gamma.
 - An alternative non-conjugate prior (that appears to have better performance) is a scaled correlation matrix with an LKJ prior on the correlation matrix (see Stan documentation for more).
 - The latter is now preferred (especially by the Stan programmers)

• Our prior on Σ_{β} starts by decomposing Σ_{β} into a diagonal scaling matrix $\operatorname{diag}(\tau)$ and a correlation matrix Ω

$$\Sigma_eta = ext{diag}(au)\Omega ext{diag}(au)$$

- For each τ_k we'll use a Half-Cauchy distribution (Cauchy with the constraint $\tau_k > 0$)
- For Ω , we'll use the Lewandowski-Kurowicka-Joe (LKJ) distribution which defines a distribution over symmetric positive definite matrices
 - o Intuition: Similar to a beta distribution in multiple dimensions.

• Our data block

```
data{
  int N; // number of observations
  int J; // Number of groups
  int K; // number of covariates
  array[N] int J_i; // group membership indicator
  matrix[N, K] X; //matrix of covariates
  vector[N] y; //outcome
}
```

• Our parameters field

```
parameters{
  corr_matrix[K] Omega; //prior correlation
  vector<lower=0>[K] tau; //prior scale
  vector[K] mu_beta;
  array[J] vector[K] beta; //group-level coefficients
  real<lower = 0> sigma; // variance parameters
}
```

• And our **model**

```
model{
    tau ~ cauchy(0, 2.5);
    Omega ~ lkj_corr(2);
    sigma ~ inv_gamma(0.001/2, 0.001/2); //Inverse-gamma
    mu_beta ~ normal(0, 9);
    beta ~ multi_normal(mu_beta, quad_form_diag(Omega, tau));
    for (n in 1:N) {
        y[n] ~ normal(X[n] * beta[J_i[n]], sigma);
    }
}
```

• Run it in Stan

```
model_var_slope <- stan(
  model_code = varying_slopes, # Stan code
  data = varying_intercepts_data, # named list of data
  chains = 4, # number of Markov chains
  warmup = 500, # number of warmup iterations per chain
  iter = 2500, # total number of iterations per chain
  cores = 4,
  refresh=0, # no progress shown
  seed = 60637 # random seed
)</pre>
```

Summarize the fit.

Pull the y_rep from the model (in the form of a list)

```
y_ppc_varslope <- rstan::extract(model_var_slope)$y_rep</pre>
```

• Calculate the upper and lower credible intervals

```
y_ppc_ci_varslope <- t(apply(y_ppc_varslope, 2, function(x) quantile(x, c(.025, .975))))</pre>
```

What's the share that cover the truth?

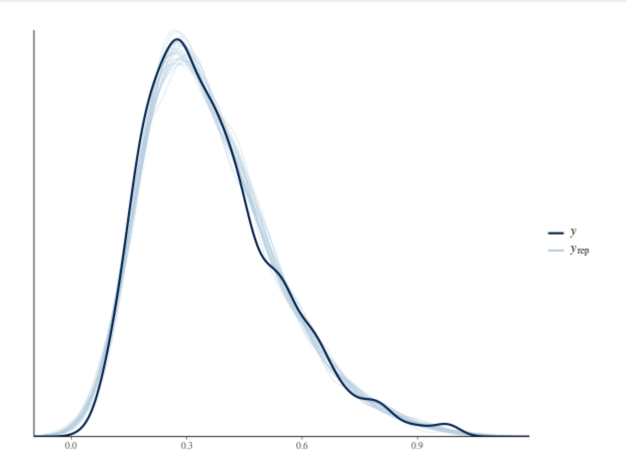
```
cover_95_varint <- y_ppc_ci_varslope[,1]<Y&Y<y_ppc_ci_varslope[,2]
mean(cover_95_varint)</pre>
```

[1] 0.966

Empirical density

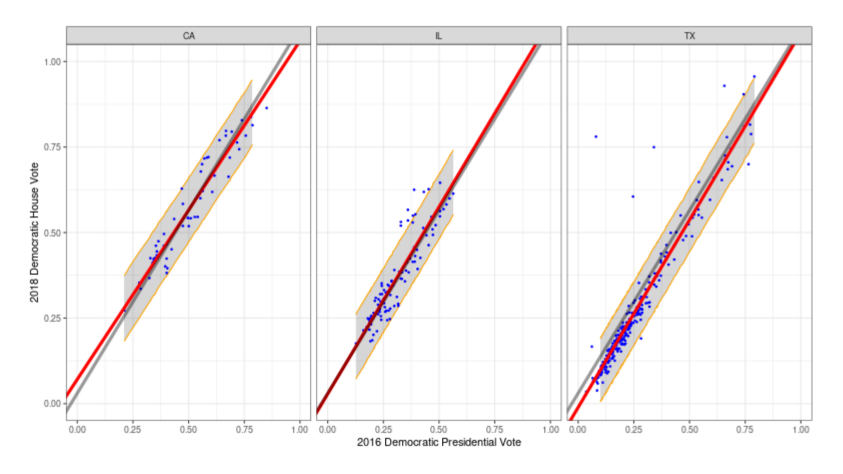
• Posterior predictions vs. empirical density

```
bayesplot::ppc_dens_overlay(y = elections_county$dem2018, yrep = y_ppc_varslope[1:25,])
```



Outcome plots

• Compare the regression lines



Leave-one-out statistic

• Calculate the leave-one-out statistic

```
library(loo)
 loo varslope <- loo(model var slope, pars="log lik")</pre>
 loo varslope$estimates
##
           Estimate
                       SF
## elpd loo 4930 170.3
## p_loo 103 15.8
## looic -9861 340.6
 loo compare(loo varslope ,loo varint, loo reg)
   elpd_diff se_diff
##
## model1 0.0 0.0
## model2 -75.3 29.1
## model3 -661.1 76.5
```