Markov Chain Monte Carlo and Stan

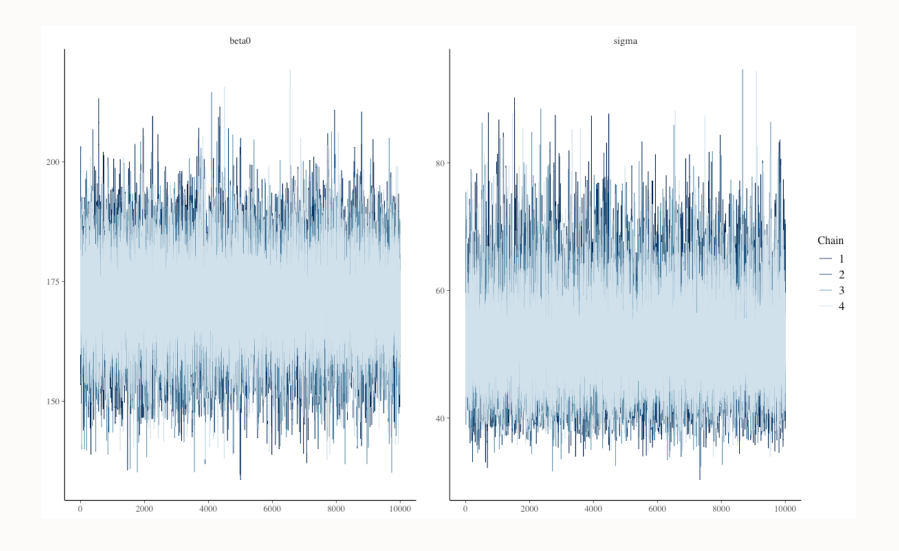
Lecture 3

https://jonathantemplin.com/bayesian-psychometric-modeling-fall-2024/

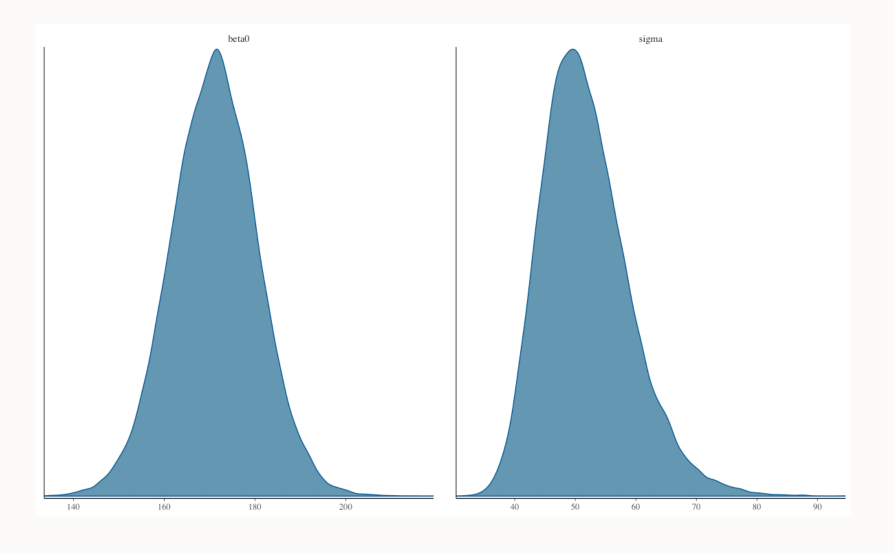
Today's Lecture Objectives

- 1. An Introduction to MCMC
- 2. An Introduction to Stan
- 3. Both with Linear Models

The Markov Chain Timeseries



The Posterior Distribution



Markov Chain Monte Carlo Estimation

Bayesian analysis is all about estimating the posterior distribution

- Up until now, we have worked with posterior distributions that fairly well-known
 - Beta-Binomial had a Beta distribution
 - In general, likelihood distributions from the exponential family have conjugate priors
 - Conjugate prior: the family of the prior is equivalent to the family of the posterior
- Most of the time, however, posterior distributions are not easily obtainable
 - No longer able to use properties of the distribution to estimate parameters

Markov Chain Monte Carlo Estimation

• It is possible to use an optimization algorithm (e.g., Newton-Raphson or Expectation-Maximization) to find maximum value of posterior distribution

- But, such algorithms may take a very long time for high-dimensional problems
- Variational Bayesian methods also attempt to approximate the center of the posterior distribution (and are faster than optimization algorithms for large models)
- Instead: MCMC "sketches" the posterior by sampling from it then use that sketch to make inferences
 - Sampling is done via MCMC

Markov Chain Monte Carlo Estimation

- MCMC algorithms iteratively sample from the posterior distribution
 - For fairly simplistic models, each iteration has independent samples
 - Most models have some layers of dependency included
 - Can slow down sampling from the posterior
- There are numerous variations of MCMC algorithms
 - Most of these specific algorithms use one of two types of sampling:
 - 1. Direct sampling from the posterior distribution (i.e. Gibbs sampling)
 - Often used when conjugate priors are specified
 - 2. Indirect (rejection-based) sampling from the posterior distribution (e.g., Metropolis-Hastings, Hamiltonian Monte Carlo)

MCMC Algorithms

- Efficiency is the main reason for so many algorithms
 - Efficiency in this context: How quickly the algorithm converges and provides adequate coverage ("sketching") of the posterior distribution
 - No one algorithm is uniformly most efficient for all models (here model = likelihood prior)
- The good news is that many software packages (stan, JAGS, MPlus, especially)
 don't make you choose which specific algorithm to use
- The bad news is that sometimes your model may take a large amount of time to reach convergence (think days or weeks)
- You can also code your own custom algorithm to make things run smoother

Commonalities Across MCMC Algorithms

- Despite having fairly broad differences regarding how algorithms sample from the posterior distribution, there are quite a few things that are similar across algorithms:
 - 1. A period of the Markov chain where sampling is not directly from the posterior
 - The burnin period (sometimes coupled with other tuning periods and called warmuup)
 - 2. Methods used to assess convergence of the chain to the posterior distribution
 - Often involving the need to use multiple chains with independent and differing starting values
 - 3. Summaries of the posterior distribution

Commonalities Across MCMC Algorithms

- Further, rejection-based sampling algorithms often need a tuning period to make the sampling more efficient
 - The tuning period comes before the algorithm begins its burnin period

MCMC Demonstration

 To demonstrate each type of algorithm, we will use a model for a normal distribution

- We will investigate each, briefly
- We will then switch over to stan to show the syntax and let stan work
- We will conclude by talking about assessing convergence and how to report parameter estimates.

Example Data: Post-Diet Weights

Example Data Link

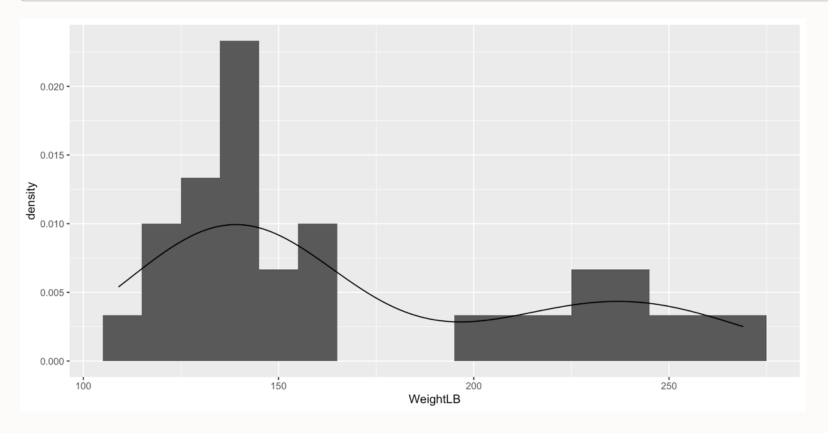
Visualizing Data: WeightLB Variable

```
DietData = read.csv(file = "DietData.csv")

ggplot(data = DietData, aes(x = WeightLB)) +

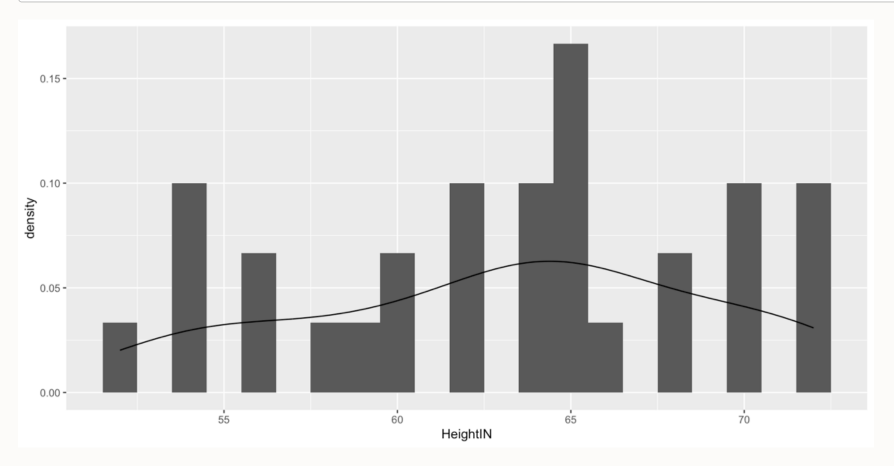
geom_histogram(aes(y = ..density..), position = "identity", binwidth = 10) +

geom_density(alpha=.2)
```



Visualizing Data: HeightIN Variable

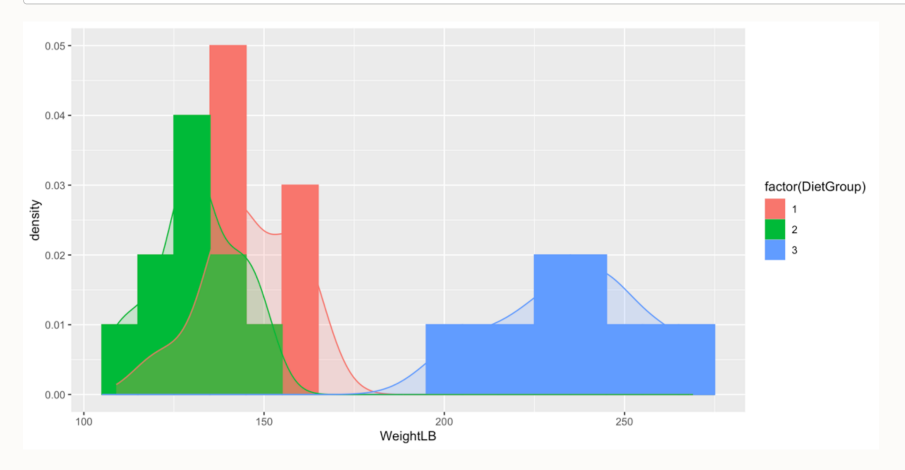
```
ggplot(data = DietData, aes(x = HeightIN)) +
geom_histogram(aes(y = ..density..), position = "identity", binwidth = 1) +
geom_density(alpha=.2)
```



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Visualizing Data: WeightLB by Group

```
ggplot(data = DietData, aes(x = WeightLB, color = factor(DietGroup), fill = factor(DietGroup))) +
geom_histogram(aes(y = ..density..), position = "identity", binwidth = 10) +
geom_density(alpha=.2)
```



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Visualizing Data: Weight by Height by Group

```
ggplot(data = DietData, aes(x = HeightIN, y = WeightLB, shape = factor(DietGroup), color = factor(Die
geom_smooth(method = "lm", se = FALSE) + geom_point()
```

Class Discussion: What Do We Do?

Now, your turn to answer questions:

- 1. What type of analysis seems most appropriate for these data?
- 2. Is the dependent variable (WeightLB) is appropriate as-is for such analysis or does it need transformed?

Linear Model with Least Squares

Let's play with models for data...

```
1 # center predictors for reasonable numbers
 2 DietData$HeightIN60 = DietData$HeightIN-60
   # full analysis model suggested by data:
   FullModel = lm(formula = WeightLB ~ 1, data = DietData)
   # examining assumptions and leverage of fit
   # plot(FullModel)
10 # looking at ANOVA table
11 # anova(FullModel)
12
   # looking at parameter summary
     summary(FullModel)
Call:
lm(formula = WeightLB ~ 1, data = DietData)
Residuals:
  Min
          10 Median
                         30
                              Max
-62.00 -36.75 -24.00 49.00 98.00
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 171.000
                         9.041 18.91 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 49.52 on 29 degrees of freedom

Path Diagram of Our Model

HeightIN60

DietGroup

DietGroup_X_HeightIN60

WeightLB

Steps in an MCMC Analysis

- 1. Specify model
- 2. Specify prior distributions for all model parameters
- 3. Build model syntax as needed
- 4. Run Markov chains (specify warmup/burnin and sampling period lengths)
- 5. Evaluate chain convergence
- 6. Interpret/report results

Specify Model

• To begin, let's start with an empty model and build up from there

• Let's examine the linear model we seek to estimate:

Where:

Questions:

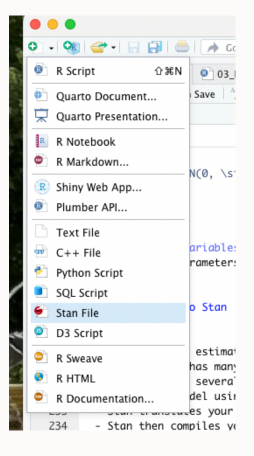
- What are the variables in this analysis?
- What are the parameters in this analysis?

Introduction to Stan

- Stan is an MCMC estimation program
 - Most recent; has many convenient features
 - Actually does severaly methods of estimation (ML, Variational Bayes)
- You create a model using Stan's syntax
 - Stan translates your model to a custom-built C++ syntax
 - Stan then compiles your model into its own executable program
- You then run the program to estimate your model
 - If you use R, the interface can be seamless

Stan and RStudio

- Stan has its own syntax which can be built in stand-alone text files
 - Rstudio will let you create one of these files in the new file menu
 - Rstudio also has syntax highlighting in Stan files
 - This is very helpful to learn the syntax
- Stan syntax can also be built from R character strings
 - Which is helpful when running more than one model per analysis



Stan Syntax

```
1 data {
     int<lower=0> N;
     vector[N] y;
   parameters {
     real beta0;
     real<lower=0> sigma;
 9 }
10
11 model {
     beta0 ~ normal(0, 1000); // prior for beta0
12
     sigma ~ uniform(0, 100000); // prior for sigma
13
14
     y ~ normal(beta0, sigma); // model for observed data
15 }
```

- Above is the syntax for our model
 - Each line ends with a semi colon
 - Comments are put in with //

Stan Syntax: R Character Object

```
1 stanModel = "
 2 data {
     int<lower=0> N;
     vector[N] y;
 5 }
   parameters {
     real beta0;
     real<lower=0> sigma;
 9
10 }
11
12 model {
13
     beta0 ~ normal(0, 1000); // prior for beta0
     sigma ~ uniform(0, 100000); // prior for sigma
14
     y ~ normal(beta0, sigma); // model for observed data
15
16 }
17 "
```

- Three blocks of syntax needed
 - Data: What Stan expects you will send to it for the analysis (using R lists)
 - Parameters: Where you specify what the parameters of the model are
 - Model: Where you specify the distributions of the priors and data

Stan Data and Parameter Delcaration

Like many compiled languages, Stan expects you to declare what type of data/parameters you are defining:

- int: Integer values (no decimals)
- real: Floating point numbers
- vector: A one-dimensional set of real valued numbers

Sometimes, additional definitions are provided giving the range of the variable (or restricting the set of starting values):

real<lower=0> sigma;

See: https://mc-stan.org/docs/reference-manual/data-types.html for more information

Stan Data and Prior Distributions

• In the model section, you define the distributions needed for the model and the priors

■ The left-hand side is either defined in data or parameters

```
y ~ normal(beta0, sigma); // model for observed datasigma ~ uniform(0, 100000); // prior for sigma
```

- The right-hand side is a distribution included in Stan
 - You can also define your own distributions

See: https://mc-stan.org/docs/functions-reference/index.html for more information

From Stan Syntax to Compilation

```
1 # compile model -- this method is for stand-alone stan files (uses cmdstanr)
2 model00.fromFile = cmdstan_model(stan_file = "model00.stan")
3
4 # or this method using the string text in R
5 model00.fromString = cmdstan_model(stan_file = write_stan_file(stanModel))
```

- Once you have your syntax, next you need to have Stan translate it into C++ and compile an executable
- This is where cmdstanr and rstan differ
 - cmdstanr wants you to compile first, then run the Markov chain
 - rstan conducts compilation (if needed) then runs the Markov chain

Building Data for Stan

```
# build R list containing data for Stan: Must be named what "data" are listed in analysis
stanData = list(
    N = nrow(DietData),
    y = DietData$WeightLB

)

# snippet of Stan syntax:
stanSyntaxSnippet = "
data {
    int<lower=0> N;
    vector[N] y;
}
```

- Stan needs the data you declared in your syntax to be able to run
- Within R, we can pass this data to Stan via a list object
- The entries in the list should correspond to the data portion of the Stan syntax
 - In the above syntax, we told Stan to expect a single integer named N and a vector named y
- The R list object is the same for cmdstanr and rstan

Running Markov Chains in cmdstanr

```
# run MCMC chain (sample from posterior)
model00.samples = model00.fromFile$sample(
data = stanData,
seed = 1,
chains = 4,
parallel_chains = 4,
iter_warmup = 10000,
iter_sampling = 10000
)
```

Running Markov Chains in rstan

```
1  rstan_options(auto_write = TRUE)
2  options(mc.cores = parallel::detectCores())
3
4  # example MCMC analysis in rstan
5  model00.rstan = stan(
6  model_code = stanModel,
7  model_name = "Empty model",
8  data = stanData,
9  warmup = 10000,
10  iter = 20000,
11  chains = 4,
12  verbose = TRUE
```

- Rstan takes the model syntax directly, then compiles and runs the chains
- The first two lines of syntax enable running one chain per thread (parallel processing)
 - As chains are independent, running them simultaneously (parallel) shortens wait time considerably
- The verbose option is helpful for detecting when things break
- The same R list supplies the data to Stan

MCMC Process

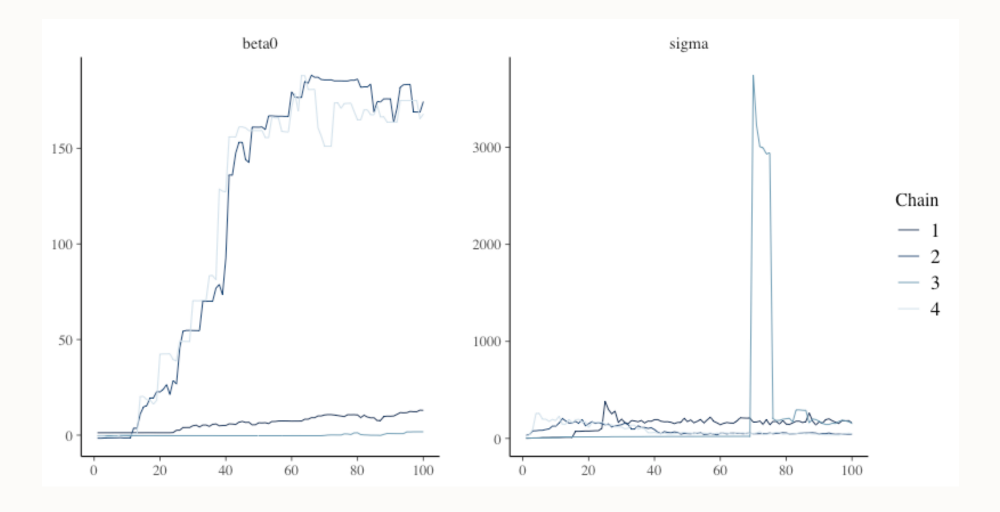
- The MCMC algorithm runs as a series of discrete iterations
 - Within each iteration, each parameter of a model has an opportunity to change its value
- For each parameter, a new parameter is sampled at random from the current belief of posterior distribution
 - The specifics of the sampling process differ by algorithm type (we'll have a lecture on this later)

MCMC Process

• In Stan (Hamiltonian Monte Carlo), for a given iteration, a proposed parameter is generated

- The posterior likelihood "values" (more than just density; includes likelihood of proposal) are calculated for the current and proposed values of the parameter
- The proposed values are accepted based on the draw of a uniform number compared to a transition probability
- If all models are specified correctly, then regardless of starting location, each chain will converge to the posterior if run long enough
 - But, the chains must be checked for convergence when the algorithm stops

Example of Bad Convergence



Examining Chain Convergence

- Once Stan stops, the next step is to determine if the chains converged to their posterior distribution
 - Called convergence diagnosis
- Many methods have been developed for diagnosing if Markov chains have converged
 - Two most common: visual in spection and Gelman-Rubin Potential Scale Reduction Factor (PSRF; quick reference)

Examining Chain Convergence

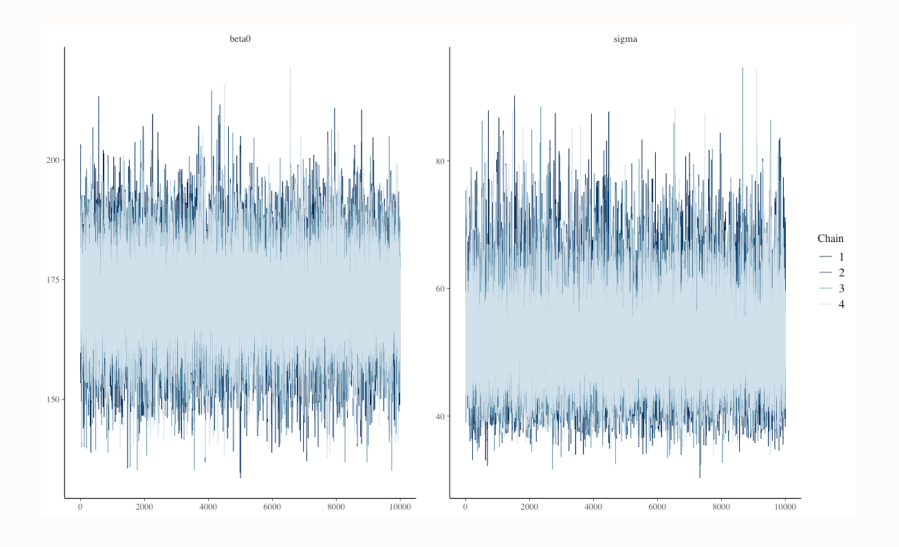
- Visual inspection
 - Want no trends in timeseries should look like a catapillar
 - Shape of posterior density should be mostly smooth
- Gelman-Rubin PSRF (denoted with)
 - For analyses with multiple chains
 - Ratio of between-chain variance to within-chain variance
 - Should be near 1 (maximum somewhere under 1.1)

Setting MCMC Options

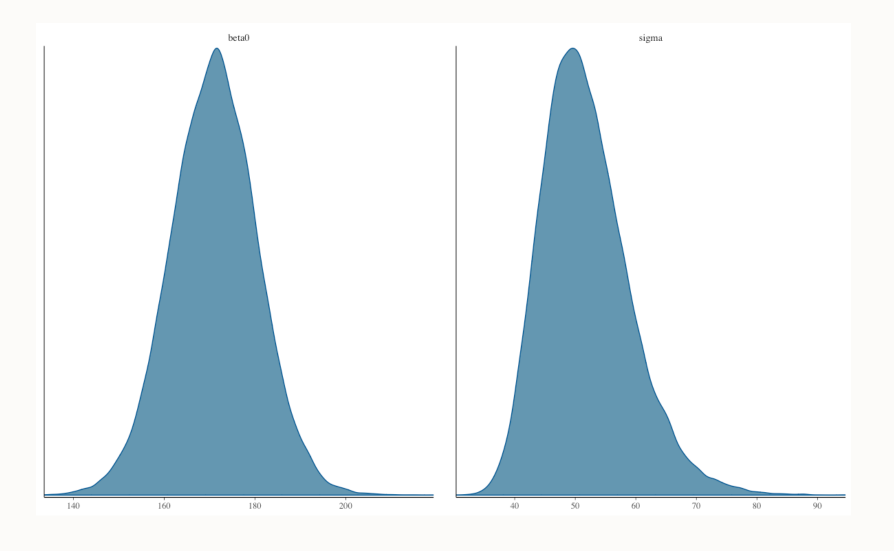
• As convergence is assessed using multiple chains, more than one should be run

- Between-chain variance estimates improve with the number of chains, so I typically use four
- Others have two; more than one should work
- Warmup/burnin period should be long enough to ensure chains move to center of posterior distribution
 - Difficult to determine ahead of time
 - More complex models need more warmup/burnin to converge
- Sampling iterations should be long enough to thoroughly sample posterior distribution
 - Difficulty to determine ahead of time
 - Need smooth densities across bulk of posterior
- Often, multiple analyses (with different settings) is what is needed

The Markov Chain Timeseries



The Posterior Distribution



Assessing Our Chains

```
1 model00.samples$summary()
# A tibble: 3 \times 10
 variable
             mean median
                                                  q95 rhat ess bulk ess tail
                                   mad
                                            q5
  <chr>
            <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
                                                                <dbl>
                                                                          <dbl>
1 lp
           -129.
                           1.02 \ 0.735 \ -131.
                                               -128.
                                                        1.00
                                                               17577.
                                                                         22523.
                                        155.
                                                       1.00
                                                                         25175.
2 beta0
            171.
                    171.
                           9.44 9.31
                                                186.
                                                               29639.
             51.8
                     51.0 7.20 6.84
                                          41.5
                                                 64.8 1.00
3 sigma
                                                               29147.
                                                                         24620.
```

- The summary function reports the PSRF (rhat)
- Here we look at our two parameters: and
- Both have , so both would be considered converged
- lp___ is posterior log likelihood—does not necessarily need examined
- ess_ columns show effect sample size for chain (factoring in autocorrelation between correlations)
 - More is better

Results Interpretation

- At long last, with a set of converged Markov chains, we can now interpret the results
 - Here, we disregard which chain samples came from and pool all sampled values to use for results
- We use summaries of posterior distributions when describing model parameters
 - Typical summary: the posterior mean
 - The mean of the sampled values in the chain
 - Called EAP (Expected a Posteriori) estimates
 - Less common: posterior median

Results Interpretation

- Important point:
 - Posterior means are different than what characterizes the ML estimates
 - Analogous to ML estimates would be the mode of the posterior distribution
 - Especially important if looking at non-symmetric posterior distributions
 - Look at posterior for variances

Results Interpretation

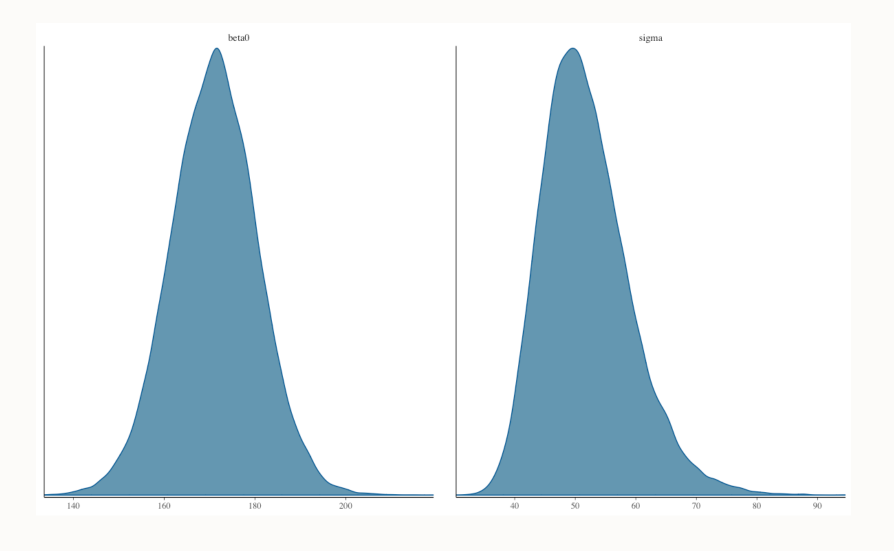
• To summarize the uncertainty in parameters, the posterior standard deviation is used

- The standard deviation of the sampled values in the chain
- This is the analogous to the standard error from ML
- Bayesian credible intervals are formed by taking quantiles of the posterior distribution
 - Analogous to confidence intervals
 - Interpretation slightly different the probability the parameter lies within the interval
 - 95% credible interval notes that parameter is within interval with 95% confidence
- Additionally, highest density posterior intervals can be formed
 - The narrowest range for an interval (for unimodal posterior distributions)

Our Results

```
1 model00.samples$summary()
# A tibble: 3 \times 10
 variable mean median
                                               q95 rhat ess bulk ess tail
                            sd
                                 mad
                                         q5
  <chr>
          <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <</pre>
                                                            <dbl>
                                                                     <dbl>
           -129. -128.
                         1.02 0.735 -131. -128.
                                                    1.00
                                                           17577.
                                                                    22523.
1 lp
           171.
                                    155.
                                            186.
                                                   1.00
                                                                    25175.
2 beta0
                  171.
                          9.44 9.31
                                                           29639.
                                              64.8 1.00
3 sigma
             51.8
                  51.0 7.20 6.84
                                       41.5
                                                                    24620.
                                                           29147.
 1 hdi(model00.samples$draws("beta0"), credMass = .9)
  lower
          upper
155.015 185.762
attr(,"credMass")
[1] 0.9
 1 hdi(model00.samples$draws("sigma"), credMass = .9)
  lower
         upper
40.2305 63.0083
attr(,"credMass")
[1] 0.9
```

The Posterior Distribution



Wrapping Up

- This lecture covered the basics of MCMC estimation with Stan
- Next we will use an example to show a full analysis of the data problem we started with today
- The details today are the same for all MCMC analyses, regardless of which algorithm is used