Bayesian Linear Regression

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Bayesian Linear Regression

- In this class, which is mostly based on chapter 4 of [McElreath, 2020], we are going to revisit the linear regression model from a Bayesian point of view.
- The idea is the same: to model the relationship of a numerical dependent variable y with m independent variables x₁, x₂,..., x_m from a dataset d.
- The response variable **y** is again modeled with a Gaussian distribution: $y_i \sim N(\mu_i, \sigma^2)$.
- We also maintain the assumption that each attribute has a linear relationship to the mean of the outcome.

$$\mu_i = \beta_0 + \beta_1 x_i + \dots \beta_m x_m$$

- However, we are not going to use least squares or maximum likelihood to obtain point estimates of the parameters.
- Instead, we are going to estimate the joint posterior distribution of all the parameters of the model:

$$f(\theta|\mathbf{d}) = f(\beta_0, \beta_1, \dots, \beta_m, \sigma|\mathbf{d})$$

Bayesian Linear Models

- The Bayesian linear regression is more flexible than least squares as it allows incorporating prior information.
- It also allows to interpret the uncertainty of the model in a much richer way.
- Notice that the parameters of the model are $\beta_0, \beta_1, \dots, \beta_m$ and σ but not μ_i .
- This is because μ_i it is determined deterministically from the linear model's coefficients.
- In order to build our posterior we need to define a likelihood function:

$$f(\mathbf{d}|\beta_0,\beta_1,\cdots,\beta_m,\sigma)=\prod_{i=1}^n f(\mathbf{d}_i|\beta_0,\beta_1,\cdots,\beta_m,\sigma)$$

- Where d_i corresponds to each data point in the dataset d containing n values for y and x_1, \ldots, x_m (IID assumption).
- The likelihood of each point is modeled with a Gaussian distribution:

$$f(d_i|\beta_0,\beta_1,\cdots,\beta_m,\sigma)=N(\mu_i,\sigma^2)$$

Bayesian Linear Models

Now we need a joint prior density:

$$f(\theta) = f(\beta_0, \beta_1, \dots, \beta_m, \sigma)$$

And the posterior gets specified as follows:

$$f(\theta|\mathbf{d}) = \frac{\prod_{i=1}^{n} f(\mathbf{d}_{i}|\beta_{0}, \beta_{1}, \cdots, \beta_{m}, \sigma) * f(\beta_{0}, \beta_{1}, \dots, \beta_{m}, \sigma)}{f(\mathbf{d})}$$

The evidence is expressed by a multiple integral:

$$f(d) = \int \int \cdots \int \prod_{i=1}^{n} f(d_{i}|\beta_{0}, \beta_{1}, \cdots, \beta_{m}, \sigma) * f(\beta_{0}, \beta_{1}, \dots, \beta_{m}, \sigma) d\beta_{0} d\beta_{1} \cdots d\beta_{m} d\sigma$$

 In most cases, the priors are specified independently for each parameter, which is equivalent to assuming:

$$f(\beta_0, \beta_1, \cdots, \beta_m, \sigma) = f(\beta_0) * f(\beta_1) * \cdots * f(\beta_m) * f(\sigma).$$



A model of height revisited

- To understand this more concretely, we will rebuild the linear model relating the height and weight of the !Kung San people using a Bayesian approach.
- We will refer to each person's height and weight as y_i and x_i respectively.
- Our probabilistic model specifying all components of a Bayesian model is defined as follows:

$$\begin{array}{ll} y_i \sim \textit{N}(\mu_i, \sigma) & \text{[likelihood]} \\ \mu_i = \beta_0 + \beta_1 x_i & \text{[linear model]} \\ \beta_0 \sim \textit{N}(150, 50) & [\beta_0 \text{ prior]} \\ \beta_1 \sim \textit{N}(0, 1) & [\beta_1 \text{ prior]} \\ \sigma \sim \text{Uniform}(0, 50) & [\sigma \text{ prior]} \end{array}$$

- Parameters β_0 and β_1 are the intercept and the slope of our linear model.
- The parameter σ is the standard deviation of all the heights.
- ullet Note that we are setting the same σ for all observations, which is equivalent to the Homoscedasticity property of the standard linear regression.

A model of height revisited

- Our priors were set independently for each parameter which implies that the joint prior density $f(\beta_0, \beta_1, \sigma)$ can be expressed as $f(\beta_0) * f(\beta_1) * f(\sigma)$.
- It should be kept in mind that the choice of priors is subjective and should be evaluated accordingly.¹
- Let's try to justify our choice a bit:
 - 1 The Gaussian prior for β_0 (intercept), centered on 150cm with a standard variation of 50, covers a huge range of plausible mean heights for human populations while giving very little chance for negative heights.
 - **2** The Gaussian prior for β_1 (slope), centered on 0 with a standard variation of 1, acts as a **regularizer** to prevent the model from **overfitting** the data by assigning extreme values to β_1 .²
 - \odot The uniform prior for the standard deviation σ between 0 and 50 prohibits obtaining negative standard deviations. The upper bound (50 cm) would imply that 95% of individual heights lie within 100cm of the average height. That's a very large range.

https://github.com/stan-dev/stan/wiki/

Prior-Choice-Recommendations

¹A guide to prior choice recommendations can be found here:

²Regularization and overfitting will be discussed later in the course.

- Now we need to fit the model to the data to build the posterior distribution.
- Grid approximation is not a valid option, as setting up a grid for 3 parameters would be too computationally expensive.
- We will use Laplace approximation instead.
- In this approach we obtain the MAP estimates for each parameter using a hill-climbing optimization method.
- Then we fit a multivariate Gaussian distribution centered on these values.
- This distribution is the multidimensional extension to the standard Gaussian.

 The multivariate Gaussian distribution in m-dimensions is defined by the following density function (PDF):

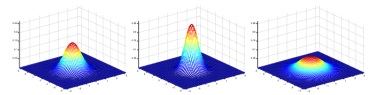
$$f_X = \frac{1}{(2\pi)^{m/2} |\Sigma|^{1/2}} \exp\left(-\frac{1}{2} (\vec{x} - \vec{\mu})^T \Sigma^{-1} (\vec{x} - \mu)\right)$$

- This density function allows working with a m-dimensional vector of random variables \vec{X} .
- The first parameter of this distributions is a mean vector $\vec{\mu} \in \mathcal{R}^m$ with the mean value of each dimension.
- The second parameter is a covariance matrix $\Sigma \in \mathbb{R}^{m \times m}$.
- This matrix contains the variance of each variable in the diagonal and the covariance of variables X_i and X_i in the other cells Σ_{i,j}:

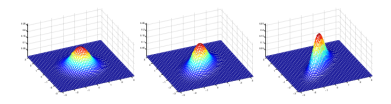
$$Cov(X) = \Sigma$$

- The matrix Σ is symmetric and positive semi-definite.
- The multivariate Gaussian $N(\vec{\mu}, \Sigma)$ is a very convenient distribution for modeling multidimensional random variables.

 Here are some examples taken from [Ng, 2008] of what the density of a multivariate Gaussian distribution looks like:



- The left-most figure shows a Gaussian with mean zero (that is, the 2x1 zero-vector) and covariance matrix Σ = I (the 2 × 2 identity matrix).
- A Gaussian with zero mean and identity covariance is also called the standard normal distribution.
- The middle figure shows the density of a Gaussian with zero mean and $\Sigma = 0.6$ *l*.
- The rightmost figure shows one with $\Sigma = 2I$.
- We see that as Σ becomes larger, the Gaussian becomes more "spread-out", and as it becomes smaller, the distribution becomes more "compressed".

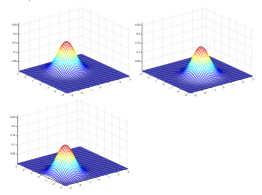


 The figures above show Gaussians with mean 0, and with covariance matrices respectively

$$\Sigma = \left[\begin{array}{cc} 1 & 0 \\ 0 & 1 \end{array} \right]; \;\; \Sigma = \left[\begin{array}{cc} 1 & 0.5 \\ 0.5 & 1 \end{array} \right]; \;\; \Sigma = \left[\begin{array}{cc} 1 & 0.8 \\ 0.8 & 1 \end{array} \right].$$

• The leftmost figure shows the familiar standard normal distribution, and we see that as we increase the off-diagonal entry in Σ , the density becomes more "compressed" towards the 45- line (given by $x_1 = x_2$).

• As our last set of examples, fixing $\Sigma = I$, by varying $\vec{\mu}$ we can also move the mean of the density around.



• The figures above were generated using $\Sigma = I$, and respectively

$$\mu = \left[\begin{array}{c} 1 \\ 0 \end{array} \right]; \;\; \mu = \left[\begin{array}{c} \text{-0.5} \\ 0 \end{array} \right]; \;\; \mu = \left[\begin{array}{c} \text{-1} \\ \text{-1.5} \end{array} \right].$$

Laplace approximation

- In Laplace approximation we assume that the joint posterior follows a multivariate Gaussian distribution $f(\theta_1, \dots, \theta_m | d) = N(\vec{\mu}, \Sigma)$.
- According to [Gelman et al., 2013], this approximation is convenient for unimodal and roughly symmetric posterior distributions.
- There are also a theoretical asymptotic argument in favor of this approximation: "if the dataset is large enough, a posterior distribution can be approximated by a Gaussian" [Gelman et al., 2013].
- The values of $\vec{\mu}$ are obtained from the posterior mode of each parameter (MAP):

$$ec{\mu} = ec{ heta}_{ extsf{MAP}}$$

Which are obtained using numeric optimization techniques.

Laplace approximation

• The values of Σ are obtained from the curvature near these values, which are obtained from the second derivatives of the posterior:

$$\Sigma = [\textit{I}(\theta_{\textit{MAP}})]^{-1}$$

where

$$I(\theta) = -\frac{d^2}{d\theta^2} \log f(\theta|d)$$

- Notice that both $\vec{\mu}$ and Σ can be calculated from the unnormalized posterior $f(d|\theta)*f(\theta)$
- We don't need to calculate the evidence f(d) to perform Laplace approximation [Srihari,].

- Laplace approximation is implemented in the quap function from the rethinking package.
- The model for height defined above can be implemented as follows:

```
library(rethinking)
data(Howell1)
d <- Howell1
d2 <- d[ d$age >= 18 , ]

b.reg1 <- quap(
    alist(
        height ~ dnorm( mu, sigma ),
        mu <- b0 + b1*weight,
        b0 ~ dnorm( 150 , 50 ),
        b1 ~ dnorm( 0 , 1) ,
        sigma ~ dunif( 0 , 50 )
        ) , data=d2 )</pre>
```

We can summarize this posterior with the command precis:

- These numbers provide Gaussian approximations for each parameter's marginal posterior distribution.
- The marginal of multivariate distribution is the univariate distribution of a single parameter θ_i after integrating (averaging) over all the other parameters $\theta_j \ \forall j \neq i$:

$$f(\theta_i|d) = \int \cdots \int f(\theta_1, \ldots, \theta_m|d) d\theta_1, \ldots, d\theta_{i-1}, d\theta_{i+1}, \ldots, d\theta_m$$

- In a multivariate Gaussian $N(\vec{\mu}, \Sigma)$, the marginal distribution of θ_i is a univariate Gaussian $N(\vec{\mu}_i, \Sigma_{ii})$.
- So, the marginal distribution of β_1 is a Gaussian distribution with $\mu=$ 0.9 and $\sigma=$ 1.9.
- The last two columns of the summary table show the lower and upper limits of a 95% equal-tailed credible interval for each parameter.
- For the case of β_1 , 95% of the posterior probability lies between 0.82 and 0.98.
- That suggests that β₁ values close to zero or greatly above one are highly incompatible with these data and this model.

• Let's compare the mean and standard deviation of β_0 and β_1 with what we obtained using least squares in previous lecture:

- These values are almost the same!
- Recall that maximum likelihood or least squares estimators are identical to MAP estimators with uniform priors.
- This indicates that our priors did not have a significant impact in the resulting posterior.

Covariance Matrix

 We can also get the covariance matrix Σ of our multivariate Gaussian obtained with Laplace approximation:

- The matrix tell us how each parameter relates to every other parameter in the posterior distribution.
- It is better to turn them into correlations for easier interpretation:

- Notice that β_0 and β_1 are almost perfectly negatively correlated.
- This means that these two parameters carry the same information: as we change the slope of the line, the best intercept changes to match it.

Centering

- In more complex models, strong correlations like this can make it difficult to fit the model to the data.
- A useful trick to avoid this problem is centering: subtract the mean of a variable from each value.

• The correlations among parameters are now all very close to zero.

Sampling from the Posterior

- In the remainder of this class we will work with samples from our multi-dimensional posterior.
- The rethinking package provides the **extract.samples** function to obtain them:

- Each sample is a different line relating height and weight.
- Bear in mind that each line is sampled in proportion to the posterior probabilities, which represent our model's beliefs.

Sampling from the Posterior

• We can see that the mean of each variable in the sample is almost identical to the corresponding MAP estimate:

```
> sapply(post,mean)
b0 b1 sigma
114.0026648 0.9023462 5.0693261
```

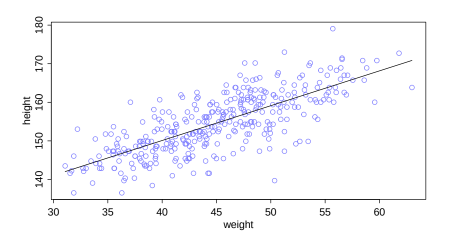
 Since our posterior was approximated with Laplace approximation, we can obtain equivalent samples directly from a multi-variate Gaussian using the mvrnorm function:

Plotting posterior inference against the data

- In truth, tables of estimates are usually insufficient for understanding the information contained in the posterior distribution.
- It's almost always much more useful to plot the posterior inference against the data.
- We're going to start with a simple version of that task, superimposing just the MAP values over the height and weight data.

```
plot( height ~ weight , data=d2 , col=rangi2 )
b0_map <- mean(post$b0)
b1_map <- mean(post$b1)
curve( b0_map + b1_map*x, add=TRUE )</pre>
```

Plotting posterior inference against the data

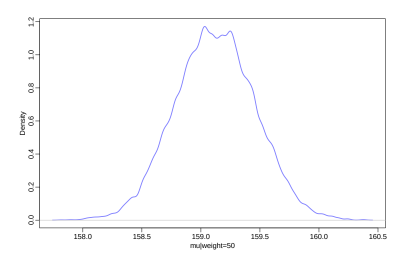


- The previous plot doesn't incorporate the uncertainty embodied in the posterior.
- This can be done by by plotting a credible interval around the MAP regression line.
- To understand how it works, let's focus for the moment on a single weight value, say 50 kilograms.
- We can quickly make a list of 10,000 values of μ for an individual who weighs 50 kilograms, by using our samples from the posterior:

```
mu_at_50 <- post$b0 + post$b1 * 50</pre>
```

• It might be helpful at this point to actually plot the density for this vector of means:

```
dens( mu_at_50 , col=rangi2 , lwd=2 , xlab="mu|weight=50" )
```



- Since the components of μ have distributions (β_0, β_1) , so too does μ .
- And since the distributions of β_0 and β_1 are Gaussian, so to is the distribution of μ .
- Adding Gaussian distributions always produces a Gaussian distribution.
- ullet Since the posterior for μ is a distribution, you can find intervals for it, just like for any posterior distribution.
- ullet To find the 95% highest posterior density interval of μ at 50 kg, we just use the HPDI command as usual:

```
> HPDI( mu_at_50 , prob=0.95 )
    |0.95     0.95|
158.4883 159.8061
```

 Now, we need to repeat the above calculation for every weight value on the horizontal axis, not just when it is 50 kg.

We first define a sequence of weights to compute predictions for:

```
weight.seg <- seg( from=25 , to=70 , by=1 )
```

- These values will be on the horizontal axis of our plot.
- Now, for each of these weights we will compute a posterior distribution of the average height μ using samples from the posterior:

```
mu.link <- function(weight) post$b0 + post$b1*weight
```

- The function above will use the 1000 different sampled regression lines to compute various values of μ for a given weight.
- We can apply this function to all our 46 weights (from 25 to 70) and obtain a matrix of 1000 × 46:

```
> mu <- sapply( weight.seq , mu.link )
> dim(mu)
[1] 10000     46
```

 Now, we can get a MAP value of μ for each weight, simply by by averaging the rows of the matrix:

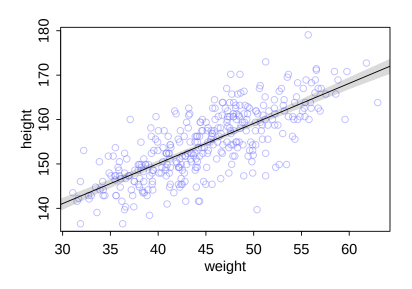
```
> mu.mean <- apply( mu , 2 , mean )
> head(mu.mean)
[1] 136.5526 137.4555 138.3583 139.2612 140.1641 141.0669
```

• And we can get our 95% HPDIs analogously:

```
mu.HPDI <- apply( mu , 2 , HPDI , prob=0.95 )
```

- The variable mu.HPDI contains 95% lower and upper bounds for each weight value.
- Now we can plot these intervals as shaded regions around the MAP estimates as follows:

```
plot( height ~ weight , data=d2 , col=col.alpha(rangi2,0.5) )
# plot the MAP line, aka the mean mu for each weight
lines( weight.seq , mu.mean )
# plot a shaded region for 95% HPPDI
shade( mu.HPDI , weight.seq )
```



 In case of working with more complex linear models (i.e., multiple attributes, interactions) we can use the link function from the rethinking package:

```
mu \leftarrow link(b.reg1, data=data.frame(weight=weight.seq), n=1000)
```

- This function will generate distributions of posterior values for μ using the formula declared in **quap** (e.g., $\mu = \beta_0 + \beta_1 * x$).
- The default behavior of link is to use the original data.
- So we have to pass it a list of new horizontal axis values we want to plot posterior predictions across.

- Now let's walk through generating prediction intervals for simulated heights \tilde{y} using the posterior predictive distribution.
- What we did before was to use samples from the posterior to visualize the uncertainty in μ_i using the linear model of the mean: $\mu_i = \beta_0 + \beta_1 * x_i$.
- But actual predictions of heights ỹ depend also upon the stochastic definition of y given in the likelihood function: y_i ~ N(μ_i, σ).
- The Gaussian distribution of the likelihood tells us that the model expects observed heights \tilde{y} to be distributed around μ , not right on top of it.
- And the spread around μ is governed by σ .
- All of this suggests we need to incorporate σ in the predictions somehow.

 The posterior predictive distribution introduced in previous class can exactly do that:

$$f(\tilde{y}|y) = \int_{\theta} f(\tilde{y}|\theta) f(\theta|y) d\theta$$

For our height regression model it would take the following form:

$$f(\tilde{\mathbf{y}}|\mathbf{y}) = \int_{\beta_0} \int_{\beta_1} \int_{\sigma} f(\tilde{\mathbf{y}}|\beta_0, \beta_1, \sigma) f(\beta_0, \beta_1, \sigma|\mathbf{y}) d\beta_0 d\beta_1 d\sigma$$

where $f(\tilde{y}|\beta_0, \beta_1, \sigma)$ is the likelihood function $N(\beta_0 + \beta_1 * \tilde{x}, \sigma^2)$ and $f(\beta_0, \beta_1, \sigma|y)$ is our posterior distribution.

 Instead of working with these complex integrals we will simulate heights using samples from the posterior.

- Here's how we will do it.
- For any unique weight value, we sample from a Gaussian distribution with the correct mean μ for that weight, using the correct value of σ sampled from the same posterior distribution.
- We do this for every sample from the posterior and for every weight value of interest.
- We end up with a collection of simulated heights that embody the uncertainty in the posterior as well as the uncertainty in the Gaussian likelihood.
- We first need to implement a function that can generate a sample of heights for a given weight w.
- The sampling should use the likelihood function $N(\beta_0 + \beta_1 * w, \sigma^2)$, using different values of β_0, β_1 and σ^2 given by our posterior samples.

```
height.weight <- function(weight)
  rnorm(
   n=nrow(post) ,
  mean=post$b0 + post$b1*weight ,
  sd=post$sigma )</pre>
```

- Notice that we are passing vectors of parameters $\vec{\mu}$ and $\vec{\sigma}$ to **rnorm** instead of scalars.
- This will result in different samples generated with different parameter values:

```
> rnorm(4,mean=c(-100,100),sd=c(1,50))
[1] -99.32946 126.17836 -100.62118 194.68311
```

 Now, we can apply this function to our sequence of weights and obtain a similar matrix than before:

```
> sim.height <- sapply( weight.seq , height.weight)
> dim(sim.height)
[1] 10000 46
```

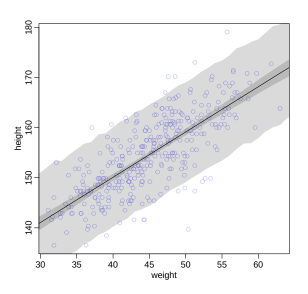
ullet This matrix is much like the earlier one, but it contains simulated heights, not distributions of plausible average height, μ .

• We can summarize these simulated heights in the same way we summarized the distributions of μ , by using apply:

```
height.HPDI <- apply( sim.height , 2 , HPDI , prob=0.95 )
```

- Now height.HPDI contains the 95% HPDIs of observable (according to the model) heights, across the values of weight in weight.seq.
- Let's plot everything we've built up: (1) the MAP line, (2) the shaded region of 95% plausible μ , and (3) the boundaries of the simulated heights the model expects.

```
# plot raw data
plot( height ~ weight , d2 , col=col.alpha(rangi2,0.5) )
# draw MAP line
lines( weight.seq , mu.mean )
# draw HPDI region for line
shade( mu.HPDI , weight.seq )
# draw HPDI region for simulated heights
shade( height.HPDI , weight.seq )
```



- The two shaded regions of the figure show different 95% plausible regions.
- The narrow shaded interval around the line is the distribution of μ .
- The wider shaded region represents the region within which the model expects to find 95% of actual heights in the population, at each weight.
- In the procedure above, we encountered both uncertainty in parameter values: β_0, β_1, σ , and uncertainty in a sampling process: $N(\mu_i, \sigma)$.
- These are distinct concepts, even though they are processed much the same way and end up blended together in the posterior predictive simulation.
- Finally, the sim function of the rethinking package allows to easily generate samples of the posterior predictive distribution of any linear model declared with quap:

```
sim.height <- sim( b.reg1 , data=list(weight=weight.seq) )</pre>
```

Conclusions

- In this class we have revisited the linear regression model using a Bayesian approach.
- The posterior distribution can be interpreted as a distribution of possible lines.
- We introduced Laplace approximation to fit these models to data.
- We also introduced new procedures for visualizing posterior distributions and posterior predictions
- In the next classes we will see more complex linear models, such as models with multiple attributes, binary attributes or interactions.
- Notice that these models can also be built with quap.
- However, a more sophisticated technique called Markov Chain Monte Carlo will allow us to sample from the posterior without making any assumptions about the posterior's shape.

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