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Introduction

This book will focus on a conceptual introduction to multilevel Bayesian models. I am going to talk about math as little as possible, so I am going to avoid giving details unless necessary. My goal is to convey some of the more important and useful concepts necessary to understand these models and to use them in your own work. I hope to set you up in a position to 'fill in the blanks' as necessary later.

I'm going to try to provide intuitive explanations for statistical models that rely on understanding the figures we use represent and interpret our data and models. The book assumes some understanding of R (only to understand the code), but none of statistics. I don't really 'explain' many basic things, for example, how to calculate a standard deviation? Why is it calculated like it is? The reason for this is that first, there are many better resources for that. And second, I am trying to make this quick, to go from 0 to understanding and interpreting Bayesian multilevel models in 10 weeks.

The book is specifically aimed for linguists only because it focuses on the sorts of research designs frequently used by linguists. However, the models and principles outlined in this book are used and useful in many domains.

If you have any comments or suggestions please let me know, and if you find any errors definitely let me know!

Inspecting a single sample of values

In this chapter I am going to present an introduction to some fundamental statistical concepts (i.e, probability, likelihood). We will discuss how to use these concepts to make inferences about our observations.

Data and research questions

Voice fundamental frequency (f0, related to pitch) is a very important cue in speech communication. It relates to phoneme identification, prosody, and to the communication of social and indexical information (speaker gender, age, . . . etc.).

We are going use a well-known data set, the Hillenbrand et al. (1995) data of Michigan English. We are going to focus on a single vector (f0), representing the f0 produced by a set of female speakers.

```
url1 = "https://raw.githubusercontent.com/santiagobarreda"
url2 = "/stats-class/master/data/h95_vowel_data.csv"
## read data from my Github page
h95 = read.csv (url(paste0 (url1, url2)))
# select the 'f0' vector, for women only (speaker type = 'w')
f0s = h95[['f0']][h95$type == 'w']
```

These speakers represent a sample from a larger population. The sample is a finite set of observations that you actually have. The population is the larger group of possible observations that you are *actually* interested in. For example, Hillenbrand et al. collected this data not to study these speakers in particular, but instead to make inferences about Michigan speakers more generally.

Similarly, we want to answer a few basic questions about the population of female speakers from Michigan, not about the sample itself:

- 1) What is the average f0 of the whole population likely to be?
- 2) Can we set bounds on likely mean f0 values based on the data we collected?

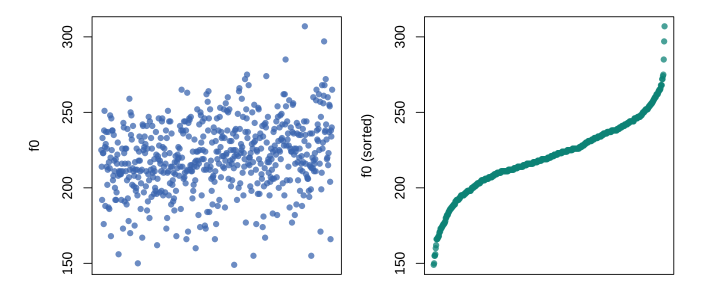
The second point is crucial. First, our sample will never exactly match the population. But it should be *representative* of it, meaning it should not be *too* far off from the real mean (otherwise, it is likely not a good sample from that population). For example, the mean f0 in the data we will discuss below is 220 Hz. This seems to suggest that, for example, a *true* mean of 100 Hz is unlikely. Is a true mean of 150 Hz also unlikely? What about 190 Hz?

Below we will discuss how to measure the spread and location of a set of numbers, and how to establish likely and credible values for the variable.

Inspecting the central location and spread of values

On the left, values are plotted according to the order they appear in the vector. This is not very useful. On the right, the same values have been ordered from low to high.

```
par (mfrow = c(1,2), mar = c(1,4,1,1))
plot (f0s, xaxt='n',xlab='', ylab = 'f0', pch = 16, col = skyblue)
plot (sort (f0s), xaxt='n',xlab='',ylab='f0 (sorted)', pch=16, col = deepgreen)
```



We can easily find descriptive statistics like the sample mean (\bar{x}) , the sample standard deviation (s_x) , and important quantiles for this sample of values. The quantiles below correspond to the values of ordered observations, like in the right plot above. The 0% quantile is the smallest observation, while 100% is the highest. Any other quantile is found by ordering the observations and selecting the observation that is higher than x% of the sample values. For example, the 50% quantile (the median) is higher than 50% of values, and the 25% quantile is higher than 1/4 of the values in the sample.

```
## calculate basic descriptive statistics
mean (f0s)

## [1] 220.401

sd (f0s)

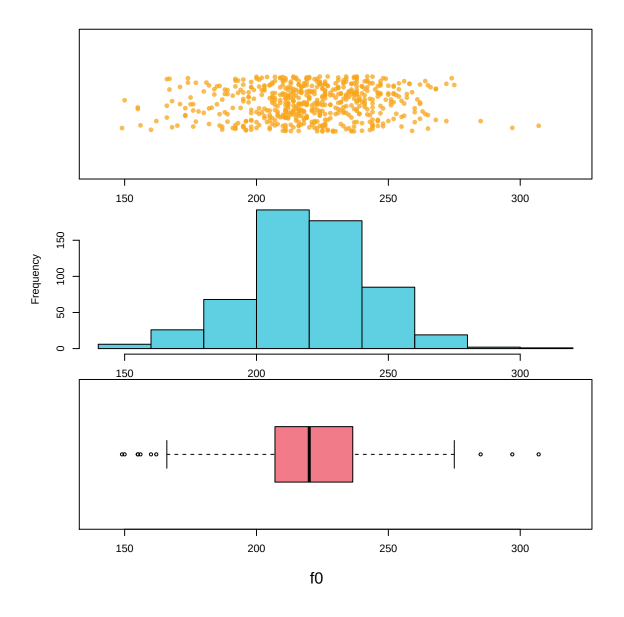
## [1] 23.22069

quantile (f0s)

## 0% 25% 50% 75% 100%

## 149.00 207.00 220.00 236.25 307.00
```

We can look at the distribution of speakers. In the top row, points indicate individual productions, and are jittered along the y axis to make them easier to see. In the middle row we see a histogram of the same data. The histogram gives you the count of observations in each bin. In the bottom row we see a box plot of the same data. The edges of the box correspond to the 25 and 75% quantiles of the distribution, and the line in the middle of it corresponds to the median. As a result, 50% of observations are contained in the box.



Probability Distributions

Histograms are particularly useful to understand because of how they relate to probability distributions. For our purposes, the probability is the number of times an event is expected to occur, out of all the other observed events and outcomes. This can also be thought of as the *percent* of times an event is expected to occur.

The total probability of all events is always equal to 1. This is like using 100 to communicate percent, its just easier that way. As a result of this convention, you know that a probability of 0.5 means something is expected to occur half the time (i.e., on 50% of trials). For example, suppose we want to know the probability of being an adult female in our sample who produces an f0 under 175 Hz. Finding the probability of observing this event is easy:

```
# the evaluation in the parenthesis will return 1 if true, 0 if false sum (f0s < 175) ## number of observations the fall below threshold
```

```
sum (f0s < 175) / length (f0s) ## divided by total number of events
```

[1] 0.03819444

```
mean (f0s < 175) ## a shortcut to calculate probability, mean = total/length
```

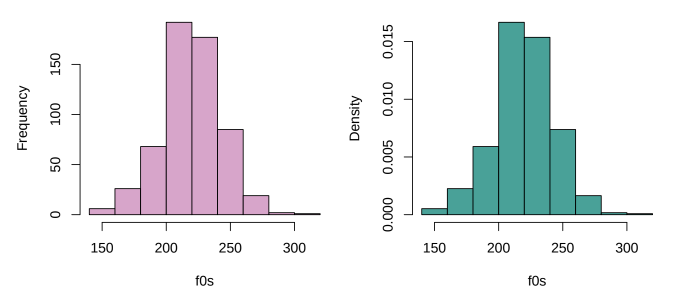
[1] 0.03819444

The top value is the frequency of the occurrence. This is not so useful because this number can mean different things given different sample sizes (e.g., 22/23, 22/10000). The middle and bottom values have been divided by the total number of observations. As a result, these now represent a proportion, or probability.

Histograms can also show this difference between total counts and probabilities. Below, the histogram on the left shows the number of observations in each bin. The histogram on the right shows *density* on the y axis. When you see *density* on the y axis, that means that y axis values have been scaled to make the area under the curve equal to 1. This has two benefits:

- 1) It lets you compare the distribution of values across different sample sizes.
- 2) It makes the histogram more comparable to a probability distribution.

```
par (mfrow = c(1,2), mar = c(4,4,1,1))
hist (f0s, main="", col = lavender)
hist (f0s, freq = FALSE, main = "", col = deepgreen)
```

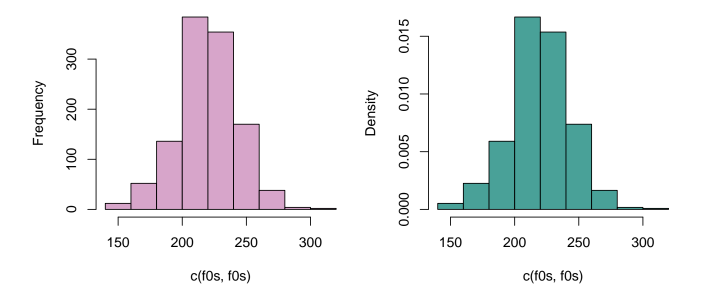


The density is just the thickness of the distribution at a certain location. In probability theory, the sum of the probabilities of all possible outcomes is 1, by definition. So, the fact that the area under the curve of a density is equal to 1 means that the density contains all your *stuff*, all the possible outcomes of the variable we are discussing.

Imagine a circle like in a Venn diagram that contains all possible productions of female f0. This circle has an area of 1 since it contains all possible instances of the variable. Imagine we spread out this circle along the x axis so that its shape reflected the relative frequencies of different values of the variable. For example, if some outcomes were 5 times more probable than others, the shape should be 5 times taller there, and so on. If we managed to do this, the height (or 'density') of this shape would exactly correspond to a probability distribution like that seen in the right plot above.

Below I've repeated the data, doubling the counts. Notice that the y axis in the right panel does not change. This is because increasing the number of observations changes your counts but not the relative frequencies of observations. For instance, increasing the number of coin flips will not change the fact that 50% will be heads, but it will change the number of heads observed.

```
par (mfrow = c(1,2), mar = c(4,4,1,1))
hist (c(f0s,f0s), breaks = 10, main = "", col = lavender)
hist (c(f0s,f0s), freq = FALSE, breaks = 10, main = "", col = deepgreen)
```



The normal distribution

The distribution of many variables (including f0) follows what is called a normal distribution. This means if you take a random sample of a variable and arrange observations into bins, they will tend to resemble the shape of a normal distribution. This distribution is also called a Gaussian distribution and has a familiar, bell-shaped curve.

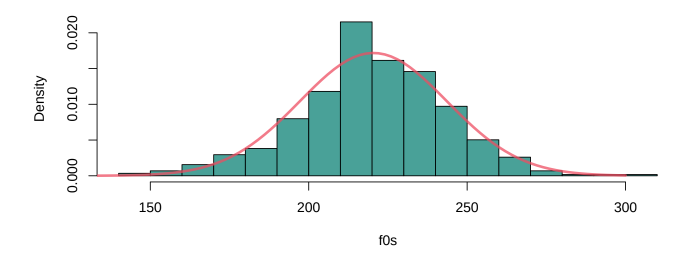
The normal distribution has the following important characteristics.

- 1. The distribution is approximately symmetrical i.e., producing a higher or lower than average f0 is about equally likely.
- 2. The probability of observing a given value decreases as you get further from the mean.
- 3. It is easy to work with, very well understood, and naturally arises in basically all domains.

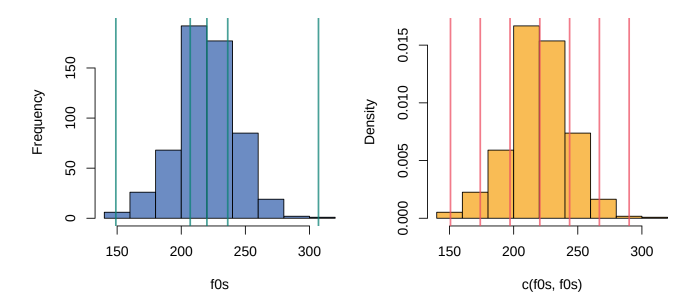
Normal distributions have two parameters. This means they vary from each other in only two ways. These parameters are:

- 1. A mean, μ , which determines where the distribution is located along the x axis. The mean is the 50% halfway point of the 'mass' of the distribution. If the distribution were an physical object, its mean would be its center of gravity.
- 2. A standard deviation, σ , that determines its *spread* along the x axis. Since every distribution has an area under the curve equal to one (they all have the same 'volume'), the smaller the variance the higher the peak of the density along the y axis must be.

Below, I compare the histogram of f0 values to the density of a normal distribution with a mean equal to our sample mean $(\mu = \bar{x})$ and a standard deviation equal to our sample standard deviation $(\sigma = s_x)$. The density was drawn using the dnorm function. This function will help draw a curve representing the shape of a theoretical normal distribution with a given mean and standard deviation.



When you are dealing with normally-distributed data, summary statistics can tell you a lot about the shape of your distribution, and about where you can expect the bulk of the density/distribution to lie. The left panel shows the locations of quantiles (0%, 25%, 50%, 75%, 100%), the right panel shows you the mean and standard deviations from the mean (-3, -2, 0, +1, +2, +3). Notice that ± 2 standard deviations enclose most of the distribution (around 95%), and ± 3 standard deviations enclose almost all of it (99%).



Referring to the normal distribution to make inferences

In general, it is impossible to know what the 'true' data distribution is, so that *perfect* inference is not possible. As a result, scientists often use theoretical probability distributions to make inferences about real-life populations and observations. Notice that our real life measurements follow the 'shape' predicted by the theoretical normal distribution. This suggests that we may be able to use the characteristics of an appropriate normal distribution to make inferences about female f0 (and other variables).

Using a normal distribution to make inferences about your data is like using a mathematical model for spheres to understand the behavior of billiard balls. In reality the balls are not perfect spheres. However, their shapes will be spherical enough to allow us to make useful predictions based on the simplified model. In general, it is useful to keep in mind that reality will never exactly conform to our model. This can result in unpredictable errors in our conclusions, which can cause errors to occur. In general, the things you don't know you don't know are the things that will cause the most problems.

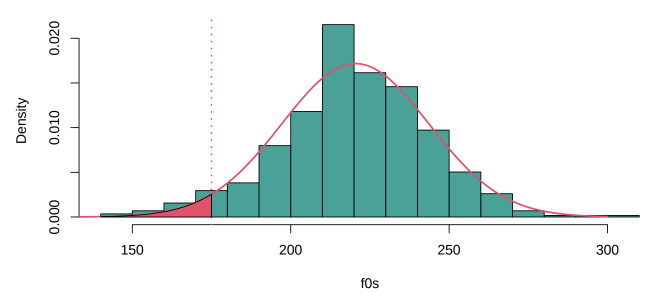
So, since we expect the distribution of f0 values to have the shape of the normal distribution, we can use the shape of the normal distribution to make inferences about the distribution of f0 values, even the ones we did not observe. For example, we can use the theoretical normal density to estimate the probability of observing a female production with an f0 of under 175 Hz, from among *all* possible observable productions of f0 in this *population*.

We do this by referring to the proportion of values expected to be less 175 Hz in the normal distribution. This can be found by finding the area under the curve of the probability density to the left of that point (the red area below). Since the *total* area is always equal to 1, the area of the red portion below corresponds to a percentage/probability.

Below, I use the function pnorm to find the proportion of values that are expected to be greater/less than 175 Hz. I use the parameters estimated form our sample to run the pnormfunction, as these are our best guesses of the population parameters. As we can see, this value is reasonably close to our empirical proportion, which was 0.038 (3.8%).

```
par (mfrow = c(1,1), mar = c(4,4,1,1))
hist (f0s, freq = FALSE, main = "", breaks = 20, col = deepgreen)
abline (v = 175, lwd = 2, col = 2, lty=3)
## plots the normal density (red line) using stats calculated form our sample.
curve (dnorm (x, mean(f0s), sd(f0s)),from=100, to=300, lwd=2, col = 2, add=TRUE)

x = c(140,seq(140,175,length.out = 100),175)
y = c(0,dnorm(seq(140,175,length.out = 100), mean (f0s), sd (f0s)),0)
polygon(x, y, col='2')
abline (v = 63.8, lwd = 2, col = 2, lty=3); abline (v = 70, lwd = 2,col=1,lty=3)
```



```
## probability of observing a production below 175 Hz
pnorm (175, mean (f0s), sd(f0s))
```

[1] 0.02527988

```
## probability of observing a production greater than 175 Hz
1 - pnorm (175, mean (f0s), sd(f0s))
```

[1] 0.9747201

Imagine you had 1 pound of clay and I asked you to make a shape **exactly** like the normal density (red curve) above with a constant depth. The 'area under the curve' would just correspond to the amount of clay in a certain area. So, if you made the density just right and I took a knife and cut the shape left of 175 Hz (the red part) and we weighed it, it should weigh 2.5% of a pound. So, the area under the curve, the probability, is just the amount of the *stuff* in the density that falls below/above a certain point, or between two points.

Since the total number of observations is always one. This helps us compare across many different actual numbers of observations. The probability above suggests the following:

```
## probability of observing a production with an f0 under 175 Hz
pnorm (175, mean (f0s), sd(f0s))

## [1] 0.02527988

## expected count
pnorm (175, mean (f0s), sd(f0s)) * length (f0s)

## [1] 14.56121

## actual count
sum (f0s < 175)</pre>
```

```
## [1] 22
```

We can also use this theoretical distribution to think about other possible outcomes:

```
min (f0s)

## [1] 149

pnorm (149, mean (f0s), sd(f0s)) # probability of observing our smallest value

## [1] 0.001052907

pnorm (140, mean (f0s), sd(f0s)) # probability of observing a smaller value

## [1] 0.0002676171

## predicted number of tokens below 175 Hz if we was had 5500 observations
pnorm (175, mean (f0s), sd(f0s)) * 5500
```

Probabilities of events and likelihoods of parameters

[1] 139.0394

We are going to switch from talking about *probabilities* to talking about *likelihoods*. A probability is the odds of observing some event/outcome, given some parameter(s). A likelihood is the odds of observing a parameter given some observed data. In both cases, these terms assume that you are referring to some probability distribution.

Every parameter for every probability distribution has a likelihood function, given some data. I am only going to talk about the likelihood of the normal mean parameter, μ , in detail.

The *likelihood function* is a curve showing the relative likelihoods of different parameter values, given a fixed set of data. The likelihood function tells you what values are *believable* given your data. If a value is very unlikely, that means that it is not supported by your data. In other words, unlikely parameter estimates represent conclusions that your data is rejecting as not viable.

Here are three useful properties of the likelihood functions of μ , the mean parameter of the normal distribution:

- 1. The likelihood function of μ will tend to be a normal distribution.
- 2. The mean (and peak) of the likelihood function of μ given some sample x is equal to the arithmetic mean of the sample (\bar{x}) .
- 3. The standard deviation of the likelihood of μ is equal to the standard deviation of the data (s_x) , divided by the square root of N (the sample size).

The first point tells us that we can use the normal distribution to make inferences about likely, and unlikely values for means, given some data.

The second point says that if you are wondering what the best (most likely) estimate of μ is given your sample, the answer is the arithmetic mean of your sample (\bar{x}) .

The third point means that the likelihood function for μ will tend to be much narrower than the distribution of our original data. This is because a mean based on, for example, 50 samples will contains many positive and negative deviations from the average that will tend to cancel out. As a result, the more data you have the more precise your estimates are, and the less uncertainty is associated with any estimate.

The left panel below shows the likelihood function for μ based on the first 10 observations of our f0 vector, shown by the blue points at the bottom of the plot. I chose this small sample just to make this example clearer. Notice

that the most likely mean values for these points like over the bulk of the sampled values. The vertical dotted lines show three possible mean values that will be highlighted.

The likelihood of any parameter estimate (e.g., $\mu = 175$ Hz in the right panel below) is equal to the product of the density of each observation in the sample, if we assume that the estimate were true. For example, to calculate the likelihood that μ , we:

- 1) assume that the data is generated by a normal distribution with a μ equal to x, and σ equal to the sample standard deviation (s_x) .
- 2) Find the height of the curve of the probability distribution (the density) over each point (indicated by lines in the right panel below).
- 3) The likelihood is the product of all of these densities. In practice, the logarithms of the individual probabilities are added together, yielding the *log-likelihood*. This is because multiplying together too many fractions can lead to numbers so small computers have a hard time representing them.

Imagine I follow the steps above for each position along the x axis, recording the likelihood values I calculate. I then plot the product of the densities for each x value: I have just plotted a likelihood function for μ given our data.

```
x = f0s[1:10] ## tiny sub sample for exampls
mean(x) ## sample mean
```

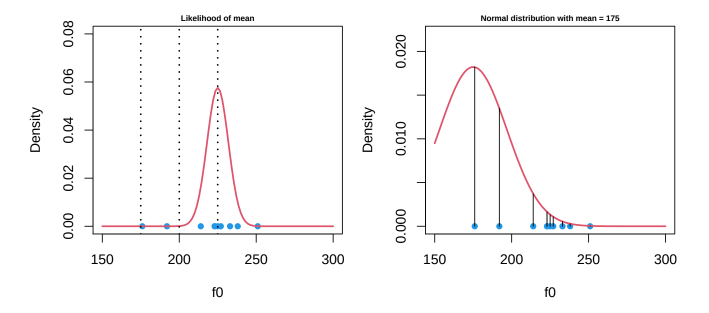
[1] 220.2

```
sd (x) ## sample standard deviation
```

```
## [1] 21.86219
```

```
par (mfrow = c(1,2), mar = c(4,4,1,1))
plot (x,rep(0,10), ylim = c(0,.08), pch=16,col=4, xlim = c(150,300),
    ylab='Density', main = 'Likelihood of mean',xlab='f0',cex.main=.6)
## here the likelihood sd is divided by the sample size
curve (dnorm (x, mean(x), 21.9 / sqrt (10)), from = c(150,300),
    add=TRUE, col = 2, lwd = 2)
abline (v = c(175, 200, 225), lwd=2,lty=3)

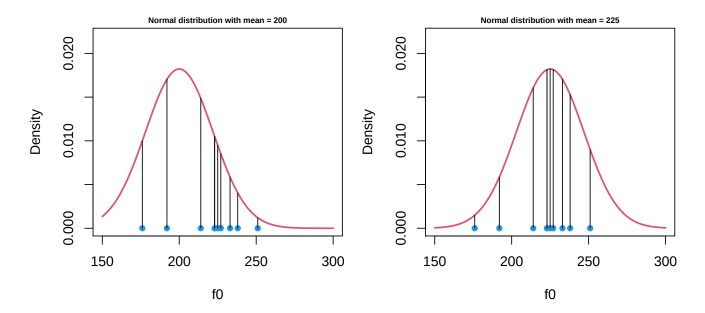
plot (x,rep(0,10), ylim = c(0,.022), pch=16,col=4,xlim = c(150,300),cex.main=.6,
    ylab='Density', main = "Normal distribution with mean = 175",xlab='f0')
## now it is centered at mean = 175 Hz
curve (dnorm (x, 175, 21.9), from = c(150,300),
    add=TRUE, col = 2, lwd = 2)
segments (x,rep(0,10),x,dnorm (x, 175, sd (x)))
```



In the right panel above we see that a normal distribution with a μ of 175 Hz is very unlikely to generate this data. Many points are extremely improbable and have densities close to zero. As a result, the product of these values (the heights of the lines) will be a very small number. This is reflected in the extremely small values in the likelihood function at 175 Hz in the left panel above.

In the left panel below, we see that a normal distribution with a μ of 200 Hz is more likely to generate this data, and the probability distribution is clearly a much better fit. However a distribution with a mean of 200 Hz is still not very likely to have generated this data.

Finally, in the right panel below we see the the maximum likelihood estimate of 225 Hz, the value representing the peak of the likelihood function (in the left panel above). When we say that 225 Hz is the most likely mean for this data, we are saying that this data is most probably the outcome of a normal distribution centered at 225 Hz, relative to the alternatives.



Making inferences using likelihoods

Previously, I mentioned using the normal distribution to make inferences. When variables are normally distributed we can use the theoretical normal distribution and functions such as pnorm to answer questions about values we expect, and don't expect, to see.

We can take this same approach to make inferences about *parameters* when their likelihood functions follow a normal distribution. For example, we can use the results of the calculations below:

```
mean (f0s) ## sample mean

## [1] 220.401

sd (f0s) ## sample standard deviation

## [1] 23.22069

length (f0s) ## sample size

## [1] 576

sd (f0s) / sqrt ( length (f0s) ) ## the standard deviation of the likelihood function

## [1] 0.9675289
```

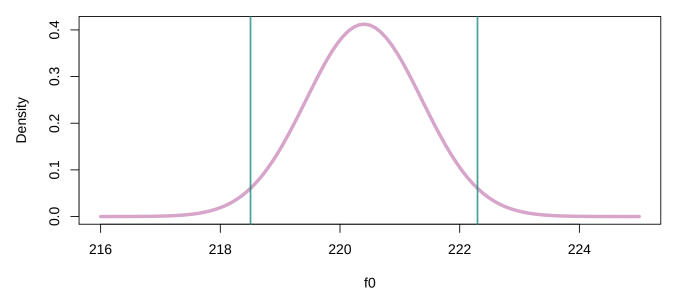
To draw the expected likelihood function for μ given our data and our model. You may be thinking, what model? It may seem too simple to be a model, but by assuming that our data can be understood as coming from a normal distribution with some given μ and σ , we have already created a simple model for our data. I'll return to this below.

We can take our model and our parameter estimates and draw the likelihood function for μ . We can then use the qnorm function to calculate quantiles for our likelihood, presented below. I added vertical lines at the 2.5% and 97.5% quantiles of our distribution. These vertical lines enclose 95% of the likelihood density, and so represent the range of values representing the 95% most likely values of μ . I chose an interval enclosing 95% of the likelihood because this is used by convention. This is a commonly-used interval but otherwise has no special significance.

```
par (mfrow =c(1,1), mar = c(4,4,1,1))
curve (dnorm (x, mean(f0s), sd(f0s)/sqrt(length(f0s))), xlim = c(216,225),
         ylab = 'Density', xlab = 'f0', col = lavender, lwd = 4)
quantiles = qnorm (c(0.025, 0.975), mean (f0s), sd (f0s) / sqrt (length (f0s) ))
quantiles
```

```
## [1] 218.5047 222.2974
```

```
abline (v = quantiles, lwd=2,col=deepgreen)
```



The likelihood tells you about the most believable/credible parameter values, given your model and data. Given the information presented in the figure above, we may conclude that the most likely parameter values fall between 218 and 222 Hz. This means that it is reasonable that the true value might be 221 Hz, as this value is very likely given our sample. Basically, maybe our sample mean is wrong and arose by accident, and 221 Hz is the true μ . This outcome is compatible with our data.

However, a value of 216 Hz is extremely *unlikely* to fit our data. It is just too far from our sample mean relative to the amount of variation in our sample. This is like if you measured the heights of 100 women in a small town (pop. 1500) and found the average was 5'4". You might accept that the actual population average is 5'5", but may find it difficult to accept that it was actually 6'0". It would mean you happened to measure all of the shortest women in the town, an extremely unlikely event.

So, since we think that 216 Hz is not a plausible mean f0 given our sample, this also means that it is very unlikely that the real μ is 216 Hz. This is because a distribution centered at 216 would be extremely unlikely to generate a sample mean of 220 Hz. Using this approach, we can rule out implausible values of μ based on the characteristics of our data.

At this point we can offer conventional responses to the research questions posed at the start of the Chapter:

- Q1) What is the average f0 of the whole population likely to be?
- A1) The most likely value for the population mean is our sample mean, 220.4 Hz.
- Q2) Can we set bounds on likely mean f0 values based on the data we collected?
- A2) Yes, there is a 95% probability that the population mean is between 218.5 222.3 Hz, given our data and model structure.

Traditional approaches to statistics (sometimes generally referred to as 'frequentist') estimate parameters by trying to find the most likely values for parameters (i.e., 'maximum likelihood estimation'). They do this by referring to

the theoretical likelihood functions such as what we plotted above. Although this works very well for simple data, it is difficult if not impossible for some of the more complicated datasets that often arise for even the simplest research questions in linguistics.

Bayesian models

In this class we are going to learn about multilevel Bayesian models. These models have many advantages over 'traditional' approaches. They provide researchers with more information, are more robust, and at worst, they are as good as traditional models. I may sound biased, but the main reason for all of these advantages is that traditional models were developed over 100 years ago. On the other hand, mathematical and technological advances have only made Bayesian multilevel models possible in the last 10+ years. It is only reasonable that the newer approaches should offer some advantages over methods developed before calculators existed.

Here, I am going to address what is meant by two aspects of the term 'Bayesian multilevel models': 'Bayesian' and 'models'.

What are regression models?

I have been referring somewhat obliquely to 'models' without really explaining what I mean by this. It's difficult to offer a precise definition because the term is so broad, but 'regression' modeling can be thought of as trying to understand variation the mean parameter (μ) of a normal distributions. Actually, you can use many other probability distributions, but for now we will focus on models based on the normal distribution.

Basically it goes like this:

- you assume that your data is well described by a normal probability distribution. This is a mathematical function $(\mathcal{N}(\mu, \sigma))$ that described what is and is not probable based on two parameters.
- the mean of this distribution is either fixed, or varies in a logical manner.
- the variation in the mean of this distribution can be understood using some other variables.

We can write this model more formally like:

$$y \sim \mathcal{N}(\mu, \sigma) \tag{1}$$

This says that we expect that the variable we are interested in is distributed according to (\sim) a normal distribution with those parameters. Basically, this just formalizes the fact that we think the *shape* of our data will be like that of a normal distribution with a mean equal to μ and a standard deviation equal to σ .

When you see this, $\mathcal{N}(\mu, \sigma)$, just picture in your mind the shape of a normal distribution, like if you see this $y = x^2$ you may imagine a parabola. $\mathcal{N}(\mu, \sigma)$ Really just represents that shape of the normal distribution, and the associated expectation about more and less probable outcomes.

The above relationship can also be presented like this:

$$y = \mu + \mathcal{N}(0, \sigma)(\#eq: 2)$$

Notice that we got rid of the \sim symbol, moved μ out of the distribution function ($\mathcal{N}()$), and that the mean of the distribution function is now 0. This breaks up our variable into two components:

- 1) A systematic component, μ , that contributes the same value to all instances of a variable.
- 2) A random component, $\mathcal{N}(0,\sigma)$, that causes unpredictable variation around μ .

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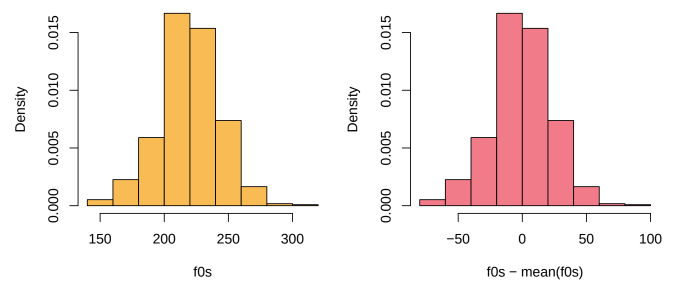
In terms of our data, I might express the distribution in either of the following ways:

$$f0 = \mathcal{N}(220.4, 23.2)(\#eq:3)$$

$$f0 = 220.4 + \mathcal{N}(0, 23.2)(\#eq:4)$$

The distribution on the left below is the original data, centered at 220.4 Hz and with a standard deviation of 23.2 Hz. On the right, the mean has been subtracted from each value. The sample now represents random variation around the sample mean, variation that our model can't explain. From the perspective of our model, this is noise, or error. This doesn't mean that it's unexplainable, it only means that we've structured our model in a way that doesn't let us explain it.

```
par (mfrow =c(1,2), mar = c(4,4,1,1))
hist (f0s, main="", freq=FALSE, col = yellow)
hist (f0s - mean (f0s), main="", freq=FALSE, col = coral)
```



In regression models, we can decompose systematic variation in μ into component parts, based on i predictor variables. The x_i . These x variables the co-vary (vary with) our y variable, and that we think help explain the variation in y. Below, I am saying that I think μ is actually equal to some combination sum of x_1 x_2 and x_3 . For example, I could think that f0 could be affected by vowel category (x_1) , the height of the speaker (x_2) , and whether the utterance is a sentence or a question (x_3) .

$$\mu = x_1 + x_2 + x_3 (\#eq : 5)$$

Actually, the mean is very unlikely to just be an equal combination of the predictors, so that a weighting of the predictors will be necessary. For example, maybe x_1 is twice as important as the other two predictors and so a is 2, while b and c are 1.

$$\mu = a * x_1 + b * x_2 + c * x_3 (\#eq : 6)$$

Decomposition of μ into sub-components makes our model something more like:

$$y = \mu + \mathcal{N}(0, \sigma)(\#eq: 7)$$

$$y = (a * x_1 + b * x_2 + c * x_3) + \mathcal{N}(0, \sigma)(\#eq : 8)$$

Often, ε is used to represent the random component, as in:

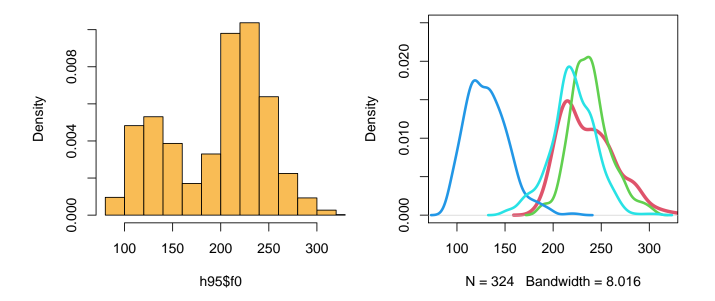
$$y = a * x_1 + b * x_2 + c * x_3 + \varepsilon (\#eq : 9)$$

When expressed in this manner, this is now a 'regression equation' or a 'regression model'. 'Fitting' a regression model basically consists of trying to guess the most likely values of a, b, and c given our data.

Notice that the above formulation means that regression models do not require that our data be normally distributed, but only that the $random\ variation$ in our data (ε) be normally distributed. For example, in the left panel below I plot the distribution of f0 from among the entire Hillenbrand et al. data, including boys, girls, men and women. The data is not normally distributed, however, we can still use a regression based on normally-distributed data to model this as long as we expect that:

- 1) There is systematic variation in the μ of f0 across different groups, speakers, conditions, etc.
- 2) The random variation around these predicted values of μ more or less follows a normal distribution.

In the right panel I plot the individual densities of boys (red), girls (green), men (blue) and women (cyan). We see that although the data is not normally distributed, the within-group variation is. This suggests a regression model is appropriate for this data.



What's 'Bayesian' about these models?

The major difference between Bayesian and traditional models is that Bayesian models rely on *posterior distributions* rather than likelihood functions. I am going to define some terms:

- prior probability distribution: the distribution of possible/believable parameter values **prior** to the *current* experiment. This *a priori* expectation can come from world knowledge, previous experiments, or some combination of the two.
- the likelihood: this is the distribution of possible/credible parameter values given the **current** data and probability model, and nothing else.
- posterior probability distribution: the distribution of possible/believable parameter values you have **after** your current experiment. You get this by combining the prior distribution and the likelihood.

Traditional models make inferences based on the likelihood functions of parameters. Bayesian models make inferences based on the posterior distributions of parameters. To do this, they have to have to actually combine information about likelihood with information about the prior probabilities of parameters.

Posterior distributions

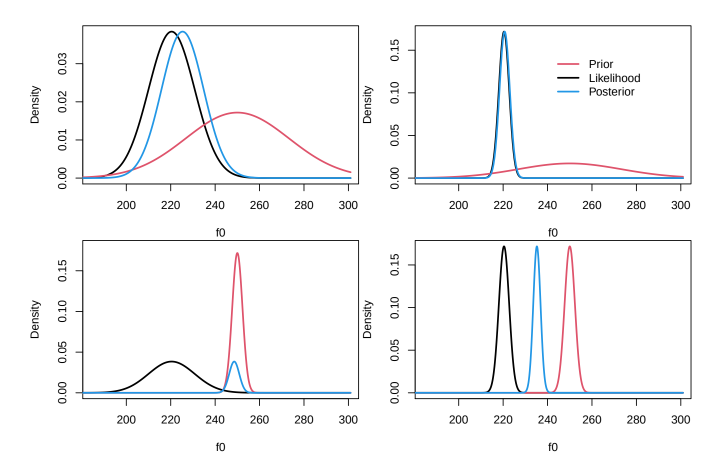
The combination of probability distributions is straightforward conceptually: you just multiply the values of the distributions at each x-axis location, and the result is the new curve. In the figure below (code at end of chapter), I combine several sets of probability distributions, showing the effects of variations in priors and likelihoods. In each plot, I scale the posterior density so that it is the same height as the likelihood. This is only to make the figures interpretable but does not affect any of the points I make below.

In the top-left panel, I plot the likelihood function for μ given a sample of size 5 with a mean of 220 Hz. I show what happens when I combine this with a relatively weak but very different prior: the standard deviation is the same as our f0 data, however the mean is much higher (250 Hz). With only 5 data points the likelihood already dominates the posterior, though the prior distribution is exerting a pull.

In the top-right panel, the posterior is almost identical to the likelihood. The likelihood represents a sample of size 100, which is actually a tiny sample in experimental linguistics work where you may have 200+ samples from each of 50+ subjects. As you might imagine, when the sample size is that large the prior exerts almost no influence on results.

In the bottom-left panel we see a situation where the prior dominates the estimate. Consider a situation where we actually have really good reasons to think that the mean is 250 Hz. If we really *know* this, why would we accept and estimate of 220 Hz based on only 5 samples? In this case, the posterior distribution is basically saying: your estimate is great, but come back when you have more evidence and I might believe you.

In the bottom-right panel we see a situation where the likelihood and the prior are equal. In this case the posterior represents compromise between new and prior knowledge.



The use of prior probabilities is often said to make Bayesian models 'subjective' but its not really a big deal. First, every model involves arbitrary decisions which can substantially affect our results. Second, a researcher will always use common sense to interpret a model. For example, before collecting my sample I can say that I expect my female average f0 to be 200 Hz or so, but think its reasonable to expect anything from 100 to 300 Hz. Based on everything we know about human speech, even these bounds are too wide, and anything outside would suggest something is very wrong. So, even if I did not use a prior, I would use my expectations to 'screen' my results, and be very wary of anything that did not meet my expectations.

A Bayesian model simply requires that you build your expectation into your model. It formalizes it, makes it definable and replicable. Also, being 'objective' does not quite make sense in many cases. Is it really being objective to ignore common sense and act as if a mean f0 of 250 is exactly as likely a priori as one of 20,000 Hz? Because not using a prior is equivalent to using a 'flat' prior and acting like almost any value is equally likely a priori, when this is hardly the case.

Sampling from the posterior

We want to understand the posterior distribution of parameters. How do we get this information? It is difficult to get this 'analytically', that is, using exact methods and solving a bunch of equations. Many traditional methods can actually be solved in this way, and that is a big part of their popularity.

Understanding the characteristics of posterior probabilities is not possible analytically for many Bayesian models. As a result, these questions are answered 'numerically', basically by using a bunch of 'guesses'. To understand the properties of posterior distributions, we use 'sampling' software that knows how to investigate these distributions.

The way these samplers work is you specify a set of data and some relationships you think are represented in your data (i.e., a model). The sampler then 'walks around' the parameter space, which is the range of possible values a parameter (or set of parameters) can take. For example, for a single parameter the parameter space is a line (like the x axis in the plots above) along which the parameter varies.

The sampler then does some variant of the following algorithm:

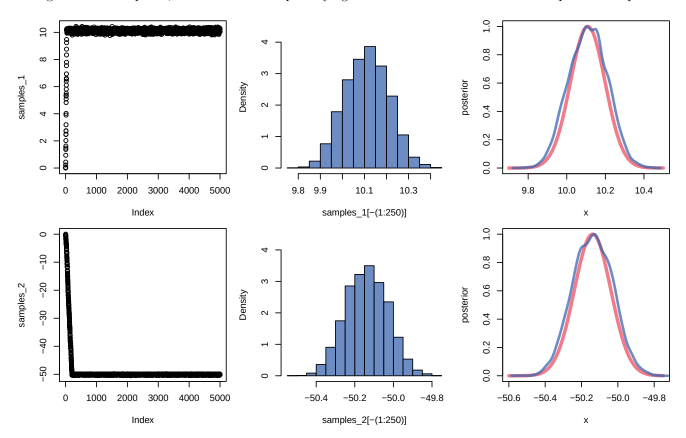
- 1) Pick a random value for the parameter (i.e., $\mu_{tmp} = 221 \text{ Hz}$).
- 2) Calculate the posterior probability for the current estimate of μ_{tmp} .
- 3) If the posterior estimate meets some criteria (e.g., it is better than the last one, it is not too low, etc.), then the value of μ_{tmp} is recorded, and becomes $\mu_{estimate}$. If not it is just discarded.
- 4) Go back to step 1.

As incredible as it may seem, under a very reasonable set of conditions if you do the above enough times, the distribution of $\mu_{estimate}$ that results from the above process will converge on the posterior distribution of μ given your data and model structure (including prior probabilities).

Below I have made a small example of this process. I use the Metropolis-Hastings algorithm, which is an algorithm to sample from probability distributions. The small example below assumes the standard deviation of the population is known, and just tries to investigate the posterior distribution of μ . It uses a very broad prior distribution ($\mu = 0$, $\sigma = 5000$) so that it will have a very weak effect on the outcomes.

```
# the function below takes a random sample, an initial mean estimate and a fixed
# standard deviation. It then takes a certain amount of samples from the
# posterior distribution of the parameter, assuming a broad prior centered at \it O
sampler_example = function (sample, mu_estimate = 0, stdev = 1, nsamples = 1000){
  # initial posterior calculation. This is the sum of the log likelihood and
  # the logarithm of the prior probability.
 prior = log (dnorm (mu_estimate[1],0, 500))
 loglik = sum (dnorm (sample, mu_estimate[1],stdev,log=TRUE))
  old_posterior = loglik + prior
 for (i in 2:nsamples){
    accept = FALSE
    ## this loop will keep proposing new steps until one gets accepted.
    while (!accept){
      ## (step 1 above)
      ## draw new proposal by randomly changing the previous mu_estimate
      mu_tmp = mu_estimate[i-1] + rnorm (1, 0, .3)
      ## (step 2 above)
      ## find prior probability for new mu_tmp proposal
      prior = log (dnorm (mu_tmp,0, 500))
      ## find log likelihood for new mu_tmp proposal
      loglik = sum (dnorm (sample, mu_tmp,stdev,log=TRUE))
      ## calculate the new posterior probability
     new_posterior = prior + loglik
      ## (step 3 above)
      ## if better accept always. If worse, accept sometimes
      if ( ( new_posterior - old_posterior ) >= log ( runif (1,0,1) ) ){
        mu_estimate[i] = mu_tmp
        ## if you accept, the new estimate becomes the current estimate
        old_posterior = new_posterior
        accept = TRUE
    }
 }
  return (mu_estimate)
```

In the plots below (code at end of chapter), you can see the algorithm begins at 0 (the initial guess) but is quickly able to find the most likely sample mean given the data (left column). In the middle, I show the distribution of the samples on the left, minus the burn-in phase (arbitrarily chosen by me). On the right, I compare our samples (blue) to the theoretical posterior distribution for the mean given the data and prior (red). I toss out the samples during the 'burn in' phase, as there are used up in trying to 'find' the correct location in the parameter space.



The results clearly coincide, but aren't perfect. But this sampler isn't very sophisticated! The samplers we will be using in this class do provide an excellent match to the posterior distribution. As a result, we can inspect the distribution of collected $\mu_{estimate}$ to understand the posterior of our parameter. We can use these distributions in the same way that we used the theoretical likelihood functions above, by using them to make statements about likely parameter values and ranges of values.

Plot Code

Plot showing combinations of different priors and likelihoods into posteriors::

PLOT CODE 23

```
## likelihood is much stronger than the prior
likelihood = dnorm (x, mean (f0s), sd (f0s) / sqrt (100))
prior = dnorm (x, 250, sd (f0s)); posterior = likelihood * prior
plot (x, likelihood / max (likelihood), type = 'l', ylab='Density', lwd=2,
      xlim = c(175, 300), ylim = c(0,1.1))
lines (x, prior / max (prior), lwd=2, col=2)
lines (x, posterior / max (posterior), lwd=2, col=4)
## prior overwhelms the likelihood
likelihood = dnorm (x, mean (f0s), sd (f0s) / sqrt (5))
prior = dnorm (x, 250, sd (f0s)/10); posterior = likelihood * prior
plot (x, likelihood / max (likelihood), type = 'l', ylab='Density',lwd=2,
      xlim = c(175, 300), ylim = c(0,1.1))
lines (x, prior / max (prior), lwd=2, col=2)
lines (x, posterior / max (posterior), lwd=2, col=4)
## prior and likelihood have about equal influence
likelihood = dnorm (x, mean (f0s), sd (f0s) / sqrt (100))
prior = dnorm (x, 250, sd (f0s)/10); posterior = likelihood * prior
plot (x, likelihood / max (likelihood), type = 'l', ylab='Density',lwd=2,
      xlim = c(175, 300), ylim = c(0,1.1)
lines (x, prior / max (prior), lwd=2, col=2)
lines (x, posterior / max (posterior), lwd=2, col=4)
```

Plot showing the output of the example sampler included above:

```
set.seed(1)
## make a random sample of data
data_1 = rnorm (100, 10, 1)
## collect samples from the likelihood of the mean
samples_1 = sampler_example (data_1,0,sd(data_1), 5000)
## do a second one to show its not a fluke
data_2 = rnorm (100, -50, 1)
## collect samples from the likelihood of the mean
samples_2 = sampler_example (data_2,0,sd(data_2), 5000)
# the left column shows the path the sampler took. the middle column shows the
# distribution of these samples, minus the burn in phase. the right column shows
# a comparison of theoretical and observed posterior distributions
par (mfrow = c(2,3), mar = c(4,4,1,1))
plot (samples_1)
hist (samples_1[-(1:250)], freq = FALSE, ylim = c(0,4.5), main="",col=4)
x = seq (9.7, 10.5, .001)
likelihood = dnorm (x, mean (data_1), sd(data_1) / sqrt (100) )
prior = dnorm (x, 0, 500)
posterior = likelihood * prior
posterior = posterior / max (posterior)
plot (x, posterior, lwd = 4, col = 2, type = 'l')
density_1 = density (samples_1[-(1:250)])
density_1$y = density_1$y / max (density_1$y)
lines (density_1, lwd = 3, col = 4)
plot (samples_2)
hist (samples_2[-(1:250)], freq = FALSE, ylim = c(0,4.5), main="",col=4)
x = seq (-50.6, -49.75, .001)
likelihood = dnorm (x, mean (data_2), sd(data_2) / sqrt (100) )
prior = dnorm (x, 0, 500)
```

```
posterior = likelihood * prior
posterior = posterior / max (posterior)
plot (x, posterior, lwd = 4, col = 2, type = 'l')
density_2 = density (samples_2[-(1:250)])
density_2$y = density_2$y / max (density_2$y)
lines (density_2, lwd = 3, col = 4)
```

Inference for a 'single group' of observations using a Bayesian multilevel model

In this chapter I am going to discuss how to use the brms package to estimate a population mean given a sample of data. For these models the data:

- can come from one speaker/subject or many speakers/subjects.
- each speaker/subject can contribute multiple data points.
- does not need to be 'balanced' or 'complete' across all subjects.

The 'traditional' designs equivalent to these models are: one-sample t-test, and repeated-measures one-way ANOVA with only two groups. However, these models won't be discussed in the chapter below.

Data and research questions

We are going to keep analyzing the female f0 data from the Hillenbrand et al. (1995) dataset.

```
url1 = "https://raw.githubusercontent.com/santiagobarreda"
url2 = "/stats-class/master/data/h95_vowel_data.csv"
h95 = read.csv (url(paste0 (url1, url2)))

# select women only
w = h95[h95$type == 'w',]
# this is unique subject numbers across all groups
w$uspeaker = factor (w$uspeaker)
# select only the vector of interest
f0s = w$f0
```

We are going to try to address the same questions we talked about last week:

- 1) What is the average f0 of the whole population likely to be?
- 2) Can we set bounds on likely mean f0 values based on the data we collected?

However, this time we are going to do this with a Bayesian multilevel model.

Estimating a single mean with the brms package

The brms Bayesian regression models package in R lets you fit Bayesian models using the STAN probabilistic programming language using R. The package is really amazing and makes Bayesian multilevel modeling easy and accessible for anyone. It also includes a lot of helper functions that making with these models very convenient.

brms should be installed in R so that the models described below will work. Make sure you have the latest version of R (and Rstudio) and the latest version of the 'brms' package installed. Sometimes using older versions can cause R to crash when fitting models.

The model formula

Model structures are expressed in R using a very specific syntax. Think of writing a model formula as writing a language within R. The good thing about learning to write models is then you can use this knowledge to describe your models in your work, and to interpret other people's models.

The model formulas resemble regression equations to some extent, but there are some differences. Remember that regression models can be thought of like this:

$$y = \mu + \varepsilon (\#eq : 21)$$

Which means that your observed variable y is the sum of some of some average value (μ) and some random error μ .

Actually, what we would really like is to understand orderly variation in μ by breaking it up into parts $(x_1, x_2, ...)$ when combined using some weights (a, b, ...).

$$y = a * x_1 + b * x_2 + ... + \varepsilon (\#eq : 22)$$

'Fitting' a regression model consists of trying to 'guess' the values of the weighing factors (a and b above), called the *model coefficients*. When we are only trying to estimate a single average, we don't have any predictors to explain variation in μ . In fact, our model structure suggests we expect no variation in μ .

Mathematically, we can't just say 'we have no predictor' since everything needs to be represented by a number. As a result, we use a 'predictor' with a value of 1 so that our regression equation is:

$$y = a * 1 + \varepsilon (\#eq : 23)$$

Now our model is trying to guess the value of a single parameter (a), and we expect this parameter to be equal to μ since it is being multiplied by a 'predictor' with a constant value of 1.

This kind of model is called an 'Intercept only' model. Regression models are really about representing differences, differences between groups and across conditions. When you are encoding differences, you need an overall reference point. For example, saying that something is 5 miles north is only interpretable given some reference point. The 'reference point' used by your model is called your 'Intercept'. Basically, our model consists only of a single reference point, the intercept a.

Generally, model formulas in R have the form:

y ~ predictor

The variable we are interested in understanding (y) goes on the left hand side, and on our predictors go on the right hand side, separated by a \sim . Notice that the random term (ε) is not included. The formula above can be read as 'y is distributed according to some predictor', which really means "we think there is systematic variation in our y variable that can be understood by considering its joint variation with our predictor variable(s).

For intercept only models, the number 1 is included in the model formula to indicate that a single constant value is being estimated. As a result, our model formula will have the form $f0 \sim 1$. This model could be said out loud like "we are trying to estimate the mean of f0" or "we are predicting mean f0 given only an intercept".

Calling the brm function

Below, I load the brms package, which contains the brm function. The brm function takes a model specification, data and some other information, and fits a model that estimates all the model parameters. Unless otherwise specified, brm assumes that the error component (ε) of my model is normally distributed. The first argument in the function call is the model formula, and the second argument tells the function where to find the data. The other argument tell the function to estimates a single set of samples (chains = 1) using a single processor on your CPU (cores = 1). These arguments will be discussed more later.

```
# To ensure predictable results in examples, I will be using the same random
# seed throughout, and resetting it before running any 'random' process.
set.seed (1)
model = brms::brm (f0 ~ 1, data = w, chains = 1, cores = 1)
## Compiling Stan program...
## Start sampling
##
## SAMPLING FOR MODEL '98dae0f1caaef07c210aac2156c73749' NOW (CHAIN 1).
## Chain 1:
## Chain 1: Gradient evaluation took 0 seconds
## Chain 1: 1000 transitions using 10 leapfrog steps per transition would take 0 seconds.
## Chain 1: Adjust your expectations accordingly!
## Chain 1:
## Chain 1:
## Chain 1: Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
## Chain 1: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 1: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 1: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 1: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 1: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 1: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 1: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 1: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 1: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 1: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 1: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 1:
## Chain 1: Elapsed Time: 0.06 seconds (Warm-up)
## Chain 1:
                           0.04 seconds (Sampling)
                           0.1 seconds (Total)
## Chain 1:
## Chain 1:
```

By default, brms takes 2000 samples, throwing out the first 1000 and returning the last 1000. The output above shows you that the sampler is working, and tells you about the progress as it works.

This is the last time I will be actually fitting a model in the code chunks. I am going to be relying on pre-fit models that you can load after downloading from the book GitHub. Models can be found in the folder corresponding to each chapter.

```
## load pre fit model
model = readRDS ('2_model.RDS')
```

Interpreting the model print statement

We can evaluate the model name to show the default brms model print statement:

model

```
##
   Family: gaussian
##
    Links: mu = identity; sigma = identity
## Formula: f0 ~ 1
##
      Data: w (Number of observations: 576)
## Samples: 1 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup samples = 1000
##
## Population-Level Effects:
##
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                           0.97
                                  218.33
                                            222.30 1.00
## Intercept
               220.40
##
## Family Specific Parameters:
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
            23.24
## sigma
                       0.69
                               21.99
                                         24.61 1.00
                                                         653
                                                                  550
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

Typing the model name into the console and hitting enter prints the information seen above. The first part just tells you technical details that we don't have to worry about for now (though some are obvious).

Next we see estimated effects for out predictors, in this case only an intercept. This is a 'population' level effect because is is shared by all observations in our sample, and not specific to any one observation.

```
Population-Level Effects:

Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
Intercept 220.40 0.97 218.33 222.30 1.00 851 557
```

The information above provides the mean (Estimate) and standard deviation (Est. Error) of the posterior distribution of μ (Intercept). The values of 1–95% CI and u–95% CI represent the upper and lower '95% credible intervals' for the posterior distribution of this parameter.

The x% credible interval for a parameter is the smallest interval that encloses x% of the distribution. This parameter has an x% chance (0.x probability) of falling inside the x% credible interval. So, this means that there is a 95% probability that μ is between 218 and 222 Hz given our data and model structure.

Notice that the parameter estimate and intervals almost exactly match the estimate and intervals we obtain by referencing the theoretical likelihood function (discussed in Chapter 1):

```
## sample mean
mean (f0s)
```

```
## [1] 220.401
```

```
## theoretical quantiles for likelihood of mean
qnorm (c(0.025, 0.975), mean (f0s), sd (f0s) / sqrt (length (f0s) )
```

```
## [1] 218.5047 222.2974
```

Our model also provides us an estimate of the error (ε), under 'Family Specific Parameters: sigma'. This estimate closely matches our sample standard deviation (s_x) estimate of 23.2. In addition, we also get a 95% credible interval for this parameter (2.5% = 21.99, 97.5% = 24.61).

Family Specific Parameters:

```
Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS sigma 23.24 0.69 21.99 24.61 1.00 653 550
```

This last section is just boilerplate and contains some basic reminders. This text will look the same after all models.

```
Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).
```

Seeing the samples

3

4

5

6

219.6445 23.22390 -2633.588

222.3286 21.93414 -2637.392

219.5626 22.87259 -2633.785

221.2339 23.08532 -2633.672

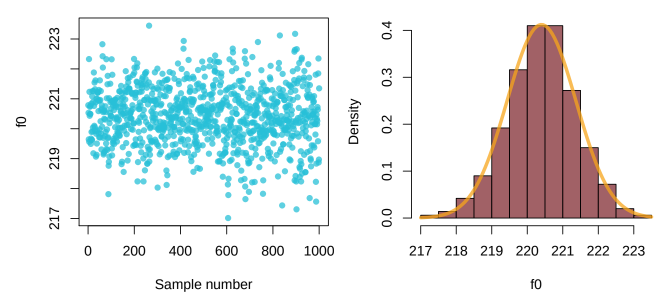
In Chapter 1 I discussed that samplers (like brm, or STAN) take samples of the posterior distributions of parameters given the data and model structure. It's helpful to see that this is quite literally what is happening, and that the print statement above just summarizes the information contained in the posterior samples.

Below I get the posterior samples from the model. We have 1000 samples, as indicated in the model output above. The first column represents the model intercept, the middle column is the error, and the third column is a statistic related to model fit.

```
## get posterior samples from model
samples = brms::posterior_samples (model)
str (samples)
## 'data.frame':
                    1000 obs. of 3 variables:
## $ b_Intercept: num
                        221 221 220 222 220 ...
   $ sigma
                        22.1 21.9 23.2 21.9 22.9 ...
                 : num
                        -2635 -2635 -2634 -2637 -2634 ...
  $ lp__
                 : num
## inspect values
head (samples)
##
     b_Intercept
                    sigma
                                lp__
## 1
        220.7860 22.06961 -2634.851
## 2
        220.5312 21.93719 -2635.160
```

I can plot the individual samples for the mean parameter on the left below. On the right I plot a histogram of the same samples, superimposed with the theoretical distribution of the likelihood. Although this is not the posterior, with so many data points we expect our posterior to be dominated by the likelihood anyways.

```
par (mfrow = c(1,2), mar = c(4,4,1,1))
plot (samples[,1], xlab = 'Sample number',ylab = 'f0',col=teal,pch=16)
hist (samples[,1], freq = FALSE, breaks = 20,main='',xlab='f0',col=maroon)
curve (dnorm (x, mean (f0s), sd (f0s) / sqrt (length (f0s) )), add = TRUE,
    lwd = 4, col = yellow)
```



Recall that our model output provides information about expected values for the mean parameter:

Population-Level Effects:

```
Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS Intercept 220.40 0.97 218.33 222.30 1.00 851 557
```

These simply correspond to the quantiles of the posterior samples!

```
quantile (samples[,1], c(.025, .5, .975))
```

```
## 2.5% 50% 97.5%
## 218.3288 220.3997 222.3001
```

There is no special status for these quantiles. We can check the values of other ones:

```
quantile (samples[,1], c(.25, .75))
```

```
## 25% 75%
## 219.8043 221.0296
```

Or even use the posterior distribution to find the probability that the mean parameter is over/under any arbitrary value:

```
mean (samples[,1] < 221)
```

```
## [1] 0.74
```

For example, given the calculation above we can say that there is a 0.74 probability (a 74% chance) that the mean f0 for female speakers in this population is under 221 Hz, given our data and model structure. We come to this conclusion by finding that 74% of the posterior samples of the parameter of interest are below 221 Hz.

Repeated measures data

The model we fit above is a reasonable starting point, but it has many weaknesses. For example, it does not consider the fact that our data was produced by a fixed number of speakers, sampled from a population. It does not consider the variation in f0 inherent between speakers, treating this as 'noise'.

Importantly, our data consists of 12 productions from each speaker in our sample, meaning we have 'repeated measures' data. Treating repeated measures data as if it were *not* repeated measures data can cause problems for our inferences. This is because it can give us a warped perspective of how much variability there really is in the sample.

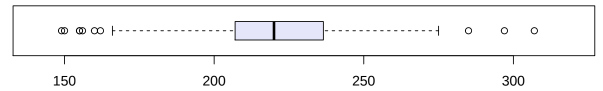
For example, if I told you I had 1,000,000 samples of speech from male speakers from Los Angeles, you may be confident that I can estimate the average f0 male speakers from Los Angeles very accurately. But what if I told you that all these samples were from only three different people? You know instinctively that this makes my data less reliable.

The reason repeated-measures data can cause problems is because the measurements are correlated: multiple measurements from the same person are obviously going to be related to each other. If you measure the height of a tall person today, they will still be tall tomorrow. Because of this general principle, although we have 12 productions from each of 48 female speakers, we do not actually have 576=48*12 totally independent observations in our data.

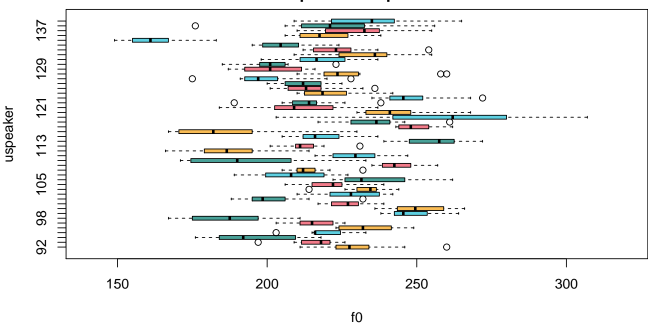
This can be seen quite clearly below. The top panel shows the distribution of all our f0 measurements. The bottom panel shows speaker boxplots (one for each speaker's data). If we were to 'push down' on the bottom panel and collapse all our boxplots into a single distribution, we would end up with the boxplot in the top panel.

These boxplots shows that each speaker has their own average f0, and that their productions tend to vary around their 'natural average. As a result, we might have closer to 46 observations (one average value per speaker) than 576. For example, the 'outliers' around 150 Hz may seem like huge 'errors' in the top plot. In the bottom plot we see that these productions all come from one speaker, and actually reflect her average f0. These are not errors but systematic between-speaker *variation*.





Speaker Boxplots

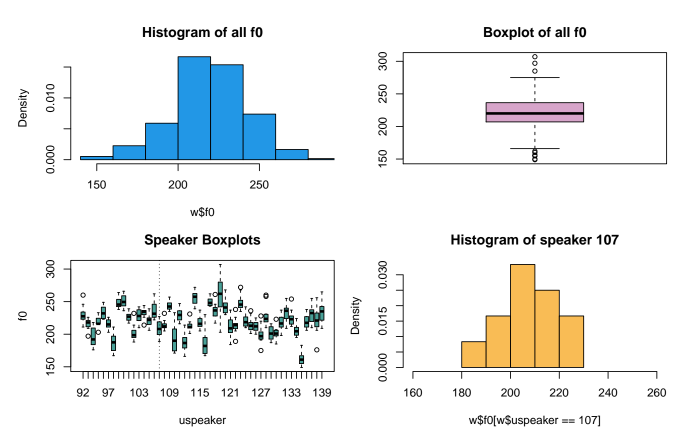


Multilevel models

In linguistics, and many other similar fields, almost all of our data is repeated measures data. The methods most commonly-used by linguists (e.g., experiments, interviews, corpora, ... etc.) yield many observations per person, and typically all involve data from multiple people/sources. As a result, the analysis of this data requires that models be able to account for within *and* between speaker variation in our data. Multilevel models address the correlated nature of repeated measures data by estimating multiple sources of variation simultaneously.

Repeated-measures data leads to random variation in parameters that is *indistinguishable* from that of our 'data'. To a large extent, whether something is a parameter or a data point depends somewhat on your perspective. For example, consider the figure below. The top left presents a histogram of all f0 measurements, while the top right presents a boxplot of the same. The bottom left presents the speaker boxplots (one per speaker), each of which resembles the overall boxplot in the top right. We can then zoom in on a single speaker's productions (bottom right) and produce a histogram that suggests a normal distribution reminiscent in shape to the overall aggregated data (top left).

If you are trying to estimate a speaker's mean f0, then the individual productions might be 'data' and the mean can be thought of as a 'parameter'. If you were instead only interested in the population average, maybe now your subject mean is actually just a single data point, and the *population* mean is actually your 'parameter'.



A multilevel model is able to simultaneously model independent variation at multiple 'levels'. For our f0 data, these are:

- The 'upper' level: Between-speaker variation in mean f0. This can be thought of like variation in $\mu_{speaker}$. Speaker's have an average f0 ($\mu_{speaker}$) that they produce over time. However, speakers are chosen randomly from a larger population, and so any given speaker's $\mu_{speaker}$ is unpredictable a priori.
- The 'lower' level: Within-speaker variation, analogous to ε . When an individual speaker produces speech, their productions will vary around their average from token to token. Our model cannot explain this and so this is 'error'

As seen in the figure above, the variation at the lower and upper levels are analogous. Just like individual speakers will rarely have an average f0 exactly like the population average, individual speakers will rarely produce f0 values exactly at their speaker average. Importantly, variation at the two levels is independent and logically distinct: within-speaker variation can be small or large independently of whether between-speaker variation is large or small.

Basically, each subject's productions form a little normal distribution around their average, and the mix of these little distributions results in the overall 'big' distribution of data across all subjects. By using multilevel models, we can estimate the effects of multiple sources of variation at the same time.

Estimating a multilevel model with brms

We are now going to fit the same model we fit above, but with a structure that reflects the repeated-measures nature of the data.

The model

To specify a multilevel model, you need to write a slightly more complicated model formula. This explanation assumes that you have a dataframe or matrix where one column contains the variable you are interested and predicting (in this case f0), and another column contains a vector containing unique labels for each speaker or source of data (in this case a unique speaker label uspeaker).

To indicate that your model contains an 'upper' level where you have clusters of data coming from different individuals, you have to put another model inside your main model!

Before, the model formula looked like this:

which meant 'predict f0 using only an intercept'. Now the model formula will look like this:

When you place a predictor in the formula in parenthesis, on the right-hand-side of a pipe, like this (| predictor), you tell brm that you expect data to be clustered according to each category represented in the grouping vector. In this case, we are telling brm that each unique speaker is a cluster of data. Whatever you put in the left-hand-side of the parentheses (in here | predictor) is the model for each subcluster!

So what does this model formula mean: f0 ~ 1 + (1 | uspeaker)? It tells brm: predict f0 based on only an intercept, and also calculate a separate intercept for each speaker. Effectively, this model formula is telling brm to figure out all the information presented in the figures above.

This regression model is now something like this:

$$y = \mu_{overall} + \mu_{speaker} + \varepsilon (\#eq: 24)$$

Recall that when we predict an intercept in a regression model, we actually perform the following decomposition $\mu = parameter*1$. This means that we are actually just fitting a parameter that we expect to equal the value of the μ we are interested in.

Below, where I have decomposed each μ into its constituent 'predictors' (1) and parameters (a, b).

$$y = a * 1 + b_{speaker} * 1 + \varepsilon (\#eq : 25)$$

In addition to the coefficient estimating the overall intercept (a), we know have another term $b_{speaker}$. This coefficient is actually a set of coefficients since it has a different value for each speaker (its a vector). It has a different value for each speaker because it will reflect variation in $\mu_{speaker}$.

However, recall that $\mu_{speaker}$ is a random variable since it reflects the random average f0 of each person drawn from the population. If $\mu_{speaker}$ behaves like a random variable, then the coefficients that reflect this value in our model $(b_{speaker})$ will behave in the same way.

This means that actually our model has two random variables. The first one is the error term:

$$\varepsilon \sim \mathcal{N}(0, \sigma_{error})(\#eq: 26)$$

which has a mean of 0 and a standard deviation which we can refer to as σ_{error} .

Our model further assumes that just like our data, our speaker-specific intercepts, or by-speaker intercepts, can be thought of as random draw from a normal distribution. This distribution is also assumed to be centered at zero, and has a population-specific standard deviation.

$$\mu_{sneaker} \sim \mathcal{N}(0, \sigma_{sneaker})(\#eq: 27)$$

Recall that I previously said that regression models encode differences. This helps explain why the distribution of speaker means is centered at 0 in our model. Our model already encodes the overall data average in the $\mu_{overall}$

parameter. Thus, the speaker-specific averages contain information about differences to this overall average, and so are centered at 0 (i.e., the average).

The initial model we fit viewed the variation in the model like this:

$$\sigma_{total} = \sigma_{error}(\#eq:28)$$

In other words, all variation was error. We don't know why values vary from the mean. Our multilevel model views the variation in our data like this:

$$\sigma_{total} = \sigma_{speaker} + \sigma_{error}(\#eq:29)$$

It sees only *some* of the variation in data as error. Basically, from the perspective of this multilevel model, the variation in the data is a combination of random (but systematic) between-speaker variation, and random within-speaker variation.

Fitting the model

We can fit a model with a formula that appropriately specifies the clustering we expect in our data. As a result, this model can estimate both between- and within-speaker variability.

```
set.seed (1)
multilevel_model =
  brm (f0 ~ (1|uspeaker), data = w, chains = 1, cores = 1)
## loand and inspect pre-fit model
multilevel_model = readRDS ("2_multilevel_model.RDS")
multilevel_model
##
   Family: gaussian
##
     Links: mu = identity; sigma = identity
## Formula: f0 ~ (1 | uspeaker)
      Data: w (Number of observations: 576)
## Samples: 1 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup samples = 1000
##
## Group-Level Effects:
##
  ~uspeaker (Number of levels: 48)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                                                 25.06 1.00
## sd(Intercept)
                    20.19
                                2.10
                                        16.65
                                                                  135
                                                                           192
##
## Population-Level Effects:
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
##
               220.39
                           3.10
                                   214.49
                                            226.13 1.03
                                                               51
                                                                       108
## Intercept
##
## Family Specific Parameters:
##
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                                         13.34 1.01
                                                         400
                                                                   700
## sigma
            12.54
                       0.39
                                11.83
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

This new model contains one new chunk its print statement:

```
Group-Level Effects:
~uspeaker (Number of levels: 48)

Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS sd(Intercept) 20.19 2.10 16.65 25.06 1.00 135 192
```

This sections contains information about the standard deviation of between-speaker averages ($\mu_{speaker}$) in the sample. We can see that the information provided by brms is quite similar to what we can estimate directly using our data. However, brms does this all for us, in addition to giving us a lot more information.

```
## find mean f0 for each speaker
speaker_means = aggregate (f0 ~ uspeaker, data = w, FUN = mean)
## find the within speaker variance. This is the within-talker 'error'.
speaker_vars = aggregate (f0 ~ uspeaker, data = w, FUN = var)

## the mean of the speaker means corresponds to our overall mean estimate
mean (speaker_means$f0)
```

[1] 220.401

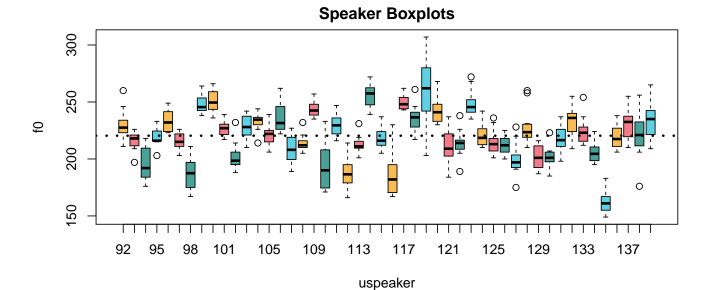
```
## sd(Intercept) in the model reflects the amount of variation in talker
## intercepts. This is the between speaker variation in our model. See how it is
## similar to the sd of the actual speaker means.
sd (speaker_means$f0)
```

```
## [1] 20.07397
```

```
## sigma in the model reflects the amount of variation in talker intercepts.
## This is the between speaker variation in our model.
sqrt (sd (speaker_vars$f0))
```

```
## [1] 12.42719
```

The overall mean f0 in our data (220.4) corresponds quite well to our model estimate of 220.4. This reflects the central location of the overall distribution below (the horizontal line in the figure below). The standard deviation of the speaker means (Intercept = 20.1) is again very similar to our model estimate (sd(Intercept) = 20.1). This reflects the average distance from each speaker's average, and the overall average. Finally, the average of the within speaker standard deviation in our data (12.4) corresponds closely to our model's error estimate (sigma = 12.5). This reflects the average spread of each speaker's data relative to their own mean, within their own little boxplot.



Checking model convergence

Remember that our model parameter estimates consist of a set of samples from the posterior distribution of a parameter. If we don't take enough of these samples, our parameter estimates will be unreliable.

For this reason, it's important to look at the ESS values (the 'expected sample size'), and the 'Rhat' values provided by brm. ESS tells you about how many independent samples you have taken from the likelihood. Bulk ESS is how many samples the sampler took in the thick part of the density, and Tail ESS reflects how much time the sampler spent in the thin part, the 'tails'. Rhat tells you about whether your 'chains' have converged (more on this later). As noted above, values of Rhat near 1 are good, and values higher than around 1.1 are a bad sign.

We haven't really taken many samples here, so we can't be confident in our parameter estimates. Ideally we would like several hundred samples (at least) for mean estimates, and thousands to be confident in the 95% confidence intervals.

To get more samples we can run the model longer, or we can use more *chains*. A chain is basically a separate set of samples for your parameter values. Just imagine you had estimated the model 4 times in a row and mixed your estimations. A model can be fit in parallel across several chains, and then the estimates can be merged across chains. When you do this across multiple cores, you can get N times as many samples when you use N cores. Since many computers these days have 4-8 (or more) cores, we can take advantage of parallel processing to fit models faster.

Below, I refit the same model from above but run it on 4 chains, and on 4 cores at once. This doesn't take any longer but it does give us a higher ESS. Just make sure you leave a couple of cores free on your computer when you fit a model!

```
set.seed (1)
multilevel_multicore =
    brm (f0 ~ 1 + (1|uspeaker), data = w, chains = 4, cores = 4)

## load and inspect pre-fit model
multilevel_multicore = readRDS ('2_multilevel_multicore.RDS')
multilevel_multicore

## Family: gaussian
## Links: mu = identity; sigma = identity
## Formula: f0 ~ 1 + (1 | uspeaker)
```

```
##
      Data: w (Number of observations: 576)
##
  Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup samples = 4000
##
##
  Group-Level Effects:
##
  ~uspeaker (Number of levels: 48)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
##
  sd(Intercept)
                    20.22
                                2.19
                                        16.54
                                                 25.19 1.02
                                                                           615
                                                                  345
##
##
  Population-Level Effects:
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
               220.47
                            2.91
                                   214.67
                                            226.14 1.02
                                                              228
                                                                       543
##
  Intercept
##
## Family Specific Parameters:
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
## sigma
            12.54
                       0.39
                                11.81
                                         13.34 1.00
                                                         3427
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

If we compare the ESS for this new model to the previous model, we see that using 4 chains has substantially increased our ESS, without taking up any more computing time.

One final tweak that I will be using going forward is to 'thin' samples. Notice that we have collected 'total post-warmup samples = 4000'. This means our model has 4000 samples for every parameter in the model. However, we have only about 400 'effective samples' to show for it for some parameters of interest. This means that a lot of our samples are basically dead weight, taking up space and slowing down computations for no good reason.

Sometimes consecutive samples can be too similar and so don't given you that much independent information. A way to fix this is to run longer chains and keep only every nth one. This lets your models be smaller while containing approximately the same information.

To do this you have to set the iter, warmup and thin parameters. You will keep every sample after the warmup is done, up to the iter maximum. So if iter=3000 and warmup=1000 you will end up with 2000 samples. After this, you keep only one every thin samples. Basically, you will end up with (iter - warmup)/thin samples per chain.

Below, I ask for 11,000 sample per chain, 10,000 post warm-up. However, since I plan to keep only 1/10, I will have 1000 per core, so 4000 samples in total. However, despite having the same number of samples as the multilevel_multicore, the ESS for this model is much higher for important parameters such as the model intercept.

```
set.seed (1)
multilevel_thinned =
  brm (f0 ~ 1 + (1|uspeaker), data = w, chains = 4, cores = 4,
       warmup = 1000, iter = 11000, thin = 10)
## load and inspect pre-fit model
multilevel_thinned = readRDS ('2_multilevel_thinned.RDS')
multilevel_thinned
##
   Family: gaussian
##
    Links: mu = identity; sigma = identity
## Formula: f0 ~ 1 + (1 | uspeaker)
##
      Data: w (Number of observations: 576)
## Samples: 4 chains, each with iter = 11000; warmup = 1000; thin = 10;
##
            total post-warmup samples = 4000
##
```

```
Group-Level Effects:
##
  ~uspeaker (Number of levels: 48)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                                2.23
                                                  24.98 1.00
                                                                 2775
                                                                          3392
##
  sd(Intercept)
                    20.06
                                        16.24
##
##
  Population-Level Effects:
##
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
               220.40
                            2.95
                                   214.57
                                            226.19 1.00
                                                             1987
                                                                      2625
  Intercept
##
  Family Specific Parameters:
##
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
            12.54
                       0.38
                                11.81
                                         13.31 1.00
                                                         3908
                                                                  3891
## sigma
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

Specifying prior probabilities

In Chapter 1 we discussed that Bayesian models require that prior probabilities be specified for all parameters. You may have noticed that to this point I haven't discussed priors at all.

If you don't specify prior probabilities for your parameters, brm will use a 'flat' prior for all parameters. When you do this, you are basically relying only on the likelihood for your analysis. You are also telling your model that, a priori, any value of average f0 is equally believable. Empirically, this is false. As a practical matter this can cause problems for samplers like brm (STAN actually). Basically, the sampler has a harder time figuring out the most likely values when you tell it to look anywhere from positive to negative infinity. Even a bit of guidance can help.

brms makes it really easy to specify prior probabilities for specific parameters or whole groups of parameters. First we figure out the overall mean and the standard deviation of the data.

```
mean(f0s)

## [1] 220.401

sd(f0s)
```

[1] 23.22069

In the example below, I use this information to set reasonable bounds on the parameters in the model. I do this by class of parameter:

- Intercept: this is a unique class, only for intercepts.
- b: this is for all the non-intercept predictors. There are none in this model.
- sd: this is for all standard deviation parameters. In our example this is sd(Intercept) for uspeaker $(\sigma_{speaker})$, and sigma (σ_{error}) .

Both priors below use a 't' distribution, which is just like a normal distribution but it is more pointy, and has more density in the outer parts of the distribution. I use this because it has good properties, but you can use normal priors, or any other priors that you think work for your model. Rather than focusing on the mathematical properties of priors, the most important thing is that their *shape* reflect the distribution of credible parameter values a priori (before you conducted your experiment).

The format for the priors looks like this student_t(nu, mean, sd), where nu is a parameter that determines how pointy the distribution is, and mean and sd are the same mean and standard deviation parameters from the

normal distribution. The nu parameter ranges from 1 to infinity, and large numbers result in a more normal-like distribution.

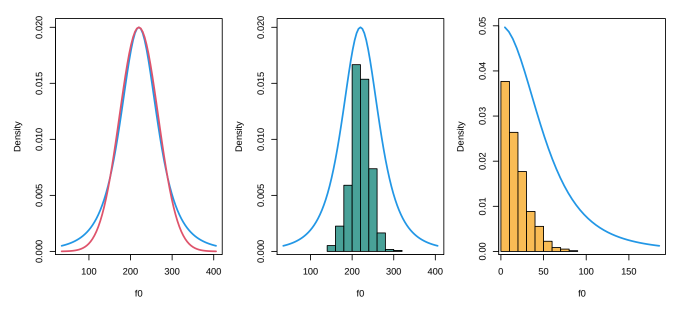
So, for the overall intercept ($\mu_{overall}$) I am using a prior with the same mean as the data mean, and a standard deviation that is twice as large as the data standard deviation. For both the model standard deviation terms (σ_{error} , $\sigma_{speaker}$) I am using a t distribution centered at 0 (explained below), with a standard deviation twice as large as the data standard deviation.

```
## load and inspect pr-fit model
multilevel_priors = readRDS ('2_multilevel_priors.RDS')
multilevel_priors
```

```
##
   Family: gaussian
    Links: mu = identity; sigma = identity
## Formula: f0 ~ 1 + (1 | uspeaker)
##
      Data: w (Number of observations: 576)
## Samples: 4 chains, each with iter = 11000; warmup = 1000; thin = 10;
##
            total post-warmup samples = 4000
##
## Group-Level Effects:
##
  ~uspeaker (Number of levels: 48)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
  sd(Intercept)
                    20.19
                                2.24
                                        16.29
                                                 25.06 1.00
                                                                 3114
                                                                          3663
##
  Population-Level Effects:
##
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
               220.42
                            2.97
                                   214.44
                                            226.38 1.00
                                                             1796
                                                                      2841
##
  Intercept
##
## Family Specific Parameters:
##
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sigma
            12.54
                       0.39
                                11.81
                                         13.34 1.00
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

In the left panel below (plot code at end of chapter) I compare the t distribution we used (blue) to the equivalent normal distribution (red). It is clear that the t distribution can tolerate more extreme values because of its 'fatter' tails. As we will discuss later, using t distributions can make our models more robust to outliers. This is really important in linguistics where subjects/speakers sometimes do weird stuff!

In the middle panel we compare this prior to the data, and see that the prior distribution is much broader (more vague) than the data distribution. The right panel compares the prior for the standard deviation parameters to the absolute value of the centered f0 data. This presentation shows how far each observation is from the mean f0 (at 220 Hz). Again, the prior distribution we have assigned for these parameters is much larger than the variation in the data. As a result, neither of these priors is going to have much of an effect on our parameter estimates.



If we compare the output of this model to multilevel_thinned, we see that specifying a prior has has no noticeable effect on our results. This is because the prior matters less and less when you have a lot of data, and because we have set wide priors that are appropriate (but vague) given our data. Although the priors may not matter much for models as simple as these, they can be very important when working with more complex data, and are a necessary component of Bayesian modeling.

Answering our research questions

Let's return again to the research questions I posed initially:

- 1) What is the average f0 of the whole population likely to be?
- 2) Can we set bounds on likely mean f0 values based on the data we collected?

And we can compare the answers provided to this question by our initial and final models.

model

```
##
   Family: gaussian
##
     Links: mu = identity; sigma = identity
  Formula: f0 ~ 1
##
##
      Data: w (Number of observations: 576)
##
  Samples: 1 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup samples = 1000
##
##
  Population-Level Effects:
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
##
  Intercept
               220.40
                           0.97
                                   218.33
                                            222.30 1.00
                                                              851
                                                                       557
##
  Family Specific Parameters:
##
##
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
            23.24
                       0.69
                                21.99
                                         24.61 1.00
  sigma
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

multilevel_priors

```
##
   Family: gaussian
##
    Links: mu = identity; sigma = identity
  Formula: f0 ~ 1 + (1 | uspeaker)
      Data: w (Number of observations: 576)
##
## Samples: 4 chains, each with iter = 11000; warmup = 1000; thin = 10;
##
            total post-warmup samples = 4000
##
## Group-Level Effects:
  ~uspeaker (Number of levels: 48)
##
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                                                 25.06 1.00
                                                                          3663
##
  sd(Intercept)
                                2.24
                                        16.29
                                                                 3114
##
## Population-Level Effects:
             Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
               220.42
                           2.97
                                   214.44
                                            226.38 1.00
                                                             1796
                                                                      2841
##
  Intercept
##
##
  Family Specific Parameters:
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
##
##
                                                                  3972
            12.54
                       0.39
                                11.81
                                         13.34 1.00
                                                         3968
  sigma
##
## Samples were drawn using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
```

Our initial model (model) and our final model (multilevel_priors) agree on what the average f0 is. However, they disagree on a credible interval for that parameter. Our initial model did not specify information about repeated measures. This causes our model to think that it has more independent observations than it does, and so it returns an overly-precise estimate.

Another difference is that the final model has a much smaller sigma parameter (12.5 vs 23.2). This indicates that the error (ε) is much smaller in the final model than in the initial model. Keep in mind that 'error' is just what your model can't explain. Our final model explains much more and so there is less error. The reduced error is a direct result of the fact that the final model splits random variation up into between-speaker and within-speaker components, estimating the between-speaker variation ($sigma_{speaker}$) to be about 20 Hz.

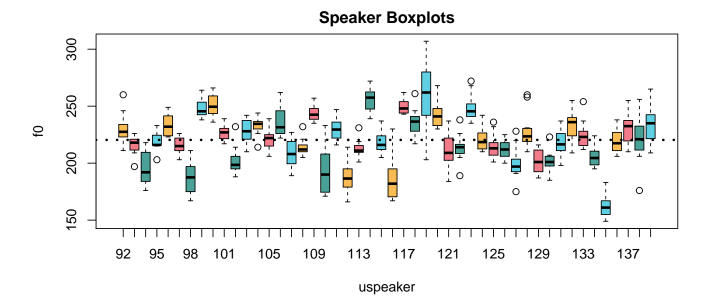
Usually, when I report parameters I provide the mean and standard deviations of the posterior distribution, in addition to the bounds of the 95% credible interval of the parameter. Based on the result of our final model, I think a thorough description of the general properties of our data might go something like:

"Based on our model the average f0 produced by adult females in Michigan is likely to be 220 Hz (s.d. = 2.97, 95% CI = 214.4, 226.4). However, consistent between-speaker variation averages about 20 Hz (s.d. = 2.24, 95% CI = 16.29, 25.06), meaning that we can expect the average f0 produced by individuals to deviate substantially from 220 Hz. Finally, the standard deviation of production error was about 12.5 Hz (s.d. = 0.39, 95% CI = 11.81, 13.34) indicating that the amount of random between speaker variation in production is about half the magnitude of the stable, between-speaker differences in f0.

I am including the speaker boxplots below because I think this image basically presents the same information as the paragraph above, but in visual form. In general, any data relationship or result can be presented in a figure, and the relationships presented in a figure can also be expressed as a mathematical model.

When you are thinking about the relationships in your data, or that you expect in your data, its a good idea to think: what kind of picture could illustrate this relationship? Conversely, if you see a figure of your results that you feel really expresses something interesting about your data you should think, how can these relationships be represented in a model?

PLOT CODE 43



Plot Code

Plot for comparison of prior distributions with f0 data

```
# get t density
x1 = seq (-4,4,.1)
y1 = dt (x1, 3); y1 = y1 / max (y1) / 50
x2 = 220+(x1*46.4)
y2 = dnorm (x2, 220, 46.4); y2 = y2 / max (y2) / 50
par (mfrow = c(1,3), mar = c(4,4,1,1))
## plot t
plot (x2, y1, type = '1', lwd = 2, ylab = 'Density', xlab = 'f0', col = 4)
## compare to equivalent normal
lines (x2, y2, lwd=2, col=2)
## plot t
plot (x2, y1, type = 'l', lwd = 2, ylab = 'Density', xlab = 'f0', col = 4)
## compare to equivalent normal
hist (f0s, add = TRUE, freq = FALSE)
## plot prior for standard deviations
x3 = x2 - mean (x2)
y3 = dt (x1, 3); y3 = y3 / max (y3) / 20
plot (x3[x3>0], y3[x3>0], type = 'l', lwd = 2, ylab = 'Density',
      xlab = 'f0', col = 4)
hist (abs (f0s - mean (f0s)), add = TRUE, freq = FALSE)
```

44INFERENCE FOR A 'SINGLE GROUP' OF OBSERVATIONS USING A BAYESIAN MULTILEVEL MODEL