# **Computational Bayesian data analysis**

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- Deriving the posterior distribution analytically is possible for only a very limited number of cases.
- The denominator, the marginal likelihood, requires us to compute a possibly intractable integral:

$$p(\Theta|y) = \frac{p(y|\Theta) \cdot p(\Theta)}{\int_{\Theta} p(y|\Theta) \cdot p(\Theta)d\Theta} \tag{1}$$

# Alternative: Deriving the posterior through sampling

# We want to derive the posterior distribution of the Cloze probability of "umbrella", $\theta$ :

- Data: a word (e.g., "umbrella") was answered 80 out of 100 times,
- · Likelihood: a binomial distribution
- Prior for  $\theta$ : Beta(a=4,b=4)

#### We sample from the posterior distribution of $\theta$ :

- · We use a probabilistic programming language,
- given enough samples we will have a good approximation of the real posterior distribution,
- say we got 20000 samples from the posterior distribution of the Cloze probability,  $\theta$ :

0.782, 0.826, 0.788, 0.671, 0.712, 0.776, 0.819, 0.75, 0.8, 0.817, 0.814, 0.863, 0.782, 0.818, 0.753, 0.743, 0.782, 0.772, 0.785, 0.724, ...

# The approximation of the posterior looks quite similar to the real posterior.<sup>1</sup>

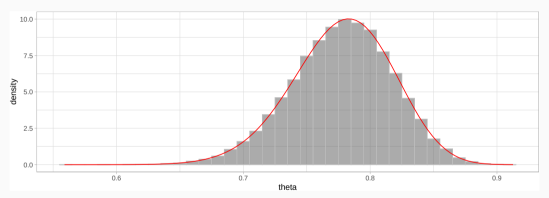


Figure 1: Histogram of the samples of  $\theta$  from the posterior distribution calculated through sampling in gray; density plot of the exact posterior in red.

<sup>&</sup>lt;sup>1</sup>The difference between the true and the approximated mean and variance are -0.0002 and 0.000005 respectively

# Computational Bayesian data analysis:

#### Why has Bayes suddenly gained in prominence since the 2000s?

- increase in computing power
- appearance of probabilistic programming languages: WinBUGS (Lunn et al. 2000), JAGS (Plummer 2016), and more recently pymc3
  (Salvatier, Wiecki, and Fonnesbeck 2016), Stan (Carpenter et al. 2017), INLA (Blangiardo and Cameletti 2015).

#### **Easier alternatives based on Stan:**

- rstanarm (Goodrich et al. 2018)
- brms (Bürkner 2019)

# **Bayesian Regression Models**

using 'Stan': brms

### Load the following:

```
set.seed(42)
library(MASS)
## be careful to load dplyr after MASS
library(dplyr)
library(tidyr)
library(purrr)
library(readr)
library(ggplot2)
library(brms)
## Save compiled models:
rstan_options(auto_write = TRUE)
## Parallelize the chains using all the cores:
options(mc.cores = parallel::detectCores())
library(bayesplot)
library(tictoc)
```

# simple linear model)

**Examples 1: A single participant** 

pressing a button repeatedly (A

We have data from a participant repeatedly pressing the space bar as fast as possible, without paying attention to any stimuli.

#### Data:

reaction times in milliseconds in each trial

#### **Question:**

How long does it take to press a key when there is no decision involved?

#### **Assumptions:**

- 1. There is a true underlying time,  $\mu$ , that the participant needs to press the space bar.
- 2. There is some noise in this process.
- 3. The noise is normally distributed (this assumption is questionable given that reaction times are generally skewed; we fix this assumption later).

#### Formal model:

#### Likelihood for each observation n:

$$rt_n \sim Normal(\mu, \sigma)$$
 (2)

#### (Bad) priors:

$$\mu \sim Uniform(0,60000)$$

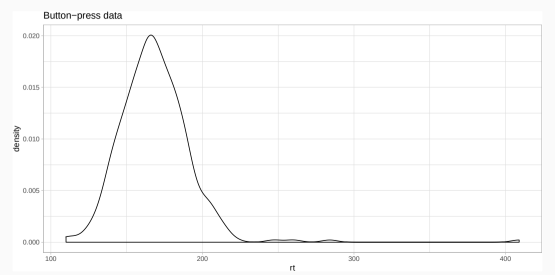
$$\sigma \sim Uniform(0,2000)$$
(3)

# Fitting the model

We'll first load the data from data/button\_press.csv:

```
df noreading data <-
  read_csv("./data/button_press.csv")
df noreading data
## # A tibble: 361 x 2
##
        rt trialn
     <dbl> <dbl>
##
## 1
      141
## 2
      138
      128
## 3
     132
## 4
## 5
      126
                5
## #
       rrith 256 mara rarra
```

```
ggplot(df_noreading_data, aes(rt)) +
  geom_density() +
  ggtitle("Button-press data")
```

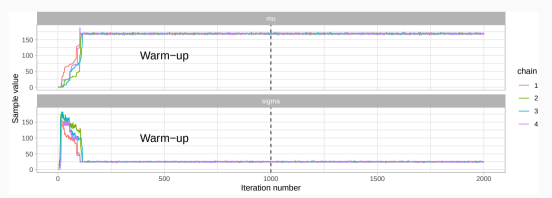


# **Specifying the model in brms**

```
fit press <- brm(rt ~ 1,
  data = df_noreading_data,
  family = gaussian(),
  prior = c(
    prior(uniform(0, 60000), class = Intercept),
    prior(uniform(0, 2000), class = sigma)
  ),
  chains = 4.
  iter = 2000.
  warmup = 1000
```

# Sampling and convergence in a nutshell

- 1. Chains start in random locations;
- 2. in each iteration they take one sample each;
- 3. samples at the beginning do not belong to the posterior distribution;
- 4. eventually, the chains end up in the vicinity of the posterior distribution;
- 5. from that point onwards the samples will belong to the posterior.



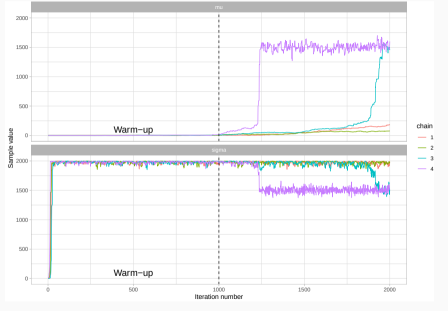
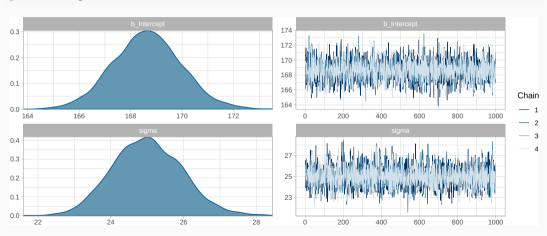


Figure 4: Trace plot of a model that did not converge.

```
posterior_samples(fit_press) %>% str()

## 'data.frame': 4000 obs. of 3 variables:
## $ b_Intercept: num 168 171 169 172 167 ...
## $ sigma : num 24.4 25.4 24 25.9 26.1 ...
## $ lp__ : num -1688 -1689 -1689 -1691 -1689 ...
```

#### plot(fit\_press)



```
fit_press
# posterior summary(fit press) is also useful
## Family: gaussian
##
  Links: mu = identity; sigma = identity
## Formula: rt ~ 1
     Data: df_noreading_data (Number of observations: 361)
##
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
           total post-warmup samples = 4000
##
##
## Population-Level Effects:
            Estimate Est.Error 1-95% CI u-95% CI Rhat
##
## Intercept 168.64 1.33 166.06 171.35 1.00
         Bulk_ESS Tail_ESS
##
## Intercept 3384
                        2336
##
## Family Specific Parameters:
        Estimate Est.Error 1-95% CI u-95% CI Rhat
##
         25.00 0.95 23.23 26.93 1.00
## sigma
        Bulk ESS Tail ESS
##
```

Notice that the Estimate is just the mean of the posterior sample, and CI are the 95% quantiles:

```
posterior_samples(fit_press)$b_Intercept %>%
 mean()
## [1] 169
posterior samples(fit press)$b Intercept %>%
  quantile(c(0.025, .975))
## 2.5% 98%
## 166 171
```

# **Important questions**

- 1. What information are the priors encoding? Do the priors make sense?
- 2. Does the likelihood assumed in the model make sense for the data?

**Prior predictive distributions** 

# **Prior predictive distributions**

We want to know the density  $p(\cdot)$  of a data vector  $y=\langle y_1,\dots,y_n\rangle$ , given a vector of priors  $\Theta$  (e.g.,  $\Theta=\langle \mu,\sigma\rangle$ )

The prior predictive density is:

$$p(y) = \int p(y|\Theta)p(\Theta) d\Theta$$
 (4)

We avoid doing the integration analytically; we will use Monte Carlo integration. We repeat the following:

- 1. Take one sample from each of the priors.
- 2. Plug those samples in the likelihood and generate a dataset

$$y_{pred} = \langle y_{pred}, \dots, y_{pred} \rangle$$
.

```
normal_predictive_distribution <- function(mu_samples, sigma_samples, N_obs) {
  # empty data frame with headers:
  df_pred <- tibble(</pre>
    trialn = numeric(0),
   rt_pred = numeric(0),
    iter = numeric(0)
  # i iterates from 1 to the length of mu samples.
  # which we assume is identical to
  # the length of the sigma samples:
  for (i in seq_along(mu_samples)) {
    mu <- mu_samples[i]</pre>
    sigma <- sigma samples[i]</pre>
    df_pred <- bind_rows(</pre>
      df pred.
      tibble(
        trialn = seq_len(N_obs), # 1, 2,... N_obs
        rt_pred = rnorm(N_obs, mu, sigma),
        iter = i
  df_pred
```

#### This approach works, but it's quite slow:

```
tic()
N_samples <- 1000
N obs <- nrow(df noreading data)
mu_samples <- runif(N_samples, 0, 60000)</pre>
sigma samples <- runif(N samples, 0, 2000)
normal_predictive_distribution(mu_samples = mu_samples,
                             sigma_samples = sigma_samples,
                              N \text{ obs} = N \text{ obs}
toc()
## # A tibble: 361,000 x 3
##
    trialn rt_pred iter
##
     <dbl> <dbl> <dbl>
## 1
     1 44587. 1
## 2 2 48271. 1
## 3 3 50291. 1
## 4 48784.
## 5 5 49350.
## # ... with 3.61e+05 more rows
## 4.762 sec elapsed
```

#### A more efficient version:

```
normal predictive distribution fast <- function(mu samples.
                                                sigma_samples,
                                                N obs) {
 # map dfr works similarly to lapply, it essentially runs
 # a for-loop, and builds a dataframe with the output.
 # We iterate over the values of mu samples and sigma samples
  # simultaneously, and in each iteration we bind a new
  # data frame with N obs observations.
 map2_dfr(mu_samples, sigma_samples, function(mu, sigma) {
   tibble(
     trialn = seq_len(N_obs),
     rt pred = rnorm(N obs. mu. sigma)
   )}. .id = "iter") %>%
    # .id is always a string and needs to be converted to a number
   mutate(iter = as.numeric(iter))
```

```
tic()
(prior_pred <- normal_predictive_distribution_fast()</pre>
 mu_samples = mu_samples,
  sigma_samples = sigma_samples,
 N obs))
toc()
## # A tibble: 361,000 x 3
##
     iter trialn rt_pred
##
    <dbl> <int> <dbl>
## 1
              1 48513.
## 2 1 2 46243.
## 3 1
              3 48812.
## 4 1 4 49650.
## 5 1
              5 49096.
## # ... with 3.61e+05 more rows
## 0.301 sec elapsed
```

```
prior_pred %>%
filter(iter <= 12) %>%
ggplot(aes(rt_pred)) +
geom_histogram() +
facet_wrap(~iter, ncol = 3)
```

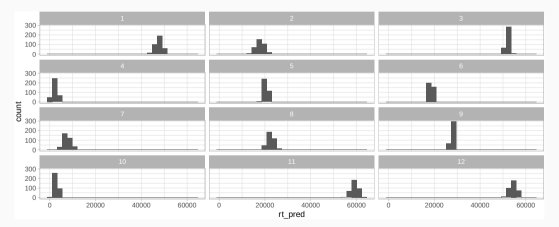


Figure 5: Eighteen samples from the prior predictive distribution.

#### **Distribution of statistics**

```
## # A tibble: 3,000 x 3
##
   iter stat
                  rt
##
    <dbl> <dbl> <dbl>
## 1
    1 min_rt 43017.
## 2 1 max_rt 52560.
## 3 1 average rt 47753.
## 4
       2 min rt
                13331.
44 5
       0 ----
                 2222
```

```
prior_stat %>%
    ggplot(aes(rt)) +
    geom_histogram(binwidth = 500) +
    facet_wrap(~stat, ncol = 1)
```

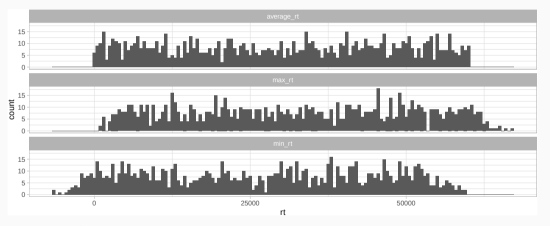


Figure 6: Prior predictive distribution of averages, maximum, and minimum values.

#### Why are our distributions so bad?

We used much less prior information than what we really had: our priors are clearly not very realistic given what we know about reaction times for such a button pressing task.

#### What priors should we have chosen?

The influence of priors:

sensitivity analysis

# **Types of priors**

- 1. Flat uninformative priors: priors as uninformative as possible.
- 2. **Regularizing priors**: priors that downweight extreme values (that is, they provide regularization), they are not very informative, and mostly let the likelihood dominate in determining the posteriors.
- 3. **Principled priors**: priors that encode all (or most of) the theory-neutral information that we do have.
- 4. **Informative priors**: There are cases where we have a lot of prior knowledge, and not much data.

# Revisiting the button-pressing example with different priors

What would happen if we use even wider priors for the model?

$$\mu \sim Uniform(-10^{10}, 10^{10}) \\ \sigma \sim Uniform(0, 10^{10})$$
 (5)

#### In brms:

```
fit_press_unif <- brm(rt ~ 1,
  data = df_noreading_data,
  family = gaussian(),
  prior = c(
      prior(uniform(-10^10, 10^10), class = Intercept),
      prior(uniform(0, 10^10), class = sigma))
)</pre>
```

#### The output of the model is virtually identical!

fit\_press\_unif

```
## Family: gaussian
##
  Links: mu = identity; sigma = identity
## Formula: rt ~ 1
##
     Data: df noreading data (Number of observations: 361)
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
           total post-warmup samples = 4000
##
##
## Population-Level Effects:
            Estimate Est.Error 1-95% CI u-95% CI Rhat
##
## Intercept 168.63 1.29 166.04 171.16 1.00
##
          Bulk_ESS Tail_ESS
## Intercept 3371
                        2578
##
## Family Specific Parameters:
        Estimate Est.Error 1-95% CI u-95% CI Rhat
##
           24.99 0.95 23.21 26.93 1.00
## sigma
        Bulk ESS Tail ESS
##
            3654
                     2897
## sigma
##
```

#### What happens if we use very informative priors and they are off?

```
\mu \sim Normal(400, 10)
\sigma \sim Normal_{+}(100, 10)
(6)
```

```
fit press inf <- brm(rt ~ 1,
  data = df noreading data,
  family = gaussian().
  prior = c(
    prior(normal(400, 10), class = Intercept),
    # brms knows that SD needs to be bounded by zero:
    prior(normal(100, 10), class = sigma)
```

# Even in this case, the new estimates are just a couple of milliseconds away from our previous estimates:

fit\_press\_inf

```
Family: gaussian
##
    Links: mu = identity; sigma = identity
## Formula: rt ~ 1
##
     Data: df noreading data (Number of observations: 361)
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
           total post-warmup samples = 4000
##
## Population-Level Effects:
            Estimate Est.Error 1-95% CI u-95% CI Rhat
##
## Intercept 172.93 1.40 170.16 175.78 1.00
         Bulk_ESS Tail_ESS
##
## Intercept 2761
                        2607
##
## Family Specific Parameters:
##
        Estimate Est.Error 1-95% CI u-95% CI Rhat
           26.07 1.07 24.11
                                      28.28 1.00
## sigma
##
        Bulk ESS Tail ESS
## sigma
            2866
                     2621
```

#### This doesn't mean that priors never matter:

- When there is enough data for a certain parameter, the likelihood will dominate
- If we are not sure about the extent to which the posterior is influenced by our priors, we can do a *sensitivity analysis* (for a published example in psycholinguistics, see Vasishth et al. 2013).
- We can use prior predictive distributions to see if we are on the right order of magnitude for our priors

**Posterior predictive distributions** 

Once we have the posterior distribution  $p(\Theta \mid y)$ , we can derive the predictions based on this distribution:

$$p(D_{pred} \mid y) = \int_{\Theta} p(D_{pred} \mid \Theta) p(\Theta \mid y) d\Theta$$
 (7)

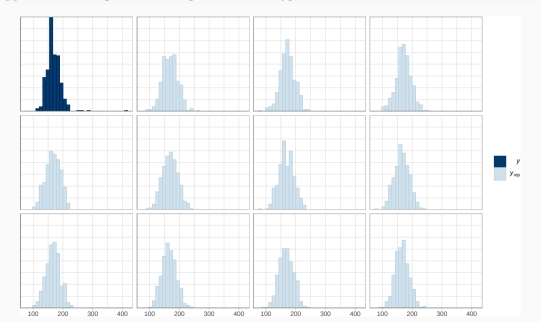
Here we can also avoid the integration, and we can even use the same function that we created before:

```
N_obs <- nrow(df_noreading_data)</pre>
mu samples <- posterior samples(fit press)$b Intercept</pre>
sigma_samples <- posterior_samples(fit_press)$sigma
(normal predictive distribution fast(
  mu_samples = mu_samples,
  sigma samples = sigma samples,
 N obs
))
## # A tibble: 1,444,000 x 3
##
      iter trialn rt_pred
##
     <dbl> <int>
                    <dbl>
## 1
                   214.
## 2
                   136
## 3
                    195.
## 4
                    125.
## 5
                5
                     159.
## # with 1 444e+06 more rows
```

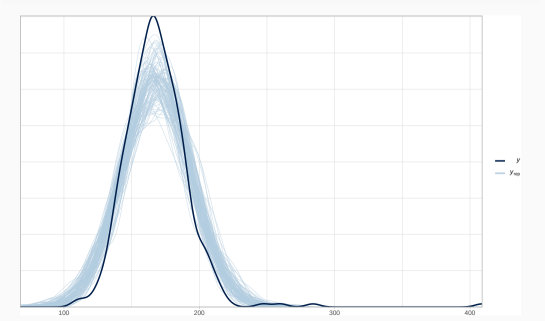
### **Descriptive adequacy/posterior predictive checks**

Could the current data have been generated by our model?

pp\_check(fit\_press, nsamples = 11, type = "hist")



pp\_check(fit\_press, nsamples = 100)



**Comparing different likelihoods:** 

The log-normal likelihood

If y is log-normally distributed, this means that  $\log(y)$  is normally distributed.<sup>2</sup>

$$\log(y) \sim Normal(\mu, \sigma)$$
 
$$y \sim \exp(Normal(\mu, \sigma))$$
 
$$y \sim LogNormal(\mu, \sigma)$$
 (8)

The log-normal distribution is again defined using  $\mu$  and  $\sigma$ , but these correspond to the mean and standard deviation of the normally distributed logarithm of the data y:  $\log(y)$ .

 $<sup>^{2}</sup>$ In fact,  $\log_{e}(y)$  or  $\ln(y)$ , but we'll write it as just log()

# Re-fitting a single participant pressing a button repeatedly with a log-normal likelihood

#### New likelihood:

$$rt_n \sim LogNormal(\mu, \sigma)$$
 (9)

#### New scale for the priors:

$$\mu \sim Uniform(0,8) \\ \sigma \sim Uniform(0,1)$$
 (10)

Because the parameters are in a different scale than the dependent variable, their interpretation changes:

- The location,  $\mu$ : In our previous linear model,  $\mu$  represented the grand mean (or the grand median, or grand mode, since in a normal distribution the three coincide). But now, the grand mean is  $\exp(\mu + \sigma^2/2)$  and the grand median is  $\exp(\mu)$ .
- The scale,  $\sigma$ : This is the standard deviation of the normal distribution of  $\log(y)$ . The standard deviation of a log-normal distribution with location  $\mu$  and scale  $\sigma$  will be  $\exp(\mu + \sigma^2/2) \times \sqrt{(\exp(\sigma^2) 1)}$ .

#### **Prior predictive distributions**

```
N samples <- 1000
N obs <- nrow(df noreading data)
mu samples <- runif(N samples, 0, 8)</pre>
sigma samples <- runif(N samples, 0, 1)
prior_pred_ln <- exp(normal_predictive_distribution_fast())</pre>
  mu samples = mu samples,
  sigma samples = sigma samples,
  N obs
```

#### **Distribution of statistics**

```
(prior_pred_stat_ln <-
prior_pred_ln %>%
group_by(iter) %>%
summarize(
  min_rt = min(rt_pred),
  max_rt = max(rt_pred),
  average_rt = mean(rt_pred),
  median_rt = median(rt_pred)
) %>%
pivot_longer(cols = ends_with("rt"), names_to = "stat", values_to = "rt"))
```

```
## # A tibble: 2,840 x 3

## iter stat rt

## <dbl> <chr> <dbl> <hr> ## 1 2.72 min_rt 106.

## 2 2.72 max_rt 154.

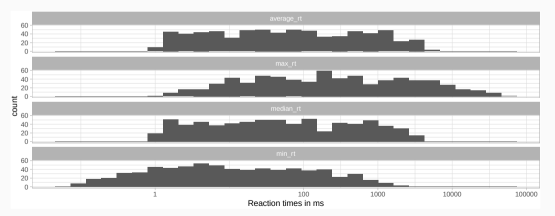
## 3 2.72 average_rt 130.

## 4 2.72 median_rt 131.

## 5 7.39 min_rt 2.99

## # ... with 2,835 more rows
```

```
prior_pred_stat_ln %%
ggplot(aes(rt)) +
scale_x_continuous("Reaction times in ms",
    trans = "log", breaks = c(0.001, 1, 100, 1000, 10000, 100000)) +
geom_histogram() +
facet_wrap(~stat, ncol = 1)
```



**Figure 9:** Prior predictive distribution of averages, maximum, and minimum value of the log-normal model; the x-axis is log-transformed.

## Better regularizing priors for the log-normal model

$$rt_n \sim LogNormal(\mu,\sigma)$$
 (11)

$$\mu \sim Normal(6, 1.5)$$
  $\sigma \sim Normal_{+}(0, 1)$  (12)

#### **Median effect for our new priors:**

```
c(
  lower = exp(6 - 2 * 1.5),
  higher = exp(6 + 2 * 1.5)
)

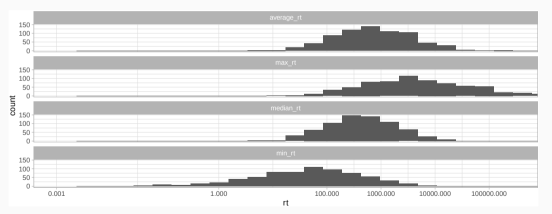
## lower higher
## 20 8103
```

#### **Prior predictive distributions**

```
N samples <- 1000
N obs <- nrow(df noreading data)
mu samples <- rnorm(N samples, 6, 1.5)
sigma samples <- rtnorm(N samples, 0, 1, a = 0)
(prior_pred_ln_better <- exp(normal_predictive_distribution_fast())</pre>
  mu samples = mu samples,
  sigma_samples = sigma_samples,
 N obs
)))
## # A tibble: 361,000 x 3
##
     iter trialn rt_pred
##
    <dbl> <dbl> <dbl>
## 1 2.72 2.72 261.
## 2 2.72 7.39 248.
## 3 2.72 20.1 441.
## 4 2.72 54.6 841.
## 5 2.72 148. 2975.
## # with 3 61e+05 more rows
```

```
(prior pred stat better ln <- prior pred ln better %>%
  group_by(iter) %>%
  summarize(
    min_rt = min(rt_pred),
    max_rt = max(rt_pred),
    average_rt = mean(rt_pred),
    median rt = median(rt pred)
  ) %>%
  pivot longer(
    cols = ends_with("rt"),
    names to = "stat", values to = "rt"
  ))
## # A tibble: 2,840 x 3
                           rt.
```

```
##
     iter stat
##
    <dbl> <chr>
                      <dbl>
## 1 2.72 min_rt
                       4.33
## 2 2.72 max_rt
                    9351.
     2.72 average_rt 734.
## 3
## 4 2.72 median rt
                     344.
## 5 7.39 min rt
                      23.8
## # ... with 2,835 more rows
```



**Figure 10:** Prior predictive distribution of averages, maximum, and minimum value of the log-normal model with better priors.

#### brms model with reasonable priors:3

```
fit_press_ln <- brm(rt ~ 1,
  data = df_noreading_data,
  family = lognormal(),
  prior = c(
    prior(normal(6, 1.5), class = Intercept),
    prior(normal(0, 1), class = sigma)
)
)</pre>
```

<sup>&</sup>lt;sup>3</sup>Notice that we need to specify that the family is lognormal()

```
## Family: lognormal
    Links: mu = identity; sigma = identity
##
## Formula: rt ~ 1
     Data: df noreading data (Number of observations: 361)
##
## Samples: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
          total post-warmup samples = 4000
##
## Population-Level Effects:
           Estimate Est.Error 1-95% CI u-95% CI Rhat
##
## Intercept 5.12
                       0.01
                               5.10
                                       5.13 1.00
##
           Bulk_ESS Tail_ESS
## Intercept 3377
                       2768
##
## Family Specific Parameters:
       Estimate Est.Error 1-95% CI u-95% CI Rhat
##
         0.13 0.01 0.13 0.15 1.00
## sigma
##
       Bulk ESS Tail ESS
## sigma
           2919
                   2201
##
## Samples were drawn using sampling(NUTS). For each parameter, Eff.Sample
## is a crude measure of effective sample size, and Rhat is the potential
```

fit press ln

#### How long does it take to press the space bar in milliseconds?

```
estimate_ms <- exp(posterior_samples(fit_press_ln)$b_Intercept)
c(mean = mean(estimate_ms), quantile(estimate_ms, probs = c(.025, .975)))
## mean 2.5% 98%
## 167 165 169</pre>
```

### **Posterior predictive checks**

pp\_check(fit\_press\_ln, nsamples = 100)

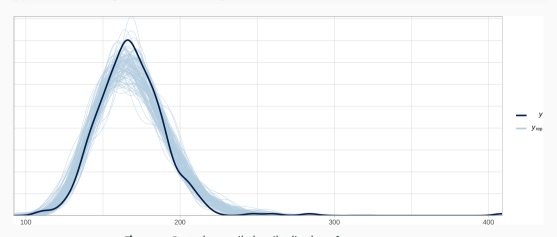


Figure 11: Posterior predictive distribution of  $fit\_noreading\_ln$ 

## Are the posterior predicted data now more similar to the real data, compared to the case where we had a Normal likelihood?

We suspect that the normal distribution would generate reaction times that are too fast (since it's symmetrical) and that the log-normal distribution may capture the long tail better than the normal model.

```
pp_check(fit_press, type = "stat", stat = "min") +
    ggtitle("Normal model")
pp_check(fit_press_ln, type = "stat", stat = "min") +
    ggtitle("Log-normal model")
```

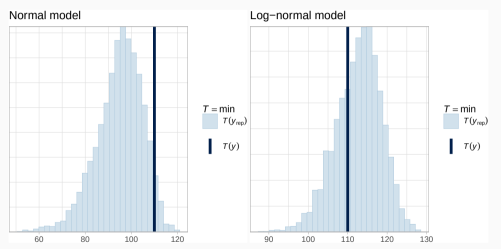


Figure 12: Distribution of minimum values in a posterior predictive check. The minimum in the data is 110 ms.

```
pp_check(fit_press, type = "stat", stat = "max") +
    ggtitle("Normal model")
pp_check(fit_press_ln, type = "stat", stat = "max") +
    ggtitle("Log-normal model")
```

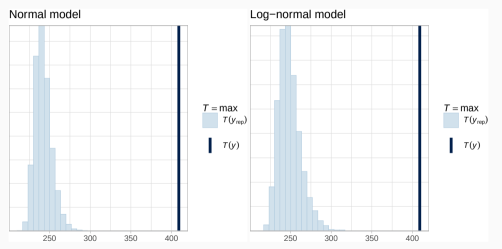


Figure 13: Distribution of maximum values in a posterior predictive check. The maximum in the data is 409 ms.

#### What did we do?

- fitted and interpreted a normal model
- looked at the effect of priors:
  - prior predictive distributions
  - sensitivity analysis
- looked at the fit of the posterior:
  - posterior predictive distribution (descriptive adequacy)
- fitted and interpreted a log-normal model
- compared a normal model with a log-normal one

#### References

Blangiardo, Marta, and Michela Cameletti. 2015. Spatial and Spatio-Temporal Bayesian models with R-INLA. John Wiley & Sons.

Bürkner, Paul-Christian. 2019. *Brms: Bayesian Regression Models Using 'Stan'*. https://CRAN.R-project.org/package=brms.

Carpenter, Bob, Andrew Gelman, Matthew D Hoffman, Daniel Lee, Ben Goodrich, Michael Betancourt, Marcus Brubaker, Jiqiang Guo, Peter Li, and Allen Riddell. 2017. "Stan: A Probabilistic Programming Language." *Journal of Statistical Software* 76 (1). Columbia Univ., New York, NY (United States); Harvard Univ., Cambridge, MA (United States).

Goodrich, Ben, Jonah Gabry, Imad Ali, and Sam Brilleman. 2018. "Rstanarm: Bayesian Applied Regression Modeling via Stan."