Introduction to Bayesian statistical modelling

A course with R, Stan, and brms

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Planning

Course n°01: Introduction to Bayesian inference, Beta-Binomial model

Course n°02: Introduction to brms, linear regression

Course n°03: Markov Chain Monte Carlo, generalised linear model

Course n°04: Multilevel models, cognitive models



Setting up a common language

```
y_i \sim \text{Normal}(\mu_i, \sigma)

\mu_i = \alpha + \beta x_i

\alpha \sim \text{Normal}(60, 10)

\beta \sim \text{Normal}(0, 10)

\sigma \sim \text{HalfCauchy}(0, 1)
```

Aim of this session: to understand this type of model.

The components of our models will always be the same and we will always follow the following three steps:

- Build the model (likelihood + priors).
- Update with data to calculate the posterior distribution.
- Interpret the model's estimates and evaluate its predictions. If necessary, modify the model.



A first model



A first model

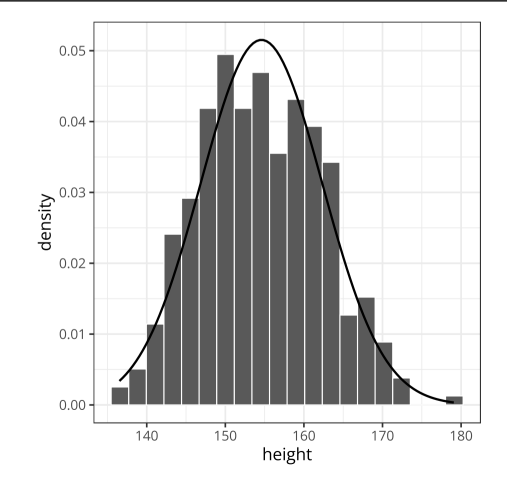
```
1 library(tidyverse)
  2 library(imsb)
  3
  4 d <- open data(howell)
  5 str(d)
 data.frame': 544 obs. of 4 variables:
 $ height: num 152 140 137 157 145 ...
 $ weight: num 47.8 36.5 31.9 53 41.3 ...
 $ age : num 63 63 65 41 51 35 32 27 19 54 ...
 $ male : int 1 0 0 1 0 1 0 1 0 1 ...
  1 d2 <- d %>% filter(age >= 18)
  2 head(d2)
   height weight age male
1 151.765 47.82561 63
2 139.700 36.48581 63
3 136.525 31.86484 65
4 156.845 53.04191 41
5 145.415 41.27687 51
6 163.830 62.99259 35
```



A first model

$h_i \sim \text{Normal}(\mu, \sigma)$

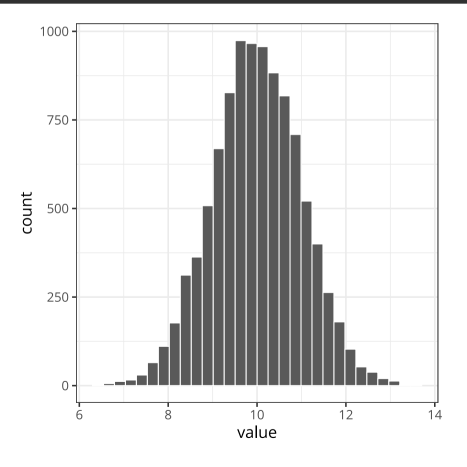
```
1 d2 %>%
2    ggplot(aes(x = height) ) +
3    geom_histogram(aes(y = ..density..), bins = 20, col = "white") +
4    stat_function(fun = dnorm, args = list(mean(d2$height), sd(d2$height) ), size = 1)
```





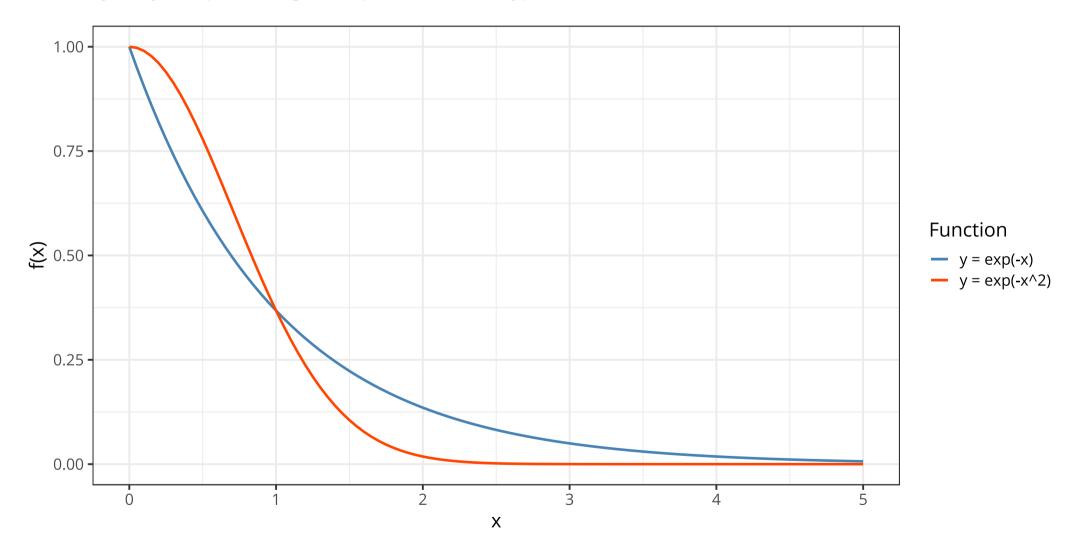
Normal distribution

$$p(x \mid \mu, \sigma) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{1}{2\sigma^2}(\mu - x)^2\right]$$





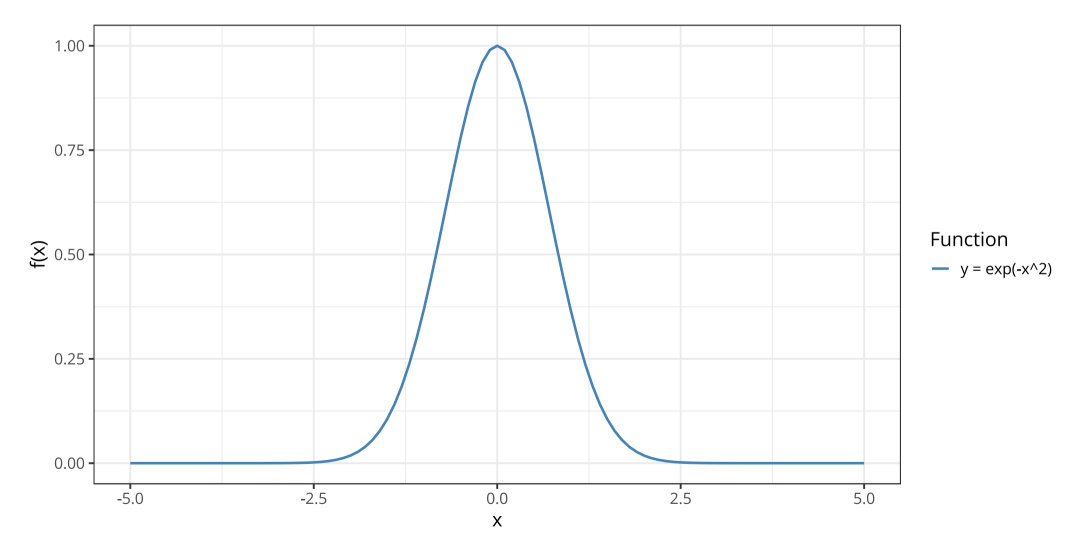
Some constraints: Some values are highly probable (around the mean μ). The further away they are, the less likely they are (following an exponential decay).





$$y = \exp\left[-x^2\right]$$

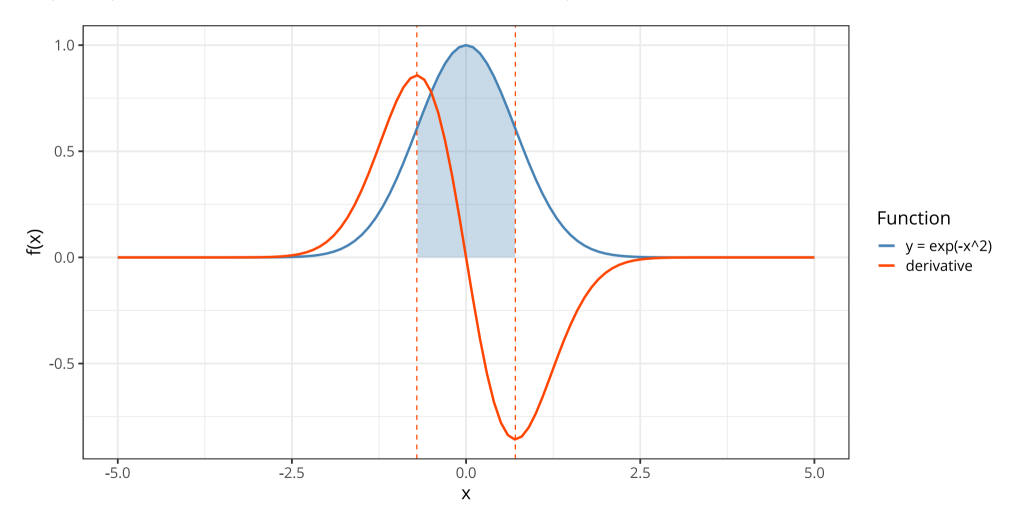
We extend our function to negative values.





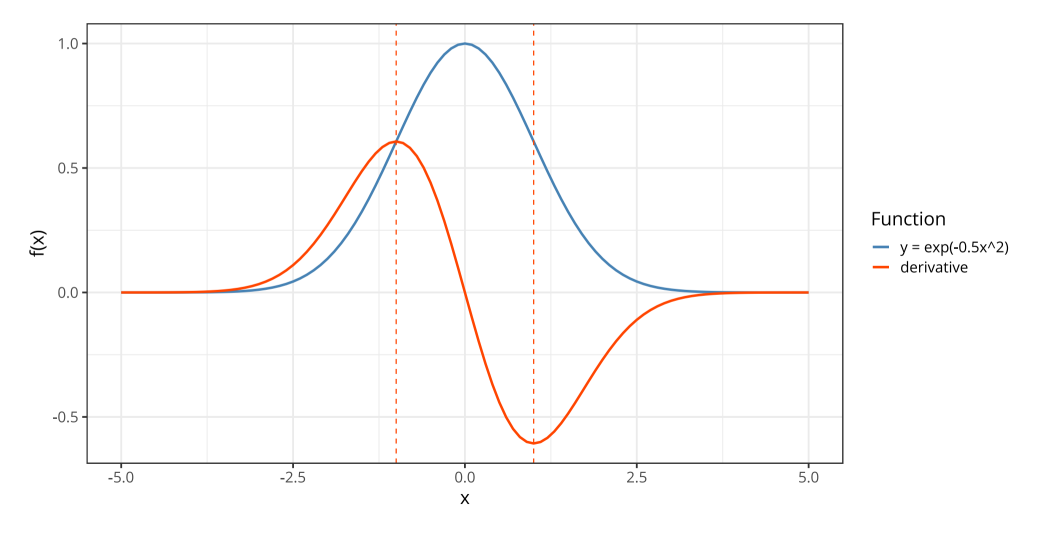
$$y = \exp\left[-x^2\right]$$

The inflection points give us a good indication of where most of the values lie (i.e., between the inflection points). The peaks of the derivative show us the inflection points.



$$y = \exp\left[-\frac{1}{2}x^2\right]$$

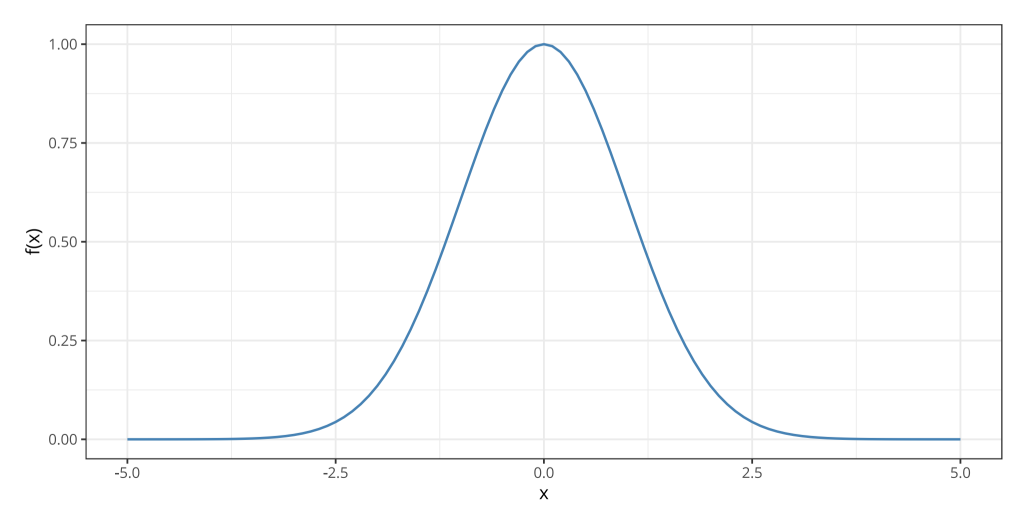
Next, we standardise the distribution so that the two inflection points are located at x = -1 and x = 1.





$$y = \exp\left[-\frac{1}{2\sigma^2}x^2\right]$$

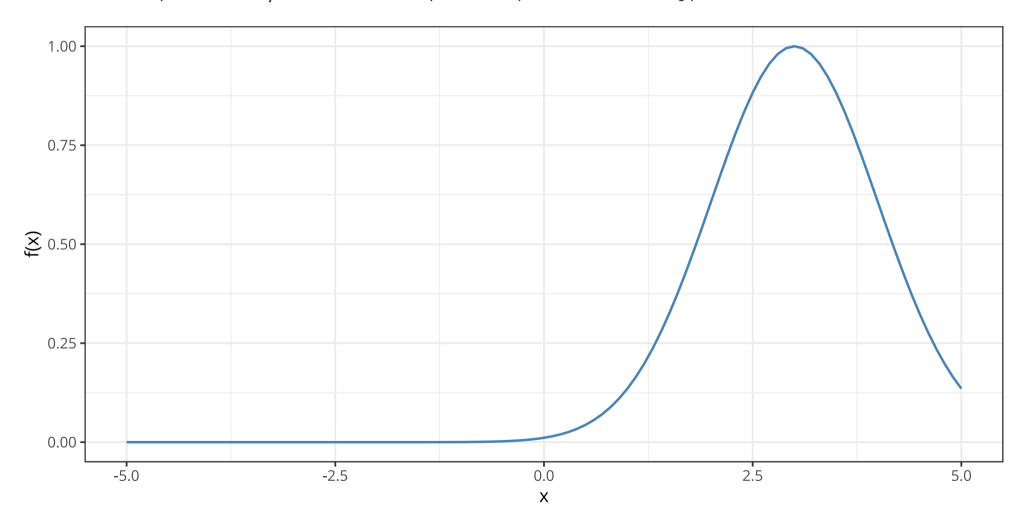
We then add a parameter σ^2 to control the distance between the inflection points.





$$y = \exp\left[-\frac{1}{2\sigma^2}(\mu - x)^2\right]$$

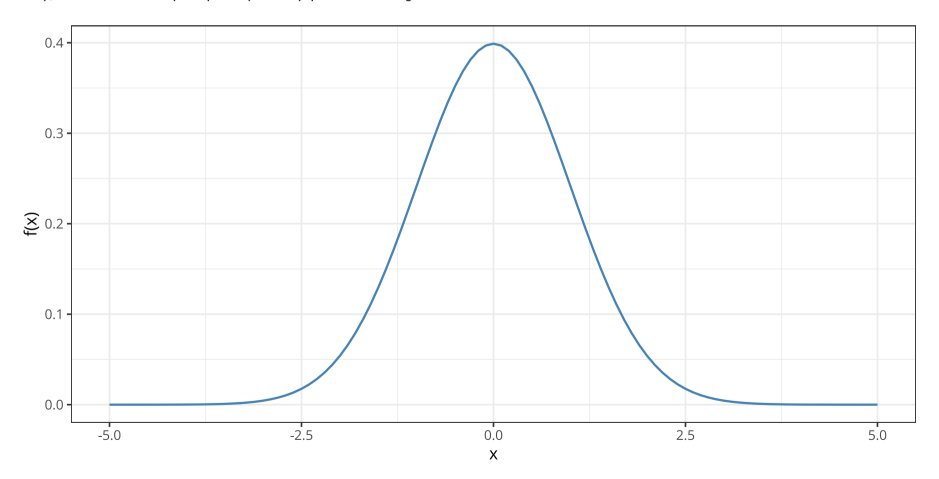
We then insert a parameter μ to control the position (central tendency) of the distribution.





$$y = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{1}{2\sigma^2}(\mu - x)^2\right]$$

But... this distribution does not integrate to 1. We therefore divide by a normalisation constant (the left-hand side), to obtain a proper (valid) probability distribution.





Gaussian model

We are going to build a regression model, but before we add a predictor, let's try to model the distribution of heights.

We want to know which model (distribution) best describes the distribution of heights. We will therefore explore all the possible combinations of μ and σ and rank them by their respective probabilities.

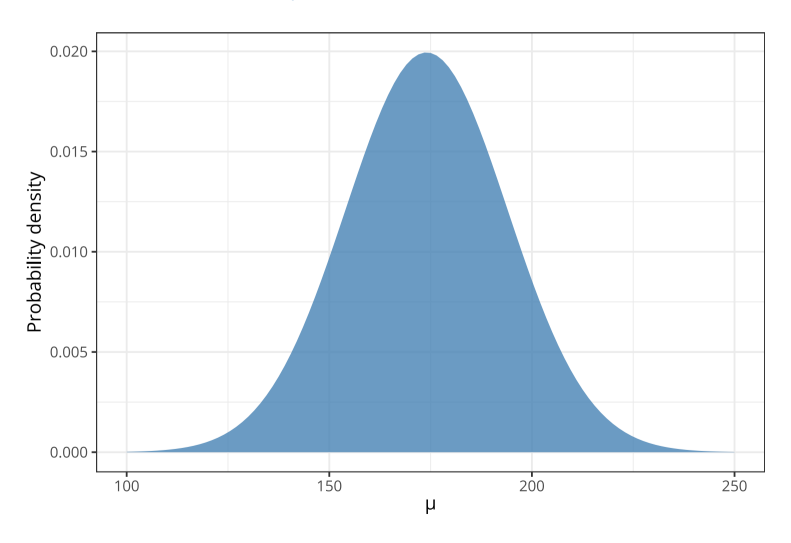
Our aim, once again, is to describe **the posterior distribution**, which will therefore be in some way **a distribution of distributions**.



Gaussian model

We define $p(\mu, \sigma)$, the joint prior distribution of all model parameters. These priors can be specified independently for each parameter, given that $p(\mu, \sigma) = p(\mu)p(\sigma)$.

$$\mu \sim \text{Normal}(174, 20)$$

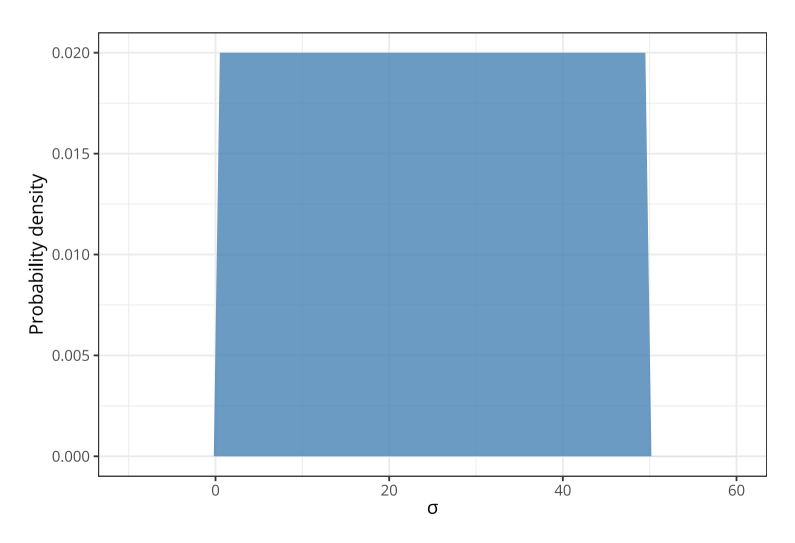




Gaussian model

We define $p(\mu, \sigma)$, the joint prior distribution of all model parameters. These priors can be specified independently for each parameter, given that $p(\mu, \sigma) = p(\mu)p(\sigma)$.

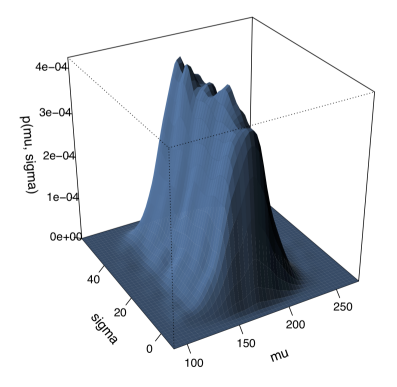
 $\sigma \sim \text{Uniform}(0, 50)$





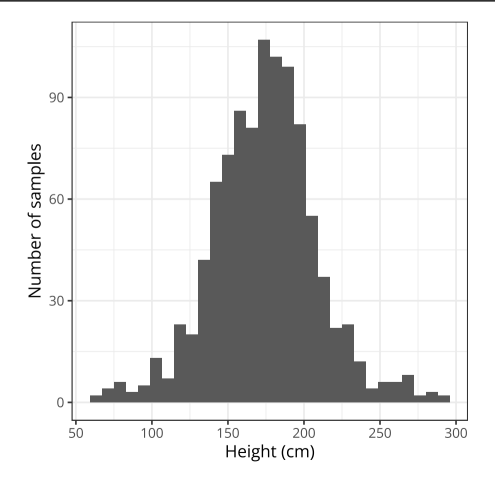
Visualiser le prior

```
1 library(ks)
2 sample_mu <- rnorm(le4, 174, 20) # prior on mu
3 sample_sigma <- runif(le4, 0, 50) # prior on sigma
4 prior <- data.frame(cbind(sample_mu, sample_sigma)) # multivariate prior
5 H.scv <- Hscv(x = prior, verbose = TRUE)
6 fhat_prior <- kde(x = prior, H = H.scv, compute.cont = TRUE)
7 plot(
8     fhat_prior, display = "persp", col = "steelblue", border = NA,
9     xlab = "\nmu", ylab = "\nsigma", zlab = "\n\np(mu, sigma)",
10     shade = 0.8, phi = 30, ticktype = "detailed",
11     cex.lab = 1.2, family = "Helvetica")</pre>
```





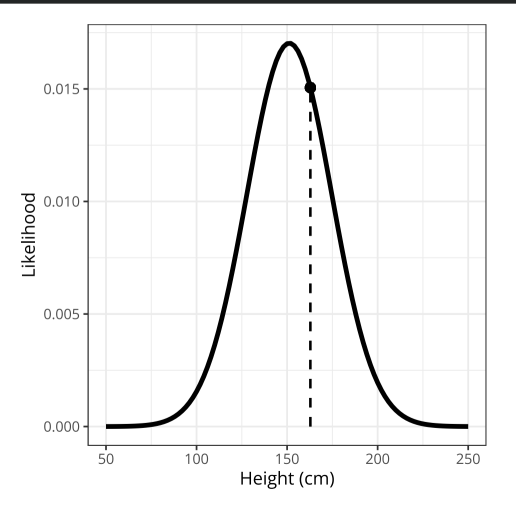
Prior predictive checking





Likelihood function

```
1 mu_exemple <- 151.23
2 sigma_exemple <- 23.42
3
4 d2$height[34] # exemplary height observation</pre>
[1] 162.8648
```





Likelihood function

We want to compute the probability of observing a certain value of height knowing certain values of μ and σ , that is:

$$p(x \mid \mu, \sigma) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{1}{2\sigma^2}(\mu - x)^2\right]$$

This **probability density** can be computed using the functions dnorm, dbeta, dt, dexp, dgamma, etc.

1 dnorm(d2\$height[34], mu_exemple, sigma_exemple)

[1] 0.01505675



Likelihood function

$$p(x \mid \mu, \sigma) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{1}{2\sigma^2}(\mu - x)^2\right]$$

Or using a custom function...

```
1 normal_likelihood <- function (x, mu, sigma) {
2
3   bell <- exp( (- 1 / (2 * sigma^2) ) * (mu - x)^2 )
4   norm <- sqrt(2 * pi * sigma^2)
5   return(bell / norm)
7   8 }
1 normal_likelihood(d2$height[34], mu_exemple, sigma_exemple)</pre>

[1] 0.01505675
```



Posterior distribution

$$p(\mu, \sigma \mid h) = \frac{\prod_{i} \text{Normal}(h_i \mid \mu, \sigma) \text{Normal}(\mu \mid 174, 20) \text{Uniform}(\sigma \mid 0, 50)}{\int \int \prod_{i} \text{Normal}(h_i \mid \mu, \sigma) \text{Normal}(\mu \mid 174, 20) \text{Uniform}(\sigma \mid 0, 50) d\mu d\sigma}$$
$$p(\mu, \sigma \mid h) \propto \prod_{i} \text{Normal}(h_i \mid \mu, \sigma) \text{Normal}(\mu \mid 174, 20) \text{Uniform}(\sigma \mid 0, 50)$$

This is the same formula from the previous course, but here considering that there are several observations of height (h_i) and two parameters to be estimated: μ and σ .

To compute the **marginal likelihood** (in green), we therefore need to integrate over two parameters: μ and σ . Here again we realise that the posterior is proportional to the product of the likelihood and the prior.



Posterior distribution - Grid approximation

```
1 # we define a grid of possible values for mu and sigma
2 mu.list <- seg(from = 140, to = 160, length.out = 200)
3 sigma.list \leftarrow seq(from = 4, to = 9, length.out = 200)
5 # we extend this grid to all possible combinations of mu and sigma
6 post <- expand.grid(mu = mu.list, sigma = sigma.list)</pre>
   # we compute the log-likelihood for each observation (under each combination of mu and sigma)
   post$LL <-
     sapply(
10
       1:nrow(post),
11
12
       function(i) sum(dnorm(
13
         d2$height,
14
        mean = post$mu[i],
         sd = post$sigma[i],
15
16
         log = TRUE))
17
18
   # we compute the (unnormalised) posterior distribution
20
   post$prod <-</pre>
21
     post$LL +
     dnorm(x = post$mu, mean = 174, sd = 20, log = TRUE) +
22
23
     dunif(x = post\$sigma, min = 0, max = 50, log = TRUE)
24
25
```



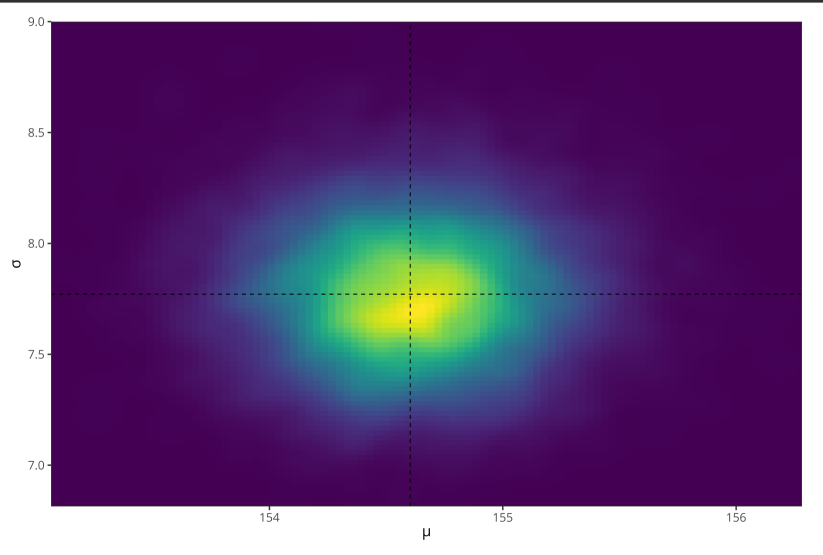
Posterior distribution - Grid approximation

```
1 # randomly selecting 20 rows from the resulting dataframe
  2 post %>% slice sample(n = 20, replace = FALSE)
               sigma
                            LL
                                    prod
                                                  prob
         mu
   155.1759 4.251256 -1418.233 -1426.502
                                         4.612132e-87
  143.2161 6.211055 -1829.978 -1838.989 3.334659e-266
  156.6834 6.462312 -1250.548 -1258.749
                                         3.297048e-14
  155.8794 6.537688 -1237.281 -1245.518
                                         1.837911e-08
  141.0050 8.221106 -1701.767 -1710.955 1.342455e-210
  159.0955 8.246231 -1273.181 -1281.286 5.378841e-24
   142.6131 4.728643 -2471.268 -2480.326 0.000000e+00
  146.3317 5.507538 -1667.237 -1676.021 1.993230e-195
  160.0000 6.788945 -1337.375 -1345.446 7.347547e-52
10 143.4171 7.517588 -1608.947 -1617.943 3.329394e-170
11 140.6030 6.437186 -2064.584 -2073.805 0.000000e+00
12 151.6583 5.005025 -1370.982 -1379.432
                                         1.276988e-66
13 156.2814 6.814070 -1236.277 -1244.496 5.108761e-08
14 153.7688 8.849246 -1226.826 -1235.164 5.767419e-04
15 144.1206 7.015075 -1615.495 -1624.437 5.033474e-173
16 143.7186 8.572864 -1506.316 -1515.289 1.271644e-125
17 154.8744 7.517588 -1219.926 -1228.210 6.039964e-01
18 146.3317 8.824121 -1379.469 -1388.252 1.886826e-70
19 140.0000 7.844221 -1828.939 -1838.211 7.264600e-266
20 159.8995 5.180905 -1478.782 -1486.857 2.832570e-113
```



Posterior distribution - Grid approximation

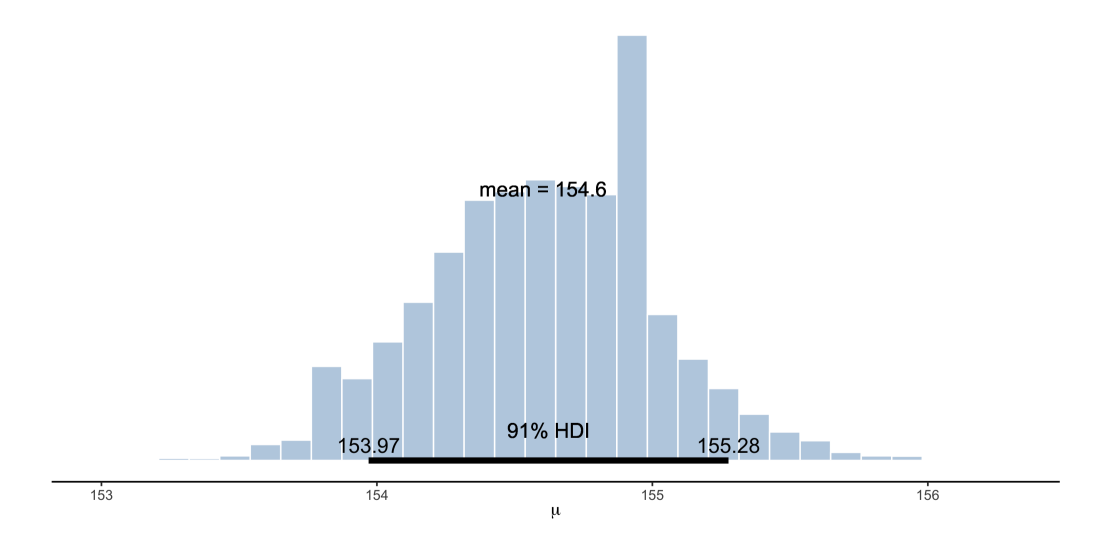
```
1 sample.rows <- sample(x = 1:nrow(post), size = 1e4, replace = TRUE, prob = post$prob)
2 sample.mu <- post$mu[sample.rows]
3 sample.sigma <- post$sigma[sample.rows]</pre>
```





Posterior distribution - Marginal distributions

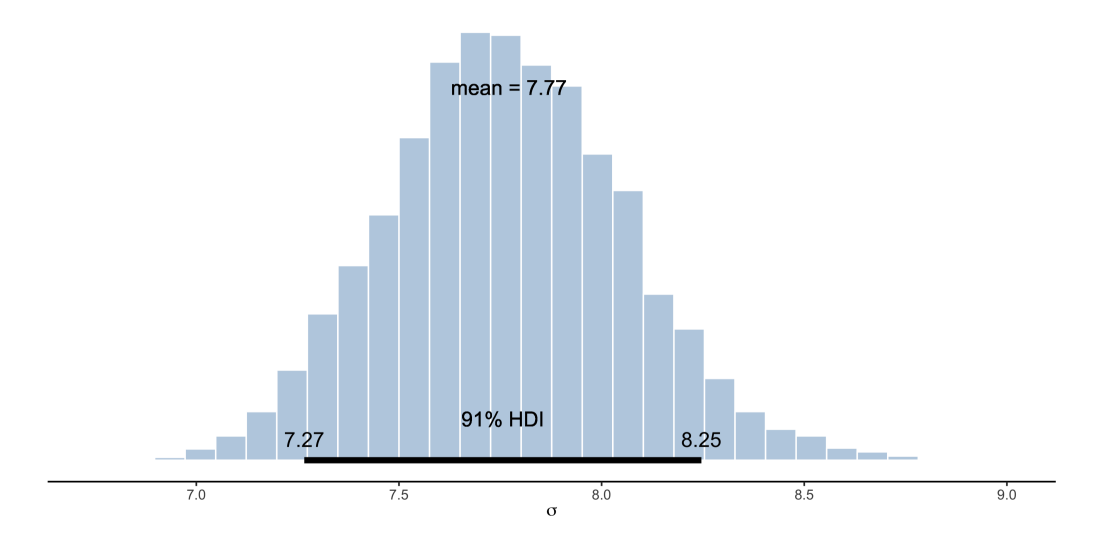
1 posterior_plot(samples = sample.mu, nbins = 30) + labs(x = expression(mu))





Posterior distribution - Marginal distributions

1 posterior_plot(samples = sample.sigma, nbins = 30) + labs(x = expression(sigma))





Introduction to brms



Introduction to brms

Under the hood: **Stan** is a probabilistic programming language written in **C++**, which implements several MCMC algorithms: HMC, NUTS, L-BFGS...

```
1 data {
     int<lower=0> J; // number of schools
     real y[J]; // estimated treatment effects
 3
     real<lower=0> sigma[J]; // s.e. of effect estimates
 4
 5
 6
   parameters {
 8
     real mu;
     real<lower=0> tau;
9
10
     real eta[J];
11
12
   transformed parameters {
14
     real theta[J];
     for (j in 1:J)
15
16
       theta[j] = mu + tau * eta[j];
17
18
19 model {
     target += normal lpdf(eta | 0, 1);
20
21
     target += normal lpdf(y | theta, sigma);
22
```



Bayesian regression models using Stan

The **brms** package (<u>Bürkner, 2017</u>) can be used to fit multilevel (or single-level) linear (or not) Bayesian regression models in **Stan** but using the intuitive syntax of **lme4** (cf. our tutorial paper, <u>Nalborczyk et al., 2019</u>).

For instance, the following model:

$$y_i \sim \text{Normal}(\mu_i, \sigma)$$

 $\mu_i = \alpha + \alpha_{\text{subject}[i]} + \alpha_{\text{item}[i]} + \beta x_i$

is specified with **brms** (as with **lme4**) as follows:

```
1 model <- brm(y ~ x + (1 | subject) + (1 | item), data = d, family = gaussian() )
```



Syntax reminders

The **brms** package uses the same syntax as the base R functions (such as lm) or functions from the lme4 package.

```
1 Reaction ~ Days + (1 + Days | Subject)
```

The left-hand side represents our dependent variable (or outcome, that is, what we are trying to predict). The **brms** package can also be used to fit multivariate models (several outcomes) by combining them with **mvbind()**.

```
1 mvbind(Reaction, Memory) ~ Days + (1 + Days | Subject)
```

The right-hand side is used to define the predictors. The intercept is is generally implicit, so that the two formulations below are equivalent.

```
1 mvbind(Reaction, Memory) ~ Days + (1 + Days | Subject)
2 mvbind(Reaction, Memory) ~ 1 + Days + (1 + Days | Subject)
```



Syntax reminders

If you want to fit a model without an intercept (because why not), you must specify it explicitly, as shown below.

```
1 mvbind(Reaction, Memory) ~ 0 + Days + (1 + Days | Subject)
```

By default **brms** assumes a Gaussian likelihood function. This assumption can be modified easily by specifying the desired likelihood via the the **family** argument.

```
1 brm(Reaction ~ 1 + Days + (1 + Days | Subject), family = lognormal())
```

In case of doubt, read the documentation (it's very exciting to read) available at ?brm.



Some useful functions

```
1 # retrieving the Stan code generated by brms
 2 make stancode(formula, ...)
 3 stancode(fit)
 5 # checking and/or defining priors
 6 get prior(formula, ...)
   set prior(prior, ...)
 9 # retrieving predictions from a brms model
10 fitted(fit, ...)
11 predict(fit, ...)
12 conditional effects(fit, ...)
13
14 # posterior predictive checking
15 pp check(fit, ...)
16
17 # model comparison and hypothesis testing
18 loo(fit1, fit2, ...)
19 bayes factor(fit1, fit2, ...)
20 model weights(fit1, fit2, ...)
21 hypothesis(fit, hypothesis, ...)
```



A first example

```
1 library(brms)
2 mod1 <- brm(height ~ 1, data = d2)

1 posterior_summary(mod1, pars = c("^b_", "sigma"), probs = c(0.025, 0.975))

Estimate Est.Error Q2.5 Q97.5
b_Intercept 154.614878 0.4224360 153.787631 155.452195
sigma 7.763965 0.2936431 7.217446 8.354354</pre>
```

These data represent the marginal (posterior) distributions of each parameter. In other words, the posterior probability μ , averaged over all possible values of sigma, is best described by a Gaussian distribution with a mean of 154.6 and a standard deviation of 0.42. The credibility interval (\neq confidence interval) indicates the 95% most plausible values of μ or σ (given the data and priors).



Using our prior

By default **brms** uses a very uninformative prior centred on the mean value of the measured variable. We can therefore refine the estimate using our knowledge of the usual distribution of heights in humans.

The **get_prior()** function is used to display a list of default priors as well as all the priors we can specify, given a certain model formula (i.e., a way of writing our model) and a set of data.



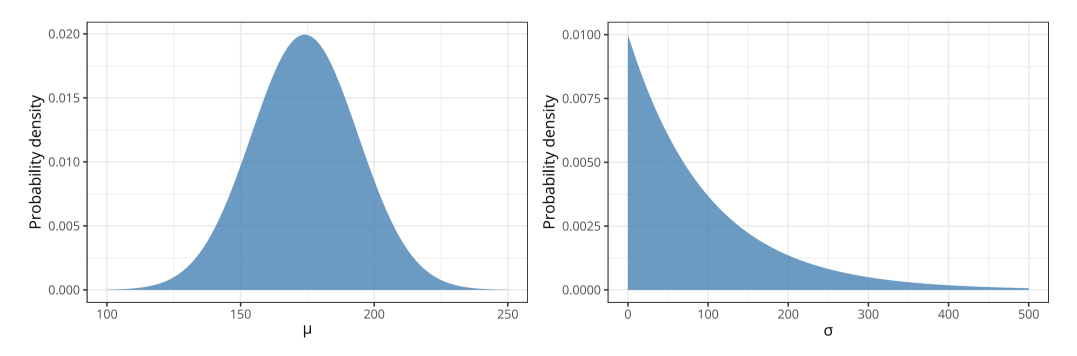
Using our prior

```
priors <- c(
prior(normal(174, 20), class = Intercept),
prior(exponential(0.01), class = sigma)

4  )

6  mod2 <- brm(
height ~ 1,
prior = priors,
family = gaussian(),
data = d2

11  )</pre>
```





Using our prior

```
1 summary(mod2)
 Family: gaussian
 Links: mu = identity; sigma = identity
Formula: height ~ 1
   Data: d2 (Number of observations: 352)
  Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
         total post-warmup draws = 4000
Population-Level Effects:
         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                       0.41 153.80 155.42 1.00
Intercept 154.60
                                                       3433
                                                                2656
Family Specific Parameters:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
         7.76
                                     8.39 1.00
                                                            2499
sigma
                   0.30
                            7.23
                                                   3032
Draws were sampled 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).
```



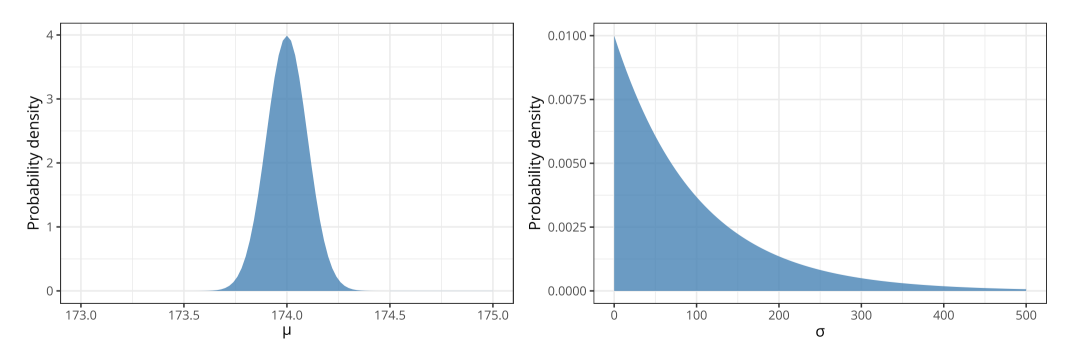
Using a more informative prior

```
priors <- c(
prior(normal(174, 0.1), class = Intercept),
prior(exponential(0.01), class = sigma)

4  )

6  mod3 <- brm(
height ~ 1,
prior = priors,
family = gaussian(),
data = d2

11  )</pre>
```





Using a more informative prior

```
1 summary(mod3)
 Family: gaussian
 Links: mu = identity; sigma = identity
Formula: height ~ 1
   Data: d2 (Number of observations: 352)
  Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
         total post-warmup draws = 4000
Population-Level Effects:
          Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                        0.10 173.65 174.03 1.00
Intercept 173.84
                                                        3108
                                                                 2389
Family Specific Parameters:
      Estimate Est. Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
         20.81
                    0.79
                            19.33
                                     22.48 1.00
                                                    2908
                                                             2312
sigma
Draws were sampled 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).
```

We note that the estimated value for μ has hardly "moved" from the prior…but we can also see that the estimated value for σ has greatly increased. What happened? We told the model that we were fairly certain of our μ value, the model then "adapted", which explains the large value of σ …



Prior precision (heuristic)

Prior distributions can generally be considered as posterior distributions obtained from previous data.

We know that the σ of a Gaussian posterior is given by:

$$\sigma_{\rm post} = 1 / \sqrt{n}$$

Which implies a **quantity of data** n=1 / σ_{post}^2 . Our prior had a $\sigma=0.1$, which implies n=1 / $0.1^2=100$.

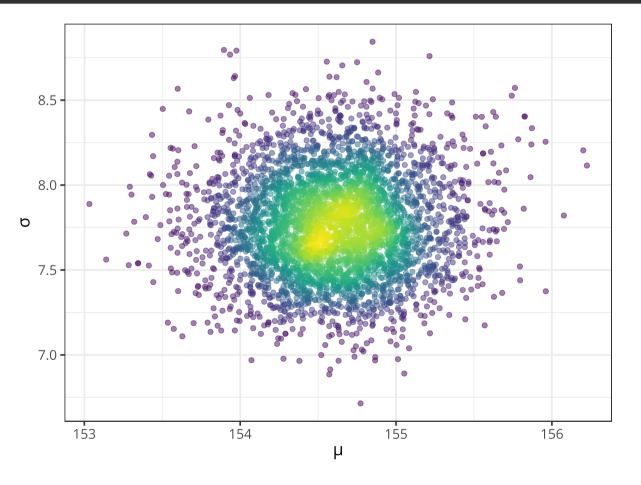
We can therefore consider that the prior Normal(174, 0.1) is equivalent to the case in which we would have observed 100 heights with mean 174.



Visualising samples from the posterior distribution

```
post <- as_draws_df(x = mod2) %>%
mutate(density = get_density(b_Intercept, sigma, n = 1e2))

ggplot(post, aes(x = b_Intercept, y = sigma, color = density)) +
geom_point(size = 2, alpha = 0.5, show.legend = FALSE) +
labs(x = expression(mu), y = expression(sigma)) +
viridis::scale_color_viridis()
```





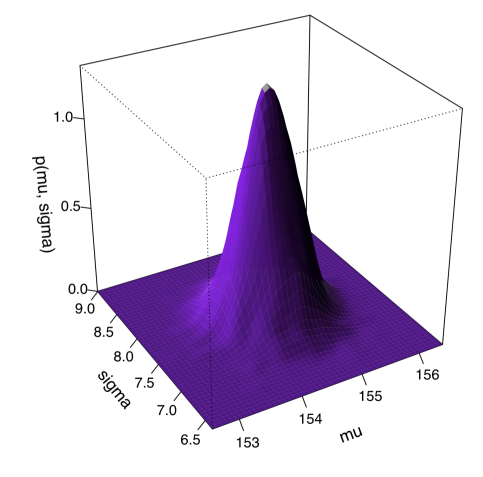
Retrieving samples from the posterior distribution

```
1 # gets the first 6 samples
  2 head(post)
# A draws df: 6 iterations, 1 chains, and 5 variables
  b Intercept sigma lprior lp density
          154
               8.2
                     -9.1 -1229
                                  0.082
                    -9.0 -1229
                                  0.150
          155
              7.2
                    -9.0 -1229
                                  0.139
         155
             8.0
                    -9.1 -1227
                                  0.582
         154
             8.3
                    -9.1 -1228
                                  0.263
         155
         155
              7.6
                    -9.0 -1228
                                  0.215
  ... hidden reserved variables { '.chain', '.iteration', '.draw'}
  1 # gets the median and the 95% credible interval
  2 t(sapply(post[, 1:2], quantile, probs = c(0.025, 0.5, 0.975))
                 2.5%
                            50%
                                     97.5%
b Intercept 153.795472 154.59952 155.418792
sigma
             7.227358
                        7.74985
                                  8.386699
```



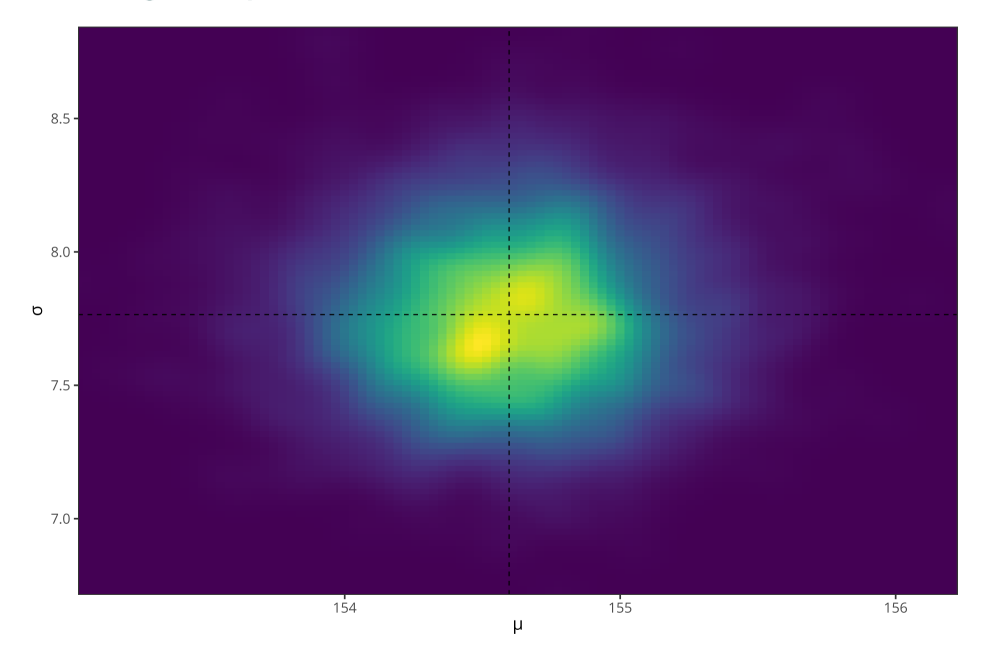
Visualising the posterior distribution

```
1 H.scv <- Hscv(post[, 1:2])
2 fhat_post <- kde(x = post[, 1:2], H = H.scv, compute.cont = TRUE)
3
4 plot(fhat_post, display = "persp", col = "purple", border = NA,
5     xlab = "\nmu", ylab = "\nsigma", zlab = "\np(mu, sigma)",
6     shade = 0.8, phi = 30, ticktype = "detailed",
7     cex.lab = 1.2, family = "Helvetica")</pre>
```





Visualising the posterior distribution

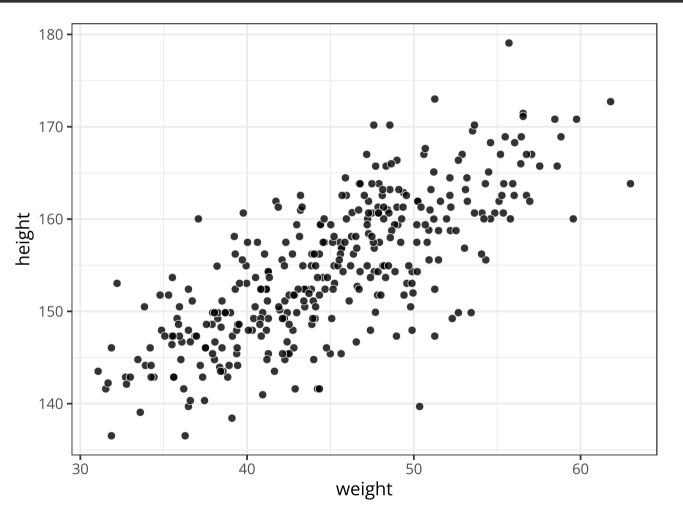




Including predictors

How does height change with weight?

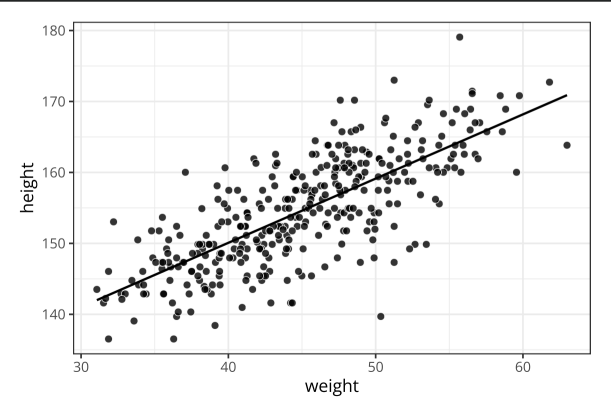
```
1 d2 %>%
2 ggplot(aes(x = weight, y = height) ) +
3 geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8)
```





$$h_i \sim \text{Normal}(\mu_i, \sigma)$$

 $\mu_i = \alpha + \beta x_i$





Notations

Let's consider a linear regression model with a single predictor, a slope, an intercept, and residuals distributed according to a normal distribution. The notation:

$$h_i = \alpha + \beta x_i + \epsilon_i$$
 avec $\epsilon_i \sim \text{Normal}(0, \sigma)$

is equivalent to:

$$h_i - (\alpha + \beta x_i) \sim \text{Normal}(0, \sigma)$$

rearranging the expression:

$$h_i \sim \text{Normal}(\alpha + \beta x_i, \sigma).$$

The above notations are equivalent, but the last one is more flexible, and will allow us to extend it more intuitively to multilevel models.



```
h_i \sim \text{Normal}(\mu_i, \sigma)

\mu_i = \alpha + \beta x_i

\alpha \sim \text{Normal}(174, 20)

\beta \sim \text{Normal}(0, 10)

\sigma \sim \text{Exponential}(0.01)
```

In this model, μ is no longer a parameter to be estimated (because μ is **determined** by α and β). Instead, we will estimate α and β .

Reminders: α is the intercept, that is, the expected height, when the weight is equal to 0. β is the slope, that is, the expected change in height when weight increases by one unit.



```
1 priors <- c(
2  prior(normal(174, 20), class = Intercept),
3  prior(normal(0, 10), class = b),
4  prior(exponential(0.01), class = sigma)
5  )
6
7  mod4 <- brm(
8  height ~ 1 + weight,
9  prior = priors,
10  family = gaussian(),
11  data = d2
12 )</pre>
```



```
1 posterior summary(mod4)
                                              02.5
                                                            097.5
                 Estimate Est.Error
b Intercept
              113.9022663 1.89453254
                                       110.2751443
                                                      117.5863999
b weight
                0.9045325 0.04172372
                                         0.8226575
                                                        0.9834808
sigma
                5.1023966 0.19607421
                                         4.7309481
                                                        5.4967173
lprior
              -12.2671372 0.01272490
                                       -12.2925817
                                                      -12.2432602
            -1083.1151428 1.14546650 -1086.1172584 -1081.7573733
lp
```

- $\alpha = 113.90, 95\%$ CrI [110.28, 117.59] represents the expected height when weight equals 0kg...
- $\beta = 0.90, 95\%$ CrI [0.82, 0.98] indicates that for an increase of 1kg, we can expect an increase of 0.90cm.



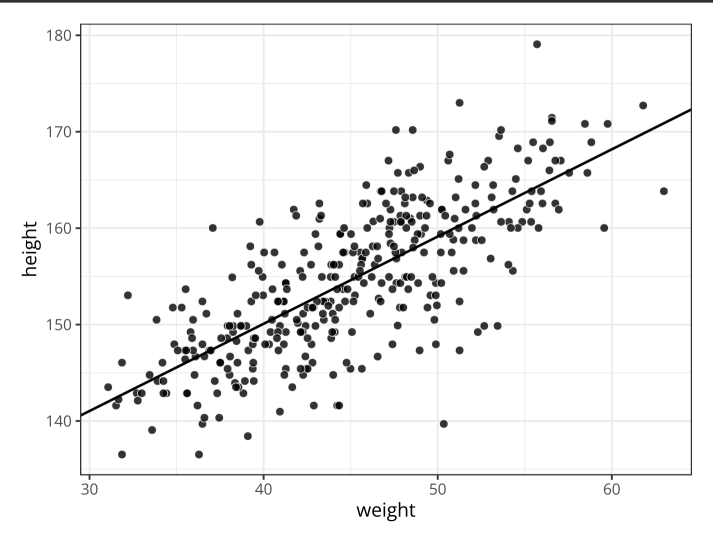
```
d2$weight.c <- d2$weight - mean(d2$weight)</pre>
  2
  3 mod5 <- brm(</pre>
      height ~ 1 + weight.c,
      prior = priors,
      family = gaussian(),
      data = d2
  8
  1 fixef(mod5) # retrieving the fixed effects estimates
             Estimate Est.Error
                                         02.5
                                                     097.5
Intercept 154.5892334 0.26511032 154.0580097 155.1021441
weight.c
            0.9048262 0.04247021
                                    0.8194467
                                                 0.9874537
```

If the predictor (weight) is centred, the intercept represents the expected value of height (in cm) when weight is at its mean value.



Visualising predictions from the model

```
1 d2 %>%
2    ggplot(aes(x = weight, y = height) ) +
3    geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4    geom_abline(intercept = fixef(mod4)[1], slope = fixef(mod4)[2], lwd = 1)
```





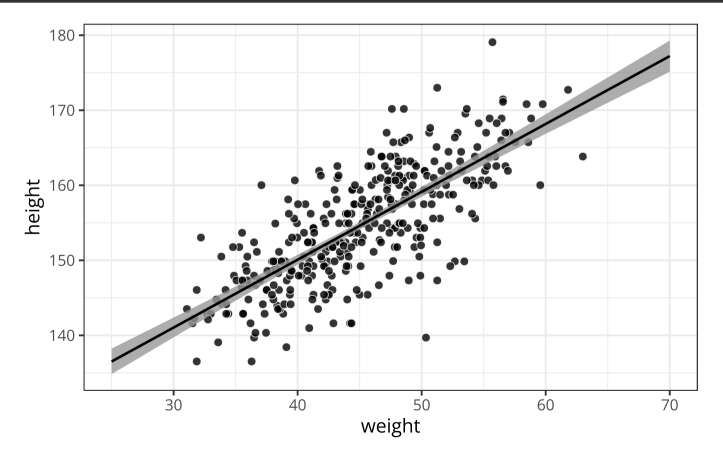
Visualising the uncertainty on μ using fitted()

```
1 # defining a grid of possible values for "weight"
  2 weight.seg <- data.frame(weight = seg(from = 25, to = 70, by = 1) )</pre>
  3
  4 # retrieving the model's predicted mus (average height) for these values of "weight"
  5 mu <- data.frame(fitted(mod4, newdata = weight.seq) ) %>% bind cols(weight.seq)
    # displaying the first 10 rows of mu
  8 head(mu, 10)
                          02.5
                                  097.5 weight
   Estimate Est.Error
 136.5156 0.8729446 134.8388 138.2282
                                            25
2 137.4201 0.8331902 135.8118 139.0699
                                            26
  138.3246 0.7936381 136.7802 139.8924
                                            27
                                            28
  139.2292 0.7543200 137.7739 140.7197
  140.1337 0.7152745 138.7559 141.5451
                                            29
  141.0382 0.6765488 139.7387 142.3767
                                            30
  141.9428 0.6382011 140.7096 143.1942
                                            31
  142.8473 0.6003039 141.6932 144.0216
                                            32
  143.7518 0.5629482 142.6682 144.8488
                                            33
10 144.6564 0.5262493 143.6445 145.6845
                                            34
```



Visualising the uncertainty on μ using fitted()

```
1 d2 %>%
2  ggplot(aes(x = weight, y = height)) +
3  geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4  geom_smooth(
5  data = mu, aes(y = Estimate, ymin = Q2.5, ymax = Q97.5),
6  stat = "identity",
7  color = "black", alpha = 0.8, size = 1
8  )
```





Prediction intervals (incorporating σ)

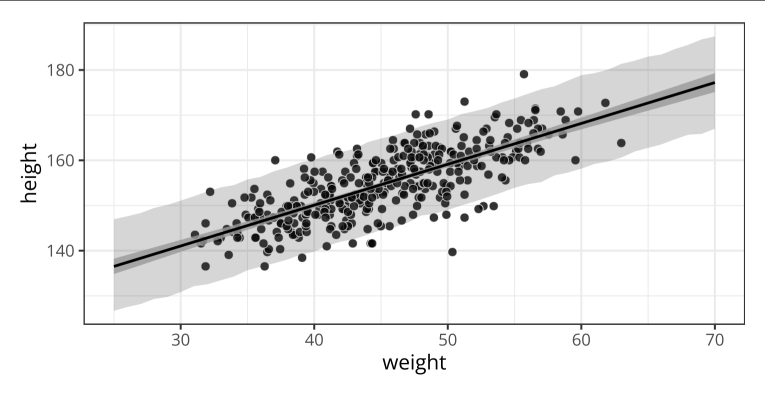
As a reminder, our model is defined as: $h_i \sim \text{Normal}(\alpha + \beta x_i, \sigma)$. For the moment, we have only represented the predictions for μ . How can we incorporate σ into our predictions?

```
1 # defining a grid of possible values for "weight"
 2 weight.seg <- data.frame(weight = seg(from = 25, to = 70, by = 1) )</pre>
  3
    # retrieving the model's predicted heights for these values of "weight"
    pred height <- data.frame(predict(mod4, newdata = weight.seq) ) %>% bind cols(weight.seq)
    # displaying the first 10 rows of pred height
 8 head(pred height, 10)
   Estimate Est.Error
                         02.5
                                 097.5 weight
  136.6003 5.241255 126.6961 146.9625
                                            25
  137.5269 5.123556 127.5401 147.6144
                                            26
  138.2405 5.176174 128.1914 148.3255
                                            27
 139.3099 5.243116 129.3009 149.4473
                                            28
  140.0189 5.168394 129.7397 150.1358
                                            29
 140.8688 5.218597 130.8407 151.0752
                                            30
  142.0739 5.152610 132.1567 152.0562
                                            31
  142.7756 5.218983 132.5639 153.1091
                                            32
  143.7965 5.244761 133.4364 154.0000
                                            33
10 144.6657 5.198395 134.3009 154.5210
                                            34
```



Prediction intervals (incorporating σ)

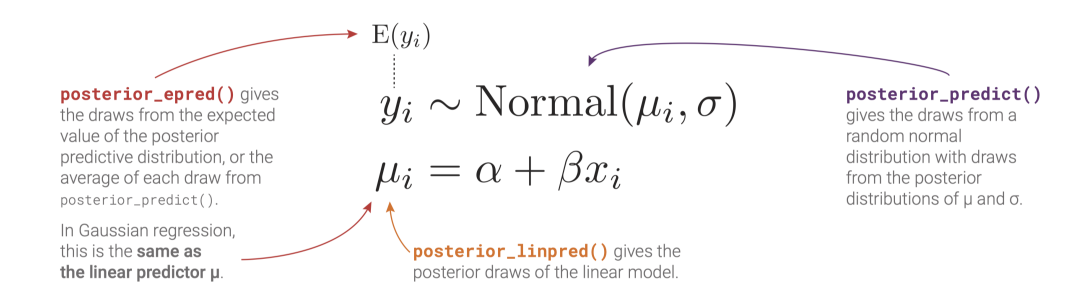
```
1 d2 %>%
2  ggplot(aes(x = weight, y = height)) +
3  geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4  geom_ribbon(
5  data = pred_height, aes(x = weight, ymin = Q2.5, ymax = Q97.5),
6  alpha = 0.2, inherit.aes = FALSE
7  ) +
8  geom_smooth(
9  data = mu, aes(y = Estimate, ymin = Q2.5, ymax = Q97.5),
10  stat = "identity", color = "black", alpha = 0.8, size = 1
11 )
```





Built-in brms functions

The brms package also includes the posterior_epred(), posterior_linpred(), and posterior_predict() functions, which can be used to generate predictions from models fitted with brms. Andrew Heiss gives a detailed description of how these functions work in this blog_post.





Reminder: Two types of uncertainty

Two sources of uncertainty in the model: uncertainty concerning the estimation of the value of the parameters but also uncertainty concerning the sampling process.

Epistemic uncertainty: The posterior distribution orders all possible combinations of parameter values according to their relative plausibility.

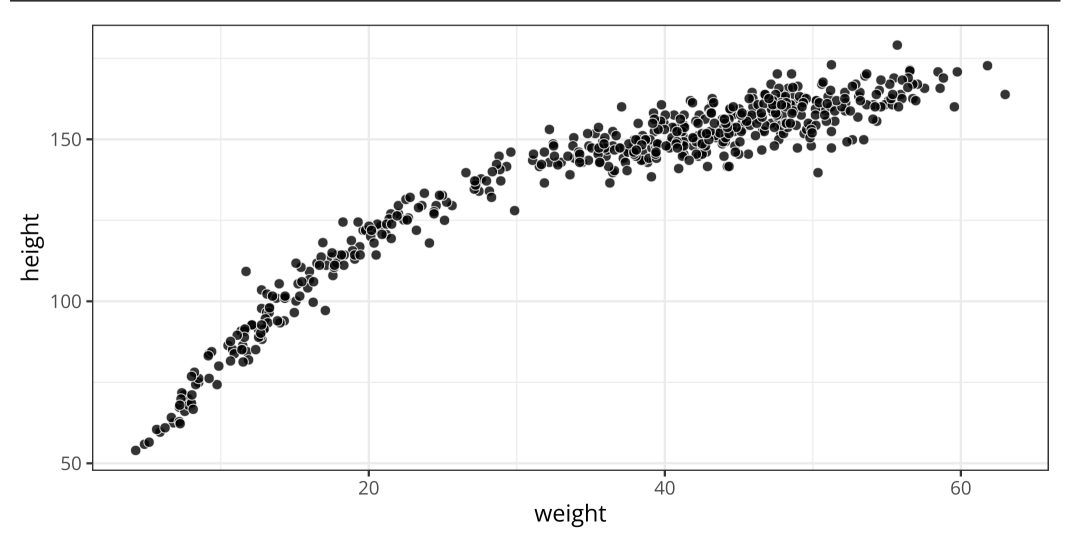
Aleatory uncertainty: The distribution of simulated data is a distribution that contains uncertainty related to a sampling process (i.e., generating data from a Gaussian distribution).

See also this short article by O'Hagan (2004).



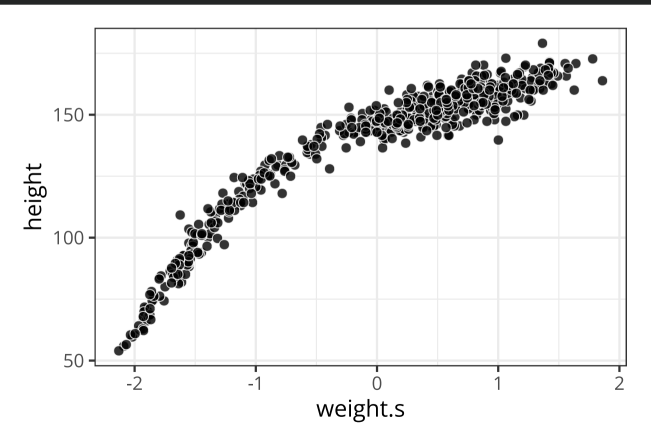
Polynomial regression

```
1 d %>% # note that we use d instead of d2
2 ggplot(aes(x = weight, y = height) ) +
3 geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8)
```





Standardised (scaled) scores





Standardised (scaled) scores

Why standardising predictors?

- **Interpretation**. Easier to compare the coefficients of several predictors. A change of one standard deviation in the predictor corresponds to to a change of one standard deviation in the response (if the response is also standardised).
- **Fitting**. When the predictors contain large values (or values that are too different from one another), this can lead to convergence problems (cf. Course n°03).



Polynomial regression - Exercise

```
h_i \sim \text{Normal}(\mu_i, \sigma)
\mu_i = \alpha + \beta_1 x_i + \beta_2 x_i^2
\alpha \sim \text{Normal}(156, 100)
\beta_1, \beta_2 \sim \text{Normal}(0, 10)
\sigma \sim \text{Exponential}(0.01)
```

It's up to you to build and fit this model using brms::brm().



Polynomial regression

```
1 priors <- c(</pre>
     prior(normal(156, 100), class = Intercept),
     prior(normal(0, 10), class = b),
 3
     prior(exponential(0.01), class = sigma)
   mod6 <- brm(</pre>
     # NB: polynomials should be written with the I() function...
     height ~ 1 + weight.s + I(weight.s^2),
 9
     prior = priors,
10
     family = gaussian(),
11
12
     data = d
13
```



Polynomial regression

```
1 summary(mod6)
 Family: gaussian
 Links: mu = identity; sigma = identity
Formula: height ~ 1 + weight.s + I(weight.s^2)
   Data: d (Number of observations: 544)
  Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
        total post-warmup draws = 4000
Population-Level Effects:
           Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
            146.65
                         0.37 145.95 147.39 1.00
Intercept
                                                        3438
                                                                 2694
weight.s
            21.40
                         0.29 20.84 21.99 1.00
                                                        3198
                                                                 2387
                         0.28 -8.96 -7.85 1.00
Iweight.sE2
              -8.41
                                                        3117
                                                                 2845
Family Specific Parameters:
     Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
                                     6.15 1.00
                                                  3352
                                                           3090
sigma
         5.78
                   0.17
                            5.46
Draws were sampled 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).
```



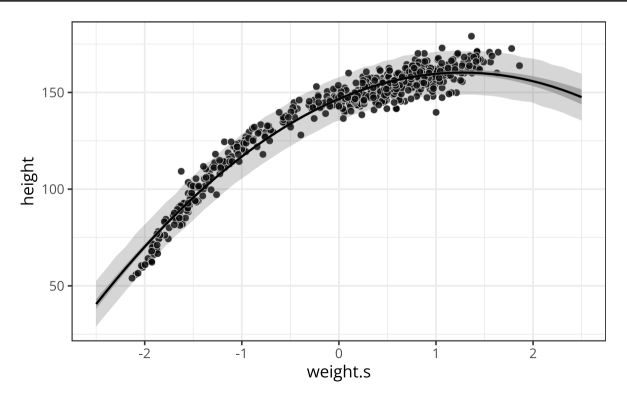
Visualising the model's predictions

```
1 # defining a grid of possible (standardised) values for "weight"
 2 weight.seg <- data.frame(weight.s = seg(from = -2.5, to = 2.5, length.out = 50))
  3
 4 # retrieving the model's predictions for these values of weight
  5 mu <- data.frame(fitted(mod6, newdata = weight.seq) ) %>% bind cols(weight.seq)
    pred height <- data.frame(predict(mod6, newdata = weight.seq) ) %>% bind cols(weight.seq)
 8 # displaying the first ten rows of pred height
  9 head(pred height, 10)
                         02.5
                                  Q97.5 weight.s
  Estimate Est.Error
  40.60063 6.010429 28.97013 52.51992 -2.500000
  47.10254 5.955673 35.32591 58.77250 -2.397959
  53.19186 6.006070 41.31088 65.14786 -2.295918
 59.29052 5.760499 47.61468 70.36520 -2.193878
  65.32630 5.963502 53.34686 76.94875 -2.091837
  70.76771 5.769429 59.88322 82.18180 -1.989796
  76.26657 5.815825 64.92816 87.47296 -1.887755
  81.64095 5.787103 70.60726 92.95473 -1.785714
 86.67635 5.665041 75.79001 97.45832 -1.683673
10 91.71143 5.880964 80.16754 103.33322 -1.581633
```



Visualising the model's predictions

```
1 d %>%
2    ggplot(aes(x = weight.s, y = height)) +
3    geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4    geom_ribbon(
5    data = pred_height, aes(x = weight.s, ymin = Q2.5, ymax = Q97.5),
6    alpha = 0.2, inherit.aes = FALSE
7    ) +
8    geom_smooth(
9    data = mu, aes(y = Estimate, ymin = Q2.5, ymax = Q97.5),
10    stat = "identity", color = "black", alpha = 0.8, size = 1
11    )
```





There are several methods for computing effect sizes in Bayesian models. Gelman & Pardoe (2006) propose a method for computing a sample-based R^2 .

Marsman & Wagenmakers (2017) and Marsman et al. (2019) generalise existing methods to compute a ρ^2 for ANOVA designs (i.e., with categorical predictors), which represents an estimate of the effect size in the population.

"

Similar to most of the ES measures that have been proposed for the ANOVA model, the squared multiple correlation coefficient ρ^2 [...] is a so-called proportional reduction in error measure (PRE). In general, a PRE measure expresses the proportion of the variance in an outcome y that is attributed to the independent variables x (Marsman et al., 2019).



$$\rho^{2} = \frac{\sum_{i=1}^{n} \pi_{i} (\beta_{i} - \beta)^{2}}{\sigma^{2} + \sum_{i=1}^{n} \pi_{i} (\beta_{i} - \beta)^{2}}$$

$$\rho^{2} = \frac{\frac{1}{n} \sum_{i=1}^{n} \beta_{i}^{2}}{\sigma^{2} + \frac{1}{n} \sum_{i=1}^{n} \beta_{i}^{2}}$$

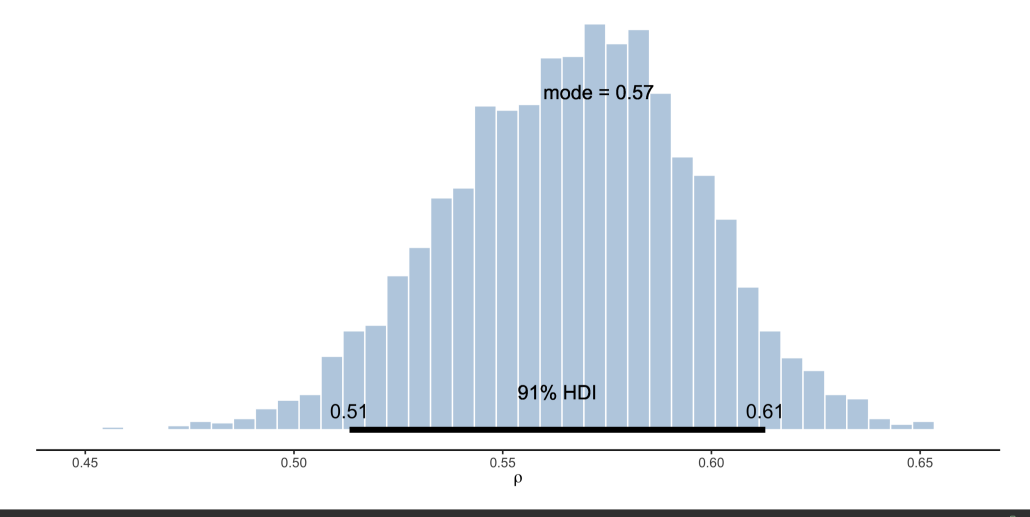
$$\rho^{2} = \frac{\beta^{2} \tau^{2}}{\sigma^{2} + \beta^{2} \tau^{2}}$$

```
post <- as_draws_df(x = mod4)
beta <- post$b_weight
sigma <- post$sigma
rho <- beta^2 * var(d2$weight) / (sigma^2 + beta^2 * var(d2$weight) )</pre>
```

Caution: if there are several predictors, it depends on the covariance structure...



```
1 posterior_plot(samples = rho, usemode = TRUE) + labs(x = expression(rho) )
```



1 summary(lm(height ~ weight, data = d2))\$r.squared
[1] 0.5696444

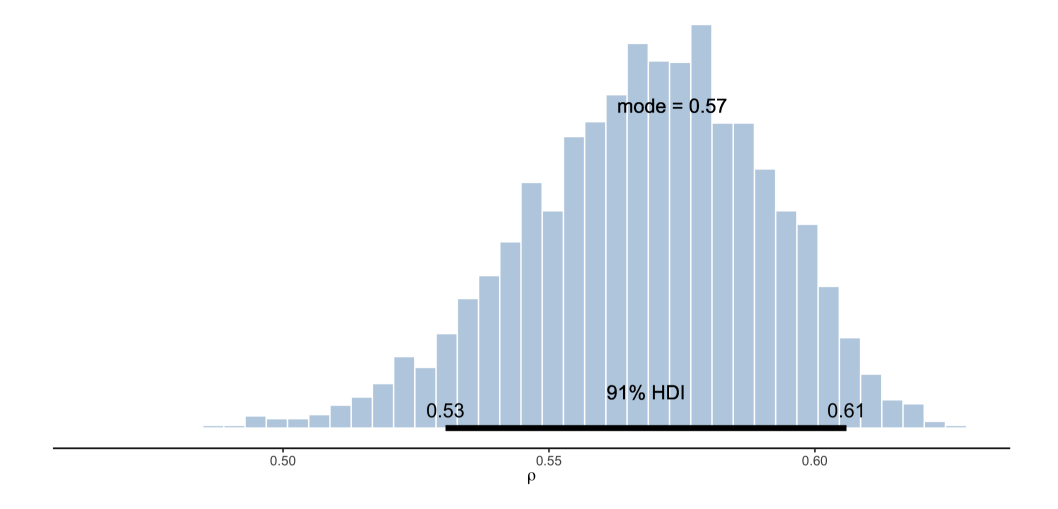


```
1 bayes_R2(mod4)

Estimate Est.Error Q2.5 Q97.5
R2 0.568025 0.02283433 0.51965 0.607488

1 bayes_R2(mod4, summary = FALSE)[, 1] %>%
2 posterior_plot(usemode = TRUE) +
3 labs(x = expression(rho))
```







Summary

A new model with two and then three parameters was presented: the Gaussian model, then Gaussian linear regression model, making it possible to relate two continuous variables.

As previously, Bayes' theorem is used to update our prior knowledge of parameter values into posterior knowledge, a synthesis between our priors and the information contained in the data.

The brms package can be used to fit all sorts of models with a syntax similar to that used by lm().

The **fitted()** function is used to retrieve the predictions of a model.

The predict() function is used to simulate data from a model fitted with brms.



Practical work - 1/2

Select all rows in the howell dataset corresponding to minors individuals (age < 18). This should result in a dataframe of 192 rows.

Fit a linear regression model using the brms::brm() function. Report and interpret the estimates from this model. For an increase of 10 units in weight, what increase in height (height) does the model predict?

Plot the raw data with weight on the x-axis and height on the y-axis. Overlay the model's regression line of the model and an 89% credibility interval for the mean. Add an 89% credibility interval for the predicted sizes.

What do you think of the 'fit' of the model? What assumptions of the model would you be willing to modify in order to improve the model's fit?



Practical work - 2/2

Let's say you've consulted a colleague who's an expert in <u>allometry</u> (i.e., the phenomena of differential organ growth) and she explains to you that it doesn't make sense to model the relationship between weight and height... because we know that it's the logarithm of weight which is (linearly) related to height!

Model the relationship between height (cm) and log weight (log-kg). Use the entire howell dataframe (all 544 rows). Fit the following model using brms::brm().

```
h_i \sim \text{Normal}(\mu_i, \sigma)

\mu_i = \alpha + \beta \cdot \log(w_i)

\alpha \sim \text{Normal}(174, 100)

\beta \sim \text{Normal}(0, 100)

\sigma \sim \text{Exponential}(0.01)
```

Where h_i is the height of individual i and w_i the weight of individual i. The function for calculating the log in R is simply log(). Can you interpret the results? Hint: plot the raw data and superimpose the model's predictions...



```
1 # we keep only the individuals younger than 18
2 d <- open data(howell) %>% filter(age < 18)</pre>
 3
4 priors <- c(
     prior(normal(150, 100), class = Intercept),
     prior(normal(0, 10), class = b),
     prior(exponential(0.01), class = sigma)
8
9
10 mod7 <- brm(
     height ~ 1 + weight,
11
12
     prior = priors,
13
     family = gaussian(),
14
     data = d
15
```



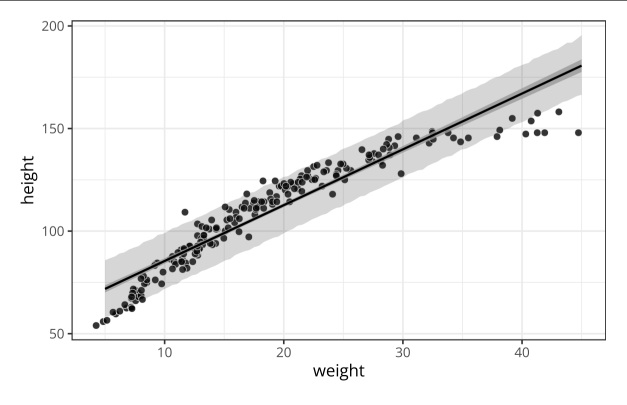
```
1 summary(mod7, prob = 0.89)
 Family: gaussian
 Links: mu = identity; sigma = identity
Formula: height ~ 1 + weight
   Data: d (Number of observations: 192)
  Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
         total post-warmup draws = 4000
Population-Level Effects:
          Estimate Est.Error 1-89% CI u-89% CI Rhat Bulk ESS Tail ESS
                               56.00
                                        60.47 1.00
Intercept
             58.22
                       1.40
                                                       3853
                                                                3011
weight
            2.72
                             2.61
                                       2.83 1.00
                        0.07
                                                       3445
                                                                2962
Family Specific Parameters:
      Estimate Est. Error 1-89% CI u-89% CI Rhat Bulk ESS Tail ESS
sigma
          8.53
                   0.45
                            7.84
                                     9.27 1.00
                                                   4283
                                                            2872
Draws were sampled 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).
```



```
1 # defining a grid of possible values for "weight"
  2 weight.seg <- data.frame(weight = seg(from = 5, to = 45, length.out = 1e2) )
  3
  4 # retrieving the model's predictions for these values fo "weight"
  5 mu <- data.frame(</pre>
      fitted(mod7, newdata = weight.seq, probs = c(0.055, 0.945))
      ) %>%
      bind cols(weight.seq)
  8
  9
 10
    pred height <- data.frame(</pre>
      predict(mod7, newdata = weight.seq, probs = c(0.055, 0.945) )
 11
 12
      bind cols(weight.seq)
 13
 14
 15 # displaying the first 6 rows of pred height
 16 head(pred height)
  Estimate Est.Error
                         05.5
                                 094.5
                                         weight
1 71.90158 8.558478 58.38327 85.76683 5.000000
2 72.80918 8.809837 58.64175 86.74542 5.404040
3 74.07137 8.558394 60.51732 87.58434 5.808081
4 75.08137 8.717473 61.08438 89.05932 6.212121
5 75.95542 8.633522 62.39603 89.61429 6.616162
6 77.39934 8.498564 63.69033 91.05000 7.020202
```



```
1 d %>%
2    ggplot(aes(x = weight, y = height)) +
3    geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4    geom_ribbon(
5     data = pred_height, aes(x = weight, ymin = Q5.5, ymax = Q94.5),
6     alpha = 0.2, inherit.aes = FALSE
7     ) +
8    geom_smooth(
9     data = mu, aes(y = Estimate, ymin = Q5.5, ymax = Q94.5),
10    stat = "identity", color = "black", alpha = 0.8, size = 1
11    )
```





```
1 # we now consider all individuals
2 d <- open_data(howell)
3
4 mod8 <- brm(
5  # we now predict height using the logarithm of weight
6 height ~ 1 + log(weight),
7 prior = priors,
8 family = gaussian(),
9 data = d
10 )</pre>
```



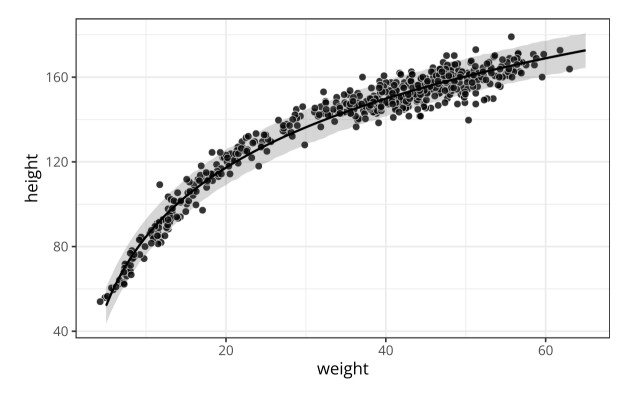
```
1 summary(mod8, prob = 0.89)
 Family: gaussian
 Links: mu = identity; sigma = identity
Formula: height ~ 1 + log(weight)
   Data: d (Number of observations: 544)
  Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
        total post-warmup draws = 4000
Population-Level Effects:
          Estimate Est.Error 1-89% CI u-89% CI Rhat Bulk ESS Tail ESS
                       1.34 -25.67 -21.42 1.00
Intercept -23.56
                                                       4356
                                                               3128
logweight 47.01
                              46.39 47.63 1.00
                       0.38
                                                       4396
                                                               2952
Family Specific Parameters:
      Estimate Est. Error 1-89% CI u-89% CI Rhat Bulk ESS Tail ESS
sigma
          5.15
                   0.15
                            4.91
                                     5.41 1.00
                                                   4641
                                                           2770
Draws were sampled 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).
```



```
1 # defining a grid of possible values for "weight"
  2 weight.seg <- data.frame(weight = seg(from = 5, to = 65, length.out = 1e2) )</pre>
  3
  4 # retrieving the model's predictions for these values of "weight"
  5 mu <- data.frame(</pre>
      fitted(mod8, newdata = weight.seq, probs = c(0.055, 0.945))
      ) %>%
      bind cols(weight.seq)
  8
  9
 10
    pred height <- data.frame(</pre>
      predict(mod8, newdata = weight.seq, probs = c(0.055, 0.945) )
 11
 12
      bind cols(weight.seq)
 13
 14
 15 # displaying the first 6 rows of "pred height"
 16 head(pred height)
  Estimate Est.Error
                         05.5
                                 094.5
                                         weight
1 52.08114 5.364763 43.64659 60.73506 5.000000
2 57.47366 5.180466 49.20607 65.72344 5.606061
3 62.29115 5.289523 53.91334 70.72491 6.212121
4 66.70296 5.104535 58.45216 74.90719 6.818182
5 70.49918 5.249748 62.17113 78.91841 7.424242
6 74.31789 5.177680 66.23217 82.75413 8.030303
```



```
1 d %>%
2    ggplot(aes(x = weight, y = height)) +
3    geom_point(colour = "white", fill = "black", pch = 21, size = 3, alpha = 0.8) +
4    geom_ribbon(
5         data = pred_height, aes(x = weight, ymin = Q5.5, ymax = Q94.5),
6         alpha = 0.2, inherit.aes = FALSE
7         ) +
8    geom_smooth(
9         data = mu, aes(y = Estimate, ymin = Q5.5, ymax = Q94.5),
10         stat = "identity", color = "black", alpha = 0.8, size = 1
11    )
```





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