Introduction to Bayesian linear regression with brms

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Random variables

- We have a question about the world, so we collect data (sample from a population).
 - \cdot y = $(y_1, y_2, y_3, y_4, ..., y_n)$
- · We want to know how the data (the sample) was generated.
- · In probability theory, data is generated by a random variable Y.

Random variables

- Y is uncertain.
 - We can describe Y as a probability distribution, expressed by a set of parameters $\Theta=(\theta_1,...,\theta_n)$.
- · Probability distributions:
 - · $Normal(\mu, \sigma)$,
 - · Binomial(n, p),
 - ..

Random variables

$$vot_i \sim Normal(\mu, \sigma)$$

$$voiced_i \sim Bernoulli(p)$$

$$DoubleDative_i \sim Poisson(\lambda)$$

Frequentist vs Bayesian view

- Parameters: μ , σ , p, λ , ...
- · Frequentist view:
 - The parameters are **fixed** (they are unknown but certain).
 - · They take on a specific value.
- · Bayesian view:
 - The parameters are random variables (they are unknown and uncertain).
 - We describe each parameter as a probability distribution, expressed by a set of hyperparameters.

Continous random variable

$$\begin{aligned} vot_i \sim Normal(\mu, \sigma) \\ \mu \sim Normal(\mu_1, \sigma_1) \\ \sigma \sim HalfCauchy(x_0, \gamma) \end{aligned}$$

Bayes' Theorem

$$P(\theta \mid d) = \frac{P(d \mid \theta) P(\theta)}{P(d)}$$

Bayes' Theorem

$$posterior\ probability = \frac{likelihood \times prior}{marginal\ likelihood}$$

Priors

- We can incorporate previous knowledge about the hyperparameters as priors (prior distributions).
- Priors are chosen based on expert knowledge, previous studies, pilot data...
 - Priors must **not** be chosen based on the data to be analysed.

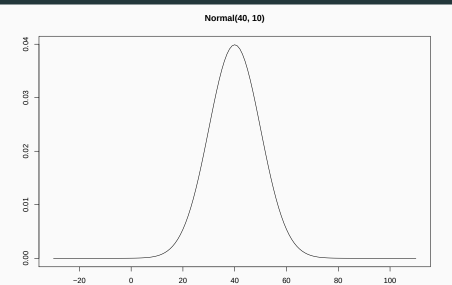
Priors

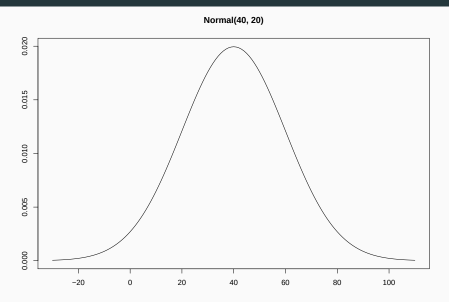
- · Informative and weakly informative priors.
- · Uninformative or diffuse priors.
 - · Uniform distribution.
- · Regularising priors.

Normal prior

[empirical rule]

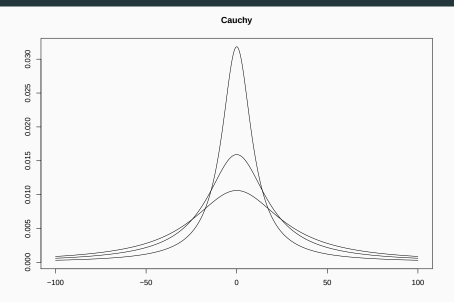
- Previous literature on VOT in Italian (Esposito, 2002; Stevens & Hajek, 2010) report VOT values for voiceless stops in the range of 20–60 ms.
 - We can express this knowledge with the prior Normal(40,10).
 - This is a somewhat strongly informative prior.





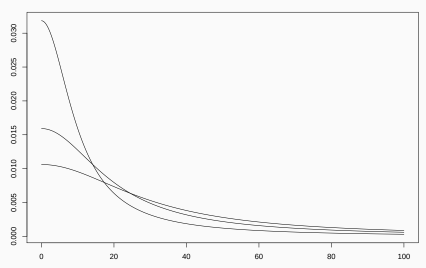
$$\begin{aligned} vot_i \sim Normal(\mu, \sigma) \\ \mu \sim Normal(40, 10) \\ \sigma \sim HalfCauchy(x_0, \gamma) \end{aligned}$$

Cauchy prior



Cauchy prior





$$\begin{aligned} vot_i \sim Normal(\mu, \sigma) \\ \mu \sim Normal(40, 10) \\ \sigma \sim HalfCauchy(0, 10) \end{aligned}$$

- We have a model which incorporates (some of) our knowledge about VOT (through the priors for μ and σ).
- · Now we want to obtain the **posterior distributions** of μ and σ .
 - The posterior distribution is the prior distribution conditioned on the data.
- brms R package: Bayesian Regression Models using Stan (Bürkner, 2018).

brms

- · Stan (Stan Development Team, 2017).
 - Statistical programming language written in C++ for fitting Bayesian models (calculate posterior distributions).
 - Calculation can be complex and/or impossible, so we take many samples from the data and from the possible parameter values to find the posterior distributions of the hyperparameters.
 - Markov Chain Monte Carlo (MCMC) sampling using the No-U-Turn sampler (NUTS).
- · brms is an interface between R and Stan.
- brm() function from brms.
 - · lme4 syntax $(y \sim x + (1|w))$.
 - · Creates a Stan model, which is compiled and run.

```
library(brms)
vot1 <- brm(
  <model_formula>,
  <family>,
  <pri>>,
  <data>,
  chains = 4,
  iter = 2000
```

```
library(brms)
vot1 <- brm(
 vot ~ 1,
  family = gaussian(),
  <pri>>,
  data = ita_egg,
  chains = 4,
  iter = 2000
```

Get prior

get_prior(
 vot ~ 1,

2 student t(3, 0, 14) sigma

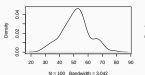
Set prior

Prior predictive checks

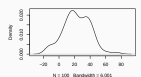
```
nsim <- 1000
nobs <- 100
y <- matrix(rep(NA, nsim * nobs), ncol = nobs)</pre>
mu <- rnorm(nsim, 40, 10)
sigma <- rhcauchy(nsim, 10)</pre>
for (i in 1:nsim) {
  v[i,] <- rnorm(nobs, mean = mu[i], sd = sigma[i])</pre>
```

Prior predictive checks

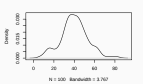
```
op \leftarrow par(mfrow = c(3, 3))
rand_sample <- sample(1:nsim, 9)</pre>
for (i in rand_sample) {
   plot(density(y[i,]), main = "")
}
  0.020
Density
                         Density
                                                  Density
  0.010
                           0.02
```



N = 100 Bandwidth = 5.72



N = 100 Bandwidth = 2.422



N = 100 Bandwidth = 1.571

Run the model

```
vot1 <- brm(
  vot ~ 1,
  family = gaussian(),
  prior = priors,
  data = ita_egg,
  chains = 4,
  iter = 2000,
  file = "./cache/vot1"
```

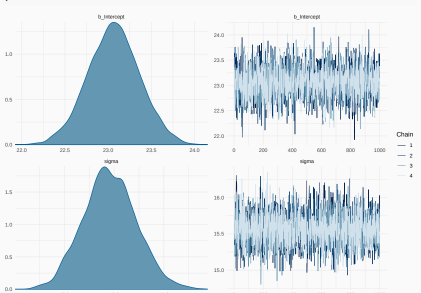
Model summary

vot1

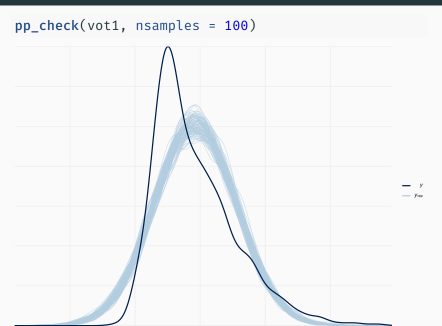
```
Family: gaussian
##
     Links: mu = identity; sigma = identity
##
## Formula: vot ~ 1
      Data: ita egg (Number of observations: 2624)
##
## Samples: 4 chains, each with iter = 2000; warmup = 1000;
            total post-warmup samples = 4000
##
##
## Population-Level Effects:
                  Estimate Est.Error 1-95% CI u-
##
95% CI Rhat Bulk ESS Tail ESS
## Intercept 23.06 0.30 22.47 23.65 1.00
                                                   3219
##
                                                     29
## Family Spacific Darameters.
```

Plot model

plot(vot1)



Posterior predictive check



Adding predictors

$$\begin{split} vot_i &\sim Normal(\mu_i, \sigma) \\ \mu_i &= \alpha + \beta_1 \times coronal_i + \beta_2 \times velar_i \\ \alpha &\sim Normal(\mu_1, \sigma_1) \\ \beta_1 &\sim Normal(\mu_2, \sigma_2) \\ \beta_2 &\sim Normal(\mu_3, \sigma_3) \\ \sigma &\sim HalfCauchy(x_0, \gamma) \end{split}$$

Adding predictors

$$\begin{split} vot_i &\sim Normal(\mu_i, \sigma) \\ \mu_i &= \alpha + \beta_1 \times coronal_i + \beta_2 \times velar_i \\ \alpha &\sim Normal(25, 10) \\ \beta_1 &\sim Normal(10, 10) \\ \beta_2 &\sim Normal(20, 10) \\ \sigma &\sim HalfCauchy(0, 10) \end{split}$$

Get prior

Set prior

Run the model

```
vot2 <- brm(
 vot ~ 1 + c1_place,
  family = gaussian(),
  prior = priors,
  data = ita_egg,
  chains = 4,
  iter = 2000,
  file = "./cache/vot2"
```

$$\begin{aligned} vot_i \sim Normal(\mu_i, \sigma) \\ \mu_i = \alpha + \alpha_{speaker[i]} + (\beta_1 + \beta_{1speaker[i]}) \times coronal_i + (\beta_2 + \beta_{2speaker[i]}) \times velar_i \end{aligned}$$

$$\begin{bmatrix} \alpha_{speaker} \\ \beta_{1speaker[i]} \\ \beta_{2speaker[i]} \end{bmatrix} \sim Normal(\begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}, S)$$

$$\alpha \sim Normal(25, 10)$$

$$\beta_{1} \sim Normal(10, 10)$$

$$\beta_{2} \sim Normal(20, 10)$$

$$\sigma_{\alpha speaker} \sim Normal(0, \sigma_{speaker})$$

$$\sigma_{\beta_{1}speaker} \sim HalfCauchy(0, 10)$$

```
##
               prior
                       class
                                      coef group resp dpa
## 1
                                     b
## 2
                                   b c1 placecoronal
                                    b
                                        c1_placevelar
## 3
                     lkj(1)
## 4
                                   cor
## 5
                                              speaker
                            cor
## 6
      student t(3, 19, 14) Intercept
     student t(3, 0, 14)
## 7
                                    sd
## 8
                             sd
                                              speaker
                          sd c1_placecoronal speaker
## 9
                           sd c1_placevelar speaker
## 10
## 11
                            sd
                                   Intercept speaker
       student_t(3, 0, 14)
## 12
                                sigma
```

```
vot3 <- brm(</pre>
  vot ~ 1 + c1_place + (1 + c1_place | speaker),
  family = gaussian(),
  prior = priors,
  data = ita egg,
  chains = 4,
  iter = 2000.
  file = "./cache/vot3"
```

References

Bürkner, Paul-Christian. 2018. Advanced Bayesian multilevel modeling with the R package brms. The R Journal 10(1). 395–411. doi: $10.32614/\mathrm{RJ}$ -2018-017.

Esposito, Anna. 2002. On vowel height and consonantal voicing effects: Data from Italian. *Phonetica* 59(4). 197–231. doi:10.1159/000068347.

Stan Development Team. 2017. Stan: A C++ library for probability and sampling, version 2.14.0. http://mc-stan.org/.

Stevens, Mary & John Hajek. 2010. Post-aspiration in standard Italian: some first cross-regional acoustic evidence. Paper presented at Interspeech, 26-30 September 2010, Makuhari, Chiba, Japan.