

# Regression with a Categorical Independent Variables

## General Principles

To study the relationship between a categorical independent variable and a continuous dependent variable, we use a *Categorical model* which applies *stratification*.

*Stratification* involves modeling how the  $k$  different categories of the independent variable affect the target continuous variable by performing a regression for each  $k$  category and assigning a regression coefficient for each category. To implement stratification, categorical variables are often encoded using one-hot encoding or by converting categories to indices .

## Considerations

### Note

- We have the same considerations as for [Regression for a Continuous Variable](#).
- As we generate regression coefficients for each  $k$  category, we need to specify a prior with a shape equal to the number of categories  $k$  in the code (see comments in the code).
- To compare differences between categories, we need to compute the distribution of the differences between categories, known as the contrast distribution. **Never compare confidence intervals or p-values directly.**

## Example

Below is an example of code that demonstrates Bayesian regression with an independent categorical variable using the Bayesian Inference (BI) package. The data consist of one continuous dependent variable (*kcal\_per\_g*), representing the caloric value of milk per gram, a categorical

independent variable (*index\_clade*), representing species clade membership, and a continuous independent variable (*mass*), representing the mass of individuals in the clade. The goal is to estimate the differences in milk calories between clades. This example is based on McElreath (2018).

## Python

```
from main import*

# Setup device-----
m = bi(platform='cpu')

# Import Data & Data Manipulation -----
# Import
from importlib.resources import files
data_path = files('BI.resources.data') / 'milk.csv'
m.data(data_path, sep=';')
m.index(["clade"]) # Manipulate
m.scale(['kcal_per_g']) # Scale
m.data_to_model(['kcal_per_g', "index_clade"]) # Send to model (convert to jax array)

# Define model -----
def model(kcal_per_g, index_clade):
    a = m.bi.dist.normal(0, 0.5, shape=(4,), name = 'a') # shape based on the number of clades
    b = m.bi.dist.normal(0, 0.5, shape=(4,), name = 'b')
    s = m.bi.dist.exponential( 1, name = 's')
    mu = a[index_clade]+b[index_clade]*mass
    m.normal(mu, s, obs=kcal_per_g)

# Run mcmc -----
m.fit(model) # Optimize model parameters through MCMC sampling

# Summary -----
m.summary()
```

## R

```
library(BI)
m=importbi(platform='cpu')
```

```

# Load csv file
m$data(paste(system.file(package = "BI"), "/data/milk.csv", sep = ''), sep=';')
m$scale(list('kcal.per.g')) # Manipulate
m$index(list('clade')) # Scale
m$data_to_model(list('kcal_per_g', 'index_clade')) # Send to model (convert to jax array)

# Define model -----
model <- function(kcal_per_g, index_clade){
  # Parameter prior distributions
  beta = bi.dist.normal( 0, 0.5, name = 'beta', shape = c(4)) # shape based on the number o
  sigma =bi.dist.exponential(1, name = 's')
  # Likelihood
  m$normal(beta[index_clade], sigma, obs=kcal_per_g)
}

# Run mcmc -----
m$fit(model) # Optimize model parameters through MCMC sampling

# Summary -----
m$summary() # Get posterior distributions

```

### 🔥 Caution

For R users, when working with indices you have to ensure 1) that indices are integers (i.e. `as.integer(index_clade)`) and, 2) that indices start at 0 (i.e. `as.integer(index_clade)-1`).

## Mathematical Details

### *Frequentist formulation*

We model the relationship between the categorical input feature (X) and the target variable (Y) using the following equation:

$$Y_i = \alpha + \beta_k X_i + \sigma$$

Where:

- $Y_i$  is the dependent variable for observation  $i$ .

- $\alpha$  is the intercept term.
- $\beta_k$  are the regression coefficients for each  $k$  category.
- $X_i$  is the encoded categorical input variable for observation  $i$ .
- $\sigma$  is the error term.

We can interpret  $\beta_i$  as the effect of each category on  $Y$  relative to the baseline (usually one of the categories or the intercept).

### **Bayesian formulation**

In the Bayesian formulation, we define each parameter with priors . We can express the Bayesian regression model accounting for prior distributions as follows:

$$Y \sim \text{Normal}(\alpha + \beta_K X, \sigma)$$

$$\alpha \sim \text{Normal}(0, 1)$$

$$\beta_K \sim \text{Normal}(0, 1)$$

$$\sigma \sim \text{Exponential}(1)$$

Where:

- $Y_i$  is the dependent variable for observation  $i$ .
- $\alpha$  is the intercept term, which in this case has a unit-normal prior.
- $\beta_K$  are slope coefficients for the  $K$  distinct independent variables categories, which also have unit-normal priors.
- $X_i$  is the encoded categorical input variable for observation  $i$ .
- $\sigma$  is a standard deviation parameter, which here has a Exponential prior that constrains it to be positive.

## Notes

### Note

- We can apply multiple variables similarly to [Chapter 2: Multiple Continuous Variables](#).
- We can apply interaction terms similarly to [Chapter 3: Interaction between Continuous Variables](#).

## Reference(s)

McElreath, Richard. 2018. *Statistical Rethinking: A Bayesian Course with Examples in r and Stan*. Chapman; Hall/CRC.