# GLM(M)s for binary data

Bayesian statistics 7 - generalized linear models for binary data

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## Some things that we learned the last time

- Poisson models good for rare events generating small counts
- Classical *link function* is the log-link so that  $Y_i|\epsilon_i \sim \mathcal{P}(\exp(a + bx_i + [\text{stuff}] + \epsilon_i))$  (with extra dispersion)
- Posterior predictive checks

#### The Binomial distribution: reminders and GLM

Let  $U_i \sim \text{Bernoulli}(p)$  a coin toss with probability p.

- Then  $Y = \sum_{i=1}^{n} U_i \sim \mathcal{B}(n, p)$
- Converges to normal distribution for large  $np \ (\geq 10)$  as n grows
- $\mathbb{E}(Y) = np$  and  $\mathbb{E}(Y) = np(1-p)$
- Conjugate prior for p = Beta distribution.

Two ways to specify a GLM:

- $Y_i \sim \mathcal{B}(n, p)$  with  $logit(p) = a + bx_i + [stuff]$  in which case the data resembles c(31, 14, 5, 0, 19)
- $U_i \sim \mathcal{B}(p)$  with  $logit(p) = a + bx_i + [stuff]$  in which case the data resembles c(0,0,1,0,1,1,0)

#### Environnementally-driven turtle sex determination



Figure 1: Green turtle, Malaysia. Bernard Dupont. Licence: CC BY SA 2.0

First described in 1966 by Madeleine Charnier in a lizard, subsequent work on turtles (Chelonia) and Crocodylia, see Janzen & Paukstis QRB 1991.

# $\mathbb{P}(\text{female hatchling}) = \text{f}(\text{temperature})$



Figure 2: Stephan Hunt. Hatching green turtle, Ascension Island. CC BY 3.0

# Empirical target

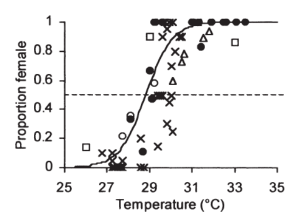
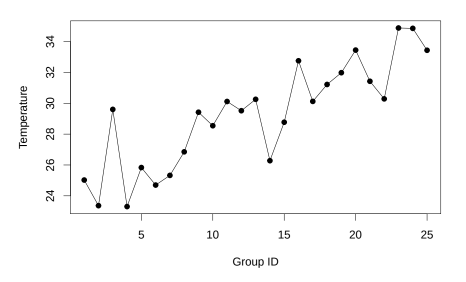


Figure 3: Temperature-dependent sex determination of Ascension Island green turtles, by Goldley et al. Marine Ecology Progress Series Vol. 226: 115–124, 2002.

# Simulating data I

We have got 25 groups i of size 50 or less (i.e., how many baby turtles are hatched at one beach location  $z_i$ ), with a different temperature in each group.

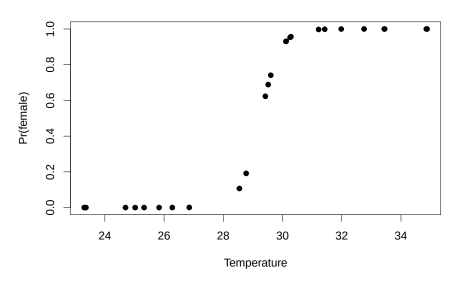
# Simulating data II



## Binomial sampling I

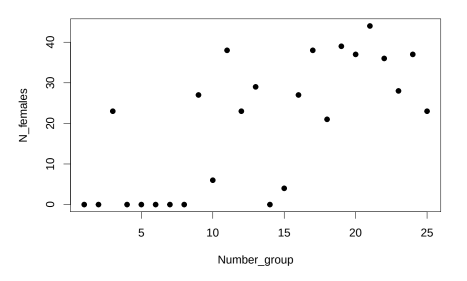
Females are born with probability  $p_i$  in group i, which itself depends on temperature.

# Binomial sampling II



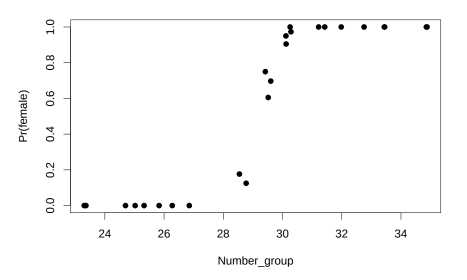
#### The data I

#### The data II



#### Empirical proportion estimates I

# Empirical proportion estimates II



#### Prior predictive checks

We have seen posterior predictive distribution = the distribution of imaginary data under the fitted model (given a posterior distribution).

Prior predictive distribution = the distribution of imaginary data under the priors.

Let's say we have proportion  $\theta \sim \mathrm{Beta}(\alpha,\beta)$  which is the prior for a very simple  $Y_i \sim \mathcal{B}(n,\theta)$  model where n is known. Then simulating data under the prior predictive distribution can be done as

```
for (rep in 1:nrep){
  theta[rep] = rbeta(1,alpha,beta)
  Yrep[rep] = rbinom(1,size=n,prob=theta[rep])
}
```

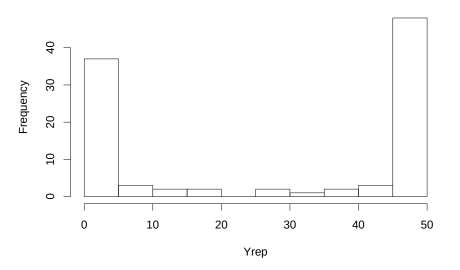
(it's a Beta-Binomial distribution)

# Prior predictive checks (practice) I

```
nrep = 100
n = 50
alpha=0.1
beta=0.1
Yrep = theta = rep(0,nrep)
for (rep in 1:nrep){
   theta[rep] = rbeta(1,alpha,beta)
   Yrep[rep] = rbinom(1,size=n,prob=theta[rep])
}
par(cex=1.5,pch=19)
hist(Yrep)
```

# Prior predictive checks (practice) II

#### **Histogram of Yrep**

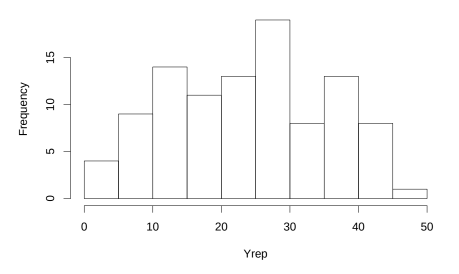


# Prior predictive checks - better prior I

```
nrep = 100
n = 50
alpha=2
beta=2
Yrep = theta = rep(0,nrep)
for (rep in 1:nrep){
   theta[rep] = rbeta(1,alpha,beta)
     Yrep[rep] = rbinom(1,size=n,prob=theta[rep])
}
par(cex=1.5,pch=19)
hist(Yrep)
```

### Prior predictive checks – better prior II

#### **Histogram of Yrep**



# Fitting the binomial model I

Now we fit that model which writes mathematically like

$$y_i \sim \mathcal{B}(z_i, p(\mathsf{temp}_i))$$

# Fitting the binomial model II

```
m11.data \leftarrow list(N = n_groups, y = Y, temp = temperature,
                 z = sample_size_per_group)
cat(file="logistic.regression.txt","
model {
  mu_temp ~ dnorm(2, 0.1) ## prior of the mean temp
  gamma ~ dnorm(1, 0.1) ## prior of the slope
  for (k in 1:N){
  y[k] ~ dbin(p[k],z[k]) ## likelihood
  logit(p[k])<-gamma*(temp[k]-mu temp)</pre>
  }
```

### Running the model I

```
# Inits function
inits <- function(){list(gamma = rnorm(1, 0, 1),</pre>
                         mu temp = rnorm(1,0,1))
# Parameters to estimate
params <- c("gamma", "mu temp")
# MCMC settings
nc <- 3 ; ni <- 2000 ; nb <- 1000 ; nt <- 2
# Call JAGS, check convergence and summarize posteriors
out <- jags(m11.data, inits, params, "logistic.regression.txt", n.thin = nt,
            n.chains = nc. n.burnin = nb. n.iter = ni)
## Compiling model graph
##
     Resolving undeclared variables
      Allocating nodes
##
## Graph information:
##
      Observed stochastic nodes: 25
     Unobserved stochastic nodes: 2
##
##
     Total graph size: 156
##
## Initializing model
```

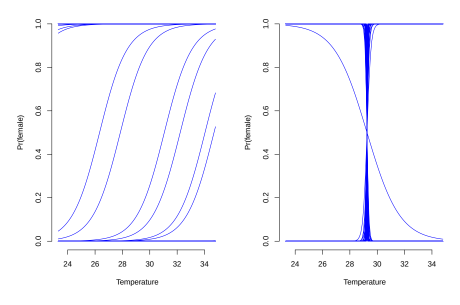
# Running the model II

```
print(out, dig = 3)  # Bayesian analysis
## Inference for Bugs model at "logistic.regression.txt", fit using jags,
   3 chains, each with 2000 iterations (first 1000 discarded), n.thin = 2
##
   n.sims = 1500 iterations saved
           mu.vect sd.vect 2.5% 25% 50% 75% 97.5% Rhat n.eff
##
             3.017 0.319 2.415 2.791 3.014 3.232 3.661 1.005
                                                                    400
## gamma
## mu_temp 29.226 0.061 29.103 29.185 29.227 29.270 29.338 1.001 1500
## deviance 38.512 2.080 36.530 37.042 37.860 39.221 44.403 1.005 1100
##
## For each parameter, n.eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
##
## DIC info (using the rule, pD = var(deviance)/2)
## pD = 2.2 and DIC = 40.7
## DIC is an estimate of expected predictive error (lower deviance is better).
```

# Priors on $\gamma$ and $\mu_{\mathsf{temp}}$ I

```
mu_temp = rnorm(100,2, 100) # prior of the mean temp
gamma = rnorm(100,1, 100) # prior on the slope
x=seq(min(temperature), max(temperature), by=0.1)
par(mfrow=c(1,2))
plot(0, bty = 'n', pch = '', ylab = "Pr(female)",
     xlab = "Temperature", ylim=c(0,1),
     xlim=c(min(temperature),max(temperature)))
for (kprior in 1:100) {
  prob = 1/(1+exp(-1*(x-mu_temp[kprior])))
  lines(x,prob,type="l",col="blue")}
plot(0, bty = 'n', pch = '', ylab = "Pr(female)",
     xlab = "Temperature", ylim=c(0,1),
     xlim=c(min(temperature), max(temperature)))
for (kprior in 1:100) {
  prob = 1/(1+exp(-gamma[kprior]*(x-mean(temperature))) )
  lines(x,prob,type="l",col="blue")}
```

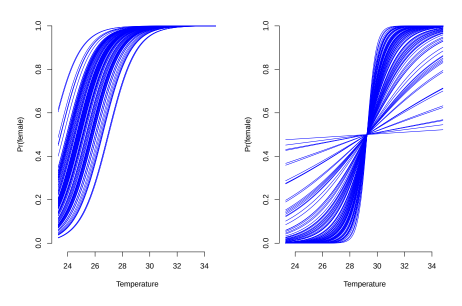
# Priors on $\gamma$ and $\mu_{\rm temp}$ II



### Better priors I

```
### Better priors
mu temp = rnorm(100,25, 1) # prior of the mean temp
gamma = rnorm(100,1, 1) # prior on the slope
par(mfrow=c(1,2))
plot(0, bty = 'n', pch = '', ylab = "Pr(female)",
    xlab = "Temperature", ylim=c(0,1),
    xlim=c(min(temperature), max(temperature)))
for (kprior in 1:100) {
 prob = 1/(1+exp(-1*(x-mu_temp[kprior])))
  lines(x,prob,type="l",col="blue")}
plot(0, bty = 'n', pch = '', ylab = "Pr(female)",
    xlab = "Temperature", ylim=c(0,1),
    xlim=c(min(temperature), max(temperature)))
for (kprior in 1:100) {
 prob = 1/(1+exp(-abs(gamma[kprior])*(x-mean(temperature))) )
  lines(x,prob,type="l",col="blue")}
```

# Better priors II



# Weakly informative priors I

#### What are they?

- Priors that provide regularization or shrinkage
- In practice, often  $\mathcal{N}(0,[\text{small}])$  instead of  $\mathcal{N}(0,[\text{huge}])$ , e.g. dnorm(0,1) or dnorm(0,0.1) instead of dnorm(0,0.0001) in JAGS.

#### A more detailed explanation

### A detour on identifiability, convergence, and priors

Borrowed from Mc Elreath's Statistical rethinking

We consider the (obviously wrong) model

$$Y_i \sim \mathcal{N}(\mu, \sigma^2)$$

$$\mu = \alpha_1 + \alpha_2$$

$$lpha_{j} \sim \mathsf{Unif}(\text{-10000,10000}) \; \mathsf{i.i.d}$$

$$\sigma \sim \mathsf{Exp}(1/10)$$

for 100 data points simulated as  $\mathcal{N}(0,1)$ .

# Coding the stupid model

```
gaussian.data = list(y=rnorm(100,0,1),N=100)
cat(file="stupid.model.txt","
model {
    # Priors
    alpha[1] ~ dunif(-10000,10000) #dnorm(0,0.00001)
    alpha[2] ~ dunif(-10000,10000) #dnorm(0,0.00001)
    sigma \sim dunif(0,1000) \#dexp(0.01) \#dexp(1)
    tau<-pow(sigma,-2)
    # Likelihood
    mu<-alpha[1]+alpha[2]
    for (i in 1:N){
    y[i] ~ dnorm(mu,tau)
```

### Fitting the stupid model I

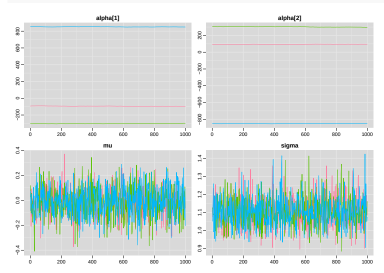
```
# Initial nalnes
inits <- function(){list(alpha=rnorm(2,0,1000))}</pre>
# Parameters to estimate
params <- c("alpha", "sigma", "mu")
# MCMC settings
nc <- 3 ; ni <- 2000 ; nb <- 1000 ; nt <- 2
# Call JAGS, check convergence and summarize posteriors
out <- jags(gaussian.data, inits, params, "stupid.model.txt", n.thin = nt,
            n.chains = nc, n.burnin = nb, n.iter = ni)
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 100
##
     Unobserved stochastic nodes: 3
##
     Total graph size: 112
##
## Initializing model
print(out, dig = 3)  # Bayesian analysis
```

#### Fitting the stupid model II

```
## Inference for Bugs model at "stupid.model.txt", fit using jags,
   3 chains, each with 2000 iterations (first 1000 discarded), n.thin = 2
## n.sims = 1500 iterations saved
            mu.vect sd.vect 2.5% 25% 50%
                                                       75%
                                                             97.5%
                                                                     Rhat.
##
## alpha[1] 151.616 502.361 -304.515 -299.895 -94.776 850.569 853.539 554.310
## alpha[2] -151.631 502.356 -853.536 -850.553 94.716 299.901 304.515 554.217
## mu
         -0.015 0.114 -0.239 -0.091 -0.010 0.064
                                                             0.214
                                                                    1.005
## sigma 1.103 0.080 0.960 1.049 1.098 1.156 1.276 1.002
## deviance 301.876 2.163 299.812 300.348 301.205 302.722 307.411 1.006
##
           n.eff
## alpha[1]
              .3
## alpha[2] 3
## mu
             660
## sigma 1100
## deviance 580
##
## For each parameter, n.eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
##
## DIC info (using the rule, pD = var(deviance)/2)
## pD = 2.3 and DIC = 304.2
## DIC is an estimate of expected predictive error (lower deviance is better).
```

#### **Traceplots**

```
library(mcmcplots)
traplot(as.mcmc(out),parms=c("alpha","mu","sigma"))
```



# Less stupid model

```
cat(file="less.stupid.model.txt","
model {
    # Priors
    alpha[1] \sim dnorm(0,0.1)
    alpha[2] \sim dnorm(0,0.1)
    sigma ~ dexp(1)
    tau<-pow(sigma,-2)
    # Likelihood
    mu<-alpha[1]+alpha[2]
    for (i in 1:N)
    y[i] ~ dnorm(mu,tau)
```

Actually even a much smaller precision on  $\alpha_i$ 's prior would work

#### Fitting the less stupid model I

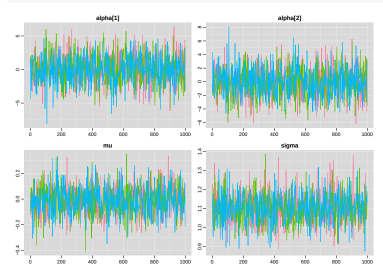
```
# Initial nalnes
inits <- function(){list(alpha=rnorm(2,0,1))}</pre>
# Parameters to estimate
params <- c("alpha", "sigma", "mu")
# MCMC settings
nc <- 3 ; ni <- 2000 ; nb <- 1000 ; nt <- 2
# Call JAGS, check convergence and summarize posteriors
out <- jags(gaussian.data, inits, params, "less.stupid.model.txt",
            n.thin = nt, n.chains = nc, n.burnin = nb, n.iter = ni)
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 100
##
     Unobserved stochastic nodes: 3
##
     Total graph size: 111
##
## Initializing model
print(out, dig = 3)  # Bayesian analysis
```

#### Fitting the less stupid model II

```
## Inference for Bugs model at "less.stupid.model.txt", fit using jags,
   3 chains, each with 2000 iterations (first 1000 discarded), n.thin = 2
## n.sims = 1500 iterations saved
         mu.vect sd.vect 2.5% 25% 50% 75% 97.5% Rhat n.eff
##
## alpha[1] 0.061 2.109 -3.999 -1.331 0.057 1.510 4.201 1.003
                                                                800
## alpha[2] -0.072 2.109 -4.231 -1.526 -0.062 1.334 4.034 1.002 830
## mu
     -0.011 0.108 -0.219 -0.083 -0.008 0.061 0.204 1.000
                                                               1500
## sigma 1.098 0.078 0.957 1.044 1.094 1.149 1.266 1.001
                                                               1500
1500
##
## For each parameter, n.eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
##
## DIC info (using the rule, pD = var(deviance)/2)
## pD = 1.9 and DIC = 303.7
## DIC is an estimate of expected predictive error (lower deviance is better).
```

#### Traceplots again

library(mcmcplots)
traplot(as.mcmc(out),parms=c("alpha","mu","sigma"))



## Now a real binary data example: bald eagles I



Figure 4: Bald eagle with salmon, Alaska. KJ Gill aka Gillfoto. CC BY 2.0

## Now a real binary data example: bald eagles II

Records of (160!) salmon-pirating attempts by one Bald eagle on another Bald eagle (not always the same!). Also borrowed from McElreath's Statistical Rethinking p. 330.

```
library(MASS)
data(eagles)
head(eagles)
## y n P A V
```

```
## 1 17 24 L A L
## 2 29 29 L A S
## 3 17 27 L I L
## 4 20 20 L I S
## 5 1 12 S A L
## 6 15 16 S A S
```

## Now a real binary data example: bald eagles III