

Binomial and negative binomial distribution

Parametrisation

The Binomial distribution is

$$\text{Prob}(y) = \binom{n}{y} p^y (1-p)^{n-y}$$

for responses $y = 0, 1, 2, \dots, n$, where

n : number of trials.

p : probability of success in each trial.

The negative binomial distribution is

$$\text{Prob}(n) = \binom{n-1}{y-1} p^y (1-p)^{n-y}$$

for given $y = 1, 2, \dots$ and response $n - y = 0, 1, 2, \dots$

Link-function

The mean and variance of y are given in the binomial case as

$$\mu = np \quad \text{and} \quad \sigma^2 = np(1-p)$$

and the probability p is linked to the linear predictor by

$$p(\eta) = \frac{\exp(\eta)}{1 + \exp(\eta)}$$

Hyperparameters

None.

Hyperparameter spesification and default values

doc The Binomial likelihood

hyper

survival FALSE

discrete TRUE

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Specification

- family = **binomial**
- Required arguments: y and n (keyword **Ntrials**)
- Optional argument: **variant=0** for binomial (default), and **variant=1** for the negative binomial.

Expert version

There is also an “expert” version where you are supposed to know what you are doing. Here, we allow y and n to be non-integers (whatever that means), however, the condition $0 \leq y \leq n$ apply. The normalizing constant is computed as above using the integer part of y and n . This is similar to using `floor(y)` and `floor(n)` in R. The marginal likelihood estimate will in this case make less sense.

- `family = xbinomial`
- Required arguments: y and n (keyword `Ntrials`)
- Optional argument: `scale=q`, which scales the probability with $0 < q \leq 1$ into p' , where

$$p' = qp(\eta).$$

By default, $q = 1$. Note that “fitted values” will still be $p(\eta)$.

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Examples

In the following example we estimate the parameters in a simulated example with binomial responses.

```
## binomial
n=100
a = 1
b = 1
z = rnorm(n)
eta = a + b*z
formula <- y ~ 1 + z
prob = exp(eta)/(1 + exp(eta))

Ntrials = sample(1:10, size=n, replace=TRUE)
y = rbinom(n, size = Ntrials, prob = prob)
data = data.frame(y, z, Ntrials)
r = inla(formula, family = "binomial", data = data, Ntrials=Ntrials)
summary(r)

## negative binomial
y = sample(1:3, size=n, replace=TRUE)
Ntrials = y + rnbino(n, size = y, prob = prob)
r = inla(formula,
          family = "binomial",
          control.family = list(variant = 1),
```

```

      Ntrials = Ntrials,
      data = data.frame(y, x, Ntrials))
summary(r)

```

In the following example we estimate the parameters in a simulated example with binomial responses using the `scale`-argument as well. This requires the use of the expert-version “`xbinomial`”.

```

n <- 10000
x <- rnorm(n, sd = 1)
q <- runif(n)
eta <- 0.88 + 0.77*x
p <- q * 1.0/(1+exp(-eta))
ntrials <- sample(1:25, size=n, replace=TRUE)
y <- rbinom(n = n, size=ntrials, prob = p)
r <- inla(y ~ 1 + x,
          family = "xbinomial",
          Ntrials = ntrials,
          scale = q,
          data = data.frame(y, x, q, ntrials))
summary(r)

```

Notes

- If the response is a **factor** it must be converted to $\{0,1\}$ before calling `inla()`, as this conversion is not done automatically (as for example in `glm()`).
- This version of the negative binomial mimics the binomial distribution, and the “data” kind of enter in the `Ntrials` argument (as `y` is pre-determined) which both can appear, and should appear, strange. There is also an alternative implementation, `family="nbinomial"`, which mimics the Poisson distribution.