Random effect multinomial model

A. J. Rominger

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The model

Given the number of sequencing reads assigned to each of S species, we want to estimate the number of individuals for each species that went into the sequencing run, call that vector x. We know the total number of reads N_{reads} , the number of species S, the total number of individuals N that were sequenced, and the vector of number of reads per species x_{read} . We assume that stochastic evolutionary processes have led to some species having a greater propensity to be sequenced. These stochastic processes we summarize in a random effect nu. We further assume that within orders these random effects are constant, thus each species within the same order gets the same random effect.

Thus our model for the number of reads is:

$$x_{reads} \sim \text{multinom}\left(N_{reads}, \frac{x\nu}{\sum_{i} x_{i}\nu_{i}}\right)$$

Thus the vector of probabilities for the multinomial distribution is proportional to the unknown abundance times the unknown random effect ν .

We could choose to put a multinomial prior on x because $N = \sum x$ is fixed, or could approximate x_i with a continuous prior and assume N is large enough such that $Cov(x_i, x_j) \approx 0$. Allowing x_i to be iid and continuous allows us to better estimate the relative contribution of x and ν in determining x_{reads} , so we opt for the prior:

$$\log(x_i) \sim norm\left(\frac{N}{S}, \sigma_x^2\right)$$

Given that ν is constant within orders, varies across orders, and must be positive, we model it as

$$\log(\nu_{order=j}) \sim \text{norm}(0, \sigma_{\nu}^2)$$

That is, for order j the log of the random effect is distributed normally with mean 0 and variance σ_{ν}^2 .

Running the model in JAGS

First we set up the simulation

```
# number of orders
nOrd <- 8

# number of species
nspp <- nOrd * 10

# vector identifying which species are in which orders
ordID <- rep(1:nOrd, each = nspp/nOrd)

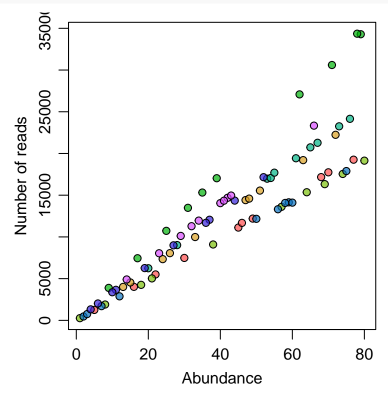
# simulate random effect
ordSD <- 0.25
set.seed(1)</pre>
```

```
ordEffect <- exp(rnorm(nOrd, mean = 0, sd = ordSD))

# generate true abundances for each species
set.seed(1)
x <- sample(seq(1, 80, length.out = nspp))

# simulate number of reads for each species
totReads <- 10^6
set.seed(1)
xreads <- rmultinom(1, totReads, x*ordEffect[ordID])[, 1]</pre>
```

A quick plot of what those simulated data look like (colors correspond to order identity)



Define the model in JAGS using rjags

```
## Loading required package: coda
## Linked to JAGS 4.2.0
```

library(rjags)

Loaded modules: basemod, bugs

```
# write the model out
cat('model{
        # reads model
        xreads[1:S] ~ dmulti(alpha[1:S], totReads)
        # priors
        sigX ~ dunif(0, 100)
        tauX <- 1/(sigX^2)
        sigCopy ~ dunif(0, 100)
        tauCopy <- 1/(sigCopy^2)</pre>
        # species-level effect
        for(i in 1:S) {
            logX[i] ~ dnorm(N/S, tauX)
            x[i] <- exp(logX[i])</pre>
        }
        # order-level random effect
        for(j in 1:nOrd) {
            logNu[j] ~ dnorm(0, tauCopy)
            nu[j] <- exp(logNu[j])</pre>
        # define multinom param
        for(i in 1:S) {
            alpha[i] <- x[i] * nu[ordID[i]]</pre>
    }',
    file = 'multiRandEffect.jag' )
```

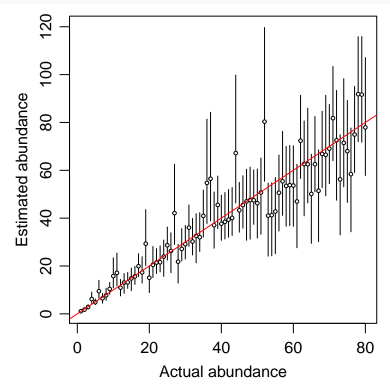
Compile the model and configure it for MCMC:

```
## Resolving undeclared variables
## Allocating nodes
## Graph information:
## Observed stochastic nodes: 1
## Unobserved stochastic nodes: 90
## Total graph size: 385
##
## Initializing model
```

Now plot real versus estimated abundance. Error bars are 95% credible intervals, red line is 1:1 line:

```
xest <- sum(x) * samp[, grep('x', colnames(samp))] /
    rowSums(samp[, grep('x', colnames(samp))])
xestCI <- apply(xest, 2, quantile, prob = c(0.025, 0.975))

par(mar = c(3, 3, 0, 0) + 0.5, mgp = c(2, 0.75, 0))
plot(x, colMeans(xest), cex = 0.5,
    panel.first = {
        segments(x0 = x, y0 = xestCI[1, ], y1 = xestCI[2, ])
        points(x, colMeans(xest), col = 'white', pch = 16, cex = 0.5)
    },
    ylim = range(xestCI),
    xlab = 'Actual abundance', ylab = 'Estimated abundance')
abline(0, 1, col = 'red')</pre>
```



Plot σ_{order} , red line is true value:

