

```
for (i in 1:n_branches) {
        bl[i] \sim dnExponential(10.0)
topology \sim dnUniformTopology(taxa)
psi := treeAssembly(topology, bl)
alpha1 <- v(1,1,1,1,1,1)
alpha2 <- v(1,1,1,1)
er^- \sim dnDirichlet(alpha1)
pi \sim dnDirichlet(alpha2)
Q := fnGTR(er, pi)
m_{alpha} < ln(2.0)
sd_alpha <- 0.587405
alpha \sim dnLognormal( m_alpha, sd_alpha)
sr := fnDiscretizeGamma( alpha, alpha, 4, false )
p_{inv} \sim dnBeta(1,1)
seq \sim dnPhyloCTMC( tree=psi, Q=Q, pInv=p_invar,
                    siteRates=sr, type="DNA" )
seq.clamp( data )
```