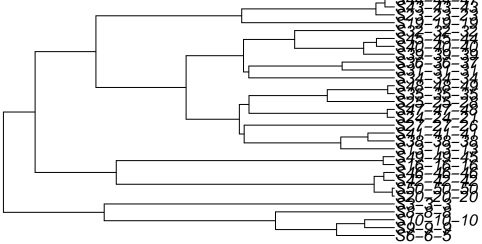
R Notebook

Speciation long:

```
results <- list()
params_a <- flechette::create_params(</pre>
  speciation_initiation_rate = 0.2,
  speciation_completion_rate = 5.0,
  extinction_rate = 0.1,
  crown_age = 15,
  crown_age_sigma = 0.01,
  sampling_method = "youngest",
  mutation_rate = 1.0,
  sequence_length = 100,
  mcmc_length = 300000,
  tree_sim_rng_seed = 43,
  alignment_rng_seed = 43,
  beast2_rnd_seed = 44
results$a <- flechette::run(parameters = params_a)</pre>
ape::plot.phylo(results$a$species_tree)
```



```
results$a$nltts_diff <- nLTT::nltts_diff(tree = results$a$species_tree, results$a$trees)
tracerer::calc_ess(results$a$estimates$posterior, sample_interval = 1000)
## [1] 300.2815
Short:</pre>
```

```
params_b <- params_a
params_b$speciation_completion_rate <- 1000000.0
results$b <- flechette::run(parameters = params_b)
ape::plot.phylo(results$b$species_tree)</pre>
```

```
$19-19-19
$25-25-25
$29-29-29
$30-30-30
$24-24-24
$22-22-22
$21-21-21
$28-28-28
$20-20-20
$27-27-27
$3-3-3
$13-13-13
$9-9-9
$11-11-11
$10-10-10
$5-5-5
$7-7-7
$8-8-8
$1-1-1
```

```
results\$b\$nltts_diff <- nLTT::nltts_diff(tree = results\$b\$species_tree, results\$b\$trees)
tracerer::calc_ess(results\$b\$estimates\$posterior, sample_interval = 1000)
```

```
## [1] 301
Plot:

df <- data.frame(
    sim = c(
        rep("a", times = length(results$a$nltts_diff)),
        rep("b", times = length(results$b$nltts_diff))
    ),
    nltts = c(results$a$nltts_diff, results$b$nltts_diff)
)
ggplot2::ggplot(
    data = df,
    ggplot2::aes(x = nltts, fill = sim)
) + ggplot2::geom_histogram(alpha = 0.5, position = "identity", binwidth = 0.01)</pre>
```

