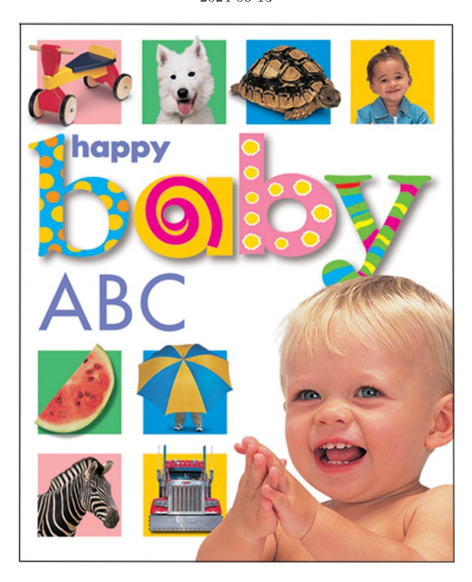
Introduction to Simulation for Biologists: Approximate Bayesian Computation (ABC) in ${\tt R}$

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Introduction

Likelihood function, which depicts the probability of observed data as a function of parameters of a statistical model, is essential in statistical inference. For many biological problems, it is challenging for us empiricists — and often even for theoreticians — to derive likelihood functions. **Approximate Bayesian Computation** (ABC) bypasses analytical evaluation of the likelihood function using simulation.

In ABC, the likelihood function is approximated by comparing simulated data (under a model with sampled parameters) with the observed data. In the simplest form of ABC with the rejection algorithm, a set of parameters are sampled from a prior distribution. Given a parameter θ , a dataset D_{sim} is simulated under a model M. Parameter θ under model M is accepted when

$$\rho(D_{sim}, D_{obs}) \le \epsilon$$

where $\rho(D_{sim}, D_{obs})$ denotes the distance (e.g. Euclidean) between D_{sim} and D_{obs} , and $\epsilon \geq 0$ is tolerance. In simple words, this means that we reject a parameter θ under model M if the simulated data D_{sim} is too different from D_{obs} .

The probability of generating a dataset D_{sim} closer to the observed dataset D_{obs} than the distance measure of ϵ typically decreases as the dimensionality of the data increases. This is problematic for ABC because it reduces the computational efficiency as would required more replicates of simulation. To deal with this issue, a set of summary statistics S(D) with lower dimensionality are used, instead of raw data D. In the Luria-Delbrück experiment, one can use the mean and variance of resistant colonies per plate instead of the frequency distribution of the number of colonies. By substituting D_{sim} and D_{obs} with $S(D_{sim})$ and $S(D_{obs})$, the acceptance criterion of the ABC rejection algorithm becomes

$$\rho(S(D_{sim}), S(D_{obs})) \le \epsilon$$

By applying this rejection algorithm to all simulated data under model M, we obtain a subset of sampled parameter values, the distribution of which can be regarded as an approximation of the posterior distribution.

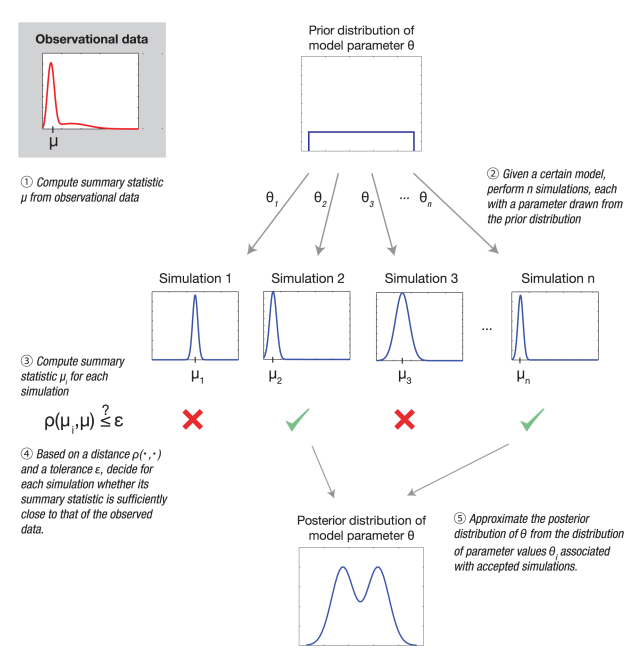


Figure 1: Parameter estimation by ABC. Adapted from Sunnåker et al. (2013)

In addition to parameter inference described above, ABC can be used for model selection (Bertorelle, Benazzo, and Mona 2010; Csilléry, François, and Blum 2012). Posterior probability of a model M_i is approximated as

$$Pr(M_i|\rho(S(D_{sim}), S(D_{obs})) \le \epsilon)$$

In simple words, the posterior probability of model M_i is approximated as the proportion of accepted simulations of model M_i out of accepted simulations of all models.

Here, we will apply ABC to the Luria-Delbrück experiment. The objectives are

- 1. to determine which of the models (i.e. induced or spontaneous mutation) is the truth;
- 2. to estimate the mutation rate under the model

In this exercise, we will apply ABC to the Luria-Delbrück experiment. First, we will determine which of models (induced and spontaneous mutation) fits better to the data. Second, we will infer the parameter (mutation rate). To this end, we will use the abc package (Csilléry, François, and Blum 2012), which implements the rejection algorithm described above in functions. Optionally, you can try other methods than the rejection algorithm implemented in abc (multinomial logistic regression and neural networks for model selection; local linear regression and neural networks for parameter inference) and ABC random forest implemented in abcrf (Raynal et al. 2019).

Installation of required packages

We will use

- abc
- abcrf
- fluctuateR

fluctuateR contains a few functions defined during the first exercise. You can use your own functions instead if you want.

To install them, run:

```
install.packages("abc")
install.packages("abcrf")
devtools::install_github("junishigohoka/fluctuateR")

To load them, run:
library("abc")
library("abcrf")
library("fluctuateR")
```

Simulation of a Luria-Delbrück experiment

Here we will run a simulation, the output of which will be used as our observation. As biologists in 2024 we know that the spontaneous mutation model is the case, so we run this simulation using simLD_spo. The mutation rate is read from a compressed file data/mu_truth.txt.gz, which we will infer with ABC later.

To be a little bit more realistic, I suggest we change the parameter values of the experiment from the previous exercise.

First, I do not want to waste my working hours waiting for cells to grow. I also do not want to leave the lab late at night or come to the lab early in the morning. Because I work 8 hours a day, overnight is 24 - 8 = 16 hours. Assuming the cells divide 3 times every hour, the number of generations overnight is $3 \times 16 = 48$. In addition, because I am lazy, I do not want to count cells before plating. So I would rather take a fixed proportion of the medium of the tube (1/1,000,000,000) by serial dilution. Let's run an experiment and store the mean and variance of the number of resistant colonies in a named vector $\mathbf{d}_{-}\mathbf{obs}$.

```
n_{sample} = n_{sample},
11
               n_plates = r
   )
13
   d_obs
    ##
            mean a
                         mean_b
                                       var_a
                                                    var b
                      40.58000
```

27.63265 1575.84041

Simulation for inference

##

28.80000

In ABC, we need to sample parameters and simulate data for models. In our Luria-Delbrück experiment, we have two models: induced and spontaneous mutation. We will simulate each model 10,000 times with mutation rates sampled from a log uniform distribution between 10^{-10} and 10^{-5} . We store sampled mutation rates in a vector mu sim.

```
nsims <- 10000
   if(file.exists("data/mu_sim.rds")){
           mu sim <- readRDS("data/mu sim.rds")</pre>
   }else{
4
           mu_sim <- 10^runif(nsims, -10, -5)</pre>
5
            saveRDS(mu_sim, "data/mu_sim.rds")
6
   }
   head(mu_sim)
```

```
## [1] 6.539262e-09 2.281259e-08 1.990270e-10 2.274115e-09 8.646574e-06 1.351128e-07
```

First, let's simulate the Luria-Delbrück experiment for these mutation rates under the spontaneous mutation model. The simulated data will be in a data frame d_sim_spo. In the code block below, I simulate the data if simulated data has not been written in data/d_sim_spo.rds and store it in data/d_sim_spo.rds.

```
if(file.exists("data/d_sim_spo.rds")){
            d_sim_spo <- readRDS("data/d_sim_spo.rds")</pre>
2
   }else{
3
            d_sim_spo <- as.data.frame(t(sapply(mu_sim,</pre>
                    function(x){
5
                             simLD_spo(T, x, 100, n_0 * (2^T)/1e9, r)
                    }
            )))
            saveRDS(d_sim_spo, "data/d_sim_spo.rds")
9
   }
10
11
12
13
   head(d_sim_spo)
```

```
##
       mean_a
                mean_b
                               var_a
                                            var_b
## 1
         8.62
                  6.70 5.832245e+00 1.037755e+01
## 2
        24.66
                 23.20 2.622898e+01 1.266939e+02
## 3
         0.10
                  0.10 9.183673e-02 9.183673e-02
## 4
         2.30
                  2.14 1.765306e+00 2.163673e+00
## 5 13099.66 11173.52 1.152745e+04 1.118872e+07
                145.34 1.071759e+02 1.489902e+03
```

Second, let's simulate the Luria-Delbrück experiment under the induced mutation model. The simulated data will be in a data frame d sim ind.

```
if(file.exists("data/d_sim_ind.rds")){
            d_sim_ind <- readRDS("data/d_sim_ind.rds")</pre>
2
   }else{
            d_sim_ind <- as.data.frame(t(sapply(mu_sim,</pre>
4
                   function(x){
5
                            simLD_ind(n_plates = r, n_sample = as.integer(n_sample), mut_rate = x)
6
                   }
            )))
8
            saveRDS(d_sim_ind, "data/d_sim_ind.rds")
9
   }
10
11
12
13
   head(d_sim_ind)
14
   ##
        mean a mean b
                              var a
                                            var b
                                      0.15061224
   ## 1
          0.16
                  0.18
                         0.1371429
   ## 2
          0.50
                  0.70
                         0.5816327
                                      0.62244898
   ## 3
          0.00
                  0.00
                         0.0000000
                                      0.00000000
   ## 4
          0.00
                  0.10
                         0.0000000
                                      0.09183673
   ## 5 245.12 243.28 136.9648980 202.94040816
          4.32
                  4.52
                          3.6506122
                                      4.37714286
   Let's concatenate the data frames into a single data frame d_sim.
   d_sim <- as.data.frame(rbind(d_sim_spo, d_sim_ind))</pre>
   head(d_sim)
   ##
           mean_a
                    mean_b
                                   var_a
                                                 var_b
   ## 1
             8.62
                      6.70 5.832245e+00 1.037755e+01
   ## 2
            24.66
                     23.20 2.622898e+01 1.266939e+02
                      0.10 9.183673e-02 9.183673e-02
   ## 3
             0.10
   ## 4
             2.30
                      2.14 1.765306e+00 2.163673e+00
   ## 5 13099.66 11173.52 1.152745e+04 1.118872e+07
                    145.34 1.071759e+02 1.489902e+03
   ## 6
           137.26
   tail(d_sim)
   ##
             mean_a mean_b
                                   var_a
                                                var_b
   ## 19995
               1.44
                      1.60
                              2.21061224 1.42857143
   ## 19996
               0.02
                      0.00
                              0.02000000 0.00000000
   ## 19997 88.30 87.58 104.78571429 82.37102041
   ## 19998
               0.04
                      0.04
                              0.03918367
                                          0.03918367
   ## 19999
               0.00
                      0.00
                              0.00000000 0.00000000
   ## 20000
               2.88
                      3.20
                              3.53632653 3.26530612
   The models of in total 20,000 simulations are recorded in a vector models.
   models <- rep(c("spo", "ind"), each = nsims)</pre>
   head(models)
   ## [1] "spo" "spo" "spo" "spo" "spo" "spo"
   tail(models)
   ## [1] "ind" "ind" "ind" "ind" "ind" "ind"
```

ABC using abc

This tutorial is based on the official vignette of abc, tailored for our Luria-Delbrück experiment. Interested readers are encouraged to read https://cran.r-project.org/web/packages/abc/vignettes/abcvignette.pdf as well as the original paper (Csilléry, François, and Blum 2012).

Cross-validation for model selection

To evaluate if ABC can, at all, distinguish between the two models, we perform a cross-validation for model selection using abc::cv4postpr, which implements a leave-one-out cross-validation. Here, we randomly take summary statistics of one simulation replicate as a pseudo-observation, and its parameter is estimated with the rejecting algorithm using all other simulations. This is repeated nval = 100 times, and numbers of classifications are summarised in a confusion matrix.

```
cv modsel rej <- cv4postpr(index = models,</pre>
                           sumstat = d_sim,
2
                           method = "rejection",
3
                           nval = 100,
                           tols = 0.05
5
  )
   summary(cv_modsel_rej)
   ## Confusion matrix based on 100 samples for each model.
   ##
   ## $to10.05
   ##
          ind spo
   ## ind 100
   ## spo 32
               68
   ##
   ##
   ## Mean model posterior probabilities (rejection)
   ##
   ## $to10.05
   ##
             ind
   ## ind 0.8288 0.1712
   ## spo 0.2361 0.7639
```

In the confusion matrix, the j-th column of the i-th row is the number of times that the i-th model were classified as the j-th model. Our confusion matrix shows that if the truth is the induced model, then you can tell that it is induced. If the truth is the spontaneous model, then you can tell that it is spontaneous 68 times out of 100. When the classified model is spontaneous, you can be sure that it is spontaneous. When the classified model is induced, the truth could still be the spontaneous model 32 times out of 132.

Model selection

Now, let's perform model selection with ABC using abc::postpr.

Call:

```
## postpr(target = as.data.frame(t(d_obs)), index = models, sumstat = d_sim,
##
       tol = 0.05, method = "rejection")
## Data:
    postpr.out$values (1000 posterior samples)
## Models a priori:
  ind, spo
##
## Models a posteriori:
##
    ind, spo
##
## Proportion of accepted simulations (rejection):
## ind spo
##
         1
##
## Bayes factors:
##
       ind spo
## ind
             0
## spo Inf
```

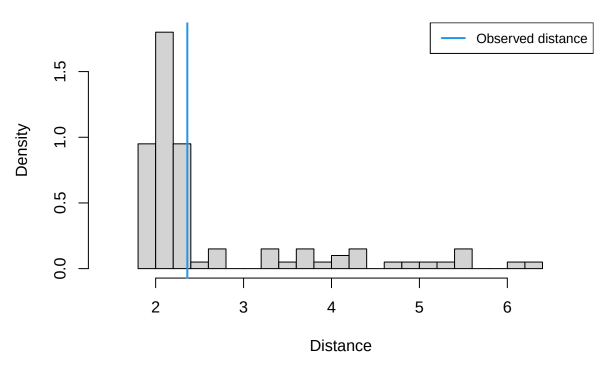
The summary shows that the posterior probability of the spontaneous model is 1 (!!!).

Goodness-of-fit

Before turning to parameter inference, it is important to check that the preferred model provides a good fit to the data. The null distribution is computed for the distance between true parameter value and the mean parameter values for accepted replicates with abc for nb.replicate = 100 randomly sampled simulation replicates. The observed distance (i.e. the distance based on the output of abc using observed data) is compared with the null distribution. If the model provides a good fit to the data, then the observed distance should be within the null distribution (say, p > 0.05).

```
gfit_abc <- gfit(target=log10(as.data.frame(t(d_obs)) + 1),</pre>
                 sumstat = log10(d_sim + 1),
2
                 tol = 0.05,
3
                 statistic=mean,
4
                 nb.replicate=100)
6
   summary(gfit_abc)
   ## $pvalue
   ## [1] 0.26
   ##
   ## $s.dist.sim
   ##
         Min. 1st Qu.
                                   Mean 3rd Qu.
                        Median
                                                    Max.
   ##
        1.891
                 2.063
                         2.182
                                  2.642
                                           2.597
                                                   6.322
   ##
   ## $dist.obs
   ## [1] 2.360076
  plot(gfit_abc)
```

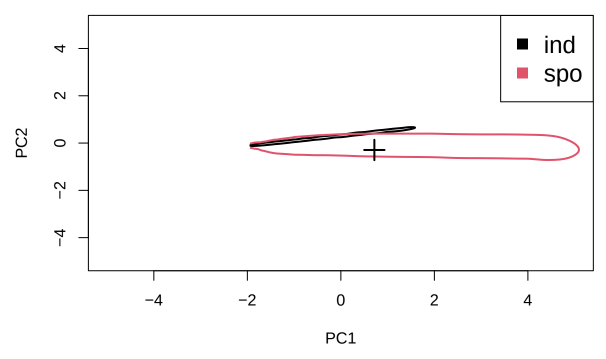
Histogramme of the null distribution



The observed distance is between mean and median of null distribution, so the model fits very well to the data.

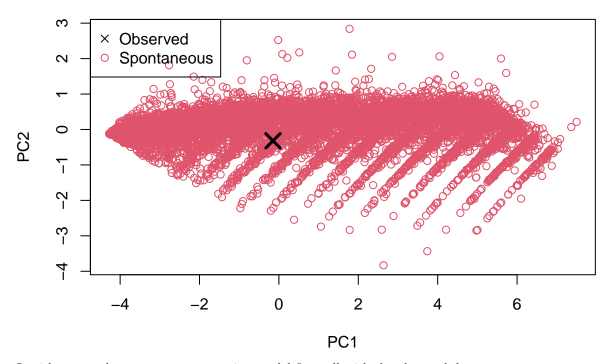
Another way to investigate the goodness-of-fit is PCA. We can visualise the PCA envelope for the considered models and plot the observed data onto it using abc::gfitpca.

```
## Warning in lfproc(x, y, weights = weights, cens = cens, base = base, geth = geth, : procv: parameter
## Warning in lfproc(x, y, weights = weights, cens = cens, base = base, geth = geth, : procv: parameter
## Warning in lfproc(x, y, weights = weights, cens = cens, base = base, geth = geth, : procv: parameter
```



Alternatively, we could also use a base R function prcomp.

```
d_sim <- as.data.frame(rbind(d_sim_spo, d_sim_ind))</pre>
2
3
   pca_both <- prcomp(log10(rbind(d_obs, d_sim_spo, d_sim_ind) + 1))</pre>
   pca_spo <- prcomp(log10(rbind(d_obs, d_sim_spo) + 1))</pre>
   plot(pca_spo$x[2:(nsims),1], pca_spo$x[2:(nsims),2], col = 2,
9
         xlab = "PC1",
10
         ylab = "PC2"
11
12
   points(pca_spo$x[1,1], pca_spo$x[1,2], pch = c(4,1), cex = 2, lwd = 3)
13
   legend("topleft",
14
           legend = c("Observed", "Spontaneous"),
15
           col = c(1, 2),
16
           pch = c(4, 1),
17
           pt.cex = c(1, 1)
18
19
```

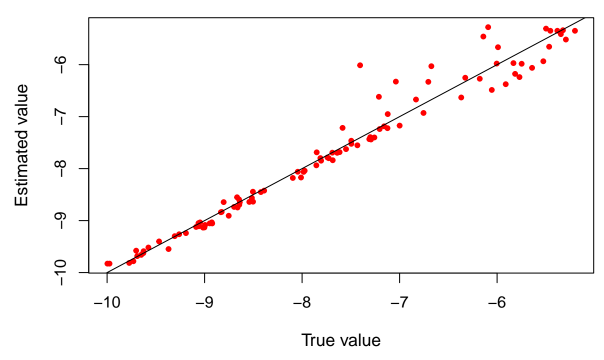


In either way, the spontaneous mutation model fits well with the observed data.

Cross-validation

As in model selection, we should check if parameter inference of ABC (implemented in abc::abc) can estimate the parameter.

log10(mut rate)



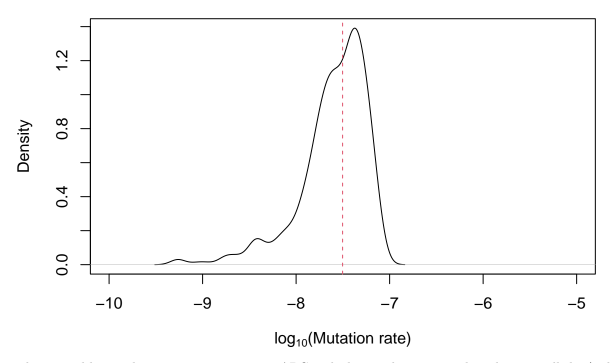
The log-transformed mutation rate can be estimated well with abc::abc.

Parameter inference

Let's estimate the mutation rate.

```
parinf_abc <- abc(target = as.data.frame(t(d_obs)),</pre>
                                                                                                              param = log10(mu_sim),
2
                                                                                                              sumstat = d_sim_spo,
                                                                                                              tol=0.05,
                                                                                                              method="rejection")
               \#\# Warning in abc(target = as.data.frame(t(d_obs)), param = log10(mu_sim), : No parameter names are given by the sum of the sum o
              plot(density(parinf_abc$unadj.value),
                                          xlim = c(-10, -5),
                                          xlab = expression(paste(log[10], "(Mutation rate)")),
                                          main = "Posterior distribution"
4
              )
              abline(v = log10(as.numeric(readLines("data/mu_truth.txt.gz"))),
6
                                                   col = 2,
                                                   lty = 2
```

Posterior distribution



The vertical line is the true mutation rate, so ABC with the simple rejection algorithm can tell the (order of) mutation rate very well.

```
10^quantile(parinf_abc$unadj.value, c(0.025, 0.5, 0.975))

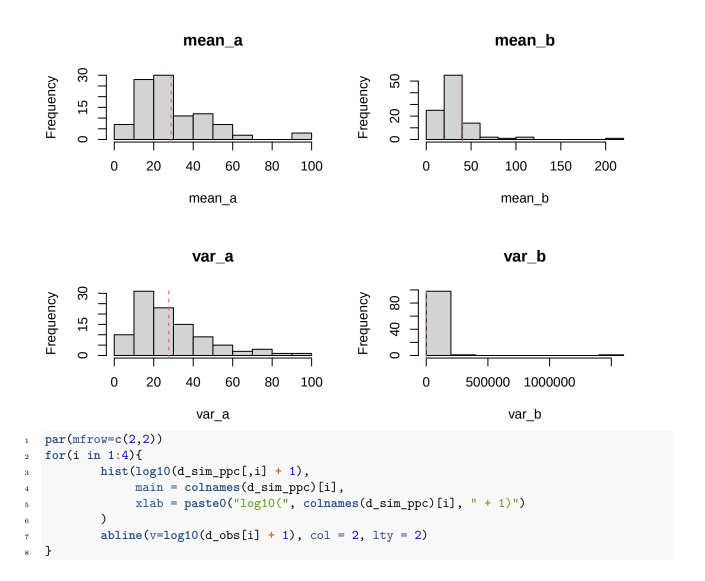
## 2.5% 50% 97.5%

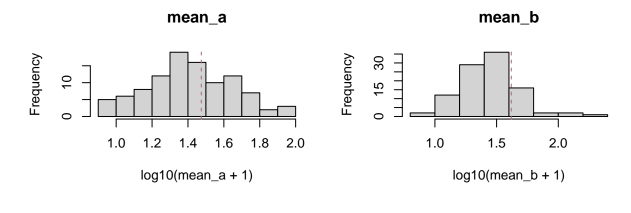
## 9.397760e-09 2.413971e-08 5.292799e-08
```

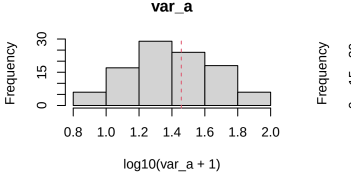
Posterior predictive check

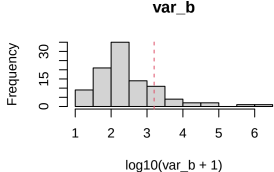
Lastly, let's check if the observed summary statistics are well represented by the summary statistics of the accepted simulations.

```
mu_posterior <- as.vector(10^ parinf_abc$unadj.values)</pre>
   d_sim_ppc <- as.data.frame(t(sapply(mu_posterior,</pre>
          function(x){
4
                   simLD_spo(T, x, 100, n_0 * (2^T)/1e9, r)
          }
   )))
   par(mfrow=c(2,2))
   for(i in 1:4){
2
           hist(d_sim_ppc[,i],
                 main = colnames(d_sim_ppc)[i],
4
                 xlab = colnames(d_sim_ppc)[i]
           abline(v=d_obs[i], col = 2, lty = 2)
   }
```









Let's get percentiles.

```
## mean_a mean_b var_a var_b
## 61 81 57 85
```

The observed data is reflected in the data simulated using parameter values taken from the (approximate) posterior distribution.

References

Bertorelle, G., A. Benazzo, and S. Mona. 2010. "ABC as a Flexible Framework to Estimate Demography over Space and Time: Some Cons, Many Pros." *Molecular Ecology* 19 (13): 2609–25. https://doi.org/10.1111/j.1365-294X.2010.04690.x.

Csilléry, Katalin, Olivier François, and Michael G. B. Blum. 2012. "Abc: An R Package for Approximate Bayesian Computation (ABC)." *Methods in Ecology and Evolution* 3 (3): 475–79. https://doi.org/10.111 1/j.2041-210X.2011.00179.x.

Raynal, Louis, Jean-Michel Marin, Pierre Pudlo, Mathieu Ribatet, Christian P Robert, and Arnaud Estoup. 2019. "ABC Random Forests for Bayesian Parameter Inference." *Bioinformatics* 35 (10): 1720–28. https://doi.org/10.1093/bioinformatics/bty867.

Sunnåker, Mikael, Alberto Giovanni Busetto, Elina Numminen, Jukka Corander, Matthieu Foll, and Christophe Dessimoz. 2013. "Approximate Bayesian Computation." *PLOS Computational Biology* 9 (1): e1002803. https://doi.org/10.1371/journal.pcbi.1002803.

Appendix

ABC random forest using abcrf

The tutorial below of abcrf is based on https://github.com/mnavascues/ABCRFtutorial.

Model selection

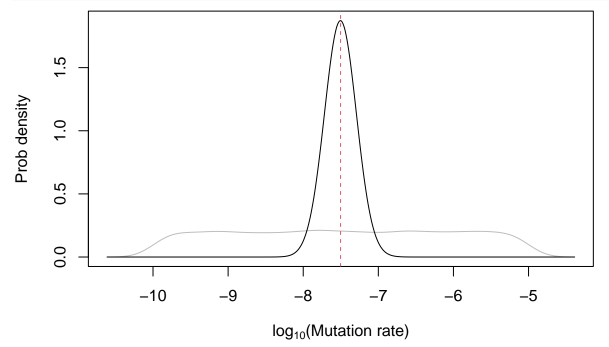
```
model_rf <- abcrf(formula = as.factor(models)~.,</pre>
                      data = d_sim,
                      ntree = 1000,
3
                      paral = TRUE,
4
                      ncores = 8
5
6
  modsel_rf <- predict(object = model_rf,</pre>
                          obs = as.data.frame(t(d_obs)),
                          training = d_sim,
3
                          ntree = 1000,
4
                          paral = TRUE,
                          paral.predict = TRUE
6
  )
  modsel_rf
        selected model votes model1 votes model2 post.proba
   ## 1
                                    0
                                               1000
                    spo
```

Parameter inference

```
model <- regAbcrf(formula = log10(mu_sim) ~ .,</pre>
                     data = d_sim_spo,
2
                     ntree = 1000,
3
                     paral = TRUE
   )
5
   posterior <- predict(object = model,</pre>
                         obs = as.data.frame(t(d_obs)),
2
                         training = d_sim_spo,
                         paral = TRUE,
4
                         rf.writes = T
   )
6
   print(posterior)
           expectation
                          median variance (post.MSE.mean) variance.cdf quantile=0.025 quantile=0.975 post.
           -7.504539 -7.495566
                                                0.001051972 0.003586242
   ## [1,]
                                                                                -7.62441
                                                                                               -7.407288
   densityPlot(object = model, obs = as.data.frame(t(d_obs)), training = d_sim_spo, paral = TRUE,
               xlab = expression(paste(log[10], "(Mutation rate)")),
               ylab = "Prob density",
               main = ""
   )
```

Warning in density.default(resp, weights = weights.std[, i], ...): Selecting bandwidth *not* using ' ## Warning in density.default(resp, weights = weights.std[, i], ...): Selecting bandwidth *not* using '

```
abline(v = log10(as.numeric(readLines("data/mu_truth.txt.gz"))),
col = 2,
lty = 2
)
```



```
smr <- c(10^posterior$med, 10^posterior$quantiles)
names(smr) <- c("median", "q2.5", "q97.5")
print(smr)</pre>
```

```
## median q2.5 q97.5
## 3.194732e-08 2.374595e-08 3.914826e-08
```

Truth

```
print(as.numeric(readLines("data/mu_truth.txt.gz")))
```

[1] 3.141593e-08