TeTame 2.1



Estimation of neutral parameters by maximum likelihood

http://www.edb.ups-tlse.fr/equipe1/tetame.htm

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Introduction

Hubbell's neutral theory of biodiversity (Hubbell, 2001) proposes a simple explanation of the maintenance of biodiversity as a result of stochastic processes of birth, death, immigration and speciation. In this model, the species relative abundances in a guild is determined by two parameters: θ , that governs the appearance of new species in the regional species pool, and m that governs immigration into local communities of individuals from the regional species pool. This model is formally analogous to a continent-island model (Wright 1931).

This model is seen as a potentially useful null model in ecology. But for this assertion to hold, an efficient estimation of these two parameters is needed. In the case where m=1 (no dispersal limitation, that is, all newborn individuals are immigrants), the likelihood of the sole parameter θ can be computed using Ewens sampling formula (Ewens, 1972), given a species abundance dataset. However, in the general case where both θ and m can take non-trivial values, the previous methods of parameter estimation consisted in sequentially estimating θ and m, leading to approximated solutions (see e.g. Hubbell 2001, McGill 2003, Volkov et al. 2003, Latimer et al. 2005).

Parameter estimation can now be rigorously performed by maximum-likelihood using the sampling formula developed by Etienne (Etienne and Olff 2005, Etienne, 2005; Etienne and Alonso 2005; Etienne et al. 2006). Etienne (2005) provided a program for this estimation in the precompiled programming language "PARI" (http://pari.math.u-bordeaux.fr/).

We here provide **TeTame2**, an easy-to-use and freely available software written in C++. This software estimates the parameters θ and *m* using Etienne's method and produces a list of likelihood values in a parameter space that can be subsequently plotted, e.g. using the R software. This software has several advantages over the version previously published by Etienne (2005), being portable, computationally fast, and simple to use.

This new version additionally computes the immigration rates in several samples belonging to the same regional pool. The underlying assumption is that the species abundances pooled among these samples reflect regional species abundances. The theory underlying this computation can be found in Jabot et al. (2008).

Thanks to Andrea Manica (U. Cambridge), Rampal Etienne (U. Groningen) and David Alonso (U Groningen) for troubleshooting the program.

Hardware and program installation

TeTame2 for Windows has been compiled using the Linux-emulated environment cygwin on Windows and its C++ GNU compiler g++. So far, it has been tested on Windows XP Professional but is expected to run on most environments.

To install **TeTame2**, download it into the same directory as your file containing the species abundance data.

Running the program

1- Formatting your data

<u>Case 1:</u> you want to analyze the species abundance distribution of a **single** sample. Save your data in a file with the extension ".txt". Your file should have the following structure:

```
abundance_species_1
abundance_species_2
...
abundance_species_n
```

Example: If your sample consists of (3 willows, 5 oaks, 1 eucalyptus), then your data entry file should read:

3 5 1

<u>Case 2:</u> you want to analyze the species abundance distribution of **multiple** samples. Save your data in a file with the extension ".txt". Your file should have the following structure:

abundance_species_1_sample1 abundance_species_2_sample1 ... abundance_species_n_sample1 & abundance_species_1_sample2 abundance_species_1_sample2 ... abundance_species_n_sample2 ... abundance_species_1_sampleD abundance_species_2_sampleD ... abundance_species_1_sampleD

<u>Example</u>: If your dataset consists of one sample of (3 willows, 5 oaks, 1 eucalyptus, 0 poplar), one sample of (1 willows, 15 oaks, 0 eucalyptus, 0 poplar), and one sample of (2 willows, 2 oaks, 0 eucalyptus, 3 poplars), then your data entry file should read:

3

5 1

0

&

1

15

<u>NB:</u> make sure you pressed enter at the end of the data file.

2- Running the program

Double-click on the executable file (tetame2.exe) and answer the questions in the console. Yes/no questions can be answered by using '0', 'n', or 'no' for no, and '1', 'y', or 'yes' for yes. You may want to test the program first with the dataset provided in the download webpage (test.txt). The typical result produced on our machine is provided in the appendix below.

Case 1: single sample analysis

Maximum-likelihood estimation

The values of the different parameters are output both in the console and in the file called [name_of_file]_out.txt created during the execution and sorted in your working directory. For the test dataset (test.txt), the output file name is thus "test_out.txt". The output file of the 'test.txt' dataset looks like:

<u>s</u>	J 112	Theta	Std_Theta	I	a 1000	Std_I	011	<u>m</u>	<u>Std_m</u>	loglike_min	Theta_Ewens
0	112	1.19528	0.383777	5.90208	e+009	5.100040+0	011	1	9.933716-007	10.7797	1.19529
6	63	2.94777	1.77646	7.04503		6.36662		0.102	0.0828011	12.2275	1.43357
6	181	21.7474	57.0798	1.24466	5	0.618653		0.006	0.00338992	13.5365	1.05382
loglike	Ewens	Theta2	Std_Theta2		I2	Std_I2	m2		Std_m2	loglike_m	in2
10.779	7	6.08536e+008	1.09707e+0	11	1.1953	0.583787	0.01	06537	0.00514788	10.7797	
13.312	9	2.94777	1.77646		7.04502	6.36661	0.10	2035	0.082801	12.2275	
13.558	1	1.0563	0.508725		9290.48	408693	0.98	0994	0.820211	13.5579	

S is the number of species in the sample, J is the total number of individuals in the sample, Theta is the maximum-likelihood estimate of θ , Std_Theta is the standard deviation of θ (under the assumption that the posterior is a Normal distribution) I is equal to m*(J-1)/(1-m) and it is a rescaled immigration rate, Std_I is the standard deviation of I, Ewens_Theta is the value of θ estimated from Ewens sampling formula (assuming that m=1), loglike_Ewens is the minimum of the opposite of the log-likelihood (assuming that m=1), loglike_min is the minimum of the opposite of the log-likelihood, and m is the immigration rate.

<u>NEW FEATURE</u> of **TeTame2**: if the programs detect a second likelihood maxima, it reports this second value too, and not just the global maximum as in the previous version of **TeTame**.

Likelihood surface plotting

It can be useful to visualize the shape of the likelihood function (see e.g. Etienne et al. 2006) for a worrisome example). Using **TeTame2**, you can output a list of log-likelihoods for values of (θ, m) on a grid. The likelihood surface can subsequently be plotted by using the freely available R software for example (and thus to have an idea of the uncertainty of the estimates). These options are provided to the user after the maximum-likelihood estimates of θ and m have been computed.

A user-supplied number of points on a grid are generated in the rectangular domain [thetamin, thetamax]*[mmin,mmax], where the four values are also user-supplied.

NB: if you have multiple samples, launch TeTame2 independently for each sample, otherwise you will only be able to draw the likelihood surface of the last sample.

Case 2: multiple sample analysis with Jabot et al (2008)'s method

Maximum-likelihood estimation

The values of the different parameters are output both in the console and in the file called [name_of_file]_out.txt created during the execution and sorted in your working directory. For the test dataset (test.txt), the output file name is thus "test outm.txt".

The output file of the 'test.txt' dataset looks like:

S	J	Ι	Std_I	m		Std_m	loglike_mir	ı
6	112	7.4197 1	5.9244	0.0626559	0.12604	48 15	.9501	
6	63	5.738 1	3.5052	0.0847088	0.18248	86 20.	.1273	
6	181	14.6712	24.470	1 0.07	53638	0.116226	15.9371	
S is	the number	er of spec	ties in the sam	ple, J is the	total numl	per of indi	viduals in the sam	mple, I is

equal to m*(J-1)/(1-m) and it is a rescaled immigration rate, Std_I is the standard deviation of I, loglike_min is the minimum of the opposite of the log-likelihood, and m is the immigration rate.

Post-processing with the R software

We suggest you to use the freely available R statistical software to plot the likelihood surface (http://www.r-project.org/). In R, you should make sure that your working directory is the one where you saved your data file (go to menu 'File', then 'Change the directory'). The R commands for plotting the likelihood surface are provided in the output file "name of file outR.txt". (in the example, this is the file test outR.txt.

Troubleshooting

- 1. For problems with the R software, there is an online help on the R website http://www.r-project.org/
- 2. The software's optimization can be stuck in a local maximum. You can try to avoid this issue by supplying initial parameter values. You may also check that you reached a global maximum by using the plotting device.
- 3. Any feedback on the software is most welcome (chave@cict.fr).

Version 1.0: J Chave 17-03-2005

Version 1.01: J Chave and R Etienne 17-05-2005

Version 1.02: J Chave 15-11-2005

Version 1.1: J Chave and F Jabot 02-02-2006

Version 2.0: J Chave and F Jabot 20-05-2008

References

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Ewens W.J., 1972. The sampling theory of selectively neutral alleles. Theor. Popul. Biol., 3, 87-112.

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McGill B.J., 2003. A test of the unified neutral theory of biodiversity. Nature, 422, 881-885.

Volkov I., Banavar J.R., Hubbell S.P. and Maritan A., 2003. Neutral theory and relative species abundance in ecology. Nature, **424**: 1035-1037.

Wright S., 1931. Evolution in Mendelian populations. Genetics, 16: 97–159.

APPENDIX

Try the test dataset (test.txt) to make sure that **TeTame2** outputs correct results:

ESTIMATING NEUTRAL PARAMETERS BY MAXIMUM LIKELIHOOD

This program can be used for:

1-estimating theta and m of Hubbell's 2001 neutral theory using Etienne's 2005 m ethod.

2-estimating m in several samples belonging to the same regional pool using Jabo t et al. 2008 method.

For more details, see the manual.

Options for answering yes/no questions: 1, y, or yes for 'yes'; 0, n, or no for 'no'.

Please enter the data file name (without'.txt') test Input file: test.txt Reading the file stats ... Number of samples: 3

TeTame detected several samples in your data file. If you want to do the simultaneous estimation of Theta and m, using Etienne (200 5)'s likelihood formula, press e and enter If you want to estimate m in each sample, assuming that they belong to the same metacommunity, using Jabot et al.(2008)'s method, press j and enter In the case you want to do the multi-samples m inference, make sure you entered the pooled-over-the-samples species abundances in the beginning of the data file.

Case 1: you enter 'e'

e In sample 1, number of species: 6 In sample 2, number of species: 6 In sample 3, number of species: 6 Start computing Stirling numbers ... Start computing ln(K(D,A)) ... Number of individuals: 112

Sample 1 Maximal abundance: 80 Compute the Ewens theta and log-likelihood ... Ewens' -log-likelihood: 10.7797 Maximizing the likelihood ... Would you like to provide initial values for the optimization procedure? 0

RESULTS (also output in file named: test_out.txt):

Theta Std Theta Std I m S JΙ Std m loglike loglike_Ewens Theta2 Std_Theta2 *Theta_Ewens* _min *I2* Std_I2 Std_m2 loglike_min2 *m*2 1.19528 112 5.96268e+0093.18884e+0111 6 0.583777 9.95571e-1.19529 6.08536e+0081.09707e+0111.1953 007 10.7797 10.7797 0.583787 0.0106537 0.00514788 10.7797

Would you like to plot the likelihood surface? 0 Number of individuals: 63

Sample 2

... it goes on for the different samples.

Case 2: you enter 'j' (after the computation, the program will close automatically so you may not see the following, but you will find the results in the file "[name_of_datafile]_outm.txt")

j In the total dataset, there are 6 species In sample 1, number of species: 6 In sample 2, number of species: 6 In sample 3, number of species: 6 Number of individuals in the pooled samples: 355 In sample 1: Number of individuals: 112 Maximal abundance: 80 Number of species present in the sample: 6 Maximizing the likelihood ...

RESULTS (also output in file named: test_outm.txt):

S	J	I ,	Std_I	m	Std_m	loglike	e_min		
6	112	7.49	668 1	6.0164	0.06326	649	0.126612	j	15.9234

In sample 2: Number of individuals: 63 Maximal abundance: 20 Number of species present in the sample: 6 Maximizing the likelihood ...

 RESULTS (also output in file named: test_outm.txt):

 S
 J
 I
 Std_I
 m
 Std_m
 loglike_min

 6
 63
 5.70176
 13.4441
 0.0842187
 0.181855
 20.1619

In sample 3: Number of individuals: 181 Maximal abundance: 81 Number of species present in the sample: 6 Maximizing the likelihood ...

 RESULTS (also output in file named: test_outm.txt):

 S
 J
 I
 Std_I
 m
 loglike_min

 6
 181
 14.6548
 24.4522
 0.0752862
 0.116161
 15.9517

Post-processing :

If you do the simultaneous estimation of θ and *m* (case 1), you can output the likelihood surface in R. TeTame2 is outputting a file named name_of_file_outR.txt. (in the example, this is the file test_outR.txt). This file contains the commands that you may use in the R software to produce graphs. You just need to paste and copy this file in the R software. (Make sure that you have changed the working directory in the R software before doing this) More details can be found in the manual of the first version of **TeTame**.