Lepidomeda aliciae dataset

POPULATION SIZE, MIGRATION, DIVERGENCE, ASSIGNMENT, HISTORY

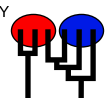
Bayesian inference using the structured coalescent

Migrate-n version 4.4.4(git:v4-series-26-ge85c6ff) [June-1-2019]

Compiled for a SYMMETRIC multiprocessors (Grandcentral)

Program started at Fri May 21 23:23:30 2021

Program finished at Sat May 22 02:57:06 2021 [Runtime:0000:03:33:36]



Options

Inheritance scalers in use for Thetas:

0.25 0.25 1.00

[The locus with a scaler of 1.0 used as reference]

Random number seed: (with internal timer) 1441484812

Start parameters:

Theta values were generated RANDOM start value from the prior

M values were generated RANDOM start value from the prior

Connection matrix:

m = average (average over a group of Thetas or M,

s = symmetric migration M, S = symmetric 4Nm,

0 = zero, and not estimated,

* = migration free to vary, Thetas are on diagonal

d = row population split off column population, D = split and then migration

Population 1 2 1 Great_Salt_Lake * 0 2 Sevier_Desert d *

Order of parameters:

 $\begin{array}{cccc} \mathbf{1} & & \Theta_1 & & \text{<displayed>} \\ \mathbf{2} & & \Theta_2 & & \text{<displayed>} \\ \mathbf{3} & & \Delta_{1->2} & & \text{<displayed>} \end{array}$

4 $\sigma_{1\rightarrow 2}$ <displayed>

Mutation rate among loci: Mutation rate is constant for all loci

Analysis strategy: Bayesian inference

-Population size estimation: Exponential Distribution

-Geneflow estimation: Exponential Distribution

-Divergence time estimation: Normal Distribution Shortcut (mean and standard dev.)

Proposal distributions for parameter

Parameter Proposal
Theta Metropolis sampling
M Metropolis sampling
Divergence Metropolis sampling
Divergence Spread Metropolis sampling
Genealogy Metropolis-Hastings

Prior distribution for parameter

| Par | ameter | | Prior | Minimum | MeanMa | iximum | Delta | Bins | UpdateFreq |
|-----|--------------------|-----|---------|----------|--------|--------|-------|------|------------|
| 1 | Theta ³ | * * | Uniform | 0.000000 | 0.050 | 0.100 | 0.010 | 1500 | 0.12500 |
| 2 | Theta ³ | * * | Uniform | 0.000000 | 0.050 | 0.100 | 0.010 | 1500 | 0.12500 |
| 3 | Splittime mean | ** | Uniform | 0.000000 | 0.250 | 0.500 | 0.050 | 1500 | 0.12500 |
| 4 | Splittime std | * * | Uniform | 0.000000 | 0.250 | 0.500 | 0.050 | 1500 | 0.12500 |

^{[* *} means priors were set globally]

Markov chain settings:

Number of chains

Long chain

1

Recorded steps [a] 10000
Increment (record every x step [b] 1000
Number of concurrent chains (replicates) [c] 2
Visited (sampled) parameter values [a*b*c] 20000000

Visited (sampled) parameter values [a*b*c] 20000000

Number of discard trees per chain (burn-in) 1000

Multiple Markov chains:

Static heating scheme 4 chains with temperatures

1000000.00 3.00 1.50 1.00

Swapping interval is 1

Print options:

Data file:
Haplotyping is turned on:
Output file:
Log file:
logfile

| Posterior distribution raw histogram file: | bayestii |
|---|----------------|
| Raw data from the MCMC run: | bayesallfile.g |
| Print data: | N |
| Print genealogies [only some for some data type]: | Nor |
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Data summary

| Data file Datatype Number | e: | | | | | | | На | infile aplotype data 3 |
|---------------------------------|-----------|-------------|---------|---------|--------------|------------------|--------|---------|------------------------------|
| Mutation | model: | | | | | | | | |
| Locus S | ublocus | Mutationn | nodel | Mu | itationmodel | parameters | | | |
| 1 | 1 | Felsenste | in 84 | [Bf:0.2 | 5 0.29 0.17 | 0.30, t/t ratio= | 2.000] | | |
| 2 | 1 | Felsenste | in 84 | - | | 0.35, t/t ratio= | _ | | |
| 3 | 1 | Felsenste | | - | | 0.31, t/t ratio= | _ | | |
| | | | | | | | | | |
| Sites pe | r locus | | | | | | | | |
| Locus | 10003 | Sites | | | | | | | |
| 1 | | 1109 | | | | | | | |
| 2 | | 849 | | | | | | | |
| 3 | | 448 | | | | | | | |
| 3 | | 440 | | | | | | | |
| Site rate | variation | and probabi | lities: | | | | | | |
| | | Region type | | nange | Probability | Patch size | | | |
| | | -5 - 71 - | | | | | | | |
| 1 | 1 | 1 | 1.000 |) | 1.000 | 1.000 | | | |
| 2 | 1 | 1 | 1.000 |) | 1.000 | 1.000 | | | |
| 3 | 1 | 1 | 1.000 |) | 1.000 | 1.000 | | | |
| Populati | on | | | | | Locus | | Gene co | opies |
| · | | | | | | | | data | (missing) |
| 1 Great_ | _Salt_Lak | ке | | | | 1 | | 10 | - |
| | | | | | | 2 | | 4 | |

(0)

(0)

(0)

2 Sevier_Desert

Total of all populations

Bayesian Analysis: Posterior distribution table

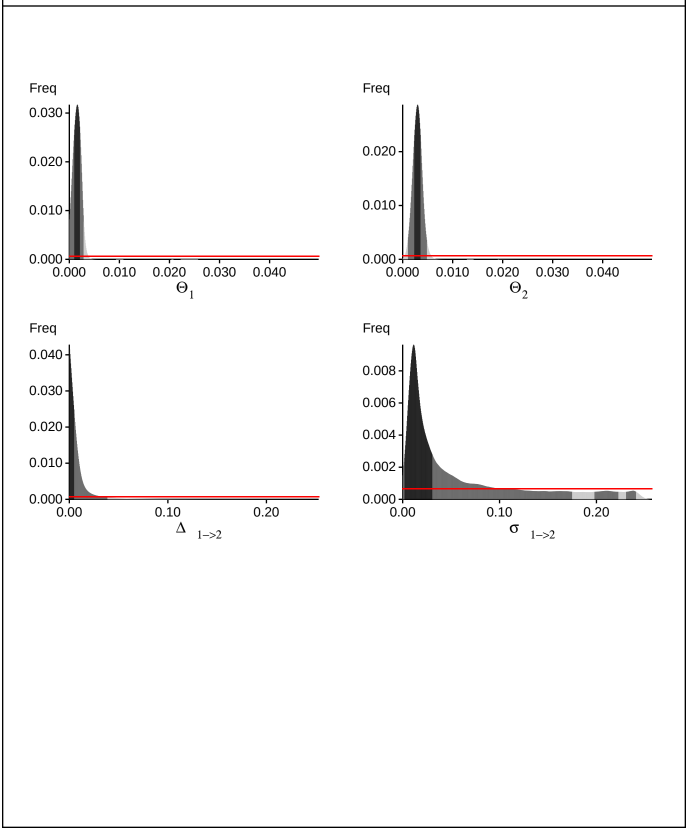
| Locus | Parameter | 2.5% | 25.0% | Mode | 75.0% | 97.5% | Median | Mean |
|-------|----------------------|---------|---------|---------|---------|---------|---------|---------|
| 1 | Θ_1 | 0.00033 | 0.00060 | 0.00090 | 0.00133 | 0.00247 | 0.00123 | 0.00127 |
| 1 | Θ_2 | 0.00080 | 0.00127 | 0.00197 | 0.00240 | 0.00433 | 0.00223 | 0.00240 |
| 1 | D _{1->2} | 0.00000 | 0.00000 | 0.00350 | 0.05333 | 0.21800 | 0.11550 | 0.15652 |
| 1 | S _{1->2} | 0.27067 | 0.27533 | 0.27583 | 0.27633 | 0.27800 | 0.27250 | 0.26362 |
| 2 | Θ_1 | 0.00000 | 0.00000 | 0.00023 | 0.00053 | 0.00167 | 0.00057 | 0.00080 |
| 2 | Θ_2 | 0.00073 | 0.00113 | 0.00170 | 0.00233 | 0.00333 | 0.00210 | 0.00228 |
| 2 | D _{1->2} | 0.00000 | 0.00000 | 0.00283 | 0.05100 | 0.13033 | 0.09750 | 0.14410 |
| 2 | S _{1->2} | 0.00033 | 0.00133 | 0.00583 | 0.02133 | 0.03967 | 0.25450 | 0.24896 |
| 3 | Θ_1 | 0.00053 | 0.00213 | 0.00310 | 0.00507 | 0.01387 | 0.00510 | 0.00672 |
| 3 | Θ_2 | 0.00200 | 0.00387 | 0.00503 | 0.00813 | 0.02127 | 0.00857 | 0.01013 |
| 3 | D _{1->2} | 0.00000 | 0.00000 | 0.00017 | 0.02000 | 0.14833 | 0.02417 | 0.08011 |
| 3 | S _{1->2} | 0.00000 | 0.00000 | 0.00283 | 0.04833 | 0.09767 | 0.08550 | 0.15474 |
| All | Θ_1 | 0.00000 | 0.00093 | 0.00163 | 0.00213 | 0.00287 | 0.00163 | 0.00158 |
| All | Θ_2 | 0.00100 | 0.00227 | 0.00297 | 0.00360 | 0.00487 | 0.00303 | 0.00299 |
| All | D _{1->2} | 0.00000 | 0.00000 | 0.00017 | 0.00500 | 0.03867 | 0.00517 | 0.01034 |
| All | S _{1->2} | 0.00000 | 0.00167 | 0.01150 | 0.03067 | 0.17500 | 0.02950 | 0.06076 |

Citation suggestions:

Beerli P., 2006. Comparison of Bayesian and maximum-likelihood inference of population genetic parameters. Bioinformatics 22:341-345

Beerli P., 2009. How to use MIGRATE or why are Markov chain Monte Carlo programs difficult to use? In Population Genetics for Animal Conservation, G. Bertorelle, M. W. Bruford, H. C. Hauffe, A. Rizzoli, and C. Vernesi, eds., vol. 17 of Conservation Biology, Cambridge University Press, Cambridge UK, pp. 42-79.

Bayesian Analysis: Posterior distribution over all loci



Log-Probability of the data given the model (marginal likelihood)

Use this value for Bayes factor calculations:

BF = Exp[In(Prob(D | thisModel) - In(Prob(D | otherModel) or as LBF = 2 (In(Prob(D | thisModel) - In(Prob(D | otherModel)) shows the support for thisModel]

| Locus | Raw thermodynamic score(1a) | Bezier approximation score(1b) | Harmonic mean(2) |
|-------|-----------------------------|--------------------------------|------------------|
| 1 | -1870.15 | -1742.75 | -1727.87 |
| 2 | -1311.18 | -1228.13 | -1217.83 |
| 3 | -715.55 | -671.55 | -664.50 |
| All | -3906.84 | -3652.38 | -3620.16 |

(1a, 1b and 2) are approximations to the marginal likelihood, make sure that the program run long enough! (1a, 1b) and (2) should give similar results, in principle.

But (2) is overestimating the likelihood, it is presented for historical reasons and should not be used (1a, 1b) needs heating with chains that span a temperature range of 1.0 to at least 100,000.

(1b) is using a Bezier-curve to get better approximations for runs with low number of heated chains [Scaling factor = -9.962152]

Citation suggestions:

Beerli P. and M. Palczewski, 2010. Unified framework to evaluate panmixia and migration direction among multiple sampling locations, Genetics, 185: 313-326.

Acceptance ratios for all parameters and the genealogies

| Parameter | Accepted changes | Ratio |
|---------------------------|------------------|---------|
| Θ_1 | 4577163/7500105 | 0.61028 |
| Θ_2 | 4684194/7497651 | 0.62475 |
| $\Delta_{1\rightarrow 2}$ | 4772707/7500000 | 0.63636 |
| $\sigma_{1\rightarrow 2}$ | 5081595/7499474 | 0.67759 |
| Genealogies | 8483724/30002770 | 0.28276 |

MCMC-Autocorrelation and Effective MCMC Sample Size

| tive Sampe Size |
|-----------------|
| 69.60 |
| 79.36 |
| 33.60 |
| 47.81 |
| 47.81 |
| - |

Potential Problems

This section reports potential problems with your run, but such reporting is often not very accurate. Whith many parameters in a multilocus analysi s, it is very common that some parameters for some loci will not be very informative, triggering suggestions (for example to increase the prior ran ge) that are not sensible. This

| gged, inspect the tables carefully and judge wether an action is required. For example, if you run a Bayesian inference with sequence data, for mac roscopic species there is rarely the need to increase the prior for Theta beyond 0.1; but if you use microsatellites it is rather common that your prior distribution for Theta should have a range from 0.0 to 100 or more. With many populations (>3) it is also very common that some migration rou tes are estimated poorly because the data contains little or no information for that route. Increasing the range will not help in such situations, reducing number of parameters may help in such situations. |
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| No warning was recorded during the run |
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