AUTO

POPULATION SIZE, MIGRATION, DIVERGENCE, ASSIGNMENT, HISTORY

Bayesian inference using the structured coalescent

Migrate-n version 5.0.0a [May-20-2017]

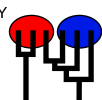
Using Intel AVX (Advanced Vector Extensions)

Compiled for PARALLEL computer architectures

One master and 100 compute nodes are available.

Program started at Sat Aug 12 15:05:04 2017

Program finished at Sat Aug 12 16:07:46 2017 [Runtime:0000:01:02:42]



Options

Datatype: DNA sequence data

Inheritance scalers in use for Thetas:

All loci use an inheritance scaler of 1.0

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

Random number seed: (with internal timer) 1565496655

Start parameters:

Theta values were generated Using a percent value of the prior

M values were generated Using a percent value of 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

1

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

Population

1 Romanshorn 0 *

Order of parameters:

1 Θ_1 <displayed>

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

Analysis strategy:

Bayesian inference

-Population size estimation: Exponential Distribution

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

Parameter Prior Minimum MeanMaximum Delta Bins UpdateFreq
1 Theta -11 Uniform 0.000000 0.050 0.100 0.010 1500 0.20000

[-1 -1 means priors were set globally]

Markov chain settings: Long chain

Number of chains1Recorded steps [a]50000Increment (record every x step [b]200Number of concurrent chains (replicates) [c]2

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

Number of discard trees per chain (burn-in) 10000

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: infile.0.4

Haplotyping is turned on:

Output file: outfile_0.4_0.5

Posterior distribution raw histogram file: bayesfile

Raw data from the MCMC run: bayesallfile_0.4_0.5

Print data:

Print genealogies [only some for some data type]:

Data summary

Data file: infile.0.4
Datatype: Sequence data
Number of loci: 100

Mutationmodel:	
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Mutation	model:			
Locus Sublocus Mut		Mutationmodel	Mutationmodel parameters	
1	1	Jukes-Cantor	[Basefreq: =0.25]	
2	1	Jukes-Cantor	[Basefreq: =0.25]	
3	1	Jukes-Cantor	[Basefreq: =0.25]	
4	1	Jukes-Cantor	[Basefreq: =0.25]	
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80	1	Jukes-Cantor	[Basefreq: =0.25]	
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99	1	Jukes-Cantor	[Basefreq: =0.25]	
100	1	Jukes-Cantor	[Basefreq: =0.25]	
Sites per	locus			
Locus		Sites		
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Locus	Sites
1	10000
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8	10000
9	10000
10	10000
11	10000
12	10000
13	10000
14	10000
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	e variation and probab				
Locus S	Sublocus Region type	Rate of change	Probability	Patch size	
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	
4	1 1	1.000	1.000	1.000	
5	1 1	1.000	1.000	1.000	
6	1 1	1.000	1.000	1.000	

7	1	1	1.000	1.000	1.000	
8	1	1	1.000	1.000	1.000	
9	1	1	1.000	1.000	1.000	
10	1	1	1.000	1.000	1.000	
11	1	1	1.000	1.000	1.000	
12	1	1	1.000	1.000	1.000	
13	1	1	1.000	1.000	1.000	
14	1	1	1.000	1.000	1.000	
15	1	1	1.000	1.000	1.000	
16	1	1	1.000	1.000	1.000	
17	1	1	1.000	1.000	1.000	
18	1	1	1.000	1.000	1.000	
19	1	1	1.000	1.000	1.000	
20	1	1	1.000	1.000	1.000	
21	1	1	1.000	1.000	1.000	
22	1	1	1.000	1.000	1.000	
23	1	1	1.000	1.000	1.000	
24	1	1	1.000	1.000	1.000	
25	1	1	1.000	1.000	1.000	
26	1	1	1.000	1.000	1.000	
27	1	1	1.000	1.000	1.000	
28	1	1	1.000	1.000	1.000	
29	1	1	1.000	1.000	1.000	
30	1	1	1.000	1.000	1.000	
31	1	1	1.000	1.000	1.000	
32	1	1	1.000	1.000	1.000	
33	1	1	1.000	1.000	1.000	
34	1	1	1.000	1.000	1.000	
35	1	1	1.000	1.000	1.000	
36	1	1	1.000	1.000	1.000	
37	1	1	1.000	1.000	1.000	
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41	1	1	1.000	1.000	1.000	
42	1	1	1.000	1.000	1.000	
43	1	1	1.000	1.000	1.000	
44	1	1	1.000	1.000	1.000	
45	1	1	1.000	1.000	1.000	
46	1	1	1.000	1.000	1.000	
47	1	1	1.000	1.000	1.000	
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50	1	1	1.000	1.000	1.000	
51	1	1	1.000	1.000	1.000	

52	1	1	1.000	1.000	1.000	
53	1	1	1.000	1.000	1.000	
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57	1	1	1.000	1.000	1.000	
58	1	1	1.000	1.000	1.000	
59	1	1	1.000	1.000	1.000	
60	1	1	1.000	1.000	1.000	
61	1	1	1.000	1.000	1.000	
62	1	1	1.000	1.000	1.000	
63	1	1	1.000	1.000	1.000	
64	1	1	1.000	1.000	1.000	
65	1	1	1.000	1.000	1.000	
66	1	1	1.000	1.000	1.000	
67	1	1	1.000	1.000	1.000	
68	1	1	1.000	1.000	1.000	
69	1	1	1.000	1.000	1.000	
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71	1	1	1.000	1.000	1.000	
72	1	1	1.000	1.000	1.000	
73	1	1	1.000	1.000	1.000	
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83	1	1	1.000	1.000	1.000	
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87	1	1	1.000	1.000	1.000	
88	1	1	1.000	1.000	1.000	
89	1	1	1.000	1.000	1.000	
90	1	1	1.000	1.000	1.000	
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92	1	1	1.000	1.000	1.000	
93	1	1	1.000	1.000	1.000	
94	1	1	1.000	1.000	1.000	
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97	1	1	1.000	1.000	1.000	
98	1	1	1.000	1.000	1.000	
99	1	1	1.000	1.000	1.000	
100	1	1	1.000	1.000	1.000	
Population		·		11000	Locus	Gene copies
	nshorn_0				1	10
- Tromai	.0.1.0111_0				2	10
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Total of all populations	1	10	
	2	10	
	3	10	
	4	10	
	5	10	
	6	10	
	7	10	
	8	10	
	9	10	
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	94	10
	95	10
	96	10
	97	10
	98	10
	99	10
1	100	10

Bayesian Analysis: Posterior distribution table

Locus	Parameter	2.5%	25.0%	Mode	75.0%	97.5%	Median	Mean
1	Θ_1	0.01593	0.02827	0.03503	0.04280	0.05013	0.03383	0.04659
2	Θ_1	0.02047	0.03607	0.04523	0.04847	0.05080	0.03777	0.05512
3	Θ_1	0.01647	0.02713	0.03297	0.04260	0.05000	0.03363	0.04606
4	Θ_1	0.01647	0.02720	0.03710	0.04247	0.05007	0.03370	0.04610
5	Θ_1	0.01640	0.02753	0.03430	0.04240	0.05007	0.03370	0.04592
6	Θ_1	0.01580	0.02740	0.03343	0.04220	0.05027	0.03377	0.04611
7	Θ_1	0.01780	0.03007	0.03823	0.04693	0.05033	0.03523	0.04929
8	Θ_1	0.02413	0.04107	0.04763	0.04907	0.05127	0.04123	0.06966
9	Θ_1	0.02307	0.03907	0.04750	0.04907	0.05113	0.04030	0.06402
10	Θ_1	0.01647	0.02647	0.03317	0.04413	0.05000	0.03377	0.04616
11	Θ_1	0.01653	0.02747	0.03597	0.04273	0.05000	0.03363	0.04577
12	Θ_1	0.02187	0.03780	0.04750	0.04893	0.05100	0.03930	0.06210
13	Θ_1	0.01820	0.03213	0.03977	0.04700	0.05047	0.03570	0.05017
14	Θ_1	0.01847	0.02913	0.03897	0.04840	0.05047	0.03583	0.05062
15	Θ_1	0.01653	0.02827	0.03210	0.04133	0.05000	0.03370	0.04605
16	Θ_1	0.01640	0.02760	0.03383	0.04240	0.05007	0.03377	0.04637
17	Θ_1	0.01647	0.02867	0.03270	0.04113	0.05007	0.03377	0.04619
18	Θ_1	0.01793	0.03260	0.03870	0.04707	0.05040	0.03537	0.04970

19	Θ_1	0.02100	0.03680	0.04603	0.04867	0.05087	0.03810	0.05601
20	Θ_1	0.01640	0.02773	0.03343	0.04207	0.05007	0.03370	0.04612
21	Θ_1	0.01633	0.02620	0.03243	0.04400	0.05000	0.03357	0.04603
22	Θ_1	0.02020	0.03713	0.04750	0.04833	0.05073	0.03750	0.05480
23	Θ_1	0.01647	0.02800	0.03463	0.04240	0.05013	0.03377	0.04621
24	Θ_1	0.01640	0.02813	0.03550	0.04360	0.05007	0.03383	0.04612
25	Θ_1	0.01647	0.02787	0.03690	0.04427	0.05007	0.03383	0.04642
26	Θ_1	0.01580	0.02800	0.03577	0.04307	0.05027	0.03383	0.04620
27	Θ_1	0.01653	0.02833	0.03557	0.04280	0.05007	0.03383	0.04599
28	Θ_1	0.01647	0.02793	0.03443	0.04260	0.05007	0.03377	0.04610
29	Θ_1	0.01627	0.02687	0.03570	0.04173	0.05007	0.03363	0.04616
30	Θ_1	0.01747	0.03127	0.04150	0.04580	0.05047	0.03530	0.04951
31	Θ_1	0.01580	0.02813	0.03383	0.04260	0.05027	0.03383	0.04596
32	Θ_1	0.01600	0.02813	0.03243	0.04320	0.05013	0.03370	0.04592
33	Θ_1	0.01647	0.02760	0.03490	0.04340	0.05007	0.03377	0.04615
34	Θ_1	0.01647	0.02807	0.03650	0.04227	0.05007	0.03377	0.04610
35	Θ_1	0.01640	0.02760	0.03443	0.04300	0.05013	0.03377	0.04598
36	Θ_1	0.01933	0.03507	0.04543	0.04813	0.05053	0.03670	0.05285
37	Θ_1	0.01960	0.03587	0.04443	0.04820	0.05067	0.03690	0.05314
38	Θ_1	0.01600	0.02753	0.03437	0.04447	0.05020	0.03383	0.04593
39	Θ_1	0.01647	0.02733	0.03603	0.04160	0.05000	0.03363	0.04599
40	Θ_1	0.01640	0.02720	0.03417	0.04400	0.05007	0.03370	0.04596
41	Θ_1	0.01600	0.02753	0.03437	0.04227	0.05020	0.03377	0.04612

_ocus	Parameter	2.5%	25.0%	Mode	75.0%	97.5%	Median	Mean
42	Θ_1	0.01993	0.03627	0.04143	0.04793	0.05067	0.03710	0.05314
43	Θ_1	0.01640	0.02180	0.03323	0.04867	0.05007	0.03370	0.04592
44	Θ_1	0.01647	0.02807	0.03377	0.04180	0.05000	0.03377	0.04604
45	Θ_1	0.01513	0.02773	0.03443	0.04293	0.05040	0.03377	0.04602
46	Θ_1	0.01633	0.02633	0.03477	0.04453	0.05007	0.03377	0.04601
47	Θ_1	0.01653	0.02787	0.03423	0.04280	0.05000	0.03377	0.04623
48	Θ_1	0.01660	0.02867	0.03330	0.04100	0.05007	0.03383	0.04601
49	Θ_1	0.01627	0.02740	0.03577	0.04287	0.05007	0.03377	0.04599
50	Θ_1	0.01640	0.02587	0.03610	0.04587	0.04993	0.03370	0.04602
51	Θ_1	0.01920	0.03647	0.04430	0.04727	0.05053	0.03663	0.05278
52	Θ_1	0.01640	0.02687	0.03337	0.04387	0.05007	0.03377	0.04599
53	Θ_1	0.01660	0.02640	0.03470	0.04400	0.05007	0.03377	0.04612
54	Θ_1	0.01847	0.03440	0.04217	0.04787	0.05067	0.03643	0.05268
55	Θ_1	0.01660	0.02713	0.03350	0.04240	0.05000	0.03370	0.04598
56	Θ_1	0.02093	0.03633	0.04743	0.04867	0.05080	0.03797	0.05599
57	Θ_1	0.01647	0.02893	0.03163	0.04167	0.05007	0.03377	0.04621
58	Θ_1	0.02247	0.03893	0.04750	0.04893	0.05107	0.03963	0.06250
59	Θ_1	0.02160	0.03780	0.04750	0.04907	0.05107	0.03937	0.06052
60	Θ_1	0.01627	0.02753	0.03350	0.04207	0.05013	0.03357	0.04613
61	Θ_1	0.01807	0.03393	0.04297	0.04767	0.05053	0.03597	0.05104

62	Θ_1	0.01647	0.02727	0.03397	0.04247	0.05007	0.03377	0.04613
63	Θ_1	0.01647	0.02800	0.03183	0.04247	0.05007	0.03377	0.04623
64	Θ_1	0.01820	0.03313	0.03730	0.04713	0.05053	0.03590	0.05079
65	Θ_1	0.01620	0.02687	0.03697	0.04307	0.05013	0.03383	0.04622
66	Θ_1	0.02533	0.04013	0.04763	0.04947	0.05127	0.04150	0.06981
67	Θ_1	0.01847	0.03393	0.04010	0.04793	0.05060	0.03617	0.05102
68	Θ_1	0.01860	0.03353	0.03997	0.04620	0.05047	0.03583	0.05013
69	Θ_1	0.01640	0.02733	0.03330	0.04280	0.04993	0.03363	0.04589
70	Θ_1	0.01647	0.02740	0.03463	0.04387	0.05013	0.03383	0.04629
71	Θ_1	0.01647	0.02760	0.03357	0.04300	0.05007	0.03383	0.04613
72	Θ_1	0.01753	0.03113	0.03870	0.04520	0.05040	0.03523	0.04932
73	Θ_1	0.01993	0.03580	0.04703	0.04840	0.05087	0.03750	0.05510
74	Θ_1	0.01613	0.02740	0.03317	0.04280	0.05020	0.03383	0.04612
75	Θ_1	0.01900	0.02307	0.04203	0.04987	0.05060	0.03643	0.05160
76	Θ_1	0.01640	0.02627	0.03110	0.04467	0.05007	0.03370	0.04595
77	Θ_1	0.01953	0.03380	0.04197	0.04820	0.05060	0.03670	0.05218
78	Θ_1	0.01820	0.03293	0.03663	0.04513	0.05033	0.03543	0.04970
79	Θ_1	0.02007	0.03700	0.04750	0.04847	0.05080	0.03757	0.05617
80	Θ_1	0.01640	0.02540	0.03397	0.04607	0.05007	0.03383	0.04631
81	Θ_1	0.02160	0.03753	0.04663	0.04867	0.05087	0.03870	0.05807
82	Θ_1	0.01600	0.02587	0.03417	0.04480	0.05013	0.03377	0.04621
83	Θ_1	0.01860	0.03340	0.04037	0.04693	0.05047	0.03583	0.05043
84	Θ_1	0.01647	0.02553	0.03243	0.04573	0.05007	0.03370	0.04616

Locus	Parameter	2.5%	25.0%	Mode	75.0%	97.5%	Median	Mean
85	Θ_1	0.02300	0.03847	0.04750	0.04907	0.05107	0.03997	0.06276
86	Θ_1	0.01607	0.02727	0.03237	0.04327	0.05020	0.03383	0.04605
87	Θ_1	0.01633	0.02573	0.03277	0.04460	0.05007	0.03370	0.04616
88	Θ_1	0.02200	0.03793	0.04757	0.04913	0.05107	0.03943	0.06110
89	Θ_1	0.01640	0.02800	0.03730	0.04153	0.05007	0.03377	0.04604
90	Θ_1	0.01640	0.02867	0.03823	0.04107	0.05007	0.03377	0.04606
91	Θ_1	0.01393	0.02707	0.03603	0.04213	0.05073	0.03377	0.04637
92	Θ_1	0.02100	0.03680	0.04750	0.04867	0.05093	0.03843	0.05678
93	Θ_1	0.01380	0.02700	0.03343	0.04273	0.05080	0.03377	0.04632
94	Θ_1	0.02347	0.03920	0.04763	0.04933	0.05120	0.04057	0.06768
95	Θ_1	0.02133	0.03760	0.04483	0.04847	0.05080	0.03830	0.05646
96	Θ_1	0.01660	0.02853	0.03250	0.04193	0.05007	0.03383	0.04640
97	Θ_1	0.01760	0.03087	0.03683	0.04747	0.05053	0.03550	0.04933
98	Θ_1	0.02273	0.03847	0.04750	0.04913	0.05113	0.03997	0.06223
99	Θ_1	0.02253	0.03467	0.04757	0.04953	0.05100	0.03963	0.06202
100	Θ_1	0.01593	0.02767	0.03523	0.04173	0.05020	0.03383	0.04643
All	Θ_1	0.03513	0.03740	0.03890	0.04053	0.04300	0.03910	0.03912

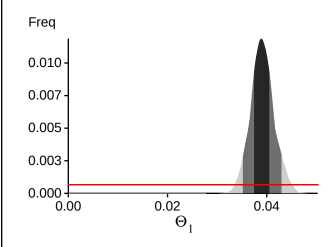
Citation suggestions:

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

Beerli P., 2007. Estimation of the population scaled mutation rate from microsatellite data, Genetics, 177:1967-1968.

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: $BF = Exp[\ ln(Prob(D \mid thisModel) - ln(\ Prob(\ D \mid otherModel)) \\ or \ as \ LBF = 2 \ (ln(Prob(D \mid thisModel) - ln(\ Prob(\ D \mid otherModel))) \\ shows the \ support for \ thisModel]$

_ocus	TI(1a)	BTI(1b)	SS(2)	HS(3)
1	-13871.66	-13727.56	-13770.82	-13864.52
2	-14049.76	-13853.24	-13895.19	-13980.85
3	-13872.41	-13728.25	-13772.07	-13865.14
4	-13873.74	-13729.47	-13771.53	-13867.46
5	-13872.43	-13728.21	-13770.53	-13864.90
6	-13873.54	-13729.25	-13773.09	-13865.90
7	-13885.38	-13741.12	-13787.31	-13877.52
8	-14142.72	-13974.05	-14023.19	-14106.10
9	-13963.28	-13809.56	-13860.99	-13945.15
10	-13872.56	-13728.65	-13772.37	-13865.18
11	-13874.11	-13730.07	-13773.47	-13866.63
12	-26734.64	-19687.66	-18483.34	-18564.46
13	-13906.30	-13752.35	-13797.63	-13886.89
14	-13907.44	-13753.89	-13798.95	-13888.70
15	-13870.91	-13726.91	-13770.05	-13863.44
16	-13872.06	-13727.95	-13771.17	-13864.37
17	-13871.84	-13727.86	-13771.92	-13864.68
18	-13889.29	-13743.51	-13788.13	-13879.68
19	-14135.88	-13911.05	-13948.13	-14034.34
20	-13874.02	-13729.68	-13773.42	-13866.31
21	-13874.55	-13730.25	-13773.50	-13866.95
22	-13916.85	-13765.89	-13812.64	-13901.05
23	-13873.68	-13729.65	-13773.15	-13866.29
24	-13872.87	-13728.72	-13771.82	-13864.99
25	-13874.21	-13730.01	-13774.07	-13866.67
26	-13870.90	-13726.67	-13768.53	-13863.55
27	-13874.19	-13730.10	-13774.36	-13866.56
28	-13870.62	-13726.62	-13770.73	-13863.24
29	-13874.32	-13730.17	-13773.94	-13866.75

Migrate 5.0.0a: (http://popgen.sc.fsu.edu) [program run on 15:05:04]

30	-13890.29	-13743.58	-13787.45	-13879.63
31	-13874.14	-13729.97	-13773.14	-13866.38
32	-13874.04	-13729.82	-13773.31	-13866.37
33	-13871.97	-13728.03	-13771.10	-13864.97
34	-13873.79	-13729.44	-13772.90	-13866.05
35	-13872.97	-13728.74	-13772.67	-13865.34
36	-13904.17	-13755.78	-13801.43	-13892.16
37	-13905.10	-13756.43	-13802.79	-13892.18
38	-13872.95	-13728.77	-13771.73	-13865.03
39	-13873.62	-13729.46	-13772.27	-13866.12
40	-13873.88	-13729.60	-13773.54	-13865.92
41	-13873.53	-13729.55	-13773.50	-13865.87
42	-13941.98	-13793.80	-13842.56	-13930.25
43	-13873.02	-13728.75	-13771.96	-13865.47
44	-13873.39	-13729.13	-13772.37	-13865.79
45	-13873.21	-13729.05	-13772.97	-13865.59
46	-13873.72	-13729.52	-13772.78	-13865.86
47	-13873.89	-13729.53	-13770.92	-13866.11
48	-13873.27	-13729.07	-13772.37	-13866.65
49	-13873.77	-13729.55	-13773.43	-13866.04
50	-13873.82	-13729.75	-13772.15	-13867.70
51	-13901.49	-13755.99	-13802.31	-13893.32
52	-13870.25	-13726.13	-13770.29	-13862.58
53	-13874.22	-13729.96	-13774.10	-13867.28
54	-13896.78	-13752.31	-13799.96	-13889.26
55	-13874.18	-13730.10	-13774.07	-13866.59
56	-13930.75	-13774.10	-13820.49	-13910.93
57	-13872.88	-13728.82	-13772.57	-13865.22
58	-20962.39	-19075.73	-18860.29	-18940.60
59	-16063.95	-14939.00	-14818.39	-14900.81
60	-13870.73	-13726.67	-13769.11	-13863.08
61	-13892.26	-13747.83	-13793.49	-13884.30
62	-13874.13	-13729.83	-13772.55	-13866.28
63	-13874.08	-13729.89	-13773.81	-13866.36
64	-13898.17	-13753.46	-13799.37	-13890.28
65	-13873.24	-13729.14	-13772.34	-13865.55
66	-14033.84	-13883.59	-13937.61	-14019.29
67	-13914.85	-13760.44	-13805.24	-13897.35
68	-13905.92	-13752.87	-13797.76	-13888.57
69	-13869.01	-13724.73	-13768.64	-13861.08
70	-13873.69	-13729.47	-13773.08	-13865.77
71	-13872.36	-13728.14	-13771.37	-13864.65
72	-13886.46	-13742.21	-13787.85	-13878.72
73	-13915.39	-13766.66	-13812.62	-13901.63
74	-13872.31	-13728.05	-13771.69	-13864.51

75	-13929.13	-13770.69	-13815.91	-13906.16
76	-13873.87	-13729.78	-13773.83	-13866.37
77	-13965.84	-13792.83	-13836.56	-13924.32
78	-13887.86	-13740.99	-13784.95	-13876.90
79	-13910.77	-13765.80	-13816.17	-13902.58
80	-13871.54	-13727.25	-13770.87	-13863.56
81	-13932.29	-13779.93	-13829.01	-13914.58
82	-13872.74	-13728.59	-13772.43	-13865.59
83	-13902.26	-13751.68	-13796.94	-13887.04
84	-13870.41	-13726.31	-13770.21	-13863.14
85	-14296.26	-14046.03	-14084.23	-14165.95
86	-13873.86	-13729.71	-13772.99	-13866.04
87	-13874.33	-13730.11	-13774.00	-13866.53
88	-13940.51	-13792.36	-13844.17	-13928.10
89	-13874.24	-13729.96	-13773.28	-13867.13
90	-13873.99	-13729.78	-13772.86	-13866.16
91	-13872.69	-13728.40	-13772.35	-13864.91
92	-13935.99	-13781.22	-13830.29	-13916.25
93	-13873.05	-13728.87	-13772.90	-13865.45
94	-14663.18	-14395.38	-14436.03	-14514.71
95	-14129.01	-13926.47	-13969.11	-14053.90
96	-13872.18	-13727.95	-13771.11	-13864.41
97	-13891.00	-13743.29	-13787.22	-13878.96
98	-14103.89	-13913.17	-13961.45	-14043.00
99	-20708.87	-17142.77	-16579.06	-16660.24
100	-13874.09	-13729.96	-13773.32	-13866.64
All	-1419905.05	-1391771.28	-1393924.77	-1403007.55

- (1a) TI: Thermodynamic integration: log(Prob(D|Model)): Good approximation with many temperatures(1b) BTI: Bezier-approximated Thermodynamic integration: when using few temperatures USE THIS!
- (2) SS: Steppingstone Sampling (Xie et al 2011)
 (3) HS: Harmonic mean approximation: Overestimates the marginal likelihood, poor variance [Scaling factor = 48.816989]

Citation suggestions:

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

Palczewski M. and P. Beerli, 2014. Population model comparison using multi-locus datasets. In M.-H. Chen, L. Kuo, and P. O. Lewis, editors, Bayesian Phylogenetics: Methods,

Algorithms, and Applications, pages 187-200. CRC Press, 2014.

Xie W., P. O. Lewis, Y. Fan, L. Kuo, and M.-H. Chen. 2011. Improving marginal likelihood estimation for Bayesian phylogenetic model selection. Systematic Biology, 60(2):150â 160, 2011.

Acceptance ratios for all parameters and the genealogies

Parameter	Accepted changes	Ratio
Θ_1 Genealogies	379373136/400006401 1015695374/1599993599	0.94842 0.63481

MCMC-Autocorrelation and Effective MCMC Sample Size

Parameter	Autocorrelation	Effective Sampe Size
Θ_1 Genealogies	0.70043 0.06691	1761887.02 8768093.52

Average temperatures during the run

4 0.00000

0.00000

3

Adaptive heating often fails, if the average temperatures are very close together try to rerun using static heating! If you want to compare models using marginal likelihoods then you MUST use static heating

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 suggestion tool will improve with time, therefore do not blindly follow its suggestions. If some parameters are fla aged inspect the tables carefully and judge wether an action is required. For example, if you run a Rayesian

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.
No warning was recorded during the run
The Warning was resorted during the run