



Nucleotide

Nucleotide

Meles meles



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Revision date

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[Meles meles](#) microsatellite DNA, locus Mel460, isolate Bap1556, clone 14_B09

181. 358 bp linear DNA

Accession: FR745722.1 GI: 315003445
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[Meles meles](#) microsatellite DNA, locus Mel459, isolate Bap1556, clone 14_B08

182. 290 bp linear DNA

Accession: FR745721.1 GI: 315003444
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[Meles meles](#) microsatellite DNA, locus Mel458, isolate Bap1556, clone 14_B07

183. 337 bp linear DNA

Accession: FR745720.1 GI: 315003443
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[Meles meles](#) microsatellite DNA, locus Mel457, isolate Bap1556, clone 14_B05

184. 572 bp linear DNA

Accession: FR745719.1 GI: 315003442
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[Meles meles](#) microsatellite DNA, locus Mel456, isolate Bap1556, clone 14_B03

185. 444 bp linear DNA

Accession: FR745718.1 GI: 315003441
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[Meles meles](#) microsatellite DNA, locus Mel455, isolate Bap1556, clone 14_B01

186. 666 bp linear DNA

Accession: FR745717.1 GI: 315003440
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Results by taxon

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- [Meles meles](#) (920)
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- [Meles leucurus](#) (45)
- [Meles anakuma](#) (11)
- [Taxidea taxus](#) (7)
- [All other taxa](#) (85)

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[Meles meles](#) microsatellite DNA, locus Mel418, isolate Bap1556, clone 12_Nucleotide

[Meles meles](#) microsatellite DNA, locus Mel468, isolate Bap1556, clone 14_Nucleotide

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Nucleotide

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Meles meles microsatellite DNA, locus Mel450, isolate Bap1556, clone 14_A04

GenBank: FR745712.1

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>gi|315003435|emb|FR745712.1| Meles meles microsatellite DNA, locus Mel450, isolate Bap1556, clone 14_A04

```
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- Meles meles microsatellite DNA, locus Mel450, isolate Bap1556, clone 14_ Nucleotide
- Meles meles microsatellite DNA, locus Mel418, isolate Bap1556, clone 12_ Nucleotide
- Meles meles microsatellite DNA, locus Mel468, isolate Bap1556, clone 14_ Nucleotide
- Meles meles microsatellite DNA, locus Mel476, isolate Bap1556, clone 14_ Nucleotide
- Meles meles microsatellite DNA, locus Mel488, isolate Bap1556, clone 14_ Nucleotide

See more...

		530	540	550	560	570	580	590	600	610	620	630	640	650	660						
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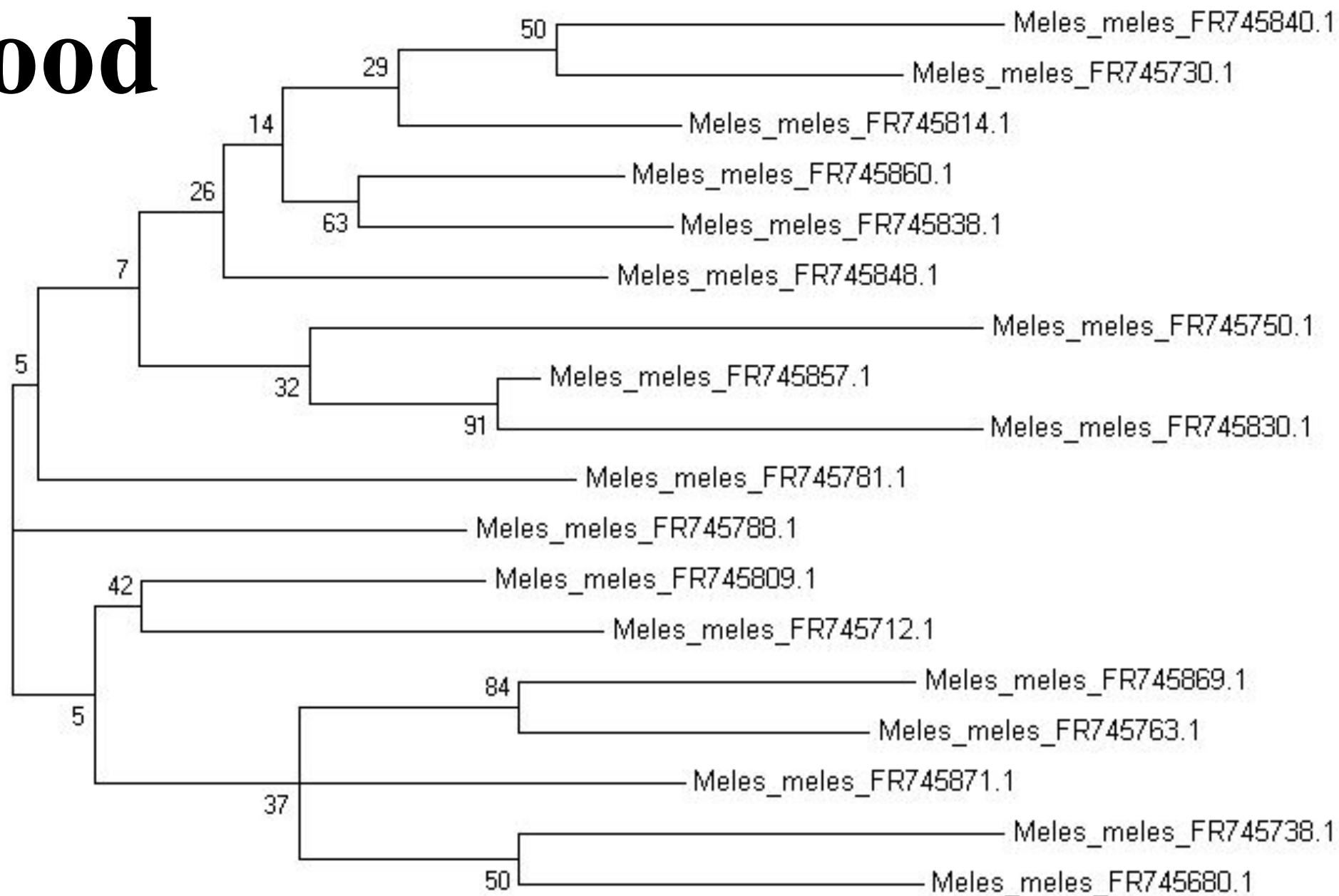
Name MEGA XL CSV UUC Phc C V Pi S 0 2 4 Special None Note

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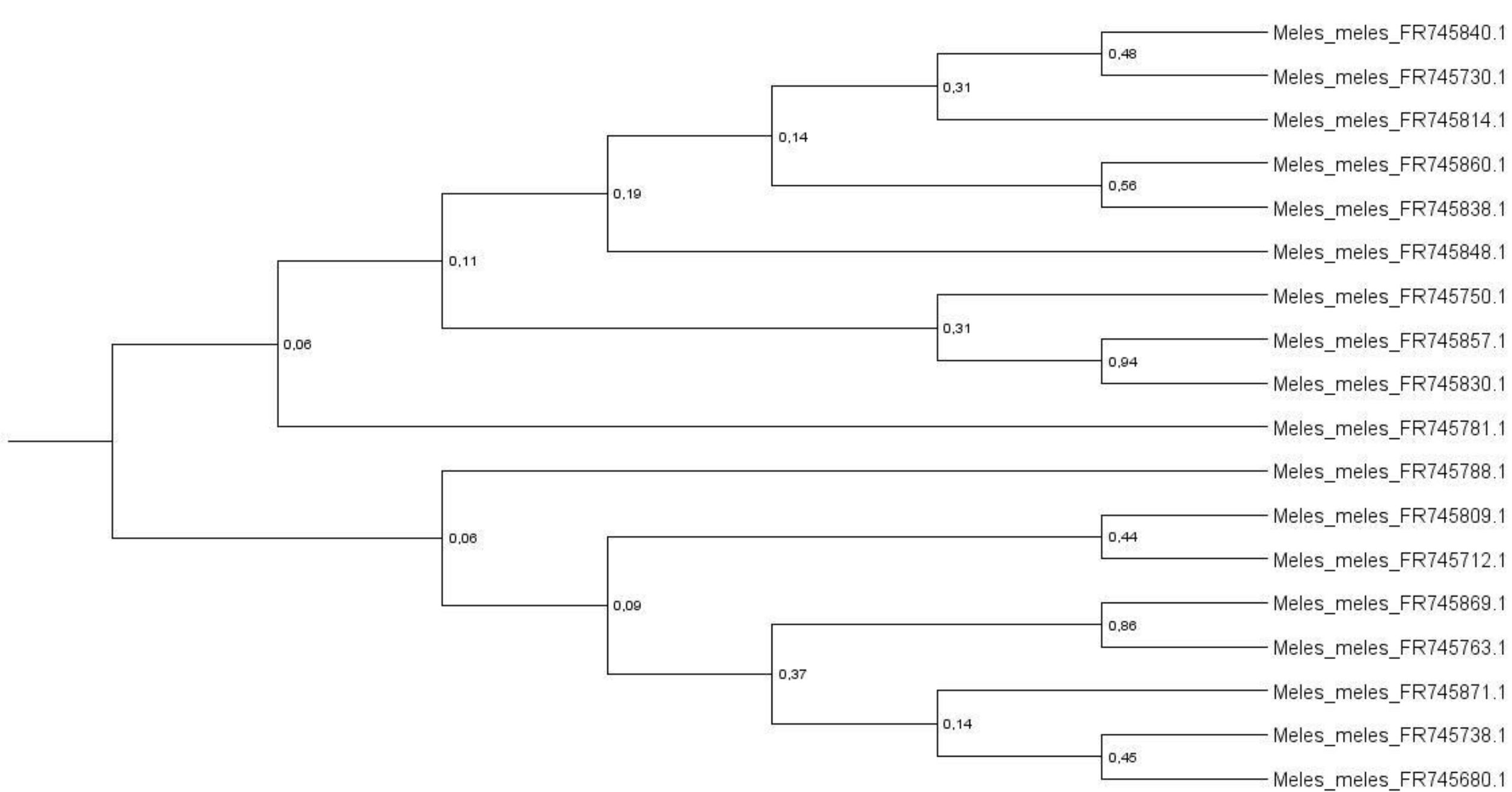
MEGA

MOLECULAR EVOLUTIONARY GENETICS ANALYSIS

Likelihood

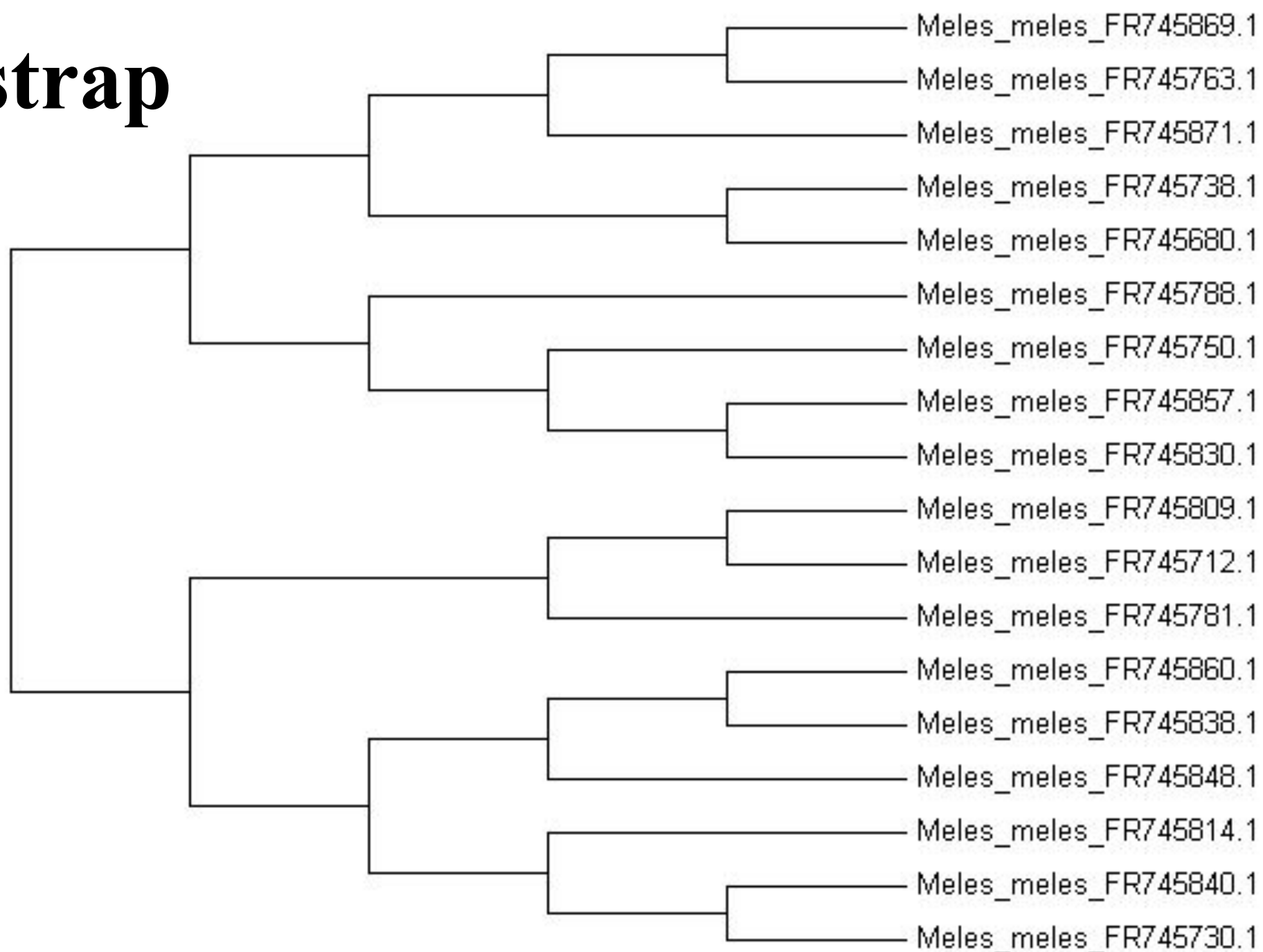


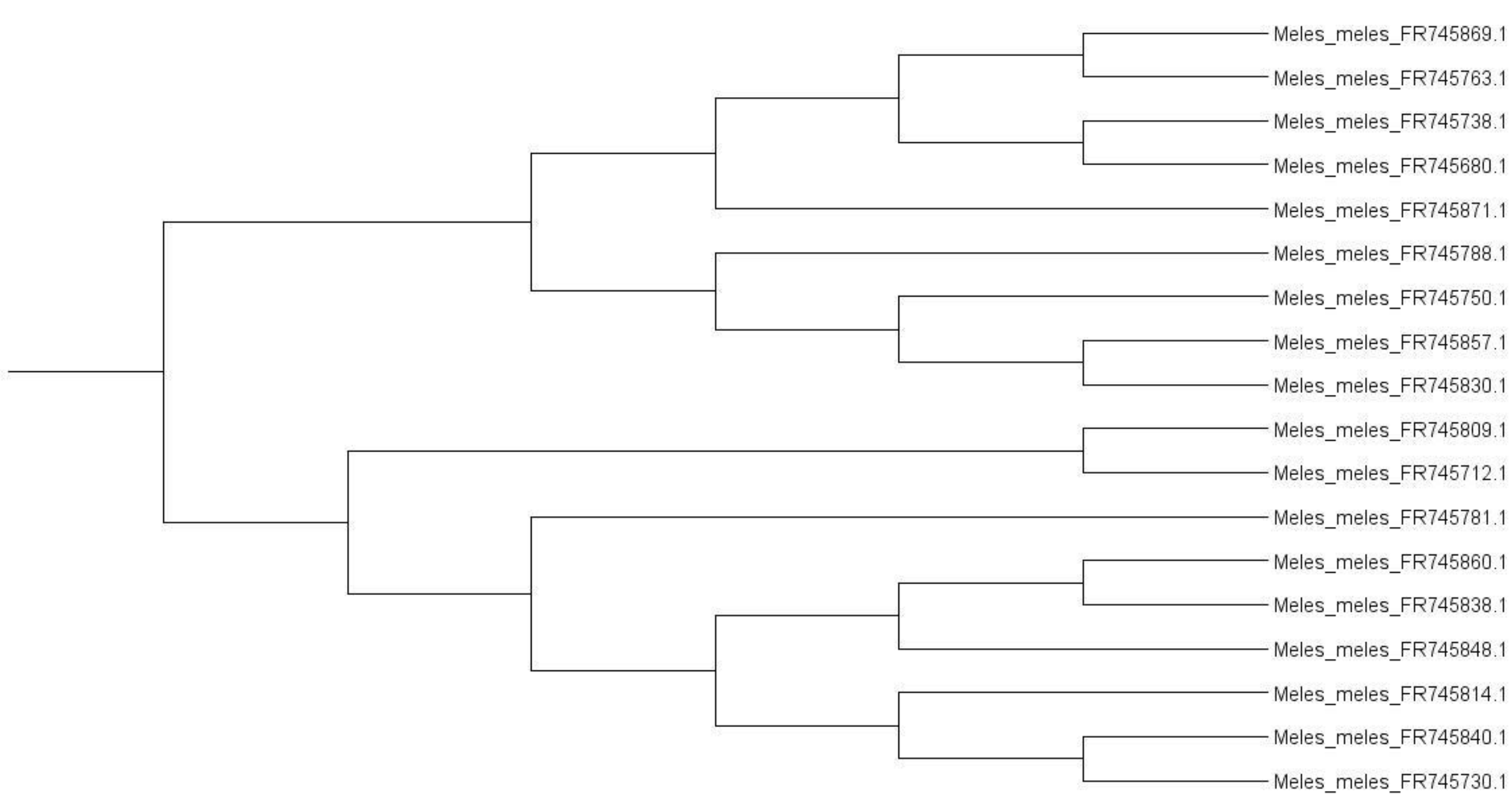
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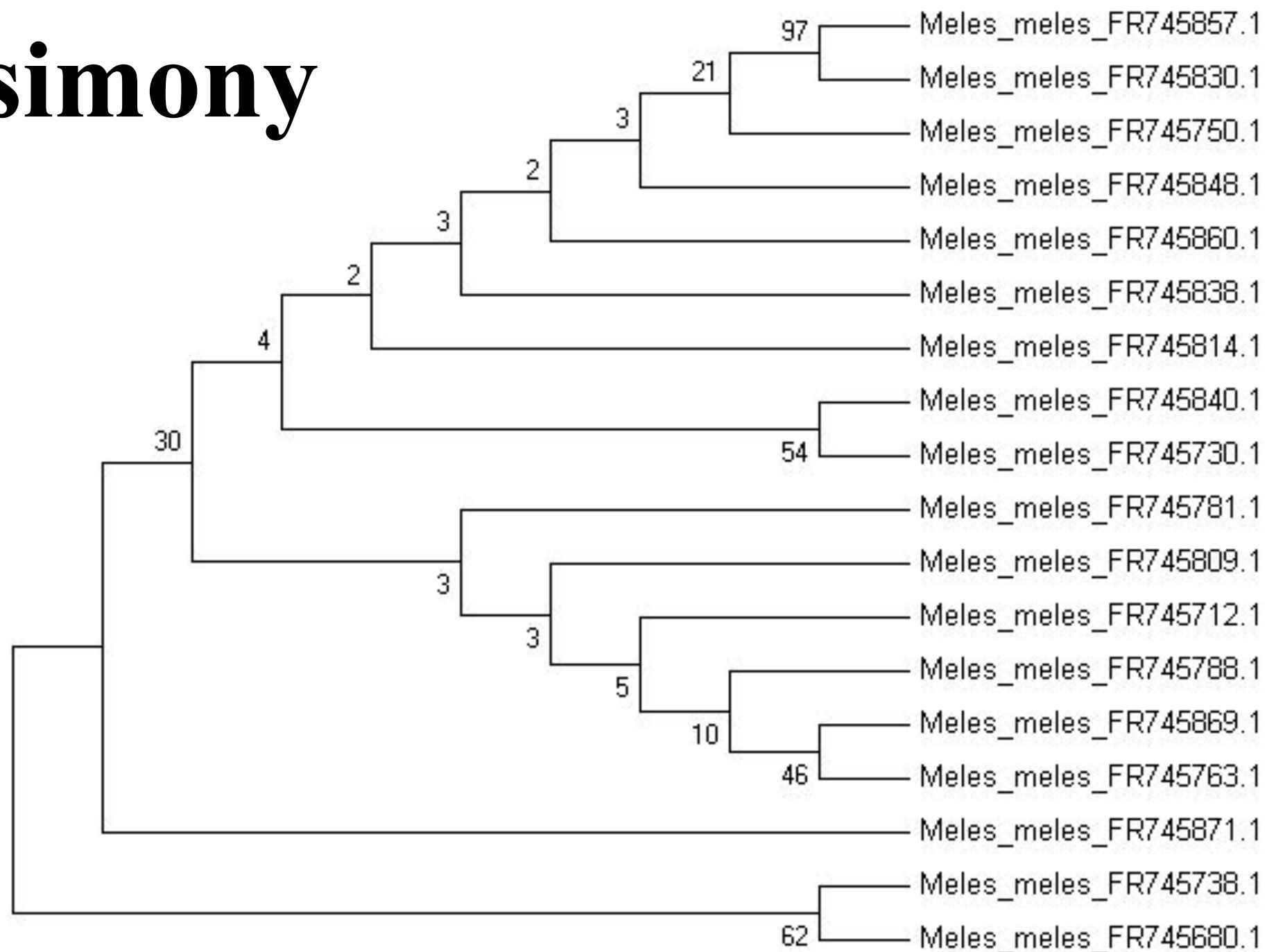
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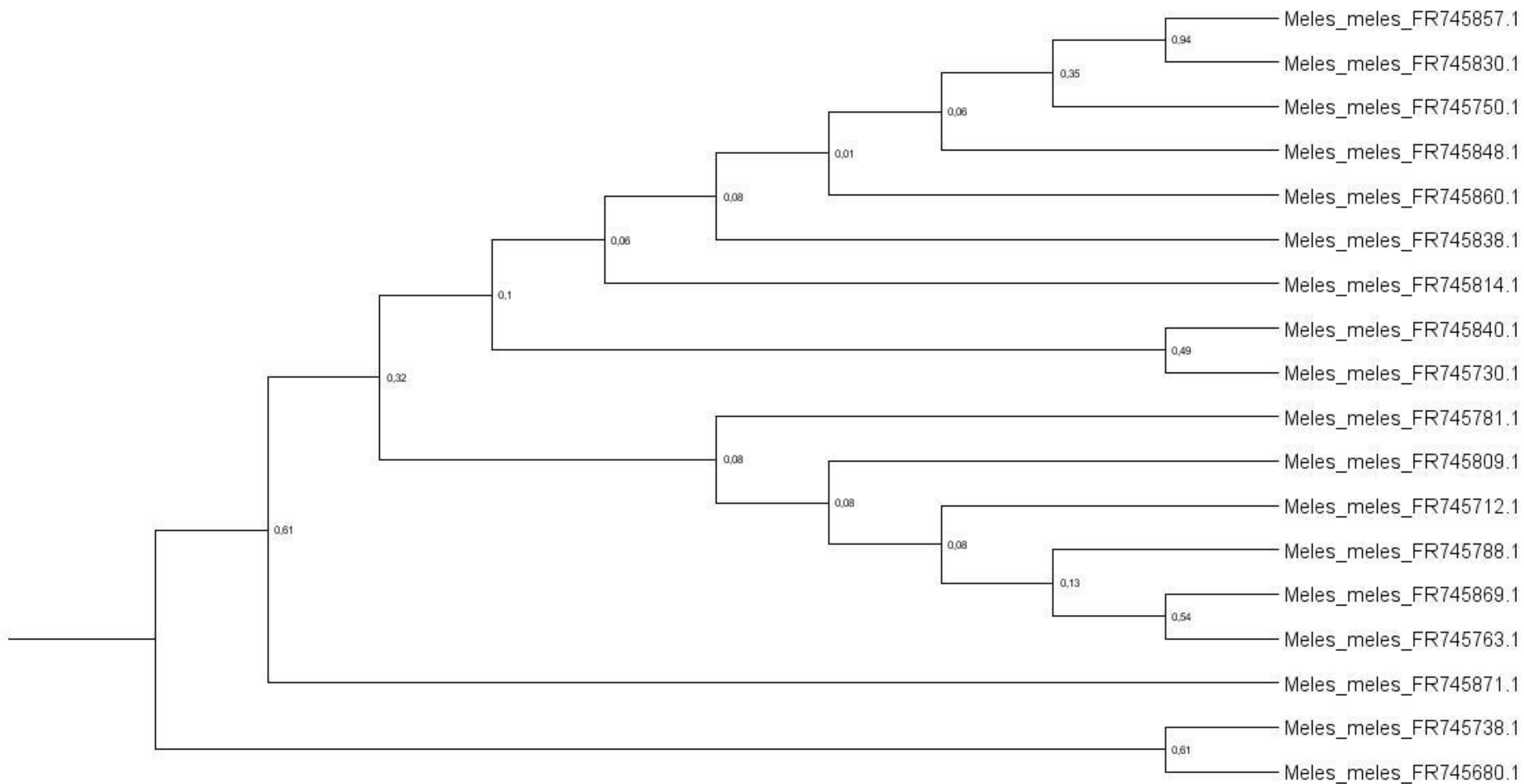
Bootstrap





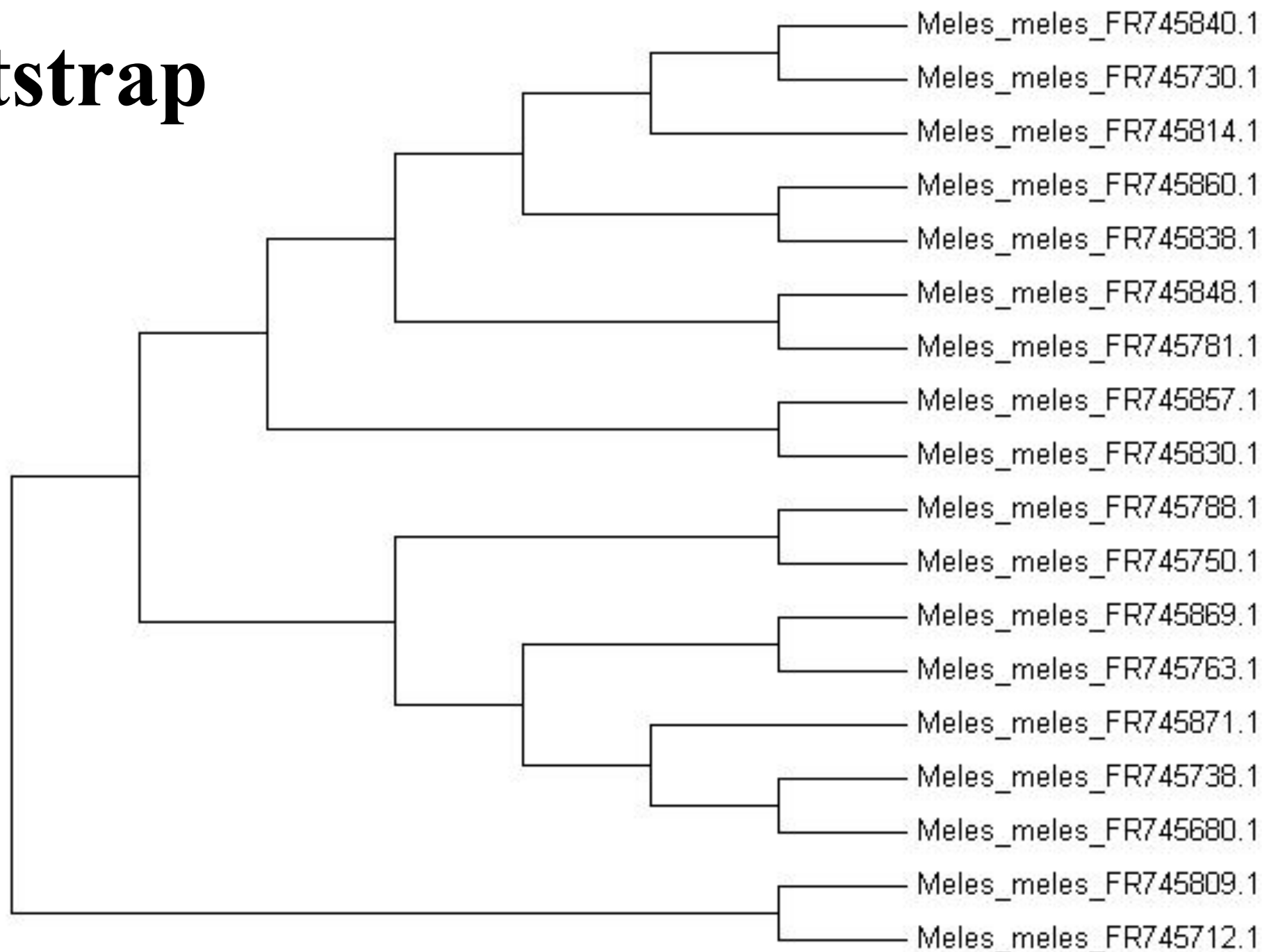
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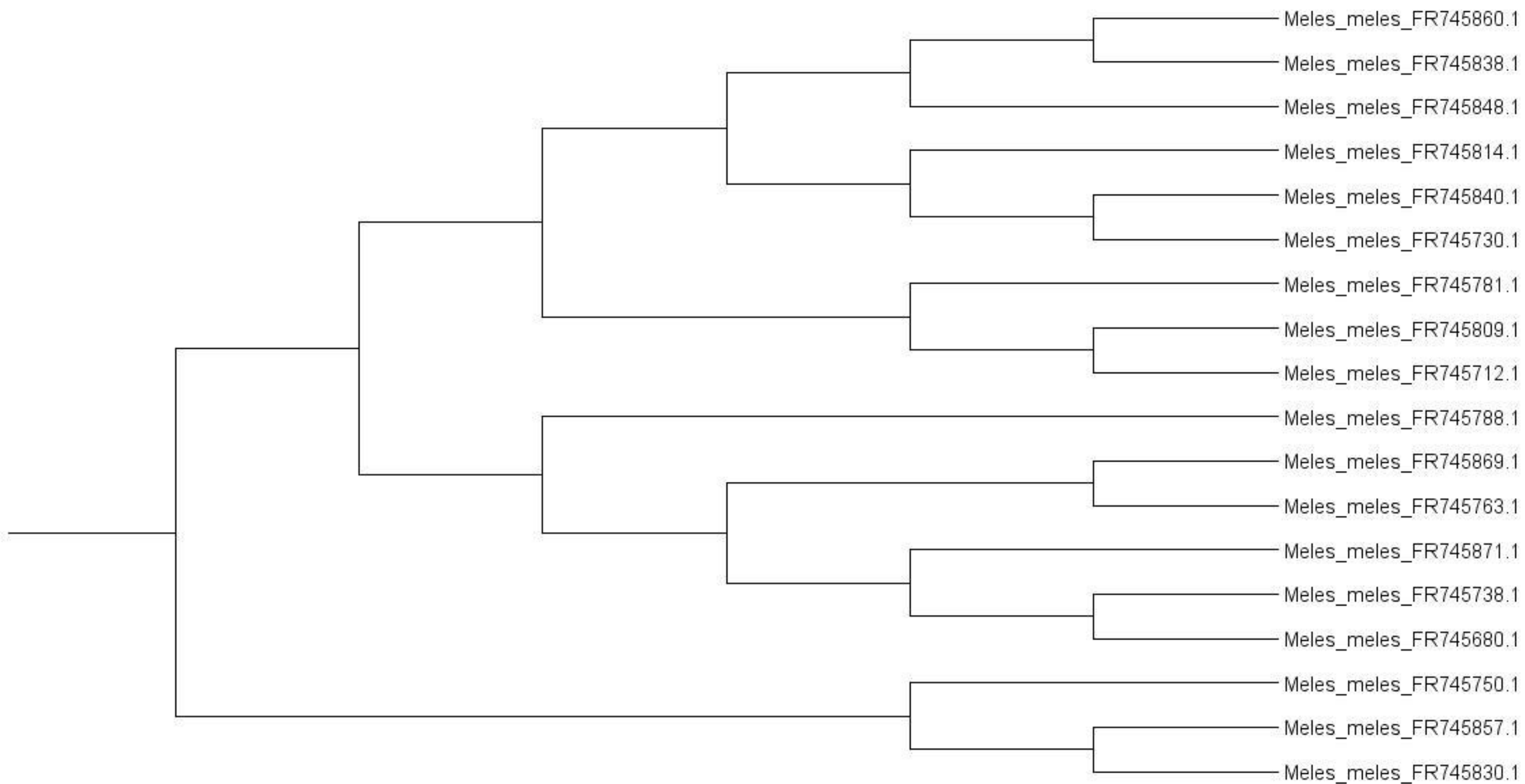




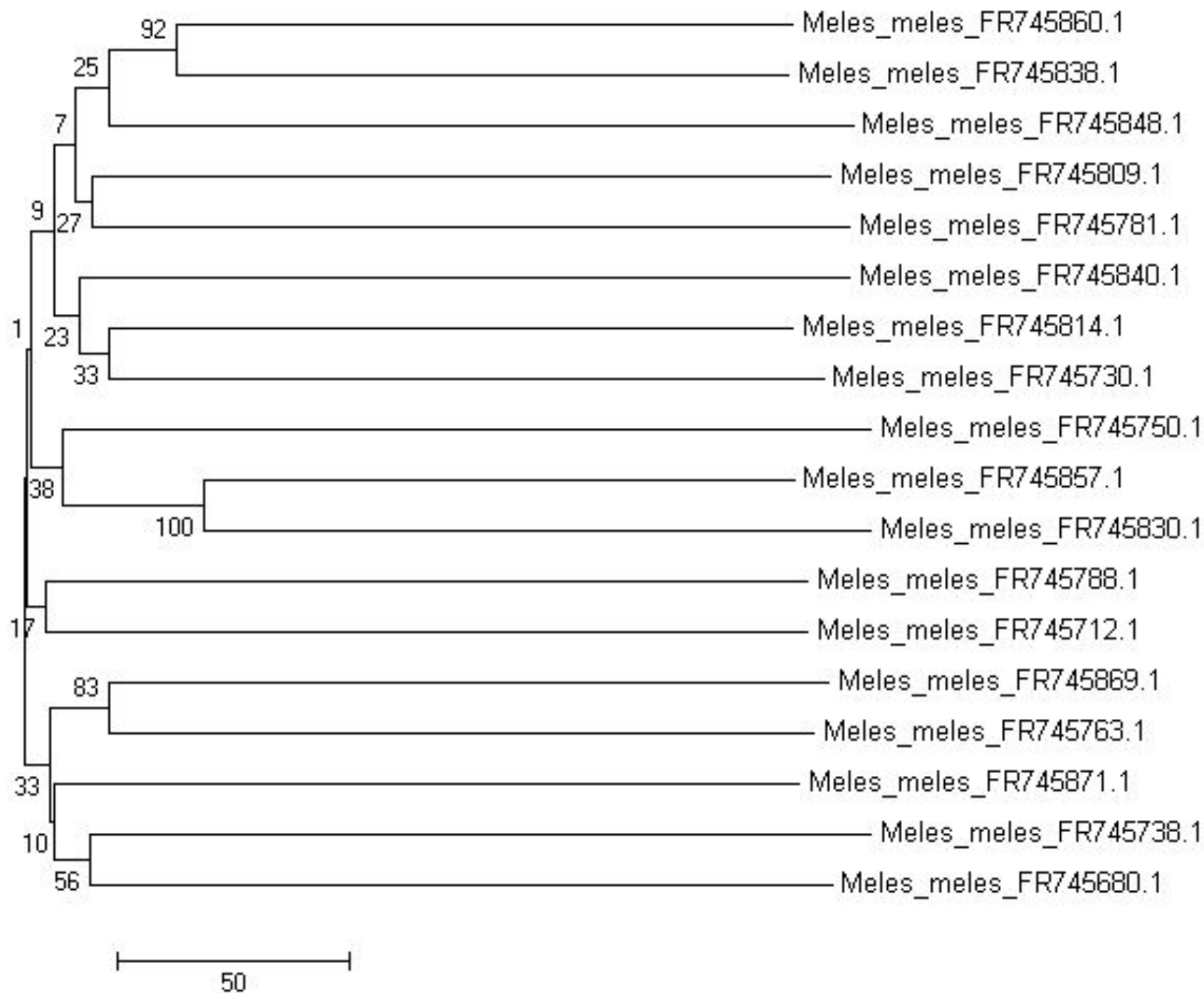
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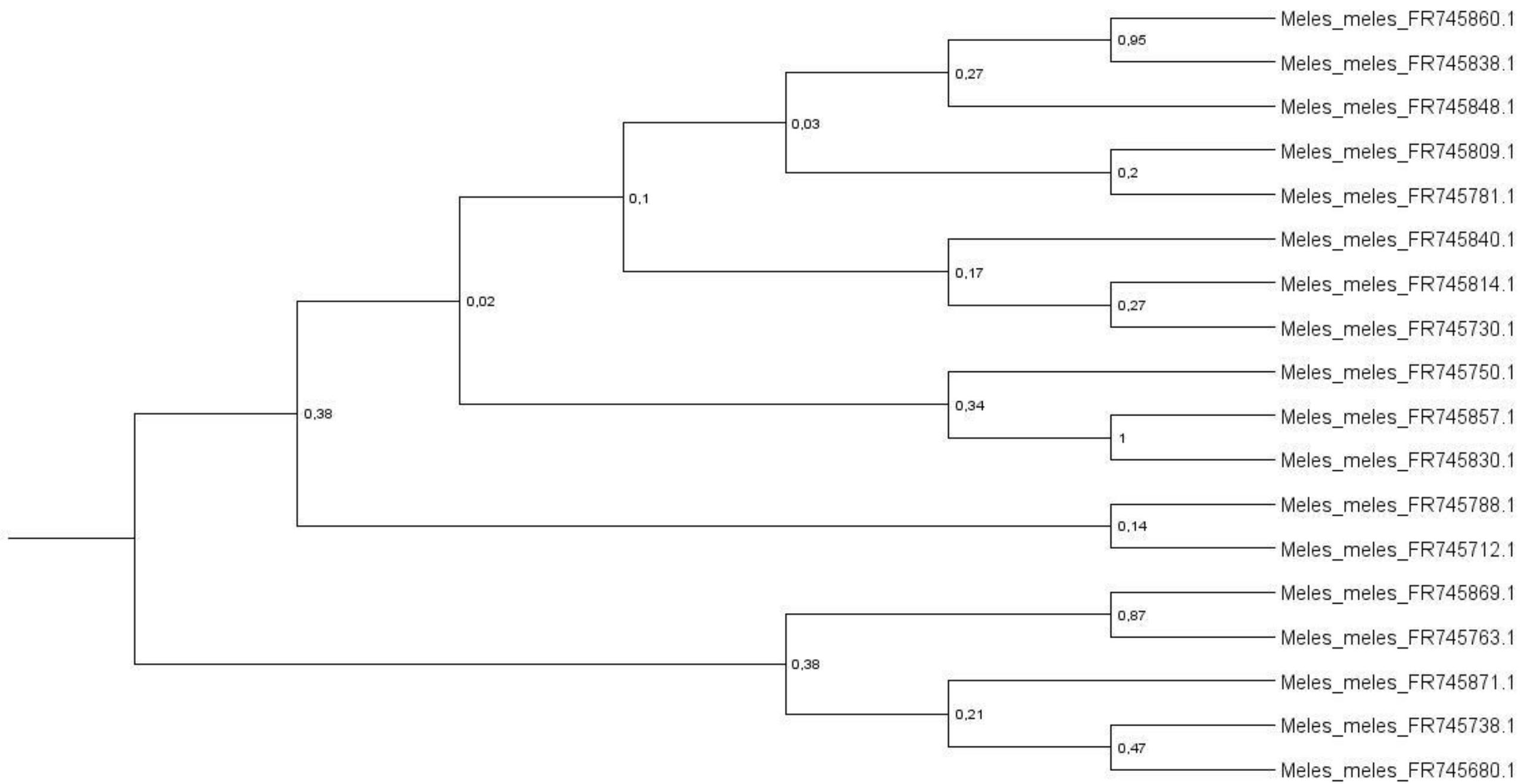
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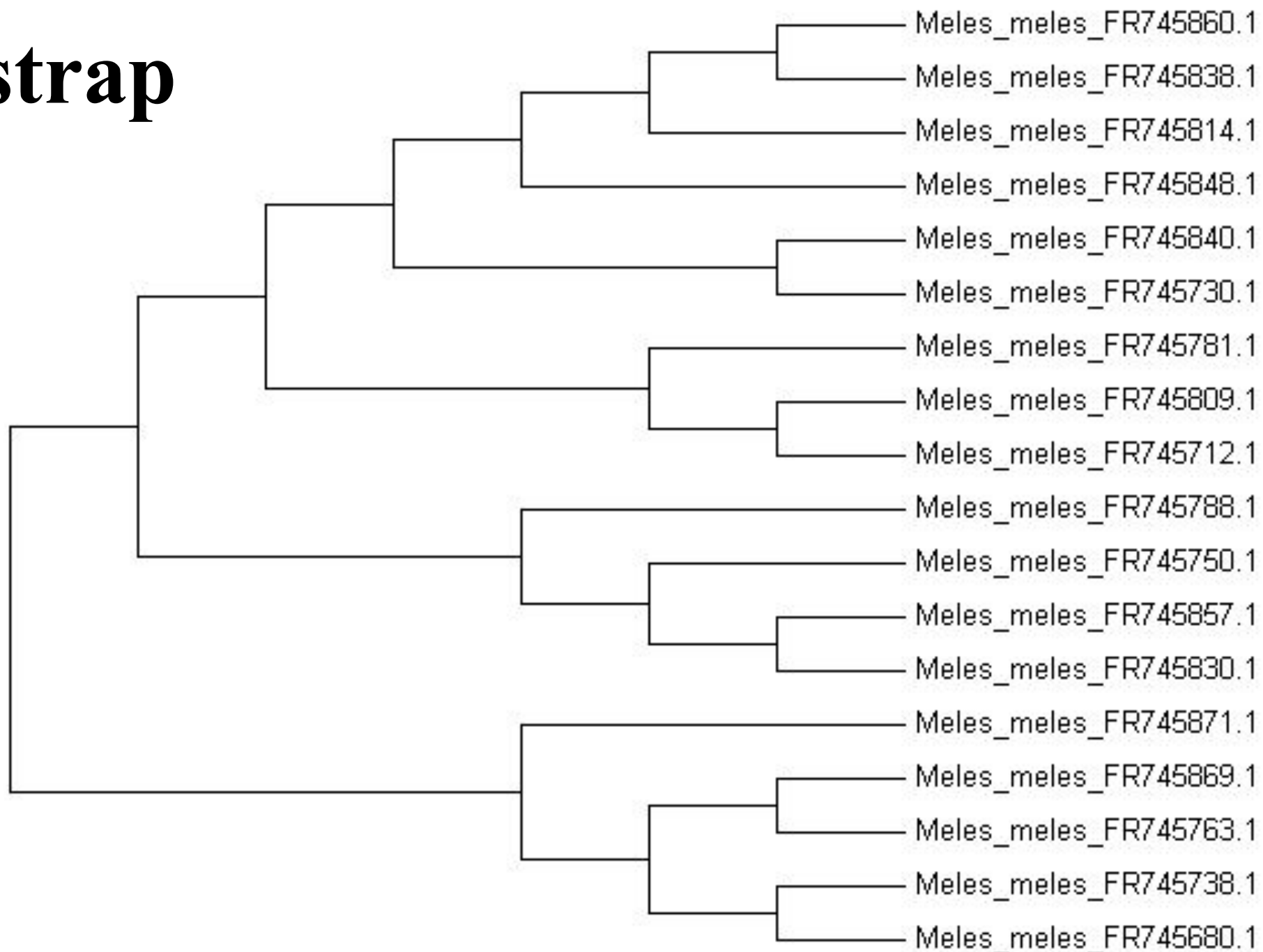
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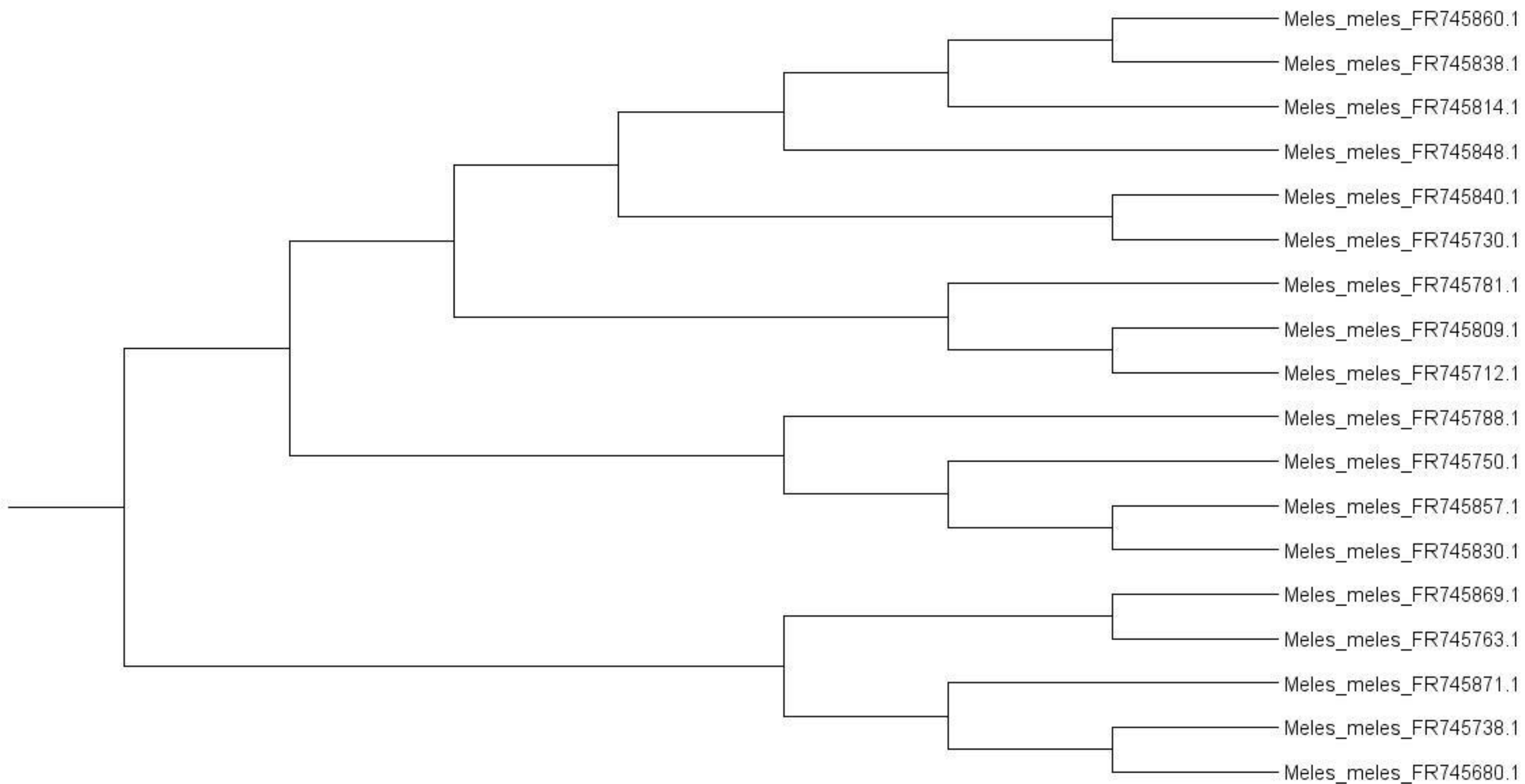




0.04

Bootstrap





**Table. Results from Tajima's Neutrality Test [1]**

<i>m</i>	<i>S</i>	<i>p_s</i>	Θ	π	<i>D</i>
18	495	1.000000	0.290735	0.681970	5.744581

NOTE.-- The analysis involved 18 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There were a total of 495 positions in the final dataset. Evolutionary analyses were conducted in MEGA6 [2].

Abbreviations: *m* = number of sequences, *n* = total number of sites, *S* = Number of segregating sites, $p_s = S/n$, $\Theta = p_s/a_1$, π = nucleotide diversity, and *D* is the Tajima test statistic (see chapter 12 in ref. [3] for details).

1. Tajima F. (1989). Statistical methods to test for nucleotide mutation hypothesis by DNA polymorphism. *Genetics* **123**:585-595.
2. Tamura K., Stecher G., Peterson D., Filipski A., and Kumar S. (2013). MEGA6: Molecular Evolutionary Genetics Analysis version 6.0. *Molecular Biology and Evolution* **30**: 2725-2729.
3. Nei M. and Kumar S. (2000). *Molecular Evolution and Phylogenetics*. Oxford University Press, New York.

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Tajima's Test

Input Data File: D:\...\Meles meles2.fas
Number of sequences: 18 Number of sequences used: 18
Selected region: 1-795 Number of sites: 795
Total number of sites (excluding sites with gaps / missing data): 495

Number of polymorphic (segregating) sites, S: 495
Total number of mutations, Eta: 1407

Average number of nucleotide differences, k: 337,57516
Nucleotide diversity, Pi: 0,68197
Theta (per sequence) from Eta: 409,06484
Theta (per site) from Eta: 0,82639

Tajima's D: -0,74808 Statistical significance: Not significant, $P > 0.10$

=====
Synonymous and NonSynonymous Substitutions
=====

No coding region defined

Calculated using the total number of mutations



Output

Pairwise No. of Differences

Input Data File: D:\...\Meles meles2.fas

Number of sequences: 18 Number of sequences used: 18

Selected region: 1-795 Number of sites: 795

Total number of sites (excluding sites with gaps / missing data): 495

Number of polymorphic (segregating) sites, S: 495

Observed values:

Nucleotide diversity, π : 0,68197Average Number of Pairwise Differences, k : 337,575Observed variance of k : 297,9960Observed C.V. of k : 0,0518Raggedness statistic r : 0,0198

Mean Absolute Error, MAE: 1,8654

Ramos-Onsins and Rozas, $R2$ statistic: 0,3183

Expected values:

Expected Total variance of k (no recombination), $V(k)$: 22931,671Expected C.V. of k (no recombination): 0,4486Expected Stochastic variance of k (no recombination), $Vst(k)$: 20340,374Expected Sampling variance of k (no recombination), $Vs(k)$: 2591,297Expected Total variance of k (free recombination), $V(k)$: 125,763Expected C.V. of k (free recombination): 0,0332Expected Stochastic variance of k (free recombination), $Vst(k)$: 112,525Expected Sampling variance of k (free recombination), $Vs(k)$: 13,238

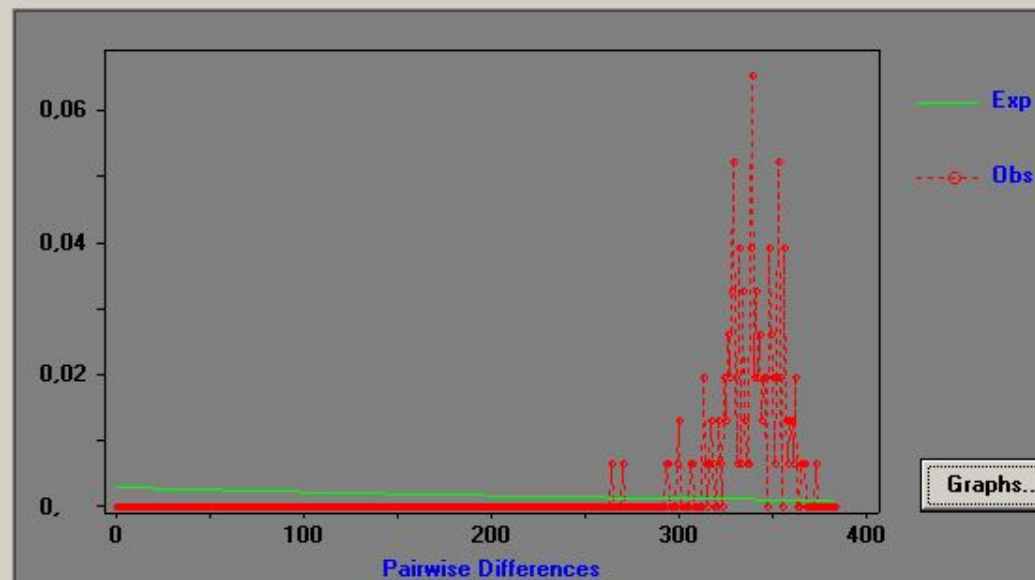
Parameters Estimation by letting Theta Final as infinite

Estimate of Theta Initial: 0,000

Estimate of Tau: 337,575

==== Expected Values for Constant Population Size =====

Graph from file: Meles meles2.fas



Output. Grid

Pairwise No. of Differences

DnaSP Graph

Differences	Freq. Obs.	Freq. Exp.
0	0,00000	0,00295
1	0,00000	0,00294
2	0,00000	0,00294
3	0,00000	0,00293
4	0,00000	0,00292
5	0,00000	0,00291
6	0,00000	0,00290
7	0,00000	0,00289
8	0,00000	0,00288
9	0,00000	0,00288
10	0,00000	0,00287
11	0,00000	0,00286
12	0,00000	0,00285
13	0,00000	0,00284
14	0,00000	0,00283
15	0,00000	0,00283
16	0,00000	0,00282
17	0,00000	0,00281
18	0,00000	0,00280
19	0,00000	0,00279



DNA Polymorphism

Input Data File: D:\...\Meles meles2.fas
Number of sequences: 18 Number of sequences used: 18
Selected region: 1-795 Number of sites: 795
Total number of sites (excluding sites with gaps / missing data): 495

Number of polymorphic (segregating) sites, S: 495
Total number of mutations, Eta: 1407

Number of Haplotypes, h: 18
Haplotype (gene) diversity, Hd: 1,000
Variance of Haplotype diversity: 0,00034
Standard Deviation of Haplotype diversity: 0,019

Nucleotide diversity, Pi: 0,68197
Theta (per site) from Eta: 0,82639

Theta (per site) from S, Theta-W: 0,29074
Variance of theta (no recombination): 0,0101528
Standard deviation of theta (no recombination): 0,10076
Variance of theta (free recombination): 0,0001708
Standard deviation of theta (free recombination): 0,01307

Finite Sites Model
Theta (per site) from Pi: 7,51841
Theta (per site) from S: 0,74902
Theta (per site) from Eta: 10,17833

Average number of nucleotide differences, k: 337,575
Stochastic variance of k (no recombination), Vst(k): 20340,374
Sampling variance of k (no recombination), Vs(k): 2591,297
Total variance of k (no recombination), V(k): 22931,671
Stochastic variance of k (free recombination), Vst(k): 112,525
Sampling variance of k (free recombination), Vs(k): 13,238
Total variance of k (free recombination), V(k): 125,763

Theta (per sequence) from S, Theta-W: 143,914
Variance of theta (no recombination): 2487,678
Variance of theta (free recombination): 41,841

DNA Sp 5.1