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1	RASP 4: ancestral state reconstruction tool for multiple genes and
2	characters
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10	Abstract: With the continual progress of sequencing techniques, genome-scale data are
11	increasingly used in phylogenetic studies. With more data from throughout the genome, the
12	relationship between genes and different kinds of characters is receiving more attention. Here,
13	we present version 4 of RASP, a software to reconstruct ancestral states through phylogenetic
14	trees. RASP can apply generalized statistical ancestral reconstruction methods to phylogenies,
15	explore the phylogenetic signal of characters to particular trees, calculate distances between
16	trees, and cluster trees into groups. RASP 4 has an improved graphic user interface and is
17	freely available from http://mnh.scu.edu.cn/soft/blog/RASP (program) and
18	https://github.com/sculab/RASP (source code).
19	Key words: Ancestral state reconstruction, Genome, Phylogeny, Phylogenetic signal
20	
21	RASP (Reconstruct Ancestral State in Phylogenies) is a software to reconstruct ancestral states
22	through phylogenetic trees. To date, the program has been used to infer biogeographic history
23	in numerous groups of animals, plants, fungi and bacteria (Blair, et al. 2015; Yu, et al. 2015;
24	Stucki, et al. 2016; Bourguignon, et al. 2018; Navaud, et al. 2018; Yan, et al. 2018). With the
25	continual progress of sequencing techniques, the data from genomes, transcriptomes and
26	proteomes have been increasingly used in phylogenetic studies (Choi and Kim 2017).
27	Additionally, morphology, ecology, and distribution data are increasingly integrated into

research (Soltis and Soltis 2016). This motivated us to add more functionality into RASP toimplement additional algorithms and tools.

The new version of RASP can analyze phylogenomic data (and other types of data), make inference on our generalized statistical method for ancestral state reconstruction (Fig 1-A and B) and summarize results under a graphical user interface (Fig 1-C). Users are also allowed to quantify phylogenetic signal of different morphological or ecological characters to particular trees (Fig 1-D), measure the fit between a tree and geography, and compute a distance matrix to cluster trees (Fig 1-E).

36 Methods to reconstruct ancestral geographical distributions using a combination of 37 phylogenetic and distributional information are increasing rapidly. In RASP 4, we implement a 38 generalized statistical method for models implemented in the R package 'BioGeoBEARS' 39 (Matzke 2014) and 'APE' (Paradis and Schliep 2018); namely our method summarizes 40 ancestral reconstructions across all input trees. The probability (p) of an ancestral range x at node *n* on the final species tree is calculated as $p(x_n) = \frac{\sum_{t \in T} [w(x_n)_t]}{g_n}$ where *T* is the set 41 42 of trees, $w(x_n)_t$ is the weight of ancestral range x at node n for tree t, and g_n is the number of 43 times node n occurs in T (see supplementary material for details). To reduce computational 44 burden, RASP applies parallel computing to all models both by taking advantage of multiple 45 threads and splitting trees into small groups. See Table S1 for a full comparison of the methods 46 of ancestral reconstruction implemented in RASP.

47 Phylogenetic signal is the tendency of related species to resemble each other in a specific 48 character more than species drawn at random from the same tree (Münkemüller, et al. 2012). 49 To test for phylogenetic signal for continuous states, RASP calculates Moran's I (Moran 1948, 50 1950), Abouheif's C_{mean} (Abouheif 1999), Pagel's λ (Pagel 1999) and Blomberg's K (Blomberg, 51 et al. 2003) using the R package 'adephylo' (Jombart, et al. 2010). For discrete states, RASP fits 52 models of trait evolution using a likelihood ratio test and calculates Pagel's λ using the R 53 package 'geiger' (Pennell, et al. 2014) (Fig 1-C). If some species have more than one state, 54 RASP will convert them to all possible combinations of single states and calculate Pagel's λ for 55 each of them. The largest Pagel's λ is used in the final result.

56 Tree distances are often used as a formal way to quantify the differences of trees inferred

57 from different genes and reconstruction methods (Sand, et al. 2014). In RASP, users can 58 compute trees distances using different methods: KC distance (Kendall and Colijn 2016), triplet 59 distance implemented in mp-est (Liu, et al. 2010), RF distance (Robinson and Foulds 1981), 60 KF distance (Kuhner and Felsenstein 1994), path differences (Steel and Penny 1993), and SPR 61 distance (de Oliveira Martins, et al. 2008; De Oliveira Martins, et al. 2014) implemented in the 62 R package 'phangorn' (Schliep 2010) (Table S2). Having the distance matrix, values can be 63 normalized using min-max normalization (Han, et al. 2006). Next, trees are clustered into 64 groups using the R package 'adegenet' (Jombart 2008) according to the distance matrix. The 65 end result may provide insight into the sources of heterogeneity among gene/species histories. 66 For example, distinct clusters of genes may indicate unique phylogenetic signatures (Kendall 67 and Colijn 2016). Additionally, the tree distance matrix and groups could be used to provide a 68 candidate species tree under the coalescent model (Liu, et al. 2010).

69 In summary, the new version of RASP 4 implements several tools for multiple gene and 70 species trees and characters while simultaneously making it easier to process trees generated 71 from different sources. We provide six tutorials to help users select appropriate methods for 72 different research questions on the our website (http://mnh.scu.edu.cn/soft/blog/RASP). We 73 will continue to develop RASP with a focus on implementing new algorithms and integrating 74 RASP Windows macOS available more tools. for and are freely from 75 http://mnh.scu.edu.cn/soft/blog/RASP (program) and https://github.com/sculab/RASP (source 76 code), and licensed under the terms of the MIT license.

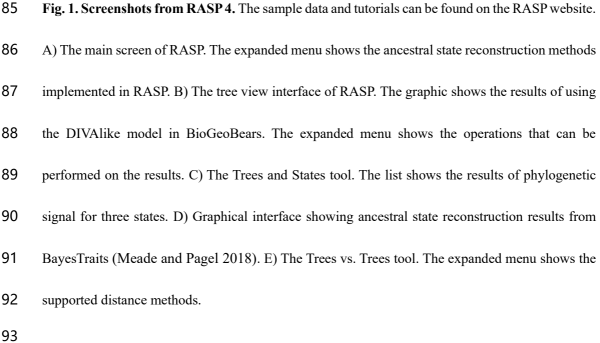
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95 References

- Abouheif E. 1999. A method for testing the assumption of phylogenetic independence in comparative
 data. Evolutionary Ecology Research 1:895-909.
- Blair C, Noonan B, Brown J, Raselimanana A, Vences M, Yoder A. 2015. Multilocus phylogenetic and
 geospatial analyses illuminate diversification patterns and the biogeographic history of Malagasy
 endemic plated lizards (Gerrhosauridae: Zonosaurinae). Journal of Evolutionary Biology 28:481-492.
- Blomberg SP, Garland Jr T, Ives AR. 2003. Testing for phylogenetic signal in comparative data:
 behavioral traits are more labile. Evolution 57:717-745.
- Bourguignon T, Tang Q, Ho SYW, Juna F, Wang Z, Arab DA, Cameron SL, Walker J, Rentz D, Evans
 TA, et al. 2018. Transoceanic Dispersal and Plate Tectonics Shaped Global Cockroach Distributions:
 Evidence from Mitochondrial Phylogenomics. Mol Biol Evol 35:970-983.
- 106 Choi J, Kim S-H. 2017. A genome tree of life for the fungi kingdom. Proceedings of the National
 107 Academy of Sciences 114:9391-9396.
- de Oliveira Martins L, Leal E, Kishino H. 2008. Phylogenetic detection of recombination with a Bayesian
 prior on the distance between trees. PLoS One 3:e2651.
- De Oliveira Martins L, Mallo D, Posada D. 2014. A Bayesian supertree model for genome-wide species
 tree reconstruction. Systematic Biology 65:397-416.
- Han J, Jian P, Michelin K. 2006. Data Mining, Southeast Asia Edition. In: San Francisco: Elsevier Inc.
- Jombart T. 2008. adegenet: a R package for the multivariate analysis of genetic markers. Bioinformatics
 24:1403-1405.
- Jombart T, Balloux F, Dray S. 2010. Adephylo: new tools for investigating the phylogenetic signal in
 biological traits. Bioinformatics 26:1907-1909.
- Kendall M, Colijn C. 2016. Mapping Phylogenetic Trees to Reveal Distinct Patterns of Evolution. Mol
 Biol Evol 33:2735-2743.
- Kuhner MK, Felsenstein J. 1994. A simulation comparison of phylogeny algorithms under equal and
 unequal evolutionary rates. Mol Biol Evol 11:459-468.
- Liu L, Yu L, Edwards SV. 2010. A maximum pseudo-likelihood approach for estimating species trees
 under the coalescent model. BMC evolutionary biology 10:302.
- Matzke NJ. 2014. Model selection in historical biogeography reveals that founder-event speciation is a
 crucial process in island clades. Systematic Biology 63:951-970.
- 125 Meade A, Pagel M. 2018. BayesTraits: a computer package for analyses of trait evolution. In: Version.
- Moran PA. 1948. The interpretation of statistical maps. Journal of the Royal Statistical Society. Series B
 (Methodological) 10:243-251.
- 128 Moran PA. 1950. Notes on continuous stochastic phenomena. Biometrika 37:17-23.
- Münkemüller T, Lavergne S, Bzeznik B, Dray S, Jombart T, Schiffers K, Thuiller W. 2012. How to
 measure and test phylogenetic signal. Methods in Ecology and Evolution 3:743-756.
- Navaud O, Barbacci A, Taylor A, Clarkson JP, Raffaele S. 2018. Shifts in diversification rates and host
 jump frequencies shaped the diversity of host range among Sclerotiniaceae fungal plant pathogens.
 Mol Ecol 27:1309-1323.
- 134 Pagel M. 1999. Inferring the historical patterns of biological evolution. Nature 401:877.
- Paradis E, Schliep K. 2018. ape 5.0: an environment for modern phylogenetics and evolutionary analyses
 in R. Bioinformatics 35:526-528.
- 137 Pennell MW, Eastman JM, Slater GJ, Brown JW, Uyeda JC, FitzJohn RG, Alfaro ME, Harmon LJ. 2014.
- geiger v2. 0: an expanded suite of methods for fitting macroevolutionary models to phylogenetic trees.

139 Bioinformatics 30:2216-2218.

- Robinson DF, Foulds LR. 1981. Comparison of phylogenetic trees. Mathematical biosciences 53:131141 147.
- Sand A, Holt MK, Johansen J, Brodal GS, Mailund T, Pedersen CN. 2014. tqDist: a library for computing
 the quartet and triplet distances between binary or general trees. Bioinformatics 30:2079-2080.
- 144 Schliep KP. 2010. phangorn: phylogenetic analysis in R. Bioinformatics 27:592-593.
- Soltis DE, Soltis PS. 2016. Mobilizing and integrating big data in studies of spatial and phylogenetic
 patterns of biodiversity. Plant Diversity 38:264-270.
- Steel MA, Penny D. 1993. Distributions of tree comparison metrics—some new results. Systematic
 Biology 42:126-141.
- Stucki D, Brites D, Jeljeli L, Coscolla M, Liu Q, Trauner A, Fenner L, Rutaihwa L, Borrell S, Luo T, et
 al. 2016. Mycobacterium tuberculosis lineage 4 comprises globally distributed and geographically
 restricted sublineages. Nat Genet 48:1535-1543.
- Yan H-F, Zhang C-Y, Anderberg AA, Hao G, Ge X-J, Wiens JJ. 2018. What explains high plant richness
 in East Asia? Time and diversification in the tribe Lysimachieae (Primulaceae). New Phytologist
 219:436-448.
- Yu Y, Harris AJ, Blair C, He X. 2015. RASP (Reconstruct Ancestral State in Phylogenies): a tool for
 historical biogeography. Molecular Phylogenetics and Evolution 87:46-49.

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