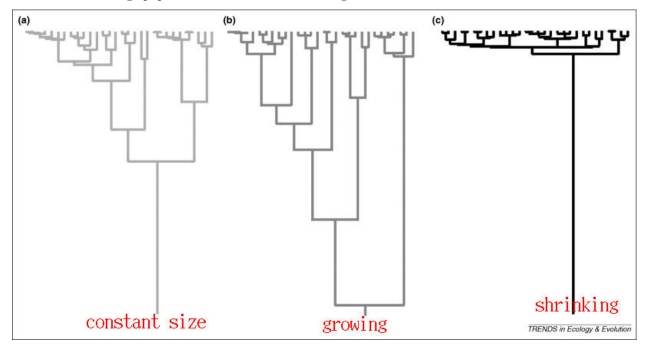
BIO4AE3 extras

Ben Bolker

coalescent methods

Basic idea: build a phylogenetic (gene) tree based on (ideally) neutral genomic samples. The **shape** of the tree (rate of branching per unit time) tells us whether the population is decreasing, increasing, or stable.

- times with small population size have more branching
- times with large population size have less branching



See Elliot and Mooers (n.d.) (intro, a little mathy, but very clear); Kuhner (2009) (intro journal article); Volz et al. (2009) (using coalescents for epidemics); Stadler et al. (2013) (ditto)

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Kuhner, Mary K. 2009. "Coalescent Genealogy Samplers: Windows into Population History." Trends in Ecology & Evolution 24 (2): 86–93. https://doi.org/10.1016/j.tree.2008.09.007.

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