Evolutionary Divergence and Population Structure of the North American Castanea Species

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Background
American chestnut (Castanea dentata) and the chinquapins (C. pumila sensu lato) are an evolutionary sister species pair that represent a promising system for studies of admixture and adaptation in wild plants. Since the 1920s, botanists and geneticists have hypothesized that hybridization between American chestnut and the chinquapins is common and that this process has been involved in the origins of some Castanea taxa and populations [1-3]. Rigorously testing these hypotheses, however, has been difficult until the recent availability of high-throughput DNA sequencing technologies and associated computational tools. Here, we present preliminary results of our analyses of whole genome sequencing data from 255 plants representing all North American Castanea species and subspecific taxa.

Research Questions
- Have American chestnut populations experienced introgression from any of the chinquapin taxa (pumila, ozarkensis, or alabamensis), particularly where they occur in sympatry?
- Has Alabama chinquapin (C. alabamensis) experienced introgression from C. dentata?
- What is the frequency of admixture between different chinquapin taxa?

Sampling
Between 2006 and 2021, we collected leaf tissue samples representing all North American Castanea taxa. We generated whole-genome sequencing (WGS) data from 255 individual plants using methods described previously [4]. Datasets containing >27,000,000 single-nucleotide polymorphisms (SNPs) were used for all analyses. Sample sites are shown below (green dot = C. dentata, purple = pumila, gold = alabamensis, light blue = ozarkensis).

Population structure analysis suggests admixture between American chestnut and the chinquapins has been rare
We inferred population structure of the North American Castanea species using ADMIXTURE software [5]. The genetic structure of our entire sample set was best characterized by a model involving four clusters (K=4): (1) Ozark chinquapins, (2) Allegheny chinquapins from Coastal Plain sites, (3) Allegheny chinquapins from sites in the Appalachians, and (4) Alabama chinquapins.

The chinquapin taxa comprise a sister group to American chestnut
We inferred a maximum-likelihood phylogenetic tree using IQ-TREE software [6]. Bootstrap support for nodes separating taxa was 100% in all cases.

D statistics (i.e., ABBA-BABA) to test for gene flow
We used Patterson’s D statistic (also known as the ABBA-BABA test) [7] to test for gene flow/admixture between different Castanea species and between different taxa within the chinquapin group. Consider the four populations (labeled P1, P2, P3, and O) and their phylogenetic tree shown below. With D statistics, we test the null hypothesis that P1 and P2 descend from a common ancestral population that split from the ancestors of P3 at an earlier time, without any gene flow between P3 and P1 or P2 after they split. The alternative hypothesis is that P3 exchanged genes with P1 or P2 after these two populations diverged. D statistics can also be applied to tests for gene flow between species, as we have done in our study.

The D statistic considers biallelic sites in the genome where we have coverage for P1, P2, P3, and O. The allele in the O population is denoted “A”, and any alternative alleles in P1, P2, or P3 are denoted “B”. The pattern “ABBA” refers to sites in the genome where P1 has the outgroup allele (A) and P2 and P3 share the derived allele (B). The pattern “BABA” refers to sites where P1 and P3 share the derived allele (B) and P2 has the outgroup allele (A). If P1 and P2 have diverged without experiencing gene flow with P3, the frequency of ABBA sites will be equal to the frequency of BABA sites, and the D statistic will not differ significantly from zero.

Hybridization between American chestnut and the chinquapins has been rare
Using the genome package ‘Discraze’ [8], we calculated Patterson’s D statistics to test for gene flow between C. dentata and chinquapin populations for which ADMIXTURE had produced evidence suggestive of hybridization. Significant test results are highlighted in green.

Hybridization between different chinquapin taxa has been more common, but still infrequent
We calculated Patterson’s D statistics to test for gene flow between chinquapin population pairs for which ADMIXTURE had produced evidence suggestive of gene flow. Significant test results are highlighted in green.

Conclusions
- Population structure analysis and D statistics provided concordant results for tests of gene flow between American chestnut and chinquapin. We found evidence of chinquapin ancestry in only one American chestnut sample site, a location in northern Alabama.
- We found no evidence to support the hypothesis that C. alabamensis has introgressed ancestry from C. dentata. Interestingly, our phylogenetic analysis indicates that C. alabamensis is a sister group to Ozark chinquapin. This is consistent with hypotheses on the relationship between Ozark chinquapin and C. alabamensis that date back to the early 20th century [9,10].
- D statistics and analysis of population structure indicate that admixture between C. alabamensis and Allegheny chinquapin is infrequent, but not rare. The same analyses indicate that admixture between Ozark and Allegheny chinquapin is also infrequent. Discrepancies between results from population structure analysis and D statistics are probably due to incomplete lineage sorting.
- We found evidence for the existence of two distinct ecotypes of Allegheny chinquapin – one from the Coastal Plain and a second from the Appalachians. These two ecotypes appear to have experienced substantial gene flow in the Atlantic Coast and eastern Gulf Coastal Plain regions. This is consistent with more limited information obtained in a recent study using genotyping-by-sequencing [11].
- Future work will include analyses of local (chromosome-scale) ancestry and tests for evidence of natural selection at introgressed ancestry tracts.

Notes
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References
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