Conflicting genomic signals affect phylogenetic inference in four species of North American pines

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Abstract. Adaptive evolutionary processes in plants may be accompanied by episodes of introgression, parallel evolution and incomplete lineage sorting that pose challenges in untangling species evolutionary history. Genus Pinus (pines) is one of the most abundant and most studied groups among gymnosperms, and a good example of a lineage where these phenomena have been observed. Pines are among the most ecologically and economically important plant species. Some, such as the pines of the southeastern USA (southern pines in subsection Australes), are subjects of intensive breeding programmes. Despite numerous published studies, the evolutionary history of Australes remains ambiguous and often controversial. We studied the phylogeny of four major southern pine species: shortleaf (Pinus echinata), slash (P. elliottii), longleaf (P. palustris) and loblolly (P. taeda), using sequences from 11 nuclear loci and maximum likelihood and Bayesian methods. Our analysis encountered resolution difficulties similar to earlier published studies. Although incomplete lineage sorting and introgression are two phenomena presumptively underlying our results, the phylogenetic inferences seem to be also influenced by the genes examined, with certain topologies supported by sets of genes sharing common putative functionalities. For example, genes involved in wood formation supported the clade echinata – taeda, genes linked to plant defence supported the clade echinata – elliottii and genes linked to water management properties supported the clade echinata – palustris. The support for these clades was very high and consistent across methods. We discuss the potential factors that could underlie these observations, including incomplete lineage sorting, hybridization and parallel or adaptive evolution. Our results likely reflect the relatively short evolutionary history of the subsection that is thought to have begun during the middle Miocene and has been influenced by climate fluctuations.

Keywords: Australes; drought tolerance; parallel evolution; phylogeny; plant defence; southern pines; wood formation.

Introduction

Plant genome structure and evolution are subjects of intensive investigations with recent milestone advances spanning whole-genome sequencing of some of the largest plant genomes, such as those of pines and spruces (Biral et al. 2013; Nystedt et al. 2013; Krutovsky et al. 2014; Neale et al. 2014; Zimin et al. 2014). These advancements bring opportunities to ask more focussed
questions about the dynamic processes that have contributed to plant genome evolution and influenced phylogenetic relationships among plant species. Duplications, from single gene to whole genome, are an intrinsic process that appears to be the major force driving genome evolution in plants (Freeling 2009; Li et al. 2015). Other processes, such as introgression, incomplete lineage sorting and parallel evolution, primarily influenced by extrinsic factors, additionally contribute to plant genome complexity and evolution (Martinsen et al. 2001; Wood et al. 2005; Willyard et al. 2009). The latter processes may also shape phenotypic and genotypic similarities among species becoming a challenge in phylogenetic studies.

Pines (genus Pinus, family Pinaceae) comprise a group of coniferous tree species occurring almost exclusively in the Northern Hemisphere, with populations often dominant across areas of the North Temperate Zone (Critchfield and Little 1966). They are keystone species of the vast boreal forest ecosystem. They provide pulp and timber products as primary commodities (Lowe et al. 1999; Borders and Bailey 2001; Alexander et al. 2002; Wear and Greis 2002; Croitoru 2007), but some additional pine forest products may include, for example, pine nuts, berries, herbs and mushrooms (Alexander et al. 2002; Bürgi et al. 2013; Nagasaka 2013). They provide a pivotal habitat for wildlife species (Brockerhoff et al. 2008), and greatly contribute to carbon storage and other ecosystem services (Goodale et al. 2002; McKinley et al. 2011). Due to their aesthetic value, they add recreational and ornamental dimensions in urban and suburban areas (Tyrväinen and Väänänen 1998; Knoth et al. 2002). These qualities extend to southeastern forest ecosystems of the USA, where pines may grow also in mixed forests with hardwoods (Sharitz et al. 1992), and where they additionally play an important role in ecosystem recovery from natural disturbances. For example, longleaf pine has developed complex adaptations to fire that allow fast stand regeneration (Outcalt 2000), and loblolly pine helps minimize soil erosion and provides watershed protection due to its fast growth (Schultz 1997).

Four major pines of the southeastern USA were investigated in this study: shortleaf (Pinus echinata), slash (P. elliottii), longleaf (P. palustris) and loblolly (P. taeda) (four of the ‘southern pines’; subsection Australes, section Trifoliae, genus Pinus). They are widely cultivated, greatly dominating the southern US forest inventory (Sternitzke and Nelson 1970), and are, therefore, a subject of breeding programmes in the region (Wear and Greis 2002; McKeand et al. 2003; Fox et al. 2007). The traditional classification (Little and Critchfield 1969) considered 11 species in this subsection with habitat stretching cumulatively from the southeastern USA, through Mexico, to the Caribbean and Central America: slash (P. elliottii), spruce (P. glabra), longleaf (P. palustris), pond (P. serotina) and loblolly (P. taeda) pines in the southeastern USA; shortleaf (P. echinata), Table Mountain (P. pungens) and pitch (P. rigida) pines in the eastern USA; Cuban pine (P. cubensis) in Cuba; West Indian pine (P. occidentalis) in the West Indies; and Caribbean pine (P. caribaea) in both the West Indies and adjacent Central America. Attempts to refine the phylogeny of Australes are ongoing, but typically have been approached in the broader context of the Pinaceae family or along with other subsections.

Several previous investigations have provided insights into the phylogenetic relationships of the four species on which we focus in our study (i.e. P. echinata, P. elliottii, P. palustris and P. taeda; Fig. 1). Adams and Jackson (1997) found a very close relationship between P. palustris and P. taeda based on 21 morphological characters, but their relationship to P. echinata and P. elliottii remained unresolved, and no statistical support was provided. Later, using RAPD markers and the neighbour-joining method, Dvorak et al. (2000) suggested a close relationship among P. echinata, P. palustris and P. taeda, in which P. echinata and P. taeda were sister lineages, and P. elliottii was sister to P. caribaea. Using a supertree approach and previously published phylogenies based on both morphological and molecular data, Grotkopp et al. (2004) also inferred a close relationship among P. echinata, P. palustris and P. taeda, but found that P. palustris was sister to P. taeda; P. elliottii was, again, sister to P. caribaea. Gerndt et al. (2005) used

![Figure 1. Cladograms for the four Australes species—P. echinata, P. elliottii, P. palustris and P. taeda—from published studies. (A) Adams and Jackson (1997). (B) Dvorak et al. (2000). (C) Grotkopp et al. (2004). (D) Gerndt et al. (2005). (E) Eckert and Hall (2006). (F) Hernández-León et al. (2013). The original studies presented the four species in a broader context along with other pines.](image-url)
two chloroplast genes in 101 pine species, but relationships among the four Australes species remained unresolved. Eckert and Hall (2006) used four chloroplast genes in 83 pine species. In their study, *P. echinata* and *P. palustris* were placed in one clade with low support, and *P. elliottii* and *P. taeda* were placed in another. Most recently, based on five chloroplast DNA (cpDNA) markers, Hernández-León et al. (2013) placed *P. echinata*, *P. elliottii* and *P. palustris* within the same clade, where *P. elliottii* and *P. palustris* were sisters, and *P. taeda* was in a separate clade, albeit with low bootstrap support (BS). Consequently, the monophyly of the four species was placed under question.

Evidence from cpDNA used in recent studies additionally questioned monophyly of the subsection Australes as defined by Little and Critchfield (1969), and suggested that the 11 Australes species may be scattered throughout a larger clade of over twice as many species (Gernandt et al. 2005; Eckert and Hall 2006; Hernández-León et al. 2013). Chloroplast genomes, however, may follow different evolutionary trajectories than nuclear genomes, with potentially confounding effects in phylogenetic studies (Rieseberg and Soltis 1991). In pines, chloroplast genomes are strictly paternally inherited, thus having smaller effective population sizes than nuclear genomes. Additionally, cpDNA experiences lower substitution rates (Willyard et al. 2007). Consequently, these factors may lead to discordant patterns of polymorphism between the two genomes (Soltis et al. 1992; Hong et al. 1993b). Moreover, cpDNA differentiation among populations of a species can be high (Hong et al. 1993a), and the cpDNA sequence variation in pines may be unevenly distributed throughout the genome requiring a representative locus sampling (Whittall et al. 2010). Finally, foreign chloroplast genomes introduced through hybridization may undergo a rapid fixation—chloroplast capture (Rieseberg and Soltis 1991; Matos and Schaal 2000; Liston et al. 2007).

Apart from the specificity of cpDNA evolution, additional problems with taxonomic classification of Australes may stem from incomplete lineage sorting, a phenomenon reported in pines (Syring et al. 2007; Willyard et al. 2009), especially considering the relatively recent evolutionary history of the subsection. Radiation within Australes is thought to have begun only 7–15 million years ago (MYA) (Willyard et al. 2007; Hernández-León et al. 2013). Large overlap in present-day areas of the four species studied here has fostered occasional or historic hybridization between some of them that may have contributed to insufficient resolution in phylogenetic studies. Natural hybridization has been observed between *P. echinata* and *P. taeda* (Zobel 1953; Mergen et al. 1965; Smouse and Saylor 1973; Edwards-Burke et al. 1997), between *P. palustris* and *P. taeda* (Chapman 1922) and between *P. elliottii* and *P. palustris* (Mergen 1958). Almost all possible hybrids for these species can be artificially generated in controlled crosses, with the exception of *P. echinata* and *P. palustris* that are not considered interfertile despite a reported hybrid case (Snyder and Squillace 1966; Campbell et al. 1969). Additionally, similar environmental constraints could lead to a parallel support for beneficial alleles and/or removal of detrimental ones. Nevertheless, sympathy could have been repeatedly disturbed in the past; for example, during the Pleistocene, *P. taeda* was likely constricted to two refugia, *P. elliottii* and *P. palustris* could have been separated, one occupying one of the two refugia, while *P. echinata*’s range was probably continuous due to its cold-hardiness (Wells et al. 1991; Schmittling and Hipkins 1998; Schmittling 2003). Each of these factors, or their combination, could confound phylogenetic inference.

The challenge with inferring relationships among Australes indicates that more work is needed. Nuclear markers are recommended to use when introgression, cytoplasmic in particular, or lineage sorting phenomena are present (Soltis et al. 1992). Consequently, we used 11 nuclear protein coding genes. Our objective was to refine and potentially clarify the phylogenetic relationships among *P. echinata*, *P. elliottii*, *P. palustris* and *P. taeda*. *Pinus pinaster* was used as an outgroup. Despite the larger dataset and application of advanced methods, untangling their phylogeny was not straightforward. We discuss potential factors that likely contributed to this problem and could explain the difficulties in inferring phylogenies using multiple genes observed in previous studies.

**Methods**

**Source of data**

Data from 32 genes recently sequenced and annotated in four pines from subsection Australes, namely *P. taeda* (Brown et al. 2004; González-Martínez et al. 2006), *P. echinata*, *P. elliottii* and *P. palustris* (Koralewski et al. 2014), were used to query the NCBI GenBank database (Benson et al. 2013) for orthologs in related pine (genus *Pinus*) taxa that could be used as appropriate outgroup species. Eleven putative orthologous genes were identified in *P. pinaster* (Table 1), which was used as an outgroup. Given that the ingroup species belong to a single subsection, it is possible that a species from a more closely related subsection could be a more optimal outgroup for phylogenetic analyses; however, *P. pinaster* is the best studied species with respect to the genes investigated in the southern pines, allowing us to utilize more sequence data in the analyses. We assumed orthology of the genes based on exon–intron structure and very high sequence similarity. For each putative ortholog, the E-value was 3E-46 or less and identity score was 95% or more in at least one comparison of an ingroup species.
with *P. pinaster*; additionally, in comparisons where the *E*-value was higher than *E*-100, the identity score was 99%. However, well-annotated, mapped and well-assembled whole-genome data are ultimately needed to validate orthology. Such data are still unavailable, although the first incomplete draft genome assembly was published recently for loblolly pine (Neale et al. 2014; Zimin et al. 2014).

### Multiple alignment

The FASTA sequences for individual genes were aligned using BioEdit (ver. 7.0.9.0) (Hall 1999) and merged into one dataset in SeaView (ver. 4.0) (Galtier et al. 1996). Conversion from FASTA to NEXUS was done in SeaView, and from FASTA to PHYLIP manually in the text editor Notepad++ (ver. 5.9.3) (Ho 2011). Coding sites were assigned based on annotation data in NCBI GenBank for each individual gene separately using DnaSP (ver. 5.10.01) (Librado and Rozas 2009). The merged NEXUS file was further manually annotated, and one of five categories was assigned to each site: a codon position (1, 2 or 3), intron or 3'UTR. All sites with missing data were removed from the analysis [see Supporting Information—Dataset S1].

### Partitioning

Partitioning schemes and models of molecular evolution were evaluated using PartitionFinder (ver. 1.1.1) (Lanfear et al. 2012). Model selection was restricted to the set of models implemented in the software in which we intended to run the phylogenetic analysis (parameter ‘model =’ in PartitionFinder). At most 42 models were evaluated simultaneously based on Akaike information criterion (AIC) and Bayesian information criterion (BIC). The parameters describing gamma distribution of rates among sites (*G* or G) and a proportion of invariable sites (*I*) were considered among the models exclusively, i.e. none of the evaluated models accounted for both *G* and *I* jointly, because of reported problems with independent optimization of these two parameters (see discussions and relevant references in Stamatakis 2008, pp. 20–21 and in Stamatakis 2014, p. 49). Additionally, in the group of models evaluated for GARLI (Zwickl 2006) (see below), one or more of the K81 models (K81, K81 + I and K81 + G, depending on the subset of data; a few data subsets were affected) caused convergence problems and were not considered. Depending on the intended phylogenetic analysis, alternative partitioning schemes were then implemented [see Supporting Information—Table S1]: (i) by-gene-site: best partitioning schemes for 39 sets corresponding to different genes and site categories within genes, or (ii) by-gene: best models identified for each of the 11 genes in the set (no partitioning within a gene). Additionally, in the cases where each gene was analysed separately, models were identified for every gene independently with sites assigned to one of the five site categories (by-site).

### Phylogenetic analysis

The combined dataset of 11 genes was subjected to two gene tree methods with partitioning by-gene-site: maximum likelihood (ML; GARLI, ver. 2.01) (Zwickl 2006) and
Bayesian inference (BI; MrBayes, ver. 3.2.2) (Ronquist et al. 2012). Species trees were reconstructed using the Bayesian method BEST (ver. 2.3) (Liu 2008) with partitioning by-gene. To account for potential polytomies, and thus potential for the star tree paradox (Suzuki et al. 2002; Lewis et al. 2005), we ran Phycas (ver. 1.2.0) (Lewis et al. 2010), also a Bayesian method, with partitioning by-gene-site. We then analysed each gene separately using GARLI and MrBayes (partitioning by-site). The MrBayes outputs were further investigated using Bayesian concordance analysis (Ané et al. 2007), as implemented in BUCKy (ver. 1.4.0, mbsum ver. 1.4.2) (Larget et al. 2001), for seven values of $\alpha$: 0.01, 0.5, 1, 2, 5, 10 and 10 000. The value of $\alpha$ corresponds to the probability that loci share the same tree (the lower $\alpha$ the higher the probability, and vice versa), thus affecting the clade support (the higher the probability the higher support). Formally, BUCKy is not considered a species-tree method; however, the resulting primary concordance tree can be generally considered comparable with species trees (for an in-depth discussion about the species methods, see Mateos et al. 2012).

All analyses were run in Windows 7 [see Supporting Information—Table S1] except for BUCKy, which was compiled and run under Cygwin. The tree figures were visualized in FigTree (ver. 1.4) (Rambaut 2013) and further edited manually to increase their readability and compactness.

**Clade support and convergence**

In order to determine the clade support, BS was calculated in GARLI using 1000 replicates [see Supporting Information—Table S1]. SumTrees (ver. 3.3.1) (Sukumaran and Holder 2010) was used to generate majority consensus trees. Posterior probability (PP) estimates were used for the BI methods. To verify that the Markov chain Monte Carlo analysis converged on a stationary distribution and that the sampling of the distribution was adequate, the following criteria were applied for MrBayes and BEST: (i) stable PP values, (ii) small and stable average standard deviation of the split frequencies of independent runs, (iii) potential scale reduction factor close to 1 and (iv) an effective sample size of at least 200 for the posterior probabilities. The conditions (i) and (iv) were evaluated also for Phycas, and additionally the split PP plot and split sojourn plot were examined. Samples prior to reaching stationarity were discarded as ‘burnin’. The conditions (i) and (iv) were evaluated in Tracer (ver. 1.5.0) (Rambaut and Drummond 2009). Average standard deviation of mean sample-wide concordance factor (CF) was examined for BUCKy, and the CF was used to determine clade support.

**Results**

**Combined evidence from all genes**

Three methods, GARLI, MrBayes and Phycas, recovered the clade *echinata–elliottii* for the concatenated matrix of 11 genes. Support from the ML method was much lower (highest BS $= 65\%$, AIC partitioning) than that from the BI methods (lowest PP $= 0.98$ in MrBayes, BIC partitioning). The BI methods additionally supported the clade *echinata–elliottii–palustris*, but support varied considerably (from PP $< 0.50$ in Phycas, AIC, to PP $= 0.94$ in MrBayes, BIC). Significant differences between the ML and BI methods in support for the clade *echinata–elliottii*, and in the case of the clade *echinata–elliottii–palustris*, also between Phycas and MrBayes (Fig. 2) are characteristic signatures of true or approximate star phylogenies (Suzuki et al. 2002; Lewis et al. 2005).

BEST supported the clade *echinata–taeda*, absent in the gene-tree methods, with PP $= 0.53$ (AIC and BIC; Fig. 3). This clade was present also in the primary concordance tree in BUCKy (CF ranging from 0.35 to 0.44, depending on $\alpha$ and model selection criterion). Another clade present in the BUCKy’s primary concordance tree was *elliottii–palustris* (CF ranging from 0.24 to 0.44, depending on $\alpha$ and model selection criterion).

**Individual gene approach**

Given the results from the analysis of the combined dataset resulting in mostly unresolved relationships among the studied species, and the conflicts found between

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**Figure 2.** Joint analysis of 11 genes in *P. echinata*, *P. elliottii*, *P. palustris* and *P. taeda* using GARLI (A), MrBayes (B) and Phycas (C). Numbers at nodes correspond to BS for GARLI (%; AIC/BIC), and PP for MrBayes and Phycas (AIC/BIC). Branch lengths are shown for the BIC partitioning schemes.
the gene-tree and species-tree methods, we looked at each gene separately using ML (GARLI) and BI (MrBayes). The two approaches produced consistent results for each gene, although the difference in clade support varied among the genes [see Supporting Information—Fig. S1]. We, therefore, clustered genes into three groups following similarity in topologies. The genes 4cl, c4h-2 and cesA3 (Group A) supported the clade echinata–taeda, the genes agp-4 and sod-chl (Group B) supported the clade echinata–elliottii and the genes dhn-2 and erd3 (Group C) supported the clade echinata–palustris. Group C genes disagreed in the placement of the species P. elliottii and P. taeda; however, we decided to focus on the similarity. Additional analysis in the R environment (ver. 2.15.2) (R Core Team 2012) supported these groupings—principal coordinate analysis (function pcoa, package ape ver. 3.0-11) (Paradis et al. 2004) run on Robinson–Foulds distance matrix for the individual gene trees (function RF.dist, package phangorn ver. 1.99-7) (Schliep 2011) placed the three groups of genes in distinct clusters [see Supporting Information—Fig. S2]. Each set was partitioned by-gene-site, and analysed using GARLI and MrBayes. In general, the BS for the clades jointly supported by genes within each group increased, the PP reached (or stayed at) 1.00 and the level of support for these clades became consistent between GARLI and MrBayes (Fig. 4) [see Supporting Information—Fig. S3]. In the case of Group B, the clade palustris–taeda, previously present only in the agp-4 gene tree, was also recovered, although with low support. In Group C, P. elliottii and P. taeda were placed as sister taxa in a clade with low BS and high PP, which reflects the conflict between the two individual gene trees.

Discussion

The discordance among various approaches applied in this study mirrors the conflicting results from previously published work (see Introduction). Comparison of the topologies supported by Group A vs. Group B vs. Group C genes shows highly consistent results within each group across different methods, yet the conclusions contradict each other among the groups. The clade echinata–taeda, supported by Group A genes, was previously recovered by Dvorak et al. (2000) using RAPD markers and the neighbour-joining method with BS = 0.90. The species in the clade palustris–taeda, supported by the gene agp-4 (Group B), were placed in one clade by Adams and Jackson (1997) based on parsimony analysis of morphological characters, and by Grotkopp et al. (2004) through a supertree approach (57% of individual trees agreed with the supertree at the node). Both clades echinata–palustris and elliottii–taeda supported by the Group C genes corresponded to grouping in Eckert and Hall (2006), who used chloroplast data.

Given life history of the subsection (see Introduction), long-distance gene flow through pollen, hybridization events that still occur today and large (but variable in the past) population sizes of the investigated species, incomplete lineage sorting could be the simplest and most straightforward explanation of the pattern observed in our study. It is a known phenomenon in pines (Syring et al. 2007; Willyard et al. 2009), and it is likely to confound phylogenetic inference in a clade as recent as Australes. Interestingly, however, the genes forming the three groups

![Figure 3](image-url) Joint analysis of 11 genes in P. echinata, P. elliottii, P. palustris and P. taeda using BEST (A) and BUCKy (cladogram; B). Numbers at nodes correspond to PP for BEST (AIC/BIC) and CFs for BUCKy (AIC/BIC; range of CFs for α values from 10 000 to 0.01).

![Figure 4](image-url) Analysis of three groups of genes: Group A (4cl, c4h-2 and cesA3; A), Group B (agp-4 and sod-chl; B) and Group C (dhn-2 and erd3; C) in P. echinata, P. elliottii, P. palustris and P. taeda using GARLI and MrBayes. Cladograms are shown. Numbers at nodes correspond to clade support: GARLI (AIC/BIC; top row) and MrBayes (AIC/BIC; bottom row). The BS for the clade palustris–taeda in Group B was < 0.50.
bound clades could have resulted from a transfer of functionally shared ancestral polymorphisms, two other hypotheses are also connected at the putative functional level (Table 2). The Group A genes all directly affect wood properties: 4cl and c4h-2 are both part of the lignin biosynthesis pathway (Whetten and Sederoff 1995; Boerjan et al. 2003), and cesA3 is involved in cellulose synthesis (Pot et al. 2005). Another gene in our dataset, comt-2, is also involved in lignin biosynthesis (Whetten and Sederoff 1995; Boerjan et al. 2003) but did not recover the echinata–taeda clade. Unlike the other three genes in Group A, it supported the clade elliottii–palustris. An additional joint analysis of all four genes, using the same methodology, showed that the support for the previously recovered clade echinata–taeda almost did not change, while the clade elliottii–palustris received a low-to-moderate support [see Supporting Information—Fig. S4]. The latter two species were previously placed in one clade (BS ≤ 0.50) by Hernández-León et al. (2013) based on an ML method applied to plastid data. Conversely, the Group B and C genes are not involved in the lignin biosynthesis pathway directly. The Group B genes (agp-4 and sod-chl) appear to contribute to plant defence, water management and other functions. Proteins from the arabinogalactan family play signalling and protective roles, and participate in xylem development, cell growth and expansion, programmed cell death and other processes (Loopstra and Sederoff 1995; Loopstra et al. 2000; Zhang et al. 2000, 2003; Showalter 2001; Seifert and Roberts 2007). Sod-chl has antioxidative properties (Huttunen and Heiska 1988; Bowler et al. 1992). It was also drought responsive (Costa et al. 1998) and hence was considered a drought-tolerance candidate gene in loblolly pine (González-Martínez et al. 2006). Dhn-2 and erd3 (Group C) are also associated with water stress recognition and response. Dhn-2, a dehydrin, is responsive to dehydration stress and also plays other protective functions (González-Martínez et al. 2006; Eveno et al. 2008). The role of erd3 is less known, but it has been found active in an early stage of dehydration stress (González-Martínez et al. 2006; Eveno et al. 2008).

Regardless of the extent of the potential effects of shared ancestral polymorphisms, two other hypotheses of parallel evolution within the functional gene groups could be pursued in follow-up studies. The functionally bound clades could have resulted from a transfer of adaptive alleles via introgressive hybridization from an adapted pine donor, followed by positive selection and subsequent purifying or balancing selection acting in both donor and acceptor populations. Alternatively, speciation could have begun while the locus was already under purifying or balancing selection, which continued working simultaneously in both populations in parallel, given common habitat locations and environmental pressures. Sympathy, shifting locale of the optimum habitat, population size changes and the recent evolutionary origin of the southern pines could have facilitated hybridization, while crucial roles of the genes belonging to the three groups could have resulted in preservation of adaptive variants, especially under similar environmental selection pressures. A tight physical linkage among the loci within each group can be excluded as an alternative possible explanation for the observed phenomenon. Most of the genes studied here are mapped to different linkage groups in P. taeda, or far from each other, although no linkage data were found for dhn-2 and erd3 [see Supporting Information—Table S2]. These two hypotheses require more assumptions to explain the observed pattern when compared with the hypothesis of incomplete lineage sorting, and therefore, the latter should be preferred, although some form of interplay of all three is certainly possible.

In order to reject the hypothesis of incomplete lineage sorting, and to pursue an alternative, limitations of our study need to be overcome. Multiple individuals sampled for each species would allow for thorough identification of shared ancestral polymorphisms and variation fixed at the species level. Including more genes per functional group would allow to examine whether the observed pattern holds also for other members of a given pathway or process, or if it is purely stochastic. Samples from all members of the clade Australes would likely help improve overall robustness of the phylogenetic inference. Additionally, to test the monophyly of the clade Australes in the traditional sense (Little and Critchfield 1969), which has been questioned by cpDNA-based studies (Gernandt et al. 2005; Eckert and Hall 2006; Hernández-León et al. 2013), species traditionally classified as Attenuatae, Leiophyllae and Oocarpae should be included in the

| Group | Genes                          | Putative function               | N    | S    |
|-------|--------------------------------|---------------------------------|------|------|
| A     | 4cl, c4h-2 and cesA3           | Wood properties                 | 2    | 10   |
| B     | agp-4 and sod-chl              | Plant defence, water management | 4    | 10   |
| C     | dhn-2 and erd3                 | Water stress recognition and response | 2    | 11   |

### Table 2. Groups of genes based on phylogenetic analysis of P. echinata, P. elliottii, P. palustris and P. taeda, their selected putative functions and numbers of nonsynonymous (N) and silent (S) nucleotide substitutions in each group.
nuclear marker-based study. Alternative outgroup species could also be considered, especially those from subsections Contortae and Ponderosae that are more closely related to the ingroup than P. pinaster. Solving these caveats, however, requires additional resources and cannot be done purely analytically. Given the limitations of the data, our primary intention was to apply a spectrum of methods to a gene sample that would maximize genome coverage.

Recent estimates of radiation within Australes suggested that it could have begun as recently as 7 MYA (P. taeda–P. radiata split), although the split between the ancestors of Ponderosae and Australes might have happened as early as 15 MYA (Willyard et al. 2007). This timeframe overlaps with the mid- to late-Miocene (about 14–15 MYA), starting at or directly following the middle Miocene climate transition (MMCT), a period of cooling and ice-sheet expansion that took place about 14 MYA (Shevenell et al. 2004). Pines experienced habitat locale shifts both before and after the MMCT, for example, during the Eocene (56–34 MYA), interpreted as the major stimulus for pine divergence at the time, and during the Pleistocene (2.6–0.01 MYA) (Millar 1993). The MMCT likely affected population sizes, species range and distribution of the allelic variation, and probably had the momentum to trigger radiation within Australes. The potential for hybridization events resulting in introgression was likely greater back in time, especially when the ancestral species were far less diverged and stressed by recurrent changes in environmental pressures and by range shifts. Range expansions could have then brought multiple genetic effects (Excoffier et al. 2009) including increase in frequency of rare (and also newly introduced) variants. This process would have happened much faster if the newly acquired alleles were advantageous.

The adaptations shared among the southern pines and shaped by the vibrant historic climate are particularly interesting in the light of the ongoing and forthcoming climate changes, amidst the discussion on assisted migration (Vitt et al. 2010; Krutovsky et al. 2012; Koralewski et al. 2015). The historical events might have led to increased standing genetic variation in these species, directly influencing their level of adaptability and making them somewhat ‘climate-change ready’. Additional inquiries directed towards the loci studied here could improve breeding strategies in the face of climate change (Krutovsky et al. 2013).

Conclusions

Incomplete lineage sorting, introgression and parallel evolution can explain inconsistencies observed in the phylogenetic analysis of the four southern pines. However, more data are needed to discriminate among these hypotheses. The conflicting signals were vigorously tested, but evidence in the current data was not robust enough to support potent claims, and thus, the simplest hypothesis of incomplete lineage sorting may be preferred, while the alternatives may be pursued in future studies. To overcome limitations of our study, additional sampling should include multiple individuals per species, additional species that form one clade with the four pines investigated here, less distant outgroup species and additional functionally related genes. Our work provided new insights into the Australes phylogeny, but their evolutionary history remains elusive.

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Contributions by the Authors

T.E.K., M.M. and K.V.K. conceived and designed the study. T.E.K. arranged the data and ran the software. T.E.K., M.M. and K.V.K. interpreted results. T.E.K. drafted the manuscript. T.E.K., M.M. and K.V.K. edited, read and approved the final version of the manuscript.

Conflict of Interest Statement

None declared.

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Supporting Information

The following additional information is available in the online version of this article —

Dataset S1. Multiple alignment in NEXUS format.

Figure S1. Separate analysis of each gene for the four Australes species: *P. echinata*, *P. elliottii*, *P. palustris* and *P. taeda*. Cladograms are shown. From top to bottom: 4cl, *c4h-2* and *cesA3* (A); *agp-4* and *sod-chl* (B); *dnh-2* and *erd3* (C); *cad*, *glyhmt*, *pp2c* and *comt-2* (D). Numbers at nodes correspond to clade support: GARLI (AIC/BIC; top row) and MrBayes (AIC/BIC; top row).

Figure S2. Principal coordinate analysis run on Robinson–Foulds distance matrix for the individual gene trees. Group A gene names (*4cl*, *c4h-2* and *cesA3*) are in green, Group B gene names (*agp-4* and *sod-chl*) are in orange and Group C gene names (*dnh-2* and *erd3*) are in blue.

Figure S3. Joint analysis of the genes from Group A (*4cl*, *c4h-2* and *cesA3*) and *comt-2* in *P. echinata*, *P. elliottii*, *P. palustris* and *P. taeda* using GARLI and MrBayes. From top to bottom: GARLI (AIC), GARLI (BIC), MrBayes (AIC) and MrBayes (BIC). Numbers at nodes correspond to clade support.

Figure S4. Joint analysis of the genes from Group A (*4cl*, *c4h-2* and *cesA3*) and *comt-2* in *P. echinata*, *P. elliottii*, *P. palustris* and *P. taeda* using GARLI and MrBayes. From top to bottom: GARLI (AIC), GARLI (BIC), MrBayes (AIC) and MrBayes (BIC). Numbers at nodes correspond to clade support.

Table S1. Partitioning schemes and software settings.

Table S2. Linkage information for the studied genes.

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