Abstract.—The placement of angiosperms and Gnetales in seed plant phylogeny remains one of the most enigmatic problems in plant evolution, with morphological analyses (which have usually included fossils) and molecular analyses pointing to very distinct topologies. Almost all morphology-based phylogenies group angiosperms with Gnetales and certain extinct seed plant lineages, while most molecular phylogenies link Gnetales with conifers. In this study, we investigate the phylogenetic signal present in published seed plant morphological data sets. We use parsimony, Bayesian inference, and maximum-likelihood approaches, combined with a number of experiments with the data, to address the morphological–molecular conflict. First, we ask whether the lack of association of Gnetales with conifers in morphological analyses is due to an absence of signal or to the presence of competing signals, and second, we compare the performance of parsimony and model-based approaches with morphological data sets. Our results imply that the grouping of Gnetales and angiosperms is largely the result of long-branch attraction (LBA), consistent across a range of methodological approaches. Thus, there is a signal for the grouping of Gnetales and angiosperms in morphological matrices, but it was swamped by convergence between angiosperms and Gnetales, both situated on long branches. However, this effect becomes weaker in more recent analyses, as a result of addition and critical reassessment of characters. Even when a clade including angiosperms and Gnetales is still weakly supported by parsimony, model-based approaches favor a clade of Gnetales and conifers, presumably because they are more resistant to LBA. Inclusion of fossil taxa weakens rather than strengthens support for a relationship of angiosperms and Gnetales. Our analyses finally reconcile morphology with molecules in favoring a relationship of Gnetales to conifers, and show that morphology may therefore be useful in reconstructing other aspects of the phylogenetic history of the seed plants.

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Introduction

The use of morphology as a source of data for reconstructing phylogenetic relationships has lost most of its ground since the advent of molecular phylogenetics, except in paleontology. However, there has recently been renewed interest in morphological phylogenetics (Lee and Palci 2015; Pyron 2015), partly because of increased focus on the phylogenetic placement of fossil taxa in trees of living organisms, stimulated by the necessity of accurate calibrations for dating the molecular trees that have become the main basis for comparative evolutionary studies. This has led to the development of methods that integrate phylogenetic placement of fossils in the dating process (Pyron 2011; Ronquist et al. 2012; Zhang et al. 2016). Another focus has been the application of statistical phylogenetics to morphological data on both a theoretical (Wright et al. 2014, 2015; O’Reilly et al. 2016) and an empirical level (Lee and Worthy 2012; Godefroit et al. 2013; Cau et al. 2015). In paleontology, for which only morphological data are available (except in the recent past), questions on the role of morphology in phylogenetics are even more critical. A major issue concerns the value of fossils in reconstructing
relationships among living organisms. Early in the history of phylogenetics, there were claims that fossils are incapable of overturning phylogenetic relationships inferred from living taxa (Patterson 1981), but also demonstrations that they can, for instance, in morphological analysis of amniote phylogeny (Gauthier et al. 1988). Whether or not fossils affect the inferred topology of living taxa, there is little doubt that they are often either useful or necessary in elucidating the homologies of novel structures (e.g., the seed plant ovule and eustele) and the order of origin of the morphological synapomorphies of extant (crown) groups (e.g., origin of secondary growth before the ovule in the seed plant line), as discussed in Doyle (2013). This is critical, because major groups, such as the now-dominant angiosperms (flowering plants), are often separated from their closest living relatives by major morphological gaps (numbers of character changes), even if the incorporation of fossils does not affect inferred relationships among living taxa (Doyle and Donoghue 1987; Donoghue et al. 1989).

Many phylogenies based on morphology have been recently published for important groups with both living and fossil representatives, including mammals (O’Leary et al. 2013), squamate reptiles (Gauthier et al. 2012), arthropods (Legg et al. 2013), and the genus Homo (Dembo et al. 2016). However, the validity and use of morphological data in reconstructing phylogeny have been severely criticized, notably by Scotland et al. (2003), based on supposed diminishing returns in the discovery of new morphological characters and the prevalence of functional convergence. The painstaking acquisition of morphological characters, which requires a relatively large amount of training and time, could turn out to be systematically worthless if the phylogenetic signal present in these data is either insufficient or misleading. Indeed, the number of characters that can be coded for morphological data sets represents a major limit to the use of morphology and its integration with molecular data, especially in the age of phylogenomics, when the ever-increasing amount of molecular signal could simply “swamp” the weak signal present in morphological data sets (Doyle and Endress 2000; Bateman et al. 2006). Morphological data may also be afflicted to a higher degree than molecules by functional convergence and parallelism (Givnish and Sytsma 1997), which could lead a morphological data set to infer a wrong phylogenetic tree. Even though the confounding effect of convergence has been formally tested in only a few studies (Wiens et al. 2003), it seems to be at the base of one of the deepest cases of conflict between molecules and morphology in the reconstruction of evolutionary history, namely, the phylogeny of placental mammals (Foley et al. 2016). In this case, the strong effect of selection on general morphology caused by similar lifestyle seems to hinder attempts to use morphology to reconstruct phylogenetic history in this group (Springer et al. 2007), and it affects even large “phenomic” data sets (Springer et al. 2013).

Another example of conflict between morphology and molecular data involves the relationships among seed plants, particularly angiosperms and the highly derived living seed plant order Gnetales. Before the advent of cladistics, some authors proposed that angiosperms and Gnetales were closest living relatives, while others argued that these two groups were strictly convergent and Gnetales were instead related to conifers (for a review, see Doyle and Donoghue 1986). However, since the earliest studies by Parenti (1980) and Hill and Crane (1982), which included only living taxa, the view that angiosperms are most closely related to Gnetales has appeared to be one of the most stable results of morphologically based parsimony analyses of seed plant phylogeny (Crane 1985a; Doyle and Donoghue 1986, 1992; Nixon et al. 1994; Rothwell and Serbet 1994; Doyle 1996, 2006; Hill and Bateman 2006; Friis et al. 2007; Rothwell et al. 2009; Rothwell and Stockey 2016; Fig. 1). The first analysis that included fossils (Crane 1985a) associated angiosperms and Gnetales with Mesozoic Bennettitales and Pentoxylon. Because all four taxa have more or less flower-like reproductive structures, this clade became known as the anthophytes, a term formerly used for angiosperms, to emphasize its implication that the flower was a synapomorphy not of angiosperms alone but rather of a larger clade to which they belong (Crane 1985b; Doyle and Donoghue 1986). Some subsequent
analyses interpolated the Mesozoic fossil *Caytonia* into this clade as the closest outgroup of angiosperms (Doyle 1996, 2006, 2008; Hilton and Bateman 2006; Friis et al. 2007). This result calls into question the original concept of anthophytes as a clade united by flowers, because *Caytonia* had large sporophylls that are unlikely to have been grouped into flower-like structures. However, all trees found in morphological analyses, with the exception of some in Doyle (2008), have agreed that Gnetales are the closest living relatives of angiosperms. Some analyses associated the clade including angiosperms and Gnetales with “Mesozoic ferns” (such as glossopterids, coryystperms, and *Caytonia*), others with “coniferophytes” (conifers, *Ginkgo*, and fossil cordaites). Inferred relationships within the clade have also varied: in some cases angiosperms and Gnetales are sister groups, in others Gnetales are linked with Bennettitales. Because some studies place taxa without flower-like structures in the clade, and molecular data are unable to distinguish such trees from anthophyte trees in the original sense, we refer to the whole class of trees in which angiosperms and Gnetales are closest living relatives as “gnetangiosperm” rather than “anthophyte” trees.

By contrast, since the advent of molecular phylogenetics, the hypothesis that angiosperms and Gnetales are closely related has lost most of its support among plant biologists. Although molecular analyses cannot directly evaluate the status of putatively related fossil taxa, they can address the relationship of angiosperms and Gnetales. Molecular data from different genomes analyzed with different approaches do not yield a Gnetales plus angiosperm clade, with the exception of few maximum parsimony (MP) and neighbor-joining analyses of nuclear ribosomal RNA or DNA (Hamby and Zimmer 1992; Stefanovic et al. 1998; Rydin et al. 2002) and one MP analysis of *rbcL* (Rydin and Källersjö 2002). The majority of molecular analyses retrieve a clade of Gnetales plus Pinaceae (Bowe et al. 2000; Chaw et al. 2000; Gugerli et al. 2001; Qiu et al. 2007; Zhong et al. 2011), conifers other than Pinaceae (cupressophytes) (Nickrent et al. 2000; Rydin and Källersjö 2002), or conifers as a whole (Wickett et al. 2014), which we refer to collectively as “gneconifer” trees. In most of these trees, angiosperms are the sister group of all other living seed plants (acrogymnosperms [Cantino et al. 2007]). The main exceptions are “Gnetales-basal” trees, in which Gnetales are sister to all other living seed plants (e.g., Albert et al. 1994; Rydin and Källersjö 2002).

Several potential issues have been identified with both sorts of data. Regarding molecules, these include limited taxonomic sampling resulting from extinction of the majority of seed plant lineages (Rothwell et al. 2009), loss of phylogenetic signal due to saturation (particularly at third codon positions), strong rate heterogeneity among sites across lineages and conflict between gene trees (Mathews 2009), composition biases among synonymous substitutions (Cox et al. 2014), as well as systematic errors and biases (Sanderson et al. 2000; Magallón and Sanderson 2002; Burleigh and Mathews 2007; Zhong et al. 2011), leading to a plethora of conflicting signals. In analyzing data sets that yielded Gnetales-basal trees, studies that have attempted to correct for these biases have generally favored trees
in which Gnetales are associated with conifers (Sanderson et al. 2000; Magallón and Sanderson 2002; Burleigh and Mathews 2007). Regarding morphology, in addition to far more complex problems in definition of characters and the role of functional convergence in confounding relationships, it has been shown that different taxon sampling strategies (which can also cause problems in molecular studies [Rydin and Källersjö 2002]), such as choice of the closest progymnosperm outgroup of seed plants (Hilton and Bateman 2006), can lead to different results concerning the rooting of the seed plants.

The conflict between molecules and morphology has led to different attitudes toward morphological data within the botanical community (Donoghue and Doyle 2000; Scotland et al. 2003; Bateman et al. 2006; Rothwell et al. 2009). Following suggestions of Donoghue and Doyle (2000), Doyle (2006, 2008) reconsidered several supposed homologies between angiosperms and Gnetales in the light of the molecular results. These studies and the analysis of Hilton and Bateman (2006) also incorporated newly recognized similarities between Gnetales and conifers, for example, in wood anatomy (Carlquist 1996), as well as new evidence on the morphology of the seed-bearing cupules in fossil taxa. Other changes involved redefinition of characters to reduce potential biases. For example, when building a morphological matrix, dissecting a character into more character states may represent an improvement by distinguishing convergent states and avoiding bias toward particular phylogenetic hypotheses during primary homology assessment (Jenner 2004; Zou and Zhang 2016), although it may be disadvantageous, because it leads to a lack of resolution when the number of states becomes excessive. In seed plants, there are many special factors that complicate character coding. Among living taxa, the assessment of homology is complicated by the plastic and modular nature of plant development (Mathews and Kramer 2012). Among fossil taxa, the mode of preservation of many key fossils has critical consequences for the amount of data available. This affects not only the number of missing characters, but also the process of primary homology assessment and character coding. Although these issues with coding are most severe in fossils preserved as compressions, such as Caytonia (Doyle 2008; Rothwell et al. 2009) and Archaefructus (Sun et al. 2002; Friis et al. 2003; Doyle 2008; Rudall and Bateman 2010; Endress and Doyle 2009; Doyle and Endress 2014), even fossil groups that are exquisitely preserved as permineralizations (e.g., Bennettitales) are not immune to conflicting interpretations (Friis et al. 2007; Rothwell et al. 2009; Crepet and Stevenson 2010; Doyle 2012: Supplemental Material; Rothwell and Stockey 2013; Pott 2016).

Despite careful reconsideration of potentially convergent traits between Gnetales and angiosperms, the conflict between morphological and molecular data appeared to persist, with most morphological parsimony analyses continuing to favor the gnetangiosperm hypothesis (Doyle 2006; Hilton and Bateman 2006; Rothwell et al. 2009). The possibility that morphological data are inadequate to resolve such a key aspect of the phylogeny of seed plants would represent a severe hindrance in understanding plant evolution, especially in the light of the small number of extant lineages that survived extinction during the Paleozoic and Mesozoic (Mathews 2009) and the great morphological gaps among the surviving lineages. However, there have been signs that the conflicts with molecular data are weakening: in the analysis of Doyle (2006), trees in which Gnetales were nested in conifers were only one step less parsimonious than gnetangiosperm trees, and in Doyle (2008), trees of the two types became equally parsimonious.

In this study, we attempt to elucidate the phylogenetic signal present in published morphological data sets of the seed plants, concentrating on the relationship of angiosperms and Gnetales. This is not the only aspect of seed plant phylogeny that varies among and between morphological and molecular analyses. Another case is whether ginkgophytes (now reduced to Ginkgo biloba) are related to conifers and cordaites, as part of a coniferophyte clade, or to cycads, as found in some molecular analyses. However, the question of angiosperms and Gnetales is probably of the broadest evolutionary interest and is especially likely to illustrate the general problem of
long-branch effects in highly derived groups. We first explore whether the possibility of convergence between angiosperms and Gnetales represents a major problem by reanalyzing the matrices that incorporated earlier homology assumptions concerning characters of the two groups (i.e., the matrices compiled before the incoming of molecular results) and later matrices that revised such assumptions (the matrices of Doyle [2006] and Hilton and Bateman [2006], and data sets derived from them) and testing whether the signal and the relative support for the gnetangiosperm and gneconifer clades changed between these two sets of matrices. After revealing a more coherent signal supporting a gneconifer clade in the more recent matrices, we investigate whether the retrieval of a gnetangiosperm topology by parsimony analyses was at least partly due to methodological biases that could be overcome by using model-based methods. Hopefully these approaches may be useful in resolving cases of conflict between morphological and molecular data in other taxa, particularly those with significant fossil representatives.

Materials and Methods

Matrices.—The matrices of Crane (1985a: version 2, in which Bennettitales and Pentoxylon were scored as having cupules potentially homologous with those of Mesozoic seed ferns), Doyle and Donoghue (1986, 1992), Nixon et al. (1994), Rothwell and Serbet (1994), and Doyle (1996, 2006, 2008) were manually coded from the respective articles. The Hilton and Bateman (2006) matrix was kindly provided by Richard Bateman. The matrices from analysis 3 of Rothwell et al. (2009) and from Rothwell and Stockey (2016) were downloaded from the supplementary materials of the respective articles. The Hilton and Bateman (2006) matrix was kindly provided by Richard Bateman. The matrices from analysis 3 of Rothwell et al. (2009) and from Rothwell and Stockey (2016) were downloaded from the supplementary materials of the respective articles.

Parsimony Analyses.—We performed parsimony analyses of all matrices with PAUP v. 4.0a136 (Swofford 2003), using the heuristic search algorithm with random addition of taxa and 1000 replicates. Bootstrap analyses were conducted with 10,000 replicates, using the “asis” addition option and keeping one tree per replicate (Müller 2005).

We also conducted analyses with a topological backbone constraint, forcing the Gnetales into a clade with the extant conifers and leaving the position of other living taxa and fossils unconstrained. Significant differences between the constrained and unconstrained topologies were evaluated using the Templeton test (Templeton 1983) as implemented in PAUP v. 4.0a136 (Swofford 2003). We investigated the effects of recoding characters by Doyle (2006, 2008) in more detail by using MacClade (Maddison and Maddison 2003) to compare the number of steps in each character on trees with Gnetales associated with angiosperms or with conifers.

Model-based Analyses.—Our model-based analyses were all conducted using the Markov k-states (Mk) model (Lewis 2001). This model assumes that characters are in one of k states, are all independent of each other, and change stochastically along branches with equal rates for all possible transitions, with all changes being independent of one another (as a Markov process). Some of these assumptions have been criticized for being unrealistic when applied to morphological change (Lewis 2001; Wright et al. 2014). For example, the model is fully symmetrical; that is, the probability of change from 0 to 1 is equal to the probability of change from 1 to 0, an assumption that is violated by Dollo characters (i.e., losses of complex structures that are unlikely to be regained). Even though some of these assumptions can be theoretically relaxed, and implementations of these relaxed models already exist in a Bayesian framework (Wright et al. 2014), we used the standard version of the model to simplify the analyses and allow a closer comparison with the maximum-likelihood (ML) implementation.

Maximum Likelihood.—ML analyses were conducted using RaxML v. 8.2.10 (Stamatakis 2014). Matrices were modified by recoding all ambiguities (e.g., 0/1 in a three-state character) as missing data, because the method cannot cope with ambiguous characters. Topology is inferred using branch lengths, which are estimated as the expected number of state changes per character on that particular branch. We conducted 1000 bootstrap replicates with a gamma-distributed rate
variation, which models different rates across characters by employing a multiplier drawn from a discretized gamma distribution.

**Bayesian Inference (BI).**—Bayesian analyses relied on MrBayes v. 3.2.3 (Ronquist et al. 2012) under the Mk model. For each matrix, we conducted two analyses, one with an equal rate of evolution among characters and another with gamma-distributed rate variation. In both cases, we used the Mkprinf correction for parsimony informative characters. The analyses were run for 5,000,000 generations, sampling every 1000th generation. The first 10,000 runs were discarded as burn-in. Posterior traces were inspected using Tracer (Rambaut and Drummond 2007).

**Model Testing and Rate Variation.**—We also conducted stepping-stone analyses (Xie et al. 2011; Ronquist et al. 2012) to evaluate the most appropriate model of rate variation among characters (equal rates vs. gamma-distributed rates). These analyses allow us to estimate the marginal likelihood for different models with better accuracy than other measures (e.g., harmonic mean estimator). We used four independent runs with two chains with the default MrBayes parameters, run for 5,000,000 generations and sampling every 1000th generation. Using the marginal likelihoods from the stepping-stone analysis, we then calculated the support for the two models using Bayes factors (BFs) (Kass and Raftery 1995).

**Exploring Conflict in the Data.**—To explore phylogenetic conflict in the data, we employed the software SplitsTree4 (Huson and Bryant 2006). We used this program to visualize conflicts among the bootstrap replicates from the MP and ML analysis and among the posterior tree samples found with Bayesian inference. The software summarizes the sets of trees using split networks, which allows us to visualize all possible conflicting hypotheses. These diagrams should not be confused with networks derived from distance-based neighbor-joining analyses. A consensus network (Holland et al. 2004) was built using the “count” option, with the cutoff for visualizing the splits set at 0.05.

**Long-Branch Attraction (LBA) Tests.**—We modified the matrices to perform tests for LBA, following the suggestions of Bergsten (2005). Two matrices were created to test the potentially destabilizing effect of the two long-branched groups suspected to create this artifact, angiosperms and Gnetales, by alternately removing each of them (long-branch extraction analysis [LBE]). If the association of angiosperms and Gnetales is indeed a result of LBA, then the removal of one of them should significantly alter the placement of the other. To test further the hypothesis of an LBA artifact exerted by angiosperms, we followed a similar approach to the sampling experiment in Rota-Stabelli et al. (2010): another matrix was created to elongate the branch subtending angiosperms by removing the three fossil taxa most commonly identified as angiosperm outgroups (*Pentoxylon*, Bennettitales, and *Caytonia*) (branch elongation [BE] analysis). In the presence of an LBA artifact, the support for the node including the two long branches (angiosperms and Gnetales) should increase with such an “elongation” of one of the two branches. To test the effect of including fossil data in the matrices, we created a set of matrices in which all fossil taxa were removed (extant experiment [EX]). Because this should lead to elongation of the branches subtending the living groups, this situation should result in the worst possible condition for long-branch artifacts and thus lead to the strongest apparent support for the node including the two long branches.

**Morphospace Analysis.**—To visualize morphological patterns in the different matrices, we conducted principal coordinates (PCO) analyses. We employed the maximum observed rescaled distance between all pairs of taxa to generate the ordination as obtained using the `MorphDistMatrix` function of the R package ‘Claddis’ (Lloyd 2016). PCO analysis was conducted using the `cmdscale` function from the ‘stats’ package (R Core Team 2017). The taxa were then plotted on the first two PCO axes.

**Results**

Our reanalyses of the historical morphological matrices of seed plants with parsimony resulted in trees identical to the published trees
The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1). The MP trees and the consensus trees always show a gnetangiosperm clade (with or without *Caytonia*), with the exception of trees based on the Doyle (2008) matrix, in which gnetangiosperm and gneconifer topologies are equally parsimonious. Constraining Gnetales and conifers to form a clade always results in trees longer than the most parsimonious trees, except with the Doyle (2008) matrix (Table 1).
the gnetangiosperms to a clade of Gnetales and conifers. This is illustrated by a split network consensus based on the Rothwell and Stockey (2016) matrix (Fig. 3C), in which Gnetales are linked with conifers, and Glossopteris, Caytonia, and Petriellaea (a Triassic fossil not included in earlier analyses that is now better known vegetatively thanks to work of Bomfleur et al. [2014]) are the closest outgroups of angiosperms.

Our first test of the hypothesis that the gnetangiosperm topology is the result of LBA consists of LBE experiments (Fig. 4A, B). These involved separate removal of the two potential long-branch taxa: angiosperms and Gnetales. The removal of the angiosperms has different effects on the pre- and post-2000 matrices. With the Crane (1985a) version 2 matrix analyzed here, a topology with Bennettitales, Pentoxylon, and the Gnetales diverging after Lyginopteris and before the other taxa becomes as parsimonious as the topology with the gnetangiosperms nested among Mesozoic seed ferns that was retrieved with the full matrix. The new tree corresponds to the most parsimonious tree that Crane (1985a) found with his version 1 matrix, which differed in that Bennettitales and Pentoxylon were scored as not having cupules potentially homologous with those of Mesozoic seed ferns. With the Doyle and Donoghue (1986) matrix, Bennettitales, Pentoxylon, and Gnetales are nested within coniferophytes. With the Doyle and Donoghue (1992) and Rothwell and Serbet (1994) matrices, the consensus tree is identical.
to the trimmed consensus derived from the full matrix. With the Nixon et al. (1994) matrix, Cordaites and Ginkgo are successive outgroups to a conifer plus gnetangiosperm clade, whereas with the full matrix, they are equally parsimoniously placed as successive outgroups to the conifers in a clade that is sister to gnetangiosperms. The inverse happens with the Doyle (1996) matrix, where the position of Ginkgo and cordaites is destabilized by the removal of the angiosperms, with these taxa being either successive outgroups to extant and fossil conifers or sister to a clade composed of other former gnetangiosperms, conifers, Peltaspermum, and Autunia. The position of the Gnaetales in a truncated gnetangiosperm clade (i.e., with Bennettitales and Pentoxylon) is maintained in all matrices.

With the post-2000 matrices, the effect of removal of the angiosperms is consistent among different matrices. With the Hilton and Bateman (2006) matrix, Gnaetales are equally parsimoniously placed within the coniferophytes, within the coniferophytes as sister to the Bennettitales, or in an anthophyte clade as sister to the conifers. In the Doyle

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**Figure 3.** Split network consensus of the posterior tree sample of the MP bootstrap analysis (A), the ML bootstrap analysis (B), and the Bayesian inference (BI) analysis of the Rothwell and Stockey (2016) matrix using gamma-distributed rate variation (C). Only splits with more than 0.15 posterior probability (PP) or 15% bootstrap are shown, and support is shown only for splits with more than 0.20 PP or 20% bootstrap. The support values for the splits within the angiosperms have been removed for clarity. If the Gnaetales-conifer clade is present and supported with all three methods, other relationships (i.e., Caytonia in a clade with angiosperms) are only supported in the BI analysis.
(2006, 2008) data sets, the resulting trees see the Gnetales nested within the coniferophytes, with or without Bennettitales. With the Rothwell et al. (2009) matrix (Fig. 4D–F), a topology with a clade of Gnetales and conifers that excludes Bennettitales and Pentoxylon becomes most parsimonious (Fig. 4E). With the Rothwell and Stockey (2016) matrix, Gnetales are sister to Taxus in a coniferophyte clade that also includes Doylea, an Early Cretaceous...
cone-like structure interpreted as consisting of seed-bearing cupules (Stockey and Rothwell 2009; Rothwell and Stockey 2016). The removal of the Gnetales has no impact at all on trees based on the Crane (1985a) and Doyle and Donoghue (1986, 1992) matrices, in which the topology is identical to the trimmed topology of the consensus in the full analysis. With the Nixon et al. (1994) matrix, the removal of the Gnetales results in trees in which coniferophytes form a clade (including Ginkgo and Cordaites), that is, eliminating most parsimonious trees in which gnetangiosperms are linked with conifers. With the Rothwell and Serbet (1994) matrix, the removal of Gnetales results in a breakup of the Caytonia–Glossopteris–corysperm clade, with the angiosperms still nested within the other gnetangiosperms.

**Figure 5.** Plot of the first two principal coordinate axes for four of the matrices analyzed. The first PCO axis mainly separates the angiosperms and the other seed plants, while the second PCO axis separates more conifer-like and more fern-like groups. These plots illustrate the effect of the reassessment of gnetalean characters between the two Doyle matrices (1996, 2008) (A, B), and the similar structure of the data in the Hilton and Bateman (2006) (C) and Rothwell and Stockey (2016) (D) matrices. Arch, Archaeopteris; Aneu, Aneurophyton; Aut, Autunia; Benn, Bennettitales; Call, Callipteris; Cayt, Caytonia; Cecr, Cecropsis; Cord, Cordaitales in A and B, Cordaixylon in C and D; Cory, Corysperms; Cyc, Cycadales in A and B, Cycadaceae in C and D; Elk, Elkinsia; Emp, Emporia; Gink, Ginkgoales; Gloss, Glossopteris in A and B, Glossopterids in C and D; Heter, Heterangium; Lyg, Lyginopteris; Medu, Medullosans in A and B, Medullosa in C and D; Meso, Mesoxylon; Pelt, Peltaspermum; Pento, Pentoxylon; Petr, Petriellales; Quaes, Quaestora; Shanx, Shanxiioxylon; Tetr, Tetraxylopteris; Thuc, Thucydia; Zam, Zamiaceae.
With the Doyle (1996) matrix, the only difference lies in the placement of the corystosperms, Autunia, and Peltaspermum, which form an unresolved clade with the coniferophyte clade in the analysis without Gnetales.

With the post-2000 matrices, the removal of the Gnetales results in trees in which the remaining gnetangiosperms (which may or may not include Caytonia) form a clade outside the coniferophytes (e.g., Fig. 4F). With the Doyle (2006, 2008) matrices, a clade including Cycadales, glossopterids, and remaining gnetangiosperms (including Caytonia) is sister to a clade of Callistophyton, Peltaspermum, Autunia, and corystosperms plus coniferophytes. The analysis of the Rothwell and Stockey (2016) matrix represents an exception, in which the placement of the remaining gnetangiosperms is not affected by the removal of Gnetales. However, the removal of Doylea in addition to Gnetales results in a pattern similar to that found with the other post-2000 matrices.

In the BE experiment, in which three fossils commonly associated with angiosperms (Bennettitales, Pentoxylon, Caytonia) were removed, we observed that MP bootstrap support for the angiosperm plus Gnetales clade increases in all matrices (Fig. 4G). This effect is even stronger in the EX matrices, in which all fossil taxa were removed, where a split including angiosperms plus Gnetales is strongly supported by the MP bootstrap in all matrices.

Bayesian analysis (BI) of the BE and EX matrices shows a less linear pattern (Fig. 4H,I). In the BE analyses, the signal for the gnetangiosperms decreases with the Doyle and Donoghue (1986, 1992) matrices, reaching less than 0.5 posterior probability (PP) in the analysis with gamma-distributed rate variation. With the Nixon et al. (1994), Rothwell and Serbet (1994), and Doyle (1996) matrices, the PP of the gnetangiosperms in the BE matrices is comparable to that from the full matrices. In the post-2000 BE matrices, BI support for the gnetangiosperms is almost null with the Hilton and Bateman (2006) and Doyle (2006) matrices (<0.07 PP) and increases with the Doyle (2008) and Rothwell et al. (2009) matrices analyzed using gamma-distributed rate variation (0.55 and 0.51 PP, respectively) and with the Rothwell and Stockey (2016) matrix (0.23 PP for the equal-rate analysis, 0.37 PP for the gamma analysis). The analyses of the EX matrices all show high to moderate support (1–0.75 PP) for the split containing angiosperms plus Gnetales. With the post-2000 matrices, the use of the gamma-distributed model recovers a higher PP for the gnetangiosperms.

The morphospace analyses (Fig. 5) provide a graphic confirmation of the morphological separation of both Gnetales and angiosperms from other seed plants and the perception that Gnetales share competing morphological similarities with both angiosperms and conifers. In the morphospace generated from most of the pre-2000 matrices, Gnetales lie closer to angiosperms. With the Doyle (1996) matrix and the post-2000 matrices, the first PCO axis appears to separate angiosperm-like and non-angiosperm-like taxa, whereas the second axis seems to represent a tendency from a seed fern-like toward a conifer-like morphology. Gnetales are always placed closer to the conifers than to the angiosperms (Fig. 5). However, in all cases, Gnetales seem to have higher levels of “angiosperm-like” morphology than do conifers, represented by their rightward placement on the first PCO axis. This position on the first axis is shared by Doylea with the Rothwell and Stockey (2016) matrix. Between the analyses of the Doyle (1996, 2008) matrices (Fig. 5A,B), there is a modest shift of Gnetales away from angiosperms and toward conifers.

Discussion

The results of our analyses help to resolve some of the main issues regarding the phylogenetic signal for the gnetangiosperm clade in morphological matrices of seed plants. Our meta-analyses of published data sets (Fig. 2) show a two-step trend: first, changes in character sampling and analysis weakened support for the gnetangiosperm hypothesis; and second, the use of model-based methods shifted the balance in favor of a relationship between Gnetales and conifers, bringing the results in line with molecular data. The effect of changes in character analysis is seen in the switch in support between matrices compiled
before the main molecular analyses of seed plant phylogeny (pre-2000) and afterward: that is, Doyle (2006) and Hilton and Bateman (2006). These two matrices, which both used Doyle (1996) as a starting point but were modified independently, with only limited discussion at later stages of the two projects, and made different choices regarding character coding, taxon sampling, and splitting of higher-level taxa, show a very similar pattern. Under the MP criterion, a gnetangiosperm topology continued to be more parsimonious, but with reduced support. By contrast, ML and the Bayesian criterion positively favor a grouping of Gnetales and conifers. The matrices descended from Doyle (2006) (i.e., Doyle 2008) and from Hilton and Bateman (2006) (i.e., Rothwell et al. 2009; Rothwell and Stockey 2016) exhibit a similar pattern, except that in Doyle (2008), gnetangiosperm and gneconifer trees were equally parsimonious. This phenomenon was already reported by Mathews et al. (2010), who reanalyzed the matrix of Doyle (2008) using BI.

**Critical Character Reassessment Weakened the Conflict between Morphology and Molecules.**—Examination of the behavior of characters on gnetangiosperm and gneconifer trees illustrates how changes in character analysis made between the studies of Doyle (1996) and Doyle (2006, 2008) increased support for gneconifer trees. Some of these changes were the result of new discoveries concerning the morphology of Gnetales and other taxa, others of critical reassessment of previous character definitions aimed at reducing bias in favor of the gnetangiosperm hypothesis. The shift of Gnetales away from angiosperms and toward conifers observed in the morphospace analyses based on the data sets of Doyle (1996, 2008) (Fig. 5A,B) is presumably the result of these changes. Especially, modifications of the latter sort illustrate general problems of analysis and definition of morphological characters, which can be far more difficult than is usually acknowledged. Because potentially homologous structures in different taxa differ to various degrees, there is often a tension between use of overly lax criteria for definition of states at the stage of primary homology assessment, which may mistake homoplasy for homology, and overly strict criteria, which may overlook real synapomorphies. Other problems can be caused by inclusion of distinct characters that are correlated for functional or developmental reasons and therefore overweight single transformations, or by decisions on whether to treat presence and absence of a structure and different forms of the structure as states of the same character or as separate characters, both of which can lead to artifacts.

Most changes of the first sort involved previously overlooked conifer-like features of Gnetales. For example, Doyle (2006) added a character for presence of a torus in the pit membranes of xylem elements in conifers and Gnetales, based on observations on Gnetales by Carlquist (1996) and studies of conifers by Bauch et al. (1972). Doyle (2006) also rescored Gnetales as having a tiered proembryo, as in conifers; two tiers of cells were illustrated by Martens (1971) and called “étages,” and by Singh (1978). This similarity may have been overlooked because of other differences related to elimination of a free-nuclear phase in the embryogenesis of Gnetales (Doyle 2006). Both characters undergo one less step on gneconifer trees than on most gnetangiosperm trees (exceptions are some trees with major rearrangements elsewhere in seed plants). In male “flowers” of Ephedra and Welwitschia, microsynangia are borne in two lateral groups, which Doyle (1996) interpreted as reduced pinnate sporophylls. Because Bennettitales, Caytonia, and many “seed fern” outgroups have pinnately organized microsporophylls, this character favored a gnetangiosperm tree by one step. However, developmental studies by Mundry and Stützel (2004) indicated that the two lateral structures are more likely branches (strobili) bearing three or four simple sporophylls. Based on these observations, Doyle (2008) rescored microsporophylls in Gnetales as simple and one-veined, as in conifers, and as a result, the character favored the gneconifer topology by one or two steps.

Doyle (2006) also made changes based on improved data on a character expressing the position of the ovule or ovules on the
sporophylls or “cupules” that bear them, which is not directly relevant to Gnetales but potentially useful for identification of gnetangiosperm outgroups. Ovules are on the abaxial surface of the sporophyll/cupule in corystosperms (Axsmith et al. 2000; Klavins et al. 2002), rather than on the adaxial surface in glossopterids (Taylor and Taylor 1992), probably Caytonia, and angiosperms (if the outer integument is a modified leaf or cupule [Doyle 2006, 2008; Kelley and Gasser 2009]). Ovules are also adaxial in the cupules of Petriellaea (Taylor et al. 1994; Bomfleur et al. 2014), which was included in the analysis of Rothwell and Stockey (2016).

Other changes were the result of doubts concerning the homology of characters that supported the gnetangiosperm hypothesis, along lines suggested by Donoghue and Doyle (2000). For example, in the apical meristem character, Doyle (1996) contrasted the presence of a tunica (an outer layer that maintains its integrity by undergoing only anticlinal cell divisions, i.e., perpendicular to the surface) of Gnetales, angiosperms, and Araucariaceae with its absence in cycads, Ginkgo, and other conifers. This character undergoes two steps when Gnetales are linked with angiosperms (the state in fossils is unknown), three when Gnetales are linked with conifers. However, the tunica consists of one layer of cells in Gnetales, but two layers in angiosperms, suggesting that it may not be homologous in the two groups. To reduce bias in favor of homology of these two conditions, Doyle (2006) split presence of a tunica into two states. The resulting three-state character undergoes three steps with Gnetales in both positions. Redefinition of the megaspore membrane character involved a shift in the limit between states, from thick versus reduced (thin or absent) to present versus absent; the megaspore membrane is thin in Gnetales but absent in angiosperms, Caytonia, and probably Bennettitales. In compressions of Bennettitalean seeds prepared by oxidative maceration, Harris (1954) observed no megaspore membrane, but Wieland (1916) and Stockey and Rothwell (2003) reported a thin layer around the megagametophyte in perminalized seeds. However, as noted by Harris (1954), there is no evidence that this layer is a true megaspore membrane (i.e., consisting of exinous material). These changes in character definition do involve a subjective element and were doubtless influenced by knowledge of the molecular evidence for a relationship of Gnetales and conifers, but the new definitions represent a shift toward greater caution in evaluating the potential homology of similar but not identical structures.

The trends seen in Figure 2 show that recognition of previously overlooked similarities between Gnetales and conifers and reconsideration of potentially convergent characters between angiosperms and Gnetales succeeded in strengthening a morphological signal associating Gnetales with conifers. This result clearly contradicts the view that morphology and molecules are in strong conflict with each other (Bateman et al. 2006; Rothwell et al. 2009) and validates arguments along these same lines that were advanced by Doyle (2006, 2008) on a parsimony basis. Indeed, in all post-2000 matrices, a topology with Gnetales linked with conifers requires the addition of only a few steps to the length of gnetangiosperm trees: for example, four in the case of Hilton and Bateman (2006) and one in Doyle (2006), and both topologies became equally parsimonious in Doyle (2008). A tendency to focus on the MP consensus tree and lack of exploration of almost equally parsimonious alternatives may have tended to inflate the perceived conflict between molecules and morphology. Among analyses since 1994, bootstrap and/or decay values were reported by Doyle (1996, 2006, 2008), Hilton and Bateman (2006), and Rothwell and Stockey (2016), but not by Nixon et al. (1994), Rothwell and Serbet (1994), and Rothwell et al. (2009). Our analyses show that the signal retrieved using MP is more correctly characterized as profoundly ambiguous.

**Contribution of Model-based Methods.**—By contrast, ML and especially Bayesian analyses of all post-2000 matrices converge on a similar result, unambiguously favoring placement of Gnetales in a coniferophyte clade that includes Ginkgoales, cordaites, and extinct and extant conifers. Stronger support is obtained in BI analyses in which gamma-distributed rate variation among sites is implemented in the
model. With ML, the difference in relative support for the two hypotheses appears smaller, but a gneconifer arrangement is consistently favored with all data sets. These results of model-based analyses of post-2000 morphological matrices have interesting implications regarding stem relatives of the angiosperms. Indeed, most post-2000 matrices are broadly congruent in attaching *Pentoxylon*, glossopterids, Bennettitales, and *Caytonia* to the stem lineage of the angiosperms. To these, the analysis of Rothwell and Stockey (2016) adds the Triassic genus *Petriellaea* (Taylor et al. 1994; Bomfleur et al. 2014) (Fig. 3), which has simple reticulate laminar venation, as in *Caytonia*, and cupules containing adaxial ovules. This may be consistent with the view that these fossils shed light on evolution of the complex reticulate venation and bitegmic ovules of angiosperms (Doyle 2006, 2008).

A cautionary note on the results of our Bayesian analyses is necessary. The differences between bootstrap support values in the MP and ML analyses and posterior probabilities in the BI analyses could be due to the very different nature of these support metrics. It has been shown that the relationship between character support and increase in PP is far from linear, and PP can easily sway results toward a hypothesis that is supported by only a few characters (Zander 2004). The strong PP support for groupings (like *Caytonia* or *Petriellaea* plus angiosperms) that receive weak or non-existent support on a character basis (MP and ML bootstrap; Fig. 3A,B) could indicate either the ability of Bayesian inference to pick up a significant signal in an otherwise noisy background or the possibility that this method can be led astray by a few potentially unimportant characters.

**The Conflict between Morphology and Molecules Is Partially Due to Long-Branch Attraction.**—Our results also add new empirical evidence on debates concerning the strengths and weaknesses of morphological data in reconstructing phylogenetic relationships, the phylogenetic importance of fossils, and the best methods to analyze morphological data (Wright and Hillis 2014; O’Reilly et al. 2016; Puttick et al. 2017b). A well-known cause of phylogenetic conflict is the presence of long branches in the tree, which can lead to LBA phenomena (Felsenstein 1978; Bergsten 2005). Analyses based on simulated matrices and real data have repeatedly shown that probabilistic, model-based approaches are more robust to LBA than parsimony (Swofford et al. 2001; Brinkmann et al. 2005; and references in both papers). LBA is most commonly discussed as a confounding factor in molecular studies, as in the case of Gnetales-basal trees found with molecular data (Sanderson et al. 2000; Magallón and Sanderson 2002; Burleigh and Mathews 2007), but here it is morphology that is potentially affected: the BI trees show that both angiosperms and Gnetales are situated on very long morphological branches, especially in the post-2000 matrices.

After following suggestions by Bergsten (2005) and other methodologies (Rota-Stabelli et al. 2011), we conclude that LBA is responsible at least in part for the continuing support for the gnetangiosperm clade in MP analyses of the post-2000 matrices. First, BI recovers a gneconifer topology with higher probability than a topology with Gnetales linked with angiosperms, thus favoring a topology that separates the long branches over a topology that unites them. Second, more complex and better-fitting models recover a higher PP for the topology in which angiosperms and Gnetales are separated (Fig. 2C). Third, removing Gnetales or angiosperms results in a rearrangement of the MP topologies in which the other long branch “flies away” from its original position. Fourth, support for Gnetales plus angiosperms increases with decreased sampling of fossil taxa on the branch leading to the angiosperms, and still more with the removal of all fossils (Fig. 4G–I). It has been suggested that molecular analyses may be incorrect about the relationship of angiosperms and Gnetales because they ignore the great diversity of extinct seed plant taxa (e.g., Rothwell et al. 2009). This reasoning seems to assume that addition of fossils would strengthen the gnetangiosperm hypothesis, but in fact our results indicate that the opposite is true.

To our knowledge, this represents the first reported case of LBA in a morphological
analysis that is supported by multiple tests (Bergsten 2005), with much stronger support than in previously reported cases (Wiens and Hollingsworth 2000; Lockhart and Cameron 2001). These analyses also support the view that model-based methods can overcome the shortcomings of parsimony in such cases. It is also noteworthy that the impact of LBA can be easily visualized with the PCO analysis (Fig. 5), in which the presumed close relationship between Gnetales and conifers and the convergence of Gnetales with the angiosperms are effectively congruent with the positions of the three taxa in the plot of the first two PCO axes. This tool could represent an interesting option for exploring the structure of the data in future phylogenetic analyses.

Less intensive examination of our results suggests that there are fewer conflicts between relationships obtained with parsimony and model-based approaches in other parts of the seed plant tree, suggesting that MP is not necessarily misleading when long-branch effects are lacking. Even when morphological parsimony analyses vary in the arrangement of extant seed plant lines, they are more consistent about relationships below the crown group, with “progymnosperms,” hydrasperman “seed ferns,” Lyginopteris, and medullosans diverging successively below the crown group, and our model-based trees show similar relationships. Another consistent result is the association of traditional coniferophyte groups, namely ginkgos, cordaites, and conifers, setting aside whether this clade also includes Gnetales or (in some morphological analyses) gnetangiosperms. Relationships among cycads and Permian and Mesozoic “seed ferns” (peltasperms, corystosperms, glossopterids, Caytonia) are more variable among parsimony analyses, possibly because of the smaller proportion of preserved characters in the fossils and/or the low number of changes on short internal branches between these lines. Assuming that molecular and model-based morphological results are correct, these considerations suggest that parsimony may perform well when branch lengths are moderate, and it would be unwarranted to reject results out of hand because they are based on parsimony.

The conclusion that similarities between angiosperms and Gnetales are the result of convergence should not be difficult to accept, because many aspects of the morphology of Gnetales can be explained in terms of a Paleozoic conifer prototype (which had female branch systems with secondary short shoots bearing sterile and fertile appendages; cf. Rothwell and Stockey 2013). However, removal of Gnetales from the former gnetangiosperm clade introduces new problems, notably by implying that similarities in seed morphology and anatomy in Gnetales and Bennettitales emphasized by Friis et al. (2009) are also convergences. Some of these similarities have been questioned or reduced by subsequent studies of Bennettitales (Rothwell et al. 2009; Doyle 2012; Rothwell and Stockey 2013; Pott 2016), but others remain. These similarities could be homologous if Bennettitales and Gnetales formed a clade within conifers, but it is much less plausible to interpret Bennettitales as modified conifers, considering their cycad-like leaf morphology, wood anatomical features, and pinnate microsporophylls.

Conclusions

The main lesson of our analyses may be that, contrary to previous impressions, morphological data do not present a strong conflict with the results of molecular analyses regarding the position of angiosperms and Gnetales. This strongly suggests that morphology carries a phylogenetic signal that is consistent with molecular data and may therefore be useful in reconstructing other aspects of the phylogenetic history of the seed plants, most notably the position of fossils relative to living taxa. The supposed conflict between the two sorts of data on the major aspect of the phylogeny of seed plants emphasized here seems to be due to a combination of difficult problems in character analysis and limitations of phylogenetic methods. Because data from the fossil record are particularly important for resolving the evolutionary history of seed plants, due to the wide gaps that separate extant groups and the potential biases in analysis of such sparsely sampled taxa (Burleigh and Mathews 2007; Mathews 2009; Rothwell et al. 2009; Magallón
et al. 2013), our results give new hope for the possibility of integrating fossils and molecules in a coherent way. This is even more important in light of new fossil discoveries (e.g., Rothwell and Stockey 2013, 2016), some of which show similarities to fossils previously associated with angiosperms (e.g., the Triassic Petriellaea plant, which shares leaf and cupule features with Caytonia [Bomfleur et al. 2014]).

The absence of deep convergence problems also opens the possibility of combining morphological and molecular data sets in a total-evidence analysis. Such an approach has been rarely employed in data sets with fossil and extant plants (Magallon 2010), but it has proven to be useful in resolving some controversial relationships (i.e., in the Cycadales [Coiro and Pott 2017]). However, especially with the recent expansion in the amount of available molecular data, both marker selection and taxon choice would have to be carefully considered to set up a successful analysis. It is possible that the ever-increasing amount of sequence data used to infer phylogenetic relationships could swamp the signal present in the many fewer morphological characters, in which case the result would not differ from that found with use of a molecular backbone constraint tree.

An important general message that emerges from our study is the importance of including an exploration of the signal in all phylogenetic analyses involving morphology. The overreliance on single consensus trees, as discussed in Brown et al. (2017) and Puttick et al. (2017a), has been a major driver of the perceived conflict in seed plant phylogeny; another factor has been the lack of support statistics in many studies. Among methods of signal dissection, consensus networks and distance-based neighbor-nets (even if these suffer from the general shortcomings associated with distance-based methods) present promising avenues for the exploration of morphological data sets (Bryant and Moulton 2004) and have proven their power in understanding the history of different groups of fossil and extant taxa at different taxonomic scales (Denk and Grimm 2009; Bomfleur et al. 2017; Grimm 2017).

Although most phylogenetic analyses based on morphology are still conducted in a parsimony framework, some authors have already underlined the potential of model-based approaches in this field (Lee and Worthy 2012; Lee et al. 2014). Our analyses show that BI yields more robust results under different taxon sampling strategies, and although parsimony and BI usually give congruent results, BI appears to be effective in correcting errors of parsimony analyses caused by long-branch effects. Our study converges with previous work indicating that the use of model-based techniques could allow the successful integration of taxa with a high proportion of missing data (Wiens 2005; Wiens and Tiu 2012), which is a prime consideration when dealing with the paleobotanical record.

Supplementary Materials

Data available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.q1c1j71

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Literature Cited

Albert, V. A., A. Backlund, K. Bremer, M. W. Chase, J. R. Manhart, B. D. Mishler, and K. C. Nixon. 1994. Functional constraints and rbcL evidence for land plant phylogeny. Annals of the Missouri Botanical Garden 81:534–567.
Axsmith, B. J., E. L. Taylor, T. N. Taylor, and N. R. Cuneo. 2000. New perspectives on the Mesozoic seed fern order Corystospermales based on attached organs from the Triassic of Antarctica. American Journal of Botany 87:757–768.

Bateman, R. M., J. Hilton, and P. J. Rudall. 2006. Morphological and molecular phylogenetic context of the angiosperms: contrasting the “top-down” and “bottom-up” approaches used to infer the likely characteristics of the first flowers. Journal of Experimental Botany 57:3471–3503.

Bauch, J., W. Liese, and R. Schultze. 1972. The morphological variability of the bordered pit membranes in gymnosperms. Wood Science and Technology 6:165–184.

Bergsten, J. 2005. A review of long-branch attraction. Cladistics 21:163–193.

Bomfleur, B., A.-L. Decombeix, A. B. Schwendemann, I. H. Escapa, E. L. Taylor, T. N. Taylor, and S. McLoughlin. 2014. Habit and ecology of the Petriellales, an unusual group of seed plants from the Triassic of Gondwana. International Journal of Plant Sciences 175:1062–1075.

Brinkmann, H., M. van der Giezen, Y. Zhou, G. P. de Raucourt, and O. M. Vargas. 2004. Neighbor-net: an agglomerative constructions of morphological traits are not discordant when taking phylogenetic methods. Journal of Human Evolution 97:17–26.

Denk, T., and G. W. Grimm. 2009. The biogeographic history of beechn trees. Review of Palaeobotany and Palynology 158:83–100.

Donoghue, M. J., and J. A. Doyle. 2000. Seed plant phylogeny: demise of the anthophyte hypothesis? Current Biology 10:R106–R109.

Donoghue, M. J., J. A. Doyle, J. Gauthier, A. G. Kluge, and T. Rowe. 1989. The importance of fossils in phylogeny reconstruction. Annual Review of Ecology and Systematics 20:431–460.

Doyle, J. A. 1996. Seed plant phylogeny and the relationships of the Gnetales. International Journal of Plant Sciences 157(6, Suppl.) 52–539.

——. 2006. Seed ferns and the origin of the angiosperms, Journal of the Torrey Botanical Society 133:169–209.

——. 2008. Integrating molecular phylogenetic and paleobotanical evidence on origin of the flower. International Journal of Plant Sciences 169:816–843.

——. 2012. Molecular and fossil evidence on the origin of angiosperms. Annual Review of Earth and Planetary Sciences 40:301–326.

——. 2013. Phylogenetic analyses and morphological innovations in land plants. In B. A. Ambrose, and M. Purugganan, eds. The evolution of plant form. Annual Plant Reviews 45:1–50. Wiley-Blackwell, Oxford.

Doyle, J. A., and M. J. Donoghue. 1986. Seed plant phylogeny and the origin of angiosperms: an experimental cladistic approach. Botanical Review 52:321–431.

——. 1987. The importance of fossils in elucidating seed plant phylogeny and macroevolution. Review of Palaeobotany and Palynology 50:63–95.

——. 1992. Fossils and seed plant phylogeny revisited. Brittonia 44:89–106.

Doyle, J. A., and P. K. Endress. 2000. Morphological phylogenetic analysis of basal angiosperms: comparison and combination with molecular data. International Journal of Plant Sciences 161 (6, Suppl.) S121–S153.

——. 2014. Integrating Early Cretaceous fossils into the phylogeny of living angiosperms: ANITA lines and relatives of Chloranthaceae. International Journal of Plant Sciences 175:555–600.

Endress, P. K., and J. A. Doyle. 2009. Reconstructing the ancestral angiosperm flower and its initial specializations. American Journal of Botany 96:22–66.

Felsenstein, J. 1978. Cases in which parsimony or compatibility methods will be positively misleading. Systematic Zoology 27:401–410.

Foley, N. M., M. S. Springer, and E. C. Teeling. 2016. Mammal madness: is the mammal tree of life not yet resolved? Philosophical Transactions of the Royal Society of London B 371:20150140.

Friis, E. M., J. A. Doyle, P. K. Endress, and Q. Leng. 2003. Archaefractus—angiosperm precursor or specialized early angiosperm? Trends in Plant Science 8:369–373.

Friis, E. M., P. R. Crane, K. R. Pedersen, S. Bengtson, P. J. C. Donoghue, G. W. Grimm, and M. Stampanoni. 2007. Phase-contrast X-ray microtomography links Cretaceous seeds with Gnetales and Bennettitales. Nature 450:549–552.
Frisi, E. M., K. R. Pedersen, and P. R. Crane. 2009. Early Cretaceous mesofossils from Portugal and eastern North America related to the Bennettitales-Erdtmannithecales-Gnetales group. American Journal of Botany 96:252–283.

Gauthier, J., A. G. Kluge, and T. Rowe. 1988. Amniote phylogeny and the importance of fossils. Cladistics 4:105–209.

Gauthier, J., M. Kearney, J. A. Maisano, O. Rieppel, and A. D. B. Behlke. 2012. Assembling the squamate tree of life: perspectives from the phenotype and the fossil record. Bulletin of the Peabody Museum of Natural History 53:3–308.

Givnish, T. J., and K. J. Sytsma. 1997. Molecular evolution and adaptive radiation. Cambridge University Press, Cambridge.

Goderforth, P., A. Cau, D.-Y. Hu, F. Escuillié, W. Wu, and G. Dyke. 2013. A Jurassic avialan dinosaur from China resolves the early phylogenetic history of birds. Nature 498:359–362.

Grimm, G. 2017. Should we try to infer trees on tree-unlikely matrices? The Genealogical World of Phylogenetic Networks. http://phylonetworks.blogspot.com/2017/07/should-we-try-to-infer-trees-on.html, accessed 5 July 2017.

Gugerli, F., C. Sperisen, U. Büchler, I. Brunner, S. Brodbeck, J. D. Palmer, and Y.-L. Qi. 2001. The evolutionary split of Pinaceae from other conifers: evidence from an intron loss and a multigene phylogeny. Molecular Phylogenetics and Evolution 21:167–175.

Hamby, R. K., and E. A. Zimmer. 1992. Ribosomal RNA as a phylogenetic tool in plant systematics. Pp. 50–91 in P. S. Soltis, D. E. Soltis, and J. J. Doyle, eds. Molecular systematics of plants. Chapman and Hall, New York.

Harriss, T. M. 1954. Mesozoic seed cuticles. Svensk Botanik Tidskrift 48:281–291.

Hill, C. R., and P. R. Crane. 1982. Evolutionary cladistics and the origin of angiosperms. In K. A. Joysey, and A. E. Friday, eds. Problems of phylogenetic reconstruction. Systematics Association Special Volume 21:269–361. Academic Press, London.

Hilton, J., and R. M. Bateman. 2006. Pteridosperms are the backbone of seed plant phylogeny. Journal of the Torrey Botanical Society 133:119–168.

Holland, B., K. T. Huber, V. Moulton, and P. J. Lockhart. 2004. Using consensus networks to visualize contradictory evidence for species phylogeny. Molecular Biology and Evolution 21:1459–1461.

Huson, D. H., and D. Bryant. 2006. Application of phylogenetic networks in evolutionary studies. Molecular Biology and Evolution 23:254–267.

Jenner, R. A. 2004. Accepting partnership by submission? Morphological phylogenetics in a molecular millennium. Systematic Biology 53:333–359.

Kass, R. E., and A. E. Raftery. 1995. Bayes factors. Journal of the American Statistical Association 90:773–795.

Kelley, D. R., and C. S. Gasser. 2009. Ovule development: genetic trends and evolutionary considerations. Sexual Plant Reproduction 22:229–234.

Klaven, S. D., T. N. Taylor, and E. L. Taylor. 2002. Anatomy of Umkomasia (Corystospermales) from the Triassic of Antarctica. American Journal of Botany 89:664–676.

Lee, M. S., and T. H. Worthy. 2012. Likelihood reinstates from discrete morphological character data. Systematic Biology 50:913–925.

Lloyd, G. T. 2016. Estimating morphological diversity and tempo with discrete character-taxon matrices: implementation, challenges, progress, and future directions. Biological Journal of the Linnean Society 118:131–151.

Lockhart, P. J., and S. A. Cameron. 2001. Trees for bees. Trends in Ecology and Evolution 16:84–88.

Maddison, D. R., and W. P. Maddison. 2003. MacClade 4: analysis of phylogeny and character evolution, version 4.06. Sinauer, Sunderland, Mass.

Magallon, S. 2010. Using fossils to break long branches in molecular dating: a comparison of relaxed clocks applied to the origin of angiosperms. Systematic Biology 59:384–399.

Magallon, S., and M. J. Sanderson. 2002. Relationships among seed plants inferred from highly conserved genes: sorting conflicting phylogenetic signals among ancient lineages. American Journal of Botany 89:1991–2006.

Magallon, S., K. W. Hilu, and D. Quandt. 2013. Land plant evolutionary timeline: gene effects are secondary to fossil constraints in relaxed clock estimation of age and substitution rates. American Journal of Botany 90:556–573.

Martens, P. 1971. Les Gnétophytes. Encyclopedia of plant anatomy 12(2). Borntraeger, Stuttgart.

Mathews, S. 2009. Phylogenetic relationships among seed plants: persistent questions and the limits of molecular data. American Journal of Botany 96:228–236.

Mathews, S., and E. Kramer. 2012. The evolution of reproductive structures in seed plants: a re-examination based on insights from developmental genetics. New Phytologist 194:910–923.

Mathews, S., M. D. Clements, and M. A. Beistein. 2010. A duplicate gene root of seed plants and the phylogenetic position of flowering plants. Philosophical Transactions of the Royal Society of London B 365:383–395.

Müller, K. F. 2005. The efficiency of different search strategies for estimating parsimony, jackknife, bootstrap, and Bremer support. BMC Evolutionary Biology 5:58.

Mundry, R., and T. Stützel. 2004. Morphogenesis of the reproductive shoots of Welwitschia mirabilis and Ephedra distachya (Gnetales), and its evolutionary implications. Organisms Diversity and Evolution 4:91–108.

Nickrent, D. L., C. L. Parkinson, J. D. Palmer, and R. J. Duff. 2000. Multigene phylogeny of land plants with special reference to bryophytes and the earliest land plants. Molecular Biology and Evolution 17:1885–1895.

Nixon, K. C., W. L. Crepet, D. W. Stevenson, and E. M. Friss. 1994. A reevaluation of seed plant phylogeny. Annals of the Missouri Botanical Garden 81:484–533.

Murry, N., and J. L. Scott. 2004. Epifonia from the Carnian (Upper Triassic) of Lunz, Austria: more evidence for a unitegmic seed coat in early Bennettitales. International Journal of Plant Sciences 177:771–791.
Rothwell, G. W., and R. Serbet. 1994. Lignophyte phylogeny and Rota-Stabelli, O., E. Kayal, D. Gleeson, J. Daub, J. L. Boore, M. J. Rothwell, G. W., and R. A. Stockey. 2013. Evolution and phylogeny of Gnetophytes: evidence from the anatomically preserved seed cone of Williamsonia (Williamsoniaceae): evidence for bennettiteal reproduction in the Late Cretaceous of western North America. Systematic Biology 60:457–604.

Springer, M. S., R. W. Meredith, E. C. Teeling, and W. J. Murphy. 2015. Post-molecular systematics and the future of phylogenetics. Cladistics 30:1132–1313.

Stefanovic, S., M. Jager, J. Deutsch, J. Broutin, and M. Masselet. 1998. Phylogenetic relationships of conifers inferred from partial 28S rRNA gene sequences. American Journal of Botany 85:688–697.

Stockey, R. A., and G. W. Rothwell. 2003. Anatomically preserved Williamsonia (Williamsoniaceae): evidence for bennettiteal reproduction in the Late Cretaceous of western North America. International Journal of Plant Sciences 164:251–262.

2009. Distinguishing angiosperms from the earliest angiosperms: a Lower Cretaceous (Valanginian-Hauterivian) fruit-like reproductive structure. American Journal of Botany 96:323–335.

Sun, G., Q. Ji, D. L. Dilcher, S. Zheng, K. C. Nixon, and X. Wang. 2002. Archaeofructaceae, a new basal angiosperm family. Science 296:899–904.

Swofford, D. L. 2003. PAUP*: Phylogenetic Analysis Using Parsimony (* and other methods), Version 4. Sinauer, Sunderland, Mass.

Taylor, E. L., and T. N. Taylor. 1992. Reproductive biology of the Permian Glossospermites and their suggested relationship to flowering plants. Proceedings of the National Academy of Sciences USA 89:11495–11497.

Taylor, T. N., G. M. Del Fuego, and E. L. Taylor. 1994. Permineralized seed fern cupules from the Triassic of Antarctica: implications for cupule and carpel evolution. American Journal of Botany 81:666–677.

Templeton, A. R. 1983. Phylogenetic inference from restriction endonuclease cleavage site maps with particular reference to the evolution of humans and the apes. Evolution 37:221–244.

Wickett, N. J., S. Mirarab, N. Nguyen, T. Warnow, E. Carpenter, N. Matsuci, S. Ayyampalayam, M. S. Barker, J. G. Burleigh, M. A. Gitzendanner, B. R. Ruhfel, E. Wafula, J. P. Der, S. W. Graham, S. Mathews, M. Melkonian, D. E. Soltis, P. S. Soltis, N. W. Miles, C. J. Rothfels, L. Polkorny, A. J. Shaw, L. DeGironimo, D. W. Stevenson, B. Surek, J. C. Villarreal, B. Boure, H. Philipp, C. W. dePamphilis, T. Chen, M. K. Deyholos, R. S. Baucom, T. M. Kutchan, M. M. Augustin, J. Wang, Y. Zhang, Z. Tian, Z. Yan, X. Wu, X. Sun, G. K. S. Wong, and J. Leebens-Mack. 2014. Phylogenomic analysis of the origin and early diversification of land plants. Proceedings of the National Academy of Sciences USA 111:E4859–E4868.

Wieland, G. R. 1916. American fossil cycads, Vol. 2. Taxonomy. Carnegie Institution of Washington, Washington, D.C.

Wiens, J. J. 2005. Can incomplete taxa rescue phylogenetic analyses from long-branch attraction? Systematic Biology 54:731–742.

Wiens, J. J., and B. D. Hollingsworth. 2000. War of the iguanas: conflicting phylogenies, long-branch attraction, and disparate

Pyron, R. A. 2011. Divergence time estimation using fossils as terminal taxa and the origins of Lissamphibia. Systematic Biology 60:466–481.

——. 2015. Post-molecular systematics and the future of phylogenetics. Trends in Ecology and Evolution 30:384–389.

Rambaut, A., and A. J. Drummond. 2007. Tracer: MCMC trace analysis tool, Version 1.4.1. http://tree.bio.ed.ac.uk/software.

R Core Team 2017. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org.

Rothewell, G. W., and R. Serbet. 1994. Lignophyte phylogeny and the evolution of spermatophytes: a numerical cladistic analysis. Systematic Botany 19:443–482.

Rothwell, G. W., and R. A. Stockey. 2013. Evolution and phylogeny of Gnetophytes: evidence from the anatomically preserved seed cone Protodrosera canesii gen. et sp. nov. and the seeds of several bennettitelean species. International Journal of Plant Sciences 174:511–529.

——. 2016. Phylogenetic diversification of Early Cretaceous seed plants: the compound seed cone of Doylea tetrahedrasperma. American Journal of Botany 103:923–937.

Rothwell, G. W., W. L. Crepet, and R. A. Stockey. 2009. Is the anthophyte hypothesis alive and well? New evidence from the reproductive structures of Bennettitales. American Journal of Botany 96:296–322.

Rudall, P. J., and R. M. Bateman. 2010. Defining the limits of flowers: the challenge of distinguishing between the evolutionary products of simple versus compound strobili. Philosophical Transactions of the Royal Society of London B 365:397–409.

Rydin, C., and M. Källersjö. 2002. Taxon sampling and seed plant phylogeny. Cladistics 18:484–513.

Rydin, C., M. Källersjö, and E. M. Friis. 2002. Seed plant relationships and the systematic position of Gnetales based on nuclear and chloroplast DNA: conflicting data, rooting problems, and the monophyly of conifers. International Journal of Plant Sciences 163:197–214.

Sanderson, M. J., M. F. Wojciechowski, J.-M. Hu, T. Sher Khan, and S. G. Brady. 2000. Error, bias, and long-branch attraction in data for two chloroplast photosystem genes in seed plants. Molecular Biology and Evolution 17:782–797.

Scotland, R. W., R. G. Olmstead, and J. R. Bennett. 2003. Phylogeny reconstruction: the role of morphology. Systematic Biology 52:539–548.

Singh, H. 1978. Embryology of gymnosperms (Handbuch der Pflanzenanatomie 10(2) Bornträger, Berlin.

Springer, M. S., A. Burk-Herrick, R. Meredith, E. Eizirik, E. Teeling, J. S. O’Brien, and W. J. Murphy. 2007. The adequacy of morphology for reconstructing the early history of placental mammals. Systematic Biology 56:673–684.

Springer, M. S., R. W. Meredith, E. C. Teeling, and W. J. Murphy. 2013. Technical comment on “The placental mammal ancestor and the post-K-Pg radiation of placents. Science 341:613.

Stamatakis, A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30:1312–1313.

Taylor, E. L., and T. N. Taylor. 1992. Reproductive biology of the Permian Glossospermites and their suggested relationship to flowering plants. Proceedings of the National Academy of Sciences USA 89:11495–11497.

Taylor, T. N., G. M. Del Fuego, and E. L. Taylor. 1994. Permineralized seed fern cupules from the Triassic of Antarctica: implications for cupule and carpel evolution. American Journal of Botany 81:666–677.

Templeton, A. R. 1983. Phylogenetic inference from restriction endonuclease cleavage site maps with particular reference to the evolution of humans and the apes. Evolution 37:221–244.

Wickett, N. J., S. Mirarab, N. Nguyen, T. Warnow, E. Carpenter, N. Matsuci, S. Ayyampalayam, M. S. Barker, J. G. Burleigh, M. A. Gitzendanner, B. R. Ruhfel, E. Wafula, J. P. Der, S. W. Graham, S. Mathews, M. Melkonian, D. E. Soltis, P. S. Soltis, N. W. Miles, C. J. Rothfels, L. Polkorny, A. J. Shaw, L. DeGironimo, D. W. Stevenson, B. Surek, J. C. Villarreal, B. Boure, H. Philipp, C. W. dePamphilis, T. Chen, M. K. Deyholos, R. S. Baucom, T. M. Kutchan, M. M. Augustin, J. Wang, Y. Zhang, Z. Tian, Z. Yan, X. Wu, X. Sun, G. K. S. Wong, and J. Leebens-Mack. 2014. Phylogenomic analysis of the origin and early diversification of land plants. Proceedings of the National Academy of Sciences USA 111:E4859–E4868.

Wieland, G. R. 1916. American fossil cycads, Vol. 2. Taxonomy. Carnegie Institution of Washington, Washington, D.C.

Wiens, J. J. 2005. Can incomplete taxa rescue phylogenetic analyses from long-branch attraction? Systematic Biology 54:731–742.

Wiens, J. J., and B. D. Hollingsworth. 2000. War of the iguanas: conflicting phylogenies, long-branch attraction, and disparate
rates of molecular and morphological evolution in iguanid lizards. Systematic Biology 49:69–85.

Wiens, J. J., and J. Tiu. 2012. Highly incomplete taxa can rescue phylogenetic analyses from the negative impacts of limited taxon sampling. PLoS ONE 7:e42925.

Wiens, J. J., P. T. Chippindale, and D. M. Hillis. 2003. When are phylogenetic analyses misled by convergence? A case study in Texas cave salamanders. Systematic Biology 52:501–514.

Wright, A. M., and D. M. Hillis. 2014. Bayesian analysis using a simple likelihood model outperforms parsimony for estimation of phylogeny from discrete morphological data. PLoS ONE 9: e109210.

Wright, A. M., G. T. Lloyd, and D. M. Hillis. 2015. Modeling character change heterogeneity in phylogenetic analyses of morphology through the use of priors. Systematic Biology 65:602–611.

Xie, W., P. O. Lewis, Y. Fan, L. Kuo, and M.-H. Chen. 2011. Improving marginal likelihood estimation for Bayesian phylogenetic model selection. Systematic Biology 60:150–160.

Zander, R. H. 2004. Minimal values of reliability of bootstrap and jackknife proportions, decay index, and Bayesian posterior probability. PhyloInformatics 2:1–13.

Zhang, C., T. Stadler, S. Klopfstein, T. A. Heath, and F. Ronquist. 2016. Total-evidence dating under the fossilized birth-death process. Systematic Biology 65:228–249.

Zhong, B., O. Deusch, V. V. Goremkin, D. Penny, P. J. Briggs, R. A. Atherton, S. V. Nikforsova, and P. J. Lockhart. 2011. Systematic error in seed plant phylogenomics. Genome Biology and Evolution 3:1340–1348.

Zou, Z., and J. Zhang. 2016. Morphological and molecular convergences in mammalian phylogenetics. Nature Communications 7:12758.