PHYLOGENY AND DIVERGENCE TIME ESTIMATES FOR THE FERN GENUS AZOLLA (SALVINIACEAE)

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A phylogeny for all extant species of the heterosporous fern genus Azolla is presented here based on more than 5000 base pairs of DNA sequence data from six plastid loci (rbcL, atpB, rps4, trnL-trnF, trnG-trnR, and rps4-trnS). Our results are in agreement with other recent molecular phylogenetic hypotheses that support the monophyly of sections Azolla and Rhizosperma and the proposed relationships within section Azolla. Divergence times are estimated within Azolla using a penalized likelihood approach, integrating data from fossils and DNA sequences. Penalized likelihood analyses estimate a divergence time of 50.7 Ma (Eocene) for the split between sections Azolla and Rhizosperma, 32.5 Ma (Oligocene) for the divergence of Azolla nilotica from A. pinnata within section Rhizosperma, and 16.3 Ma (Miocene) for the divergence of the two lineages within section Azolla (the A. filiculoides + A. rubra lineage from the A. caroliniana + A. microphylla + A. mexicana complex).

Keywords: Azolla, divergence time estimates, ferns, freshwater plants, heterospory, Salviniaceae.

Introduction

Because of its great agricultural importance, Azolla Lam. (Salviniaceae) is “the most frequently studied genus of ferns in the world” (Lumpkin 1993, p. 339). Azolla has an obligate symbiotic association with the cyanobacterium Anabaena azollae Strasburger (Strasburger 1873; Peters and Meeks 1989; Plazinski et al. 1990; Baker et al. 2003). The nitrogen-fixing capabilities of this symbiosis have long been appreciated, promoting the use of Azolla as a “green manure” in rice paddies in Southeast Asia (Moore 1969; Peters and Meeks 1989; Wagner 1997).

Azolla is one of five fern genera to exhibit a heterosporous life cycle—a phenomenon observed in less than 1% of extant ferns (Pryer et al. 2004). Together with all other heterosporous ferns, Azolla is nested within leptosporangiate ferns and forms the sister lineage to a large clade comprised of tree ferns and polypods (Pryer et al. 2004; Schuettbelz et al. 2006; Schuettbelz and Pryer 2007). Molecular and morphological analyses have shown Azolla to be sister to Salvinia, together forming a clade that is sister to the three genera (Marsilea, Regnellidium, and Pilularia) that make up extant Marsileaceae (Rothwell and Stockey 1994; Hasebe et al. 1995; Pryer et al. 1995, 2004; Pryer 1999). Many authors (Tan et al. 1986; Peter and Meeks 1989; Saunders and Fowler 1993) have placed Azolla in its own monotypic family Azollaceae, but more recently, it has been recognized together with Salvinia in Salviniaceae (Smith et al. 2006).

The taxonomic history of Azolla has long been contentious and convoluted (for a thorough account, see Reid et al. 2006; but see also Dunham 1986; Dunham and Fowler 1987). Azolla has traditionally been divided into two sections, Azolla and Rhizosperma (Mettenius 1847; Tan et al. 1986; Wagner 1997; Reid et al. 2006), based primarily on reproductive structures (e.g., section Azolla is characterized by three floats in the megaspore apparatus, whereas section Rhizosperma has nine).

Section Rhizosperma is consistently considered to comprise two species (Azolla nilotica and A. pinnata; Saunders and Fowler 1992); however, little consensus has been reached on the diversity in section Azolla, which has varied from two to five species. For example, the status of A. rubra as either a distinct species or a variety of A. filiculoides has long been debated (Svenson 1944), although most recently it has been treated as a distinct species (Zimmerman et al. 1989; Saunders and Fowler 1993; Reid et al. 2006). Likewise, the taxonomic status and rank of three sympatric New World taxa (A. caroliniana, A. mexicana, and A. microphylla) have also generated considerable debate. Numerous authors maintain them as distinct species (Svenson 1944; Moore 1969; Saunders and Fowler 1993). Others have contributed evidence or reviews suggesting that they be regarded as a single species (Zimmerman et al. 1989, 1991; Evrard and Van Hove 2004) or that the extent of the taxonomic conundrum is such that some of the names may have been misapplied and will require a detailed study of type material (Dunham 1986; Dunham and Fowler 1987). Reid et al. (2006) produced the first molecular phylogeny of Azolla and hypothesized that A. caroliniana was distinct whereas A. microphylla and A. mexicana should be considered the same species.

In this study, we use a data set of six plastid loci (rbcL, atpB, rps4, trnL-trnF, trnG-trnR, and rps4-trnS) to examine the phylogeny of extant Azolla taxa; only one of the six loci (trnL-trnF) duplicates data from Reid et al. (2006), although we sequenced this locus independently. This allows us to

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Manuscript received October 2006; revised manuscript received April 2007.

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critically evaluate the traditional sectional classification and the conclusions of Evrard and Van Hove (2004) and Reid et al. (2006) regarding the A. caroliniana-microphylla-mexicana (CAR-MIC-MEX) complex. We also integrate data from Azolla’s rich fossil record to estimate divergence times for all well-supported nodes in our molecular phylogeny.

Material and Methods

Taxonomic Sampling

Twelve accessions of Azolla were studied here (table 1): three from section Rhizosperma and nine from section Azolla. Salvinia minima and Marsilea botryocarpa were used as outgroup taxa to root the topology.

DNA Isolation, Amplification, and Sequencing

Genomic DNA was extracted from silica-dried leaf material using the DNeasy kit (Qiagen, Valencia, CA) according to the manufacturer’s instructions. Six plastid loci (rbcL, atpB, rps4, trnL-trnF, trnG-trnR, and rps4-trnS) were amplified separately for each accession using the polymerase chain reaction (PCR), according to established protocols (Pryer et al. 2004). PCR products were cleaned using Montage columns (Millipore, Bedford, MA) according to the manufacturer’s protocol. Sequencing reactions were performed for both strands of all purified PCR products using Big Dye Terminator Cycle Sequencing reagents (Applied Biosystems, Foster City, CA). All sequencing reactions were completed on either an ABI 3700 or an ABI 3730XL automated sequencer (Applied Biosystems). Primer information for amplification and sequencing reactions is shown in table 2. All sequence reads were examined for contamination using the National Center for Biotechnology Information nucleotide-nucleotide BLAST (blastn) function (Altschul et al. 1997). DNA sequence chromatograms were corrected and assembled into contiguous alignments using Sequencher (Gene Codes, Ann Arbor, MI). Seventy-five sequences used in this study were generated specifically for this project and are available in GenBank (table 1).

Sequence Alignments

Sequence alignments were performed by eye using MacClade, version 4.06 (Maddison and Maddison 2003). The protein-coding rbcL, atpB, and rps4 data sets were easily aligned (without gaps) across all Azolla accessions and outgroup taxa. The noncoding trnG-trnR, trnL-trnF, and rps4-trnS data were also easily aligned but only across all Azolla accessions. Because numerous ambiguously aligned regions resulted when the outgroup sequences were added, they were not included in the noncoding locus data sets. The noncoding alignments for the Azolla taxa alone did have some insertions and deletions, but none was ambiguous; therefore, no regions were excluded, nor was any gap coding method employed.

Data Set Combinability

Using PAUP*, version 4.0b10 (Swofford 2002), separate maximum parsimony (MP) and maximum likelihood (ML) bootstrap analyses were run for each data set. The MP analyses were run for 1000 replicates with 10 random addition sequence replicates each and tree-bisection-reconnection (TBR) branch swapping. The ML analyses were run for 500 replicates, each with 10 random addition sequence replicates. In addition, four independent Bayesian Markov chain Monte Carlo (B/MCMC) analyses, each using four chains, were run for each data set, using MrBayes, version 3.1.1 (Ronquist and Huelsenbeck 2003). These analyses were run for 10 million generations, with trees being sampled every 1000 generations; the first 2.5 million generations were discarded as the burn-in phase. The 18 resultant topologies (six from MP, six from ML, and six from B/MCMC) were inspected for topological conflicts using a threshold of 70% bootstrap proportion or higher for the MP and ML analyses and a threshold of 0.95 posterior probability or higher for the B/MCMC analyses. No topological conflict was visually detected among data sets or analytical methods, and hence all six data sets were combined into a single data set.

Phylogenetic Analyses

As noted, an excess of ambiguously aligned regions resulted in the alignments of the noncoding data sets (trnG-trnR, trnL-trnF, and rps4-trnS) when outgroups were included. Rather than excluding much of the data for these three loci, we performed two sets of phylogenetic analyses to maximize the information available (similar to the approach in Nagalingum et al. 2007). Analysis 1 included all Azolla accessions, plus the outgroups, for only the protein-coding loci (atpB, rbcL, and rps4). Because this analysis unequivocally established section Rhizosperma as the monophyletic sister to the rest of the genus (see “Results”), it was then used as an outgroup in analysis 2, which focused on exploring relationships within section Azolla. Analysis 2 included only the Azolla accessions but incorporated all six loci (atpB, rbcL, rps4, rps4-trnS, trnG-trnR, and trnL-trnF). No ambiguous regions were present in these alignments, and no sequence data were excluded.

For both analyses 1 and 2, MP and ML analyses were run using PAUP*, version 4.0b10 (Swofford 2002), and B/MCMC analyses were performed using MrBayes, version 3.1.1 (Ronquist and Huelsenbeck 2003). All MP heuristic searches were for 1000 replicates with TBR branch swapping, and all MP bootstrap analyses were performed using 1000 replicates, each with 10 random addition sequence replicates and TBR branch swapping. The ML heuristic searches were run with TBR branch swapping, and the ML bootstrap analyses all included 500 replicates with 10 random addition sequence replicates each and TBR branch swapping. Each B/MCMC analysis was performed using four independent tree searches. Each of these searches used four chains, running for 10 million generations apiece. The B/MCMC analyses were performed with data partitioned by locus, flat priors, and trees being sampled every 1000 generations. The likelihood versus generation scores were plotted for all four searches, and stationarity was determined to have occurred after 2,500,000 generations in each analysis. The prestationarity trees were excluded as the burn-in period. For each B/MCMC analysis, post-burn-in trees from all four runs were pooled, and a majority-rule consensus tree with average branch lengths and posterior probabilities computed from 30,000 trees, using the “sumt” command in MrBayes.
<table>
<thead>
<tr>
<th>Taxon</th>
<th>Voucher information</th>
<th>GenBank accession nos.</th>
<th>Fern DNA database number</th>
</tr>
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<td><strong>Section Azolla:</strong></td>
<td></td>
<td>rbcL</td>
<td>atpB</td>
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<td><strong>Azolla caroliniana</strong> Willd.</td>
<td>USA: Ohio, W. Tuleke s.n. (DUKE); CA3001 in IRRI(^b)</td>
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<td><strong>A. filiculoides</strong> Lam.</td>
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<td><strong>A. mexicana Schlecht. &amp; Cham.</strong></td>
<td>USA: California, Sutter County, Sutter Basin of Sacramento Valley, D. W. Rains s.n. (DUKE)</td>
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<td><strong>A. mexicana</strong> 2</td>
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<td>EF520874</td>
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<td>EF520879</td>
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<td>Sudan: Kosti, T. Lumpkin s.n. (DUKE); NIS001 in IRRI(^b)</td>
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<td><strong>A. pinnata</strong> R. Br. 1</td>
<td>China: Shandong province, Tancheng District, Bai Kai-Ze s.n. (DUKE); PI0022 in IRRI(^b)</td>
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<td>EF520870</td>
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<td><strong>A. pinnata</strong> 2</td>
<td>Malaysia: Penang, Bumhorn Lima, Butterworth s.n. (DUKE); PI0002 in IRRI(^b)</td>
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<td>EF520877</td>
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<td><strong>Outgroup taxa:</strong></td>
<td></td>
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<td>atpB</td>
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<td><strong>Marsilea botryocarpa</strong> Ballard</td>
<td>Kenya: Tsavo East National Park, R. Faden s.n. (UC)</td>
<td>DQ643294</td>
<td>DQ643259</td>
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<td><strong>Salvinia minima</strong> Baker</td>
<td>USA: Louisiana, S. Fredericq s.n. (DUKE)</td>
<td>EF520931</td>
<td>EF520881</td>
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</tbody>
</table>

Note. NA = data not available for this taxon.
\(^a\) Permanent record numbers in Fern DNA Database at http://www.pryerlab.net/DNA_database.shtml.
\(^b\) International Rice Research Institute (IRRI), Manila.
Models of sequence evolution for all ML and B/MCMC analyses were determined using Modeltest, version 3.6 (Posada and Crandall 1998). For analysis 1, the TrN + G model was used for atpB and rbcL, HKY + G was used for rps4, and TrN + G was used for the combined data set. For analysis 2, the HKY model was used for atpB, rbcL, and trnF, and TrNef + G was used for the combined three-locus data set (TrN + G, as estimated by Modeltest). Marsilea botryocarpa was then pruned from the tree, leaving the Salvinia-Azolla node as the root node.

**Divergence Time Estimation**

The topology obtained from the Bayesian inference search of the combined data set in analysis 2 was used for the penalized likelihood (PL) divergence date analysis. Because PL divergence time estimation programs can experience difficulty with topologies that have extremely short branch lengths, we followed the placeholder approach (as recommended by Sanderson [2003]) by deleting several redundant accessions. We removed Azolla mexicana 2, A. mexicana 3, A. microphylla 1, A. microphylla 3, and A. pinnata 2. This left us with at least one accession representing each commonly recognized species in Azolla. Because of the well-supported (Pryer et al. 2004) intergeneric relationships in the heterosporous fern clade ((Salvinia, Azolla) (Marsilea (Regnellidium, Pilularia))), Salvinia minima was added to the tree as nearest sister genus to the Azolla clade, and Marsilea botryocarpa was added to the tree as nearest sister genus to the Azolla-Salvinia clade. Branch lengths for this topology were then re-estimated in PAUP*, version 4.0b10, using the combined rbcL, atpB, and rps4 data sets and the ML parameters for the combined three-locus data set (TrN + G, as estimated by Modeltest). Marsilea botryocarpa was then pruned from the tree, leaving the Salvinia-Azolla node as the root node.

Using r8s (ver. 1.60; Sanderson 2003), we implemented two fossil constraints (table 3) in conjunction with this tree topology. The Salvinia-Azolla node was fixed as a calibration point at 89 Ma, using the earliest reports of Azolla megaspores (Collinson 1991). The section Azolla node was assigned a minimum age constraint of 13.65 Ma using a megaspore fossil of A. filiculoides (Mai 2001). This fossil indicates that the first divergence in section Azolla had occurred no later than this date. The fossil was not assigned to the node representing the most recent common ancestor of A. filiculoides and A. rubra because the megaspore morphology of these two species is essentially identical. Instead, the fossil was assigned to the next-lowest node (section Azolla node) because it is at least assignable to the A. filiculoides/A. rubra lineage.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5' to 3')</th>
<th>Reference</th>
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<tbody>
<tr>
<td>trnL-trnF</td>
<td>TACGACGATCTYTCTAAACAAGC</td>
<td>Tabet et al. 1991</td>
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<tr>
<td>TRNLG</td>
<td>ATGTCACCACAAAACCGAGACTAAACG</td>
<td>Korall et al. 2006</td>
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<td>TRNL</td>
<td>ATGGCAATATTTCCGAGGATRTYA</td>
<td>Wolf 1997</td>
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<tr>
<td>TRNF</td>
<td>GTGAACTTCATGGATTCTGATTTCA</td>
<td>Nagalingum et al. 2007</td>
</tr>
<tr>
<td>rbcL</td>
<td>ATGB672F</td>
<td>TGGATACCCGGAGYCCCTCTWAGTGT</td>
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<td>rps4</td>
<td>ATPE384R</td>
<td>GAATCGGACTTCCATTTGAGATTG</td>
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<td>trnG-trnR</td>
<td>ATPS5F</td>
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<td>TRNG63R</td>
<td>ATPS4IR</td>
<td>TTTATATCCAARCTSAITAGATTCTG</td>
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<td>TRNG353R</td>
<td>ATPS4LF</td>
<td>GTCCTGKATGCGMGGACCTG</td>
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<td>TRNR22R</td>
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<td>ATPB609R</td>
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<tr>
<td>rbcL</td>
<td>ATPB609R</td>
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Note. Primers in italics were used both for amplifying and sequencing.
A r8s analysis using the three-locus data set was performed with a smoothing parameter of 1 as selected by a cross-validation procedure (Sanderson 2002). Using the procedure described in the r8s manual (Sanderson 2002), we ran 1000 bootstrap replicates. First, 1000 bootstrap data matrices were generated using PAUP*, version 4.0b10. Branch lengths for the tree used in the initial r8s analysis were then reestimated for each of the bootstrap data sets. Finally, divergence times for all the resulting bootstrap trees were calculated by r8s. Out of the 1000 replicates, 984 were successfully completed.

**Results**

**Phylogenetic Analyses**

**Analysis 1.** The MP analysis of the combined three-locus data set (atpB, rbcL, and rps4) produced two equally most parsimonious trees (916 steps, consistency index [CI] = 0.882, retention index [RI] = 0.816), and the strict consensus tree was well resolved (not shown). The ML analysis resulted in one most likely tree (−ln L = 8530.8154; tree not shown). The B/MCMC analysis returned a majority-rule consensus tree with a well-resolved backbone (fig. 1A). Analysis 1 allowed us to robustly confirm the monophyly of *Azolla* from all three measures of support—Bayesian posterior probability (PP = 1.00) and ML and MP bootstrap percentages (BPML = 99 and BPMP = 100). We also established the reciprocal monophyly of sections *Azolla* (BPMP = 100; BPML = 100; PP = 1.00) and *Rhizosperma* (BPMP = 94; BPML = 100; PP = 1.00). The well-supported position of section *Rhizosperma* as sister to the rest of the genus allowed us to use it as the outgroup in analysis 2. No well-supported conflicts were observed among the three analytical methods.

**Analysis 2.** The MP analysis of the six-locus combined data set resulted in six equally most parsimonious trees (960 steps, CI = 0.969, RI = 0.971), which yielded a well-resolved strict consensus tree (tree not shown). The ML tree search returned one most likely tree (−ln L = 13,042.541; tree not shown). The Bayesian analysis resulted in a majority-rule consensus tree with a well-resolved backbone (fig. 1B). All three analytical methods returned highly congruent topologies without conflicts. *Azolla nilotica* and the accessions of *A. pinnata* are sister taxa in a well-supported clade (BPMP = 100; BPML = 100; PP = 1.00). This group is sister to the rest of the genus. In this larger clade, *A. fliculoides* and *A. rubra* are sister taxa (BPMP = 100; BPML = 100; PP = 1.00) and form a sister group to the *A. caroliniana* clade (CAR-MIC-MEX) (BPMP = 100; BPML = 100; PP = 1.00). There is little resolution within this group, but our single accession of *A. caroliniana* is consistently sister to all of the *A. microphylla* and *A. mexicana* accessions (BPMP = 100; BPML = 100; PP = 1.00); the latter two species are both not monophyletic. The *A. mexicana* 1 accession was sister to the remaining *A. microphylla* and *A. mexicana* accessions with strong support (BPMP = 88; BPML = 86; PP = 1.00); in addition, *A. mexicana* 2 and *A. microphylla* 2 were always sister taxa (BPMP = 99; BPML = 100; PP = 1.00), as were *A. mexicana* 3 and *A. microphylla* 3.

There was no conflict between the topologies recovered in analyses 1 and 2 (fig. 1A, 1B). However, we were able to obtain a more resolved topology by conducting analysis 2, which resulted in three well-supported nodes within the CAR-MIC-MEX clade, compared with one in analysis 1 (well supported: BPMP ≥ 70; BPML ≥ 70; PP ≥ 0.95).

**Divergence Time Estimates**

The average branch lengths of the topology used for the divergence time estimation were based on the atpB, rbcL, and rps4 data and are shown in figure 1C. Age estimates from the PL analysis, as well as age estimates and standard deviations from the PL bootstrap analysis for all well-supported nodes in the Bayesian consensus tree, are presented in table 3. The age estimate data are graphically depicted in figure 2 as a chronogram plotted against the geologic timescale.

Our results from the PL r8s analysis estimate the divergence of sections *Azolla* and *Rhizosperma* at 50.7 Ma (Eocene; fig. 2, node 2). *Azolla nilotica* and *A. pinnata* (section *Rhizosperma*) are estimated to have diverged at 32.5 Ma (Oligocene; fig. 2, node 3). The *A. fliculoides-A. rubra* (FL-RUB) clade and the CAR-MIC-MEX clade (section *Azolla*) diverged at 16.3 Ma (Miocene; fig. 2, node 4), and *A. fliculoides* and *A. rubra* diverged at 3.1 Ma (Pliocene; fig. 2, node 5). *Azolla caroliniana* (CAR-MIC-MEX species complex) diverged from *A. microphylla* and *A. mexicana* at 3.9 Ma (Pliocene; fig. 2,
node 6). Accessions within the MIC-MEX complex diverged at 2.3 Ma (Pliocene; fig. 2, node 7). All estimates from the PL analysis were within 1 SD of their estimates from the bootstrap analysis (table 3).

**Discussion**

**Phylogenetic Relationships**

Our six-locus plastid data set resulted in a topology congruent with the results of Reid et al.’s (2006) analysis of nrITS, *trnL-trnF*, and *atpB-rbcL*. Previous studies suggested that *Azolla nilotica* belonged in a separate subgenus, *Tetrasporocarpia* (Saunders and Fowler 1993), due to its diploid chromosome number (2n=52 for *A. nilotica* vs. 2n=44 for all other *Azolla* species) and numerous unique morphological traits (e.g., sporocarps in groups of four and cauline multicellular trichomes). However, both Reid et al.’s (2006) study and our study confirm the monophyly of the traditional sections *Azolla* and *Rhizosperma* (Mettenius 1847; Tan et al. 1986; Wagner 1997), which form well-supported sister clades (fig. 1), each with clearly defined morphological characters (fig. 2). Morphological similarities between *A. pinnata* and section *Azolla* species are likely due to parallel evolution or to apomorphic changes in *A. nilotica*. *Azolla rubra* and *A. filiculoides* are closely related sister taxa (fig. 1B) and, as with Reid et al.’s (2006) work, little resolution is achieved within the CAR-MIC-MEX species complex, despite our data set having nearly twice as much data. Accessions of *A. microphylla* and *A. mexicana* are paraphyletic to one another and show little to no evidence of constituting separate evolutionary lineages (see fig. 1B). *Azolla caroliniana* is sister to a clade comprising both *A. microphylla* and *A. mexicana* accessions. Reid et al. (2006) showed this same relationship and also provided evidence for the monophyly of *A. caroliniana*. Based on the results of their three-locus study, Reid et al. (2006) recommended recognizing *A. caroliniana* as a separate species but suggested that *A. microphylla* and *A. mexicana* be treated as a single species. The results of our six-locus data support the conclusions of Reid et al. (2006) and do not confirm the morphological conclusion of Evrard and Van Hove (2004) that the entire CAR-MIC-MEX complex should be recognized as a single species. Future investigations into the MIC-MEX complex require greatly increased taxon sampling and the use of additional molecular tools, such as intersimple sequence repeats (Barker and Hauk 2003) and amplified fragment length polymorphisms (Perrie

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**Fig. 1** A, Analysis 1: Bayesian phylogram depicting the topology and branch lengths in *Azolla* obtained from the three-locus coding data set (*atpB*, *rbcL*, and *rps4*); *Salvinia* and *Marsilea* are outgroups. An asterisk highlights the well-supported node for *Azolla mexicana* 2 plus *A. microphylla* 2. B, Analysis 2: Bayesian phylogram depicting the topology and branch lengths in *Azolla* sect. *Azolla* topology generated using the six-locus coding and noncoding data set (*atpB*, *rbcL*, *rps4*, *rps4-trnS*, *trnG-trnK*, and *trnL-trnF*); *Azolla* sect. *Rhizosperma* is used as an outgroup. An asterisk highlights the well-supported node for *A. mexicana* 3 plus *A. microphylla* 3. C, Summary topology used for divergence time estimation; branch lengths as calculated from the three-locus data set (*atpB*, *rbcL*, and *rps4*). For all topologies shown, thickened lines indicate well-supported branches from all three measures (maximum parsimony bootstrap percentage [BPMP ≥ 70]; maximum likelihood bootstrap percentage [BPML ≥ 70]; posterior probability [PP ≥ 0.95]).
et al. 2003). It is important to note that several taxonomic and nomenclatural issues pertaining to this species complex have been identified in an unpublished thesis (Dunham 1986), as well as alluded to by Dunham and Fowler (1987). These complications will need to be carefully considered in any future study of this group.

**Divergence Time Estimates and the Fossil Record**

This study is the first to provide a temporal framework for evolution in *Azolla* based exclusively on extant lineages. *Azolla* possesses a tremendous fossil diversity, with more than 50 extinct species of *Azolla* described (Collinson 1980, 2001). Many are represented solely by megaspores and exhibit wide variation in morphology, particularly with regard to float number (Collinson 1980). The earliest stratigraphic occurrence (89 Ma) is the many-floated *A. montana* Hall in the Maastrichtian/Late Cretaceous (table 3; Collinson 1991).

Our 89-Ma fixed-age calibration on the root node is identical to the minimum age constraint on the *Salvinia/Azolla* divergence used by Pryer et al. (2004) and is similar to their molecular age estimate of 89.17 ± 0.69 Ma for the Salviniaeae. We evaluated our molecular divergence time estimates for *Azolla* against the earliest reported occurrences of extant *Azolla* taxa. We estimate the diversification within *Azolla* to have begun at 50.7 Ma (Eocene; fig. 2, node 2) with the divergence of sections *Azolla* and *Rhizosperma*. However, the first fossil record of an *Azolla* species with a collared megaspore, a trait shared by all extant *Azolla*, does not occur until 37.2–28.4 Ma (Oligocene; *A. colwellensis* Collinson; Collinson 1980).

*Azolla pinnata* is recorded from the Pliocene (1.8–5.3 Ma) (Florschutz 1945; Batten and Kovach 1990), and the oldest fossil record for *A. nilotica* is from only 550 AD (Birks 2002), whereas our results suggest that *A. pinnata* diverged from *A. nilotica* at 32.5 Ma (Oligocene; fig. 2, node 3). This discordance between the fossil and molecular ages suggests either (1) that the lineages leading to *A. pinnata* and *A. nilotica* diverged in the Oligocene while the extant species did not appear until much more recently (e.g., the Pliocene) or (2) that older fossils of extant species have not yet been found.

*Azolla filiculoides* is reported from the Langhian (13.65–15.97 Ma) (Mai 2001), and the oldest fossil records for the CAR-MIC-MEX complex are attributed to *A. caroliniana* from the Upper Miocene (5.3–11.6 Ma) (Gamerro 1981; Batten and Kovach 1990). We estimate 16.3 Ma (Miocene) for the divergence of CAR-MIC-MEX from the FIL-RUB lineage (fig. 2, node 4), which is in accordance with the *A. filiculoides* fossil from the Langhian (13.65–15.97 Ma).

Within the CAR-MIC-MEX complex, our analyses indicate the MIC-MEX lineage diverged from *A. caroliniana* at
3.9 Ma (fig. 2, node 6) and that divergences within the MIC-MEX lineage occurred as recently as 1.9 Ma (fig. 2, node 7). One fossil record for *A. microphylla* exists from 50,000 years ago (Schofield and Colinaux 1969). Given the lack of morphological distinction between the members of the recently derived MIC-MEX lineages, we do not see any conflict between our age estimates and the fossil record for this group. No fossil specimens have been assigned to *A. mexicana*, so no comparisons were possible for this species. *Azolla rubra* also does not have any described fossils, but this is likely due to the overall similarity of *A. rubra* and *A. filiculoides* megaspores.

The literature on *Azolla* fossils consistently refers to a progressive reduction in the number of megaspore floats, from 24, through 18, 15, 12, and nine, to three. Unfortunately, our study of extant species alone cannot inform us about the possible pattern of reduction in megaspore float number in *Azolla* across geologic time (Saunders and Fowler 1993). We can simply determine that there was a basal split in the genus at about 50 Ma (fig. 2, node 2), with one lineage giving rise to extant taxa that have three floats and the other lineage to taxa with nine. Explaining this megaspore float pattern will require the combined phylogenetic analysis of fossil and extant species of *Azolla*.

### Acknowledgments

We are grateful to Gerald Peters for material of nearly all *Azolla* specimens sampled in this study; Nathalie Nagalingum for material of *Azolla rubra*; Suzanne Fredericq for material of *Salvinia minima*; Michal Skakuj for laboratory assistance; David Hearn for providing a Perl script to automate the divergence time estimation bootstrap analyses; and Eric Schuettpelz, Nathalie Nagalingum, Petra Korall, and Rick Lupia for advice and assistance with analyses and for comments on earlier versions of this manuscript. We are grateful to an anonymous reviewer and to Richard Saunders, who provided valuable insight and information in review, especially with regard to persistent taxonomic problems that have been identified in an unpublished thesis by D. G. Dunham. We thank Jill Reid for allowing us to view a preprint of her research before its publication. Our research was supported by National Science Foundation grants DEB-0089909 (K. M. Pryer and H. Schneider) and DEB-0347840 (K. M. Pryer).

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