Testing the monophyly of *Spathiphyllum*, and the relationship between Asian and tropical American species.

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ABSTRACT

As part of a broader phylogenetic study of the subfamily Monsteroideae we used three plastid and one nuclear markers to test the monophyly of the genus *Spathiphyllum*. Our results support the tribe Spathiphyllum. Our results support the tribe Spathiphyllum. The monophyly of *Spathiphyllum*. The monotypic genus *Holochlamys* appears nested within *Spathiphyllum* in a clade with the Asian species and *S. cannifolium*. However, support for this clade is low. The section *Spathiphyllum* shows a high support and is sister to the remaining species in the genus. As a consequence *Spathiphyllum* remains as

the only disjunct aroid genus between the Old and New World Tropics.

INTRODUCTION

With the increased application molecular data for phylogenetic analysis, the understanding of relationships Araceae has been improved considerably (Cabrera, 2008, Cusimano, 2011, Henríquez, Additionally, new fossils 2014). biogeographical studies have helped to elucidate the historical and geographic distribution patterns in the family (Nauheimer, 2012a). As a result, classification aroid has changed dramatically; several genera have been resurrected; and new genera have been created (Cabrera, 2008, Cusimano, 2011). Nevertheless, many critical issues are still in need of more data.

Before this molecular era there were thought to be three geographically disjunct genera between the old world and new world tropics. Species in the genera Homalomena, Schismatoglottis and Spathiphyllum were found in both tropical America and tropical Asia, particularly within the Malay Archipelago and Melanesia (van Steenis, 1962, Grayum, 1990, Mayo et al., 1997). However, recent studies (Barabé, 2002, Gauthier, 2008, Wong et al., 2013) have shown evidence challenging the monophyly of disjunct Schismatoglottis and Homalomena (they are no longer disjunct in their revised circumscriptions), leaving Spathiphyllum as the only remaining aroid genus with a tropical disjunction.

The genus Schismatoglottis, with only three species in South America, and about 150 in belongs Asia, tropical to the tribe Schismatoglottidae that in its current circumscription comprises 11 genera. Several of these genera are small groups recognized thanks to molecular data (Wong et al., 2010a, 2010b, Low, 2014). The three Neotropical species previously recognized in Schismattoglotis are now in the resurrected genus Philonotion (Wong, 2010c) that belong to its own tribe Philonotieae. Equally, the genus Homalomena is considerable more diverse in Tropical Asia with about 500 species, versus only 10 species in Tropical

America (Boyce & Croat, 2013). Unlike Schismatoglottis, molecular evidence for the separation of Neotropical Homalomena is not conclusive (Barabé, 2002, Gauthier, 2008, Wong et al., 2013). The most plausible assignment options the for Neotropical Homalomena would be resurrection of the genus Adelomena, or the combination of these species Homalomena with Philodendron subgenus Pterosmischum in the genus Elopium (Wong et al., 2013). Thus, more extensive sampling and more molecular data are necessary to fully understand the relationships among subgenera Homalomena, and the of Philodendron.

Finally, we have the genus Spathiphyllum, which has a different distribution pattern. The genus is more diverse in Tropical America with about 50 species (Cardona, 2004), and only three species in Southeast Asia. Spathiphyllum commutatum Schott is widespread in Malesia and Micronesia: S. solomonense Nicolson restricted to the Solomon Islands; and S. schlechteri (Engl. & K.Krause) Nicolson is restricted to New Guinea. Neotropical species are concentrated in the northern Andes, with only a few species reaching Central America and Mexico (Cardona, 2004), and one species, S. leave reaches Cocos Island in the Pacific Ocean west of Costa Rica. No comprehensive phylogenetic studies, until now, have included more than three Spathiphyllum from Tropical America, or any of the Asian species.

Spathiphyllum together with Holochlamys becarii form the tribe Spathiphylleae in the subfamily Monsteroideae (Cabrera, Cusimano, 2011). The relationships of the tribe Spathiphylleae within Monsteroideae have been controversial, but recent studies strongly support Spathiphylleae as sister to the remaining Monsteroideae (Chartier, 2014, Henriquez, 2014). These two genera are very similar morphologically ecologically, differing mainly in a few characteristics of the flowers. Whereas Spathiphyllum species have two to three locules and ovules with axillar placentation, Holochamys has one locule and ovules with basal placentation (Bunting, 1960, Mayoet al.,1997). Nevertheless, anatomical studies suggest that the Holochlamys ovary could actually have two locules, placentation may not be basal (Eyde, 1967, Carvell, 1989).

Currently, Spathiphyllum is divided into four sections based the fusion of the perianth segments, the form and length of the pistils relative to the perianth, and the attachment the spathe to the peduncle (Bunting, 1960). Asian species together with S. cannifolium and S. leave form the section Massowia, which is recognized by having the perianth segments completely connate forming a cup around the pistil. This trait led Bunting (1960) to propose the monophyly of this group. However, Williams & Dressler (1967) and Grayum (1984, 1990) suggested this could be an artificial group with Holochlamys nested within it. Sections Amomophyllum, Dysspathiphyllum, and Spathiphyllum have free perianth segments.

As part of a broader systematic study of the subfamily Monsteroideae (Zuluaga, unpublished data) we tested the monophyly of *Spathiphyllum* and *Holochlamys*, and investigated the relationships of the Asian and American species of *Spathiphyllum*.

METHODS

as outgroups.

Taxon Sampling and DNA sequencing. During the sampling for the study of systematics of the subfamily Monsteroideae we sequenced a total of seven accessions of Spathiphyllum from Tropical genus America, S. commutatum and S. solomonense from Tropical Asia, and one accession of Holochlamys becarii. Additional sequences for phryniifolium obtained were from S. GENBANK. For this study we used representatives from all ten additional

Monsteroideae genera, plus six other aroids

DNA was extracted from silica-dried plant material using the DNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA) according to the manufacturer's specifications. samples were extracted using a CTAB method adapted from Doyle & Doyle (1987). We sequenced four plastid markers, matK, rbcL, trnC-petN spacer, and partial ycf1. PCR products were purified using ExoSap-It (Cleveland, OH). Sequencing reactions BigDye Terminator used Cycle the Sequencing Kit, using and cleaned Agencourt CleanSeq (Beverly, MA) magnetic beads. BigDye Terminator Cycle Sequencing Ready Reaction Kit (Applied Biosystems, Foster City, CA, USA). Samples were sequenced using PE-Biosystems version 3.7 of Sequencing Analysis at the University Wisconsin–Madison Biotechnology Center.

Molecular analysis. All sequences were manually edited in Geneious 6.0 (Biomatters, http://www.geneious.com). Sequences were aligned in Geneious 6.0 using a combination of Muscle (Edgar 2004) and MAFFT (Katoh 2002). Finally, all alignments were checked manually. We performed individual combined phylogenetic and analyses. Analyses for each marker/dataset were performed under Maximum Likelihood using RaxML 7.0 (Statamakis, 2006), and MrBayes 3.1.2 on XSEDE, as implemented in the CIPRES portal. For RaxML, we used the GTRGAMMA model in all cases. For Bayesian analysis models were calculated using Jmodeltest2 (Darriba, 2012).

RESULTS.

So far at least one of the four molecular markers has been successfully sequenced for S. cannifolium (two accessions), cochlearispathum, S. commutatum, S. floribundum, S. minor, S. patulinervum, S. solomonense, S. wallisii, and Holochlamys becarii. Only the trnCpetN spacer was sequenced for the Asian Spathiphyllum. The combined alignment length is 4970 bp with 36% missing data. Despite using four molecular markers the identical sites between pairs of species within Spathiphyllum is more than 97%. This low variation is common within the genera of the subfamily Monsteroideae (Unpublished data).

Trees derived from individual analyses congruent, however the are resolution is low except for a clade comprising the species of the section combined Spathiphyllum. Thus, a analysis is presented (Figure 1). individual and combined analyses show the tribe Spathiphylleae as monophyletic and sister to the remaining Monsteroideae, thus supporting the topology presented Chartier (2014) and Henríquez (2014). Section Spathiphyllum represented by S. cochlearispathum, S. phryniifolium, and S. wallisii shows strong support, and is sister to the Spathiphylleae. remaining The genus Holochlamys appears within nested Spathiphyllum in a clade with the Asian species and one accession of S. cannifolium (section Massowia). Section Amomophyllum (S. minor, S. floribundum, S. patulinervum) and one accession of S. cannifolium are sister to the containing Holochlamys. However support for the last two clades is low (Figure 1).

DISCUSSION.

A monophyletic tribe Spathiphylleae has been supported since the first molecular studies in Araceae, based on only a few genes, were published (Tam, 2004, Cabrera, 2008). However, the position of Spathiphyllleae as sister to rest of Monsteroideae is only recovered in the most recent studies with more data included

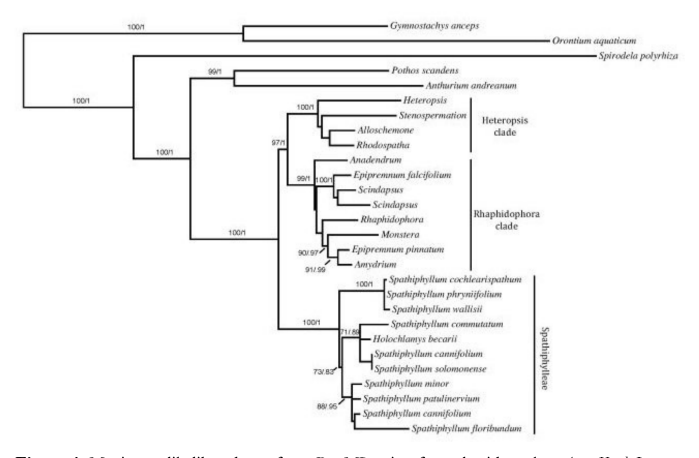


Figure 1. Maximum likelihood tree from RaxML using four plastid markers (*matK*, *rbcL*, *trnC-petN*spacer, and partial *ycf1*). Bootstrap supports >75 and Bayesian posterior probabilities >0.85 are shown on the branches.

(Chartier, 2014, Henriquez, 2014). Our results strongly support this topology (Figure 1). Most of the synapomorphies defining Monsteroideae as presented in Cusimano (2011) seem to be appropriate to be assigned to the clade all Monsteroideae containing Spathiphylleae. These synapomorphies and the alternative characters in Spathiphylleae are: pollen zonate (ring-like) (vs. pollen multiaperturate with surface exine

striate/polyplicate); spathe soon deciduous with a distinct basal abscission (vs. Spathe marcescent), large trichosclereids not in bundles (vs. smaller trichosclereids in bundles). The only synapomorphy remaining for Monsteroideae is the presence of trichosclereids.

Within the tribe Spathiphylleae, the support for the position of *H. becarii* is not conclusive, but some markers suggest this

species belong may to the genus Spathiphyllum. Holochlamys appears consistently embedded in a clade with section Massowia. These two genera are very similar in habit and morphology (Bunting,1960), stem vasculature (French & Tomlison, 1981), palynological features (Grayum,1992), and floral anatomy (Carvell, 1989). Moreover, Holochlamys shares with section Massowia the presence of fused perianth parts forming a cup around the pistil (Bunting, 1960, Carvell, 1989). The proposed differences between these two genera - a unilocular ovary, and ovules with basal placentation - have been challenged by more detailed anatomical studies (Eyde, 1967). Carvell (1989) undertook a very detailed anatomical study of Monsteroideae and Pothoideae, and reported unilocular ovaries, and basal ovules for H. becarii. However, he also reported the presence of an incomplete septum in several flowers, and the attachment of some the ovules to that septum. Additionally, Carvell (1989) found considerable intra- and interspecific variation in locule, ovules, tepals, and stamen numbers in Spathiphyllum. Despite the fact that we consider there to be no strong morphological evidence to separate Holochlamys from Spathiphyllum (especially from the section Massowia), our analysis currently lacks adequate resolution within this clade, and so we prefer to be cautious at this point in time, necessarily gathering more data before proposing any taxonomic changes for these species.

Our sampling did not allow a complete assessment of the sectional classification in

Spathiphyllum as proposed by Bunting (1960). The only section with strong support through all our analysis is the section Spathiphyllum, which is also supported by the morphology of their flowers with a long and conical pistil exceeding the perianth (Bunting, 1960). Williams and Dressler (1967), and Grayum (1984, 1990), discussed hypotheses for the origin and biogeography of Spathiphyllum and Holochlamys. We are undertaking additional molecular biogeographical analyses of the subfamily Monsteroideae, and more detailed discussion on this topic will be presented in subsequent papers.

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