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Research article

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A phylogenetic framework of the legume genus *Aeschynomene* for comparative genetic analysis of the Nod-dependent and Nod-independent symbioses

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69 SUMMARY

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• Some *Aeschynomene* legume species have the property of being nodulated by photosynthetic *Bradyrhizobium* lacking the *nodABC* genes. Knowledge of this unique Nod (factor)-independent symbiosis has been gained from the model *A. evenia* but our understanding remains limited due to the lack of comparative genetics with related taxa using a Nod-dependent process.

• To fill this gap, this study significantly broadened previous taxon sampling, including in allied genera, to construct a comprehensive phylogeny. This backbone tree was matched with data on chromosome number, genome size, low-copy nuclear genes and strengthened by nodulation tests and a comparison of the diploid species.

The phylogeny delineated five main lineages that all contained diploid species while
polyploid groups were clustered in a polytomy and were found to originate from a single
paleo-allopolyploid event. In addition, new nodulation behaviours were revealed and Noddependent diploid species were shown to be tractable.

• The extended knowledge of the genetics and biology of the different lineages in the legume genus *Aeschynomene* provides a solid research framework. Notably, it enabled the identification of *A. americana* and *A. patula* as the most suitable species to undertake a comparative genetic study of the Nod-independent and Nod-dependent symbioses.

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99 Keywords

100 *Aeschynomene*, genetics, legumes, nodulation, phylogenetics, polyploidy, symbiosis

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103 INTRODUCTION

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105 In the field of nitrogen-fixing symbiosis, scientists have a long-standing interest in the tropical 106 papilionoid legume genus Aeschynomene since the discovery of the ability of the species A. 107 afraspera to develop abundant nitrogen-fixing stem nodules (Hagerup, 1928). This nodulation 108 behavior is uncommon in legumes, being shared by very few hydrophytic species of the genera Discolobium, Neptunia and Sesbania, but it is exceptionally widespread among the 109 110 semi-aquatic Aeschynomene species (Alazard, 1985; Boivin et al., 1997; Chaintreuil et al., 111 2013). These stem-nodulating Aeschynomene species are able to interact with Bradyrhizobium 112 strains that display the unusual property of being photosynthetic (Giraud et al., 2000; Miché 113 et al., 2010). Most outstanding is the evidence that some of these photosynthetic 114 Bradyrhizobium strains lack both the nodABC genes required for the synthesis of the key 115 "Nod factors" symbiotic signal molecules and a type III secretion system (T3SS) that is 116 known in other rhizobia to activate or modulate nodulation (Giraud et al., 2007; Okazaki et 117 al., 2013, 2015). These traits revealed the existence of an alternative symbiotic process 118 between rhizobia and legumes that is independent of the Nod factors.

119 As in the legume genus Arachis (peanut), Aeschynomene uses an intercellular symbiotic 120 infection process instead of infection thread formation that can be found in other legume 121 groups (Sprent et al., 2017). This led to the suggestion that the Nod-independent process 122 might correspond to the ground state of the rhizobial symbiosis, although we cannot exclude 123 that it represents an alternative symbiotic process compared to the one described in other 124 legumes (Sprent & James, 2008; Madsen et al., 2010, Okubo et al., 2012). It is noteworthy 125 that all the Nod-independent species form a monophyletic clade within the Aeschynomene 126 phylogeny and jointly they also display striking differences in the bacteroid differentiation 127 process compared to other Aeschynomene species (Chaintreuil et al., 2013; Czernic et al., 128 2015). To decipher the molecular mechanisms of this distinct symbiosis, the Nod-independent 129 A. evenia has been used as a new model legume, because its genetic and developmental 130 characteristics (diploid with a reasonable genome size -2n=20, 415 Mb/1C-, short perennial 131 and autogamous, can be hybridized and transformed) make this species tractable for 132 molecular genetics (Arrighi et al., 2012, 2013, 2015). Functional analyses revealed that some 133 symbiotic determinants identified in other legumes (SYMRK, CCaMK, HK1 and DNF1) are 134 recruited, but several key genes involved in bacterial recognition (e.g. LYK3), symbiotic 135 infection (e.g. EPR3 and RPG), and nodule functioning (e.g. DNF2 and FEN1) were found 136 not to be expressed in A. evenia roots and nodules, based on RNAseq data (Czernic et al.,

2015; Fabre *et al.*, 2015; Chaintreuil *et al.*, 2016a; Nouwen *et al.*, 2017). This suggested that
the Nod-independent symbiosis is distinct from the Nod-dependent one.

139 Forward genetics are now expected to allow the identification of the specific molecular 140 determinants of the Nod-independent process in A. evenia (Arrighi et al., 2012; Chaintreuil et 141 al., 2016a). In addition, comparing A. evenia with closely related Nod-dependent 142 Aeschynomene species will promote our understanding how symbiosis evolved in legumes. 143 The genus Aeschynomene (restricted now to the section Aeschynomene as discussed in 144 Chaintreuil et al. (2013)) is traditionally composed of three infrageneric taxa, subgenus 145 Aeschynomene (which includes all the hydrophytic species) and subgenera Bakerophyton and 146 Rueppellia (Rudd, 1955; Gillet et al., 1971). The genus has also been shown to be 147 paraphyletic, with a number of related genera being nested within it, but together they form a 148 distinct clade in the tribe Dalbergieae (Chaintreuil et al., 2013; Rudd, 1981; Lavin et al., 149 2001; Klitgaard et al., 2005; LPWG, 2017). Within this broad clade, two groups of semi-150 aquatic Aeschynomene have been well-studied from a genetic and genomic standpoint: the A. 151 evenia group, which contains all the Nod-independent species (most of them being 2x), and 152 the A. afraspera group (all species being Nod-dependent) that appears to have a 4x origin 153 (Arrighi et al., 2014, Chaintreuil et al. 2016b, 2018). For comparative analyses, the use of 154 Nod-dependent species with a diploid structure would be more appropriate, but such 155 Aeschynomene species are poorly documented.

156 To overcome these limitations, our aim was to produce a species-comprehensive phylogenetic 157 tree supplemented with genetic and nodulation data. For this, we made use of an extensive 158 taxon sampling in both the genus *Aeschynomene* and in closely related genera to capture the 159 full species diversity of the genus and to clarify phylogenetic relationships between taxa. For 160 most species, we also documented chromosome number, genome size and molecular data for 161 low-copy nuclear genes, thus allowing the identification of diploid species as well as 162 untangling the genome structure of polyploid taxa. In addition, these species were 163 characterized for their ability to nodulate with various Bradyrhizobium strains containing or lacking *nod* genes and, finally, the diploid species were submitted to a comparative analysis 164 165 of their properties. In light of the data obtained in this study, we discuss the interest of two 166 Aeschynomene species, A. americana and A. patula, to set up a comparative genetic system to 167 complement the A. evenia model.

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170 MATERIALS & METHODS

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172 Plant material

All the accessions of *Aeschynomene* used in this study, including their geographic origin and collection data are listed in Tables S1 and S4. Seed germination and plant cultivation in the greenhouse were performed as indicated in Arrighi *et al.* (2012). Phenotypic traits such as the presence of adventitious root primordia and nodules on the stem were directly observed in the glasshouse.

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179 Nodulation tests

180 Nodulation tests were carried out using Bradyrhizobium strains ORS278 (originally isolated from A. sensitiva nodules), ORS285 (originally isolated from A. afraspera nodules), 181 ORS285 (Giraud and DOA9 (originally isolated from A. americana nodules) (Giraud et al., 182 183 2007; Teamtisong et al., 2014; Bonaldi et al., 2011). Bradyrhizobium strains were cultivated 184 at 34°C for seven days in Yeast Mannitol (YM) liquid medium supplemented with an 185 antibiotic when necessary (Howieson et al., 2016). Plant in vitro culture was performed in 186 tubes filled with buffered nodulation medium (BNM) as described in Arrighi et al. (2012). 187 Five-day-old plants were inoculated with 1 mL of bacterial culture with an adjusted OD at 188 600nm to 1. Twenty one days after inoculation, six plants were analysed for the presence of 189 root nodules. Nitrogen-fixing activity was estimated on the entire plant by measurement of 190 acetylene reducing activity (ARA) and microscopic observations were performed using a 191 stereo-macroscope (Nikon AZ100, Champigny-sur-Marne, France) as published in Bonaldi et 192 al. (2011).

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194 Molecular methods

195 Plant genomic DNA was isolated from fresh material using the classical CTAB (Cetyl 196 Trimethyl Ammonium Bromide) extraction method. For herbarium material, the method was 197 adapted by increasing the length of the incubation (90 min), centrifugation (20 min) and 198 precipitation (15 min) steps. The nuclear ribosomal internal transcribed spacer region (ITS), 199 the chloroplast *matK* gene and four low-copy nuclear genes (*CYP1*, $eiF1\alpha$, SuSy, and TIP1;1) 200 previously identified in the A. evenia and A. afraspera transcriptomes were used for 201 phylogenetic analyses (Arrighi et al., 2014; Chaintreuil et al., 2016). The genes were PCR-202 amplified, cloned and sequenced as described in Arrighi et al. (2014) (Table S2). For genomic 203 DNA extracted from herbarium specimens, a battery of primers was developed to amplify the

different genes in overlapping fragments as short as 250 bp (Table S2). The DNA sequences

205 generated in this study were deposited in GenBank (Table S3).

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207 Phylogenetic analyses and traits mapping

Sequences were aligned using MAFFT (*--localpair –maxiterate 1000*; Katoh & Standley, 209 2013). Phylogenetic reconstructions were performed for each gene as well as for concatenated 210 datasets under a Bayesian approach using Phylobayes 4.1b (Lartillot & Philippe, 2004) and 211 the site-heterogeneous CAT+F81+ Γ 4 evolution model. For each analysis, two independent 212 chains were run for 10,000 Phylobayes cycles with a 50% burn-in. Ancestral states 213 reconstruction was done through stochastic character mapping using the Phytools R package 214 (Revell, 2012) running 10 simulations for each character.

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216 Species networks and hybridizations

217 To test if the phylogeny obtained by concatenating the four low-copy nuclear genes (CYP1, 218 $eiF1\alpha$, SuSy, and TIP1;1) was most likely obtained by gene duplications followed by 219 differential losses or by a combination of duplications, losses coupled with one or several 220 allopolyploidy events involving A. patula and Soemmeringia semperflorens, the method 221 presented in To & Scornavacca (2015) was used. In short, this method computes a 222 reconciliation score by comparing a phylogenetic network and one or several gene trees. The 223 method allows allopolyploidy events at hybridization nodes while all other nodes of the 224 network are associated to speciation events; meanwhile, duplication and loss events are 225 allowed at a cost (here, arbitrarily fixed to 1) on all nodes of the gene tree.

226 Thus, the set of 4 nuclear gene trees was used to score different phylogenetic networks 227 corresponding to four different potential evolutionary histories. Two alternative networks with 228 no reticulation corresponding to the two topologies obtained either with the group A (T1) or 229 group B (T2) served to evaluate a no-allopolyploidisation hypothesis. The topology yielding 230 the best score (T2) served to generate and compare all phylogenetic networks with one or two 231 hybridization nodes, involving A. patula and/or S. semperflorens, to test successively a one-232 allopolyploidisation scenario (N1-best) and a two-allopolyploidisation evolutionary scenario 233 (N2-best).

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235 **GBS analysis**

A GBS library was constructed based on a described protocol (Oueslati *et al.*, 2017). For each
sample, a total of 150 ng of genomic DNA was digested using the two-enzyme system, PstI

(rare cutter) and Mse (common cutter) (New England Biolabs, Hitchin, UK), by incubating at
37°C for 2 h. The ligation reaction was performed using the T4 DNA ligase enzyme (New
England Biolabs, Hitchin, UK) at 22°C for 30 min and the ligase was inactivated at 65°C for
30 min. Ligated samples were pooled and PCR-amplified using the Illumina Primer 1
(barcoded adapter with PstI overhang) and Illumina Primer 2 (common Y-adapter). The
library was sequenced on an Illumina HiSeq 3000 (1x 150 pb) (at the Get-PlaGe platform in
Toulouse, France).

- 245 The raw sequence data were processed in the same way as in the study described in Garsmeur 246 et al. (2018). SNP calling from the raw Illumina reads was performed using the custom 247 python pipeline VcfHunter (available at https://github.com/SouthGreenPlatform/VcfHunter/) 248 (Guillaume Martin, CIRAD, France). For all samples, these sequence tags were aligned to the 249 A. evenia 1.0 reference genome (JF Arrighi, unpublished data). The SNP results from all the 250 samples were converted into one large file in VCF format and the polymorphism data were 251 subsequently analysed using the web-based application SNiPlay3 (Dereeper et al., 2015). 252 First, the SNP data were treated separately for each species and filtered out to remove SNP 253 with more than 10% missing data as well as those with a minor allele frequency (MAF) of 254 0.01 using integrated VCFtools. Second, an overall representation of the species diversity 255 structures was obtained by making use of the PLINK software as implemented in SNiPlay3. 256 This software is based on the multidimensional-scaling (MSD) method to produce two-257 dimensional plots. The Illumina HiSeq 3000 sequencing raw data are available in the NCBI 258 SRA (Sequence Read Archive) under the study accession number: SRP149516.
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260 Genome size estimation and chromosome counting

261 Genome sizes were measured by flow cytometry using leaf material as described in Arrighi et 262 al. (2012). Genome size estimations resulted from measurements of three plants per accession 263 and Lycopersicum esculentum (Solanaceae) cv "Roma" (2C = 1.99 pg) was used as the 264 internal standard. The 1C value was calculated and the conversion factor 1 pg DNA = 978 Mb 265 was used to express it in Mb/1C. To count chromosome number, metaphasic chromosomes 266 were prepared from root-tips, spread on slides, stained with 4',6-diamidino-2-phenylindole 267 (DAPI) and their image captured with a fluorescent microscope as detailed in Arrighi et al. 268 (2012).

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271 **RESULTS**

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273 A comprehensive phylogeny of the genus Aeschynomene and allied genera

274 To obtain an in-depth view of the phylogenetic relationships within the genus Aeschynomene 275 subgenus Aeschynomene, which contains the hydrophytic species, we significantly increased 276 previous sampling levels by the addition of new germplasm accessions and, if these were not 277 available, we used herbarium specimens. DNA was isolated for 40 out of the 41 species 278 (compared to the 27 species used in Chaintreuil et al. (2013)) included in this group in 279 taxonomic and genetic studies (Table S1) (Arrighi et al., 2014; Chaintreuil et al., 2012, 280 2016b, 2018; Rudd, 1955). In addition, to determine the phylogenetic relationship of this 281 subgenus with respect to Aeschynomene subgenera Bakerophyton and Rueppellia, unclassified 282 Aeschynomene species, as well as with the allied genera Bryaspis, Cyclocarpa, Geissaspis, 283 Humularia, Kotschya, Smithia and Soemmeringia, representatives of these 10 taxa were also 284 sampled (compared to the 5 taxa present in Chaintreuil et al. (2013)) (Rudd, 1981; Lewis, 285 2005). This added 21 species to our total samples (Table S1). The dalbergioid species Pictetia 286 angustifolia was used as outgroup (Chaintreuil et al. 2013; LPWG, 2017).

287 Phylogenetic reconstruction of all the taxa sampled was undertaken using Bayesian analysis 288 of the chloroplast *matK* gene and the nuclear ribosomal *ITS* region (Table S2 and S3). The 289 matK and ITS gene trees distinguished almost all the different Aeschynomene groups and 290 related genera (Fig. S1 and S2). The two phylogenetic trees have a very similar topology 291 although some branches of both trees can be lowly supported. Incongruences were also 292 observed for A. deamii and the genus Bryaspis, but the conflicting placements are poorly 293 supported and were thus interpreted as a lack of resolution typical of single-marker trees, 294 rather than strong incongruence. To improve the phylogenetic resolution among the major 295 lineages, the *matK* gene and the *ITS* sequence datasets were combined into a single 296 phylogenetic analysis where only well-supported nodes were considered (posterior probability 297 $(PP) \ge 0.5$ (Fig. 1). Our analysis recovered a grade of five main lineages with a branching 298 order that received robust support (PP \geq 0.92): (1) a basally branching lineage represented by 299 A. americana, (2) an A. montevidensis lineage, (3) an A. evenia lineage corresponding to the 300 Nod-independent clade (Arrighi et al., 2012, 2014), (4) a newly-identified lineage containing 301 A. patula and (5) a lineage represented by an unresolved polytomy clustering the A. afraspera 302 clade (Chaintreuil et al., 2016b) with all the remaining taxa. 303 In large part, our work also provided good species-level resolution and demonstrated that

Aeschynomene subgenus *Aeschynomene* (as currently circumscribed) is interspersed on the phylogenetic tree with the lineage containing *A. patula*, the two other subgenera of 306 Aeschynomene and a number of other genera related to Aeschynomene (Fig. 1) (Chaintreuil et 307 al., 2013; Lavin et al., 2001; LPWG, 2017; Du Puy et al., 2002). The combined analysis also 308 grouped the genus Bryaspis with the species related to A. afraspera in a highly supported 309 clade but its exact position with respect to other taxa remained inconclusive, as previously 310 observed (Fig. 1) (Chaintreuil et al., 2013). Most noticeably, several intergeneric relationships 311 are consistently recovered, notably sister-clade relationships between *Cyclocarpa* and *Smithia* 312 as well as between Aeschynomene subgenera Bakerophyton and Rueppellia together with the 313 genus Humularia (referred to as the BRH clade herein after) (Fig. 1). This clade supports 314 previous observations of a morphological continuum between Aeschynomene subgenus 315 Rueppellia and the genus Humularia and brings into question their taxonomic separation 316 (Gillett *et al.*, 1971).

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Ploidy level of the species and genomic structure of the different lineages

319 The revised Aeschynomene phylogeny was used as a backbone tree to investigate the 320 evolution of ploidy levels. Previous studies had demonstrated that the A. evenia clade is 321 mostly diploid (2n=2x=20) even if some species such as A. indica (2n=4x=40, 2n=6x=60)322 appear to be of recent allopolyploid origin (Arrighi et al., 2014; Chaintreuil et al., 2018). 323 Conversely, all the species of the A. afraspera group were found to be polyploid 324 (2n=4x=28,38,40, 2n=8x=56,76) and to have a common AB genome structure but the origin of the polyploidy event remained undetermined (Chaintreuil et al., 2016b). To assess the 325 326 ploidy levels in *Aeschynomene* species and related genera, chromosome numbers and nuclear 327 DNA content were determined (appended to labels in Fig. 2a, Table S1, Fig. S3 and S4). We 328 provide evidence that the lineages containing A. americana, A. montevidensis, A. evenia and 329 A. patula, as well as Soemmeringia semperflorens, are diploid with 2n=20, with the smallest 330 2x genome found in A. patula (0.58 pg/2C) and the largest 2x genome in A. deamii (1.93 331 pg/2C). With the exception of S. semperflorens, all the groups that are part of the polytomy 332 were characterized by higher chromosome numbers 2n=28,36,38,40 (up to 76). These 333 chromosome numbers equate to approximately twice that of the diploid species (except for 334 2=28), suggesting that the corresponding groups are most probably polyploid. Species with 335 chromosome numbers departing from 2n=40 are likely to be of disploid origin as already 336 described in the A. afraspera clade (Chaintreuil et al., 2016b). Here again, important genome 337 size variations ranging from 0.71 pg/2C for the *Geissaspis* species to 4.82 pg/2C for the 4x A. 338 schimperi highlight the genomic differentiation of the various taxa (Fig. 2a, Table S1).

339 To firmly link chromosome numbers to ploidy levels and to clarify genetic relationships 340 between the different lineages, we cloned and sequenced four nuclear-encoded low-copy 341 genes in selected species: CYP1 (Cyclophilin 1), eiF1a (eukaryotic translation initiation factor 342 α), SuSy (Sucrose Synthase) and TIP1;1 (tonoplast intrinsic protein 1;1) (Table S2). For all 343 diploid species, only one gene sequence was obtained, while for all the polyploid species, in 344 almost all cases, a pair of putative homeologues was isolated, thus confirming their genetic 345 status inferred from the karyotypic data (Table S3). In general, the duplicated copies were 346 highly divergent and nested in two different major clades in the resulting Bayesian phylogenic 347 trees generated for each gene (Fig. S5). One clade contained all the A copies (except for one 348 anomalous sequence for *Bryaspis lupulina* in the $eiF1\alpha$ tree) and the other clade gathered all 349 the B copies previously identified in A. afraspera (Chaintreuil et al., 2016b). These two 350 clades A and B did not always receive high support, however it is notable that the A copies 351 formed a monophyletic group with, or sister to, the A. patula sequence and similarly the B 352 copies with, or sister to, the *Soemmerignia semperflorens* sequence, in all gene trees (Fig. S5). 353 In an attempt to improve phylogenetic resolution, the four gene data sets were concatenated. 354 This combination resulted in a highly supported Bayesian tree that places the A copy clade as 355 the sister to the diploid A. patula (PP = 1), and the B copy clade as sister to the diploid S. 356 semperflorens (PP =1) (Fig. 2b). As a result, these phylogenetic analyses combined to 357 karyotypic data show that all the five main lineages contain diploid species. They also reveal 358 that all the polyploid groups share the same AB genome structure, with the diploid A. patula 359 and S. semperflorens species being the closest modern representatives of the ancestral donors 360 of the A and B genomes.

361 In addition, an ancestral state reconstruction analysis performed on the *ITS+matK* phylogeny 362 indicates that diploidy is the ancestral condition in the whole revised group and that 363 tetraploidy most likely evolved once in the polytomy (Fig. S6). To provide support on a 364 probable single origin of the allopolyploidy event, separate and concatenated nuclear gene 365 trees were further used for a phylogenetic network analysis. In this analysis, the two non-366 allopolyploidisation hypotheses (T1 and T2) were found to be more costly (scores of 207 and 367 196) than the two hypotheses allowing for hybridization (N1-best and N2-best with scores of 368 172 and 169, respectively) (Fig. S7a-d). The one-allopolyploidisation hypothesis (N1-best) 369 strongly indicates that a hybridization involving the lineages that contain A. patula and S. 370 semperflorens gave rise to all the polyploid groups (Fig. S7c). Although the two-371 allopolyploidisation hypothesis (N2-best) yielded the absolute best score, the score 372 improvement was very low (169 vs 172) and the resulting network included the hybridization inferred with the one-allopolyploidisation hypothesis making this latter hypothesis mostprobably the correct one (Fig. S7d).

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376 Nodulation properties of the different Aeschynomene lineages

377 Species of Aeschynomene subgenus Aeschynomene are known to be predominantly 378 amphibious and more than 15 of these hydrophytic species (found in the A. evenia and A. 379 afraspera clades, as well as A. *fluminensis*) have been described as having the ability to 380 develop stem nodules (Boivin et al., 1997; Chaintreuil et al., 2016b; Lock, 1989; Rudd, 381 1955). In A. fluminensis, these nodules are observed only on submerged stems (as also seen in 382 the legume Discolobium pulchellum), while they occur on aerial stems within the A. evenia 383 and A. afraspera clades (Fig. 3a) (Alazard & Duhoux, 1987; Chaintreuil et al., 2013; Loureiro 384 et al., 1994, 1995). Phenotypic analysis of representatives of the different lineages under 385 study revealed that they all display adventitious root primordia along the stem (Fig. 3a,b). 386 Adventitious roots are considered to be an adaptation to temporary flooding and they also 387 correspond to nodulation sites in stem-nodulating Aeschynomene species (Fig. 3b) (Alazard & 388 Duhoux, 1987). Given that the A. evenia and A. afraspera clades are now demonstrated not to 389 share the same genomic components provides a genetic argument for independent 390 developments of stem nodulation by photosynthetic bradyrhizobia. Reconstruction of 391 ancestral characters based on the ITS+matK phylogeny confirmed that the whole group was 392 ancestrally a wet ecology taxon endowed with adventitious root primordia but that the stem 393 nodulation ability evolved several times, as previously inferred (Fig. S8, S9; and S10) 394 Chaintreuil et al., 2013, 2016b).

To investigate whether the newly studied species could be nodulated by photosynthetic 395 396 bradyrhizobia, we extended the results obtained by Chaintreuil et al. (2013) by testing the 397 nodulation abilities of 22 species available (listed in Fig. 4a) for which adequate seed supply 398 was available. Three different strains of Bradyrhizobium equating to the three cross-399 inoculation (CI) groups defined by Alazard (1985) were used: DOA9 (non-photosynthetic 400 Bradyrhizobium of CI-group I), ORS285 (photosynthetic Bradyrhizobium with nod genes of 401 CI-group II) and ORS278 (photosynthetic Bradyrhizobium lacking nod genes of CI-group 402 III). These strains were used to inoculate the 22 species and their ability to nodulate them was 403 analysed at 21 dpi. For this, we recorded nodule formation and compared nitrogen fixation 404 efficiency by an acetylene reduction assay (ARA) and observation of plant vigor. Nodulation 405 was observed on all species tested except for *Smithia sensitiva* that had a problem with root 406 development, A. montevidensis and S. semperflorens. For these three species, either the
407 culture conditions or the Bradyrhizobium strains used were not appropriate (Fig. 4a).

408 The non-photosynthetic strain DOA9 displayed a wide host spectrum but was unable to 409 nodulate the Nod-independent species, A. deamii, A. evenia and A. tambacoundensis. The 410 photosynthetic strain ORS285 efficiently nodulated A. afraspera and the Nod-independent 411 Aeschynomene species (Fig 4a), as previously reported (Chaintreuil et al., 2013). 412 Interestingly, the ORS285 strain was also able to induce nitrogen-fixing nodules in A. patula 413 and ineffective nodules were observed on A. fluminensis and the genera Bryaspis, Cyclocarpa 414 and Smithia (Fig. 4a). To examine if in these species the nodulation process relies on a Nod-415 dependent or Nod-independent symbiotic process, we took advantage of the availability of a 416 Δnod mutant of the strain ORS285. None of them were found to be nodulated by 417 ORS285 Δ nod, suggesting that the nodule formation depended on a Nod signaling in these 418 species (Fig. 4a). In fact, the ORS285 Δ nod mutated strain was able to nodulate only species 419 of the A. evenia clade, similarly as to the photosynthetic strain ORS278 naturally lacking nod-420 genes (Fig. 4a). Analysis of the evolution of these nodulation abilities by performing an 421 ancestral state reconstruction on the revisited phylogeny indicated several emergences of the 422 ability to interact with photosynthetic bradyrhizobia and a unique emergence of the ability to 423 be nodulated by the *nod* gene-lacking strain as observed earlier (Fig. S11 and Fig. S12) 424 (Chaintreuil et al., 2013). From these nodulation tests, different nodulation patterns emerged 425 for the diploid Aeschynomene species (as detailed in Fig. 4b-d) with the DOA9 and ORS278 426 strains being specific to the Nod-dependent and Nod-independent groups respectively and 427 ORS285 showing a gradation of compatibility between both.

428

429 Diversity of the diploid species outside the Nod-independent clade

430 To further characterize the diploid species that fall outside of the Nod-independent clade, in 431 which A. evenia lies, they were analysed for their developmental properties and genetic 432 diversity (Fig. 5a). All species are described as annuals or short-lived perennials (Du Puy et 433 al., 2002; Lewis, 2005; Rudd, 1955). As for A. evenia, A. americana, A. villosa, A. 434 *fluminensis*, A. *parviflora* and A. *montevidensis* are robust and erect, reaching up to 2 m high 435 when mature, whilst A. patula and S. semperflorens are creeping or decumbent herbs. These 436 differences in plant habit are reflected by the important variation in seed size between these 437 two groups (Fig. 5a). This has an impact on plant manipulation, because for A. patula and S. 438 semperflorens seed scarification needs to be adapted (25 min with concentrated sulfuric acid 439 instead of 40 min for the other species) and *in vitro* plant growth takes slightly more time to 440 get a root system sufficiently developed for inoculation with *Bradyrhizobium* strains (10 days-441 post-germination instead of the 5-7 dpi for other species) (Arrighi et al., 2012). Consistent 442 flowering and seed production was observed for A. americana, A. villosa, A. patula and S. 443 semperflorens when grown under full ambient light in the tropical greenhouse in short days 444 conditions as previously described for A. evenia, making it possible to develop inbred lines by 445 successive selfing (Fig. 5a) (Arrighi et al., 2012). For A. fluminensis, A. parviflora and A. 446 *montevidensis*, flowering was sparse or not observed, indicating that favorable conditions for 447 controlled seed set were not met (Fig. 5a).

448 Five species (A. villosa, A. fluminensis, A. parviflora, A. montevidensis and S. semperflorens) 449 are strictly American while A. americana is a pantropical species and A. patula is endemic to 450 Madagascar (Du Puy et al., 2002; Lock, 1989; Rudd, 1955). Several species have a narrow 451 geographic distribution or seem to be infrequent, explaining the very limited accession 452 availability in seedbanks (Fig. 5a). This is in sharp contrast with both A. americana and A. 453 villosa that are well-collected, being widely found as weedy plants and sometimes used as 454 component of pasture for cattle (Fig. 5a) (Cook et al., 2005). To assess the genetic diversity of 455 these two species, a germplasm collection containing 79 accessions for A. americana and 16 456 accessions for A. villosa, and spanning their known distribution was used (Table S4). A 457 Genotyping-By-Sequencing (GBS) approach resulted in 6370 and 1488 high quality 458 polymorphic SNP markers for A. americana and A. villosa accessions, respectively. These two SNP datasets subsequently served for a clustering analysis based on the 459 460 multidimensional-scaling (MSD) method. The MSD analysis distinguished three major 461 groups of accessions for both A. americana and A. villosa along coordinate axes 1 and 2 (Fig. 462 5b). When mapping the accessions globally, the three groups identified for A. villosa were 463 observed together in Mexico and only group (3) extended to the northern part of South 464 America (Fig. 5c, Table S4). Converly, a clear geographical division was observed for A. 465 americana with group (1) occupying the central part of South America, group (2) being found 466 in the Caribbean area while group (3) was present in distinct regions from Mexico to Brazil 467 and in across the Palaeotropics (Fig. 5c, Table S4). A. americana is hypothesized to be native 468 in America and naturalized elsewhere (Cook et al., 2005). The observed distributions in 469 combination with the MSD analysis, accessions being tightly clustered in group (3) compared 470 to groups (1) and (2), support this idea and indicate that group (3) recently spread worldwide.

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- 472

473 **DISCUSSION**

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475 A well-documented phylogenetic framework for the legume genus *Aeschynomene*

476 We produced a new and comprehensive phylogeny of the genus *Aeschynomene* and its closely 477 related genera complemented by gene data sets, genome sizes, karyotypes and nodulation 478 assays. For plant genera, there are few for which documentation of taxonomic diversity is so 479 extensive and supported by a well-resolved, robustly supported phylogeny which reveals the 480 evolutionary history of the group (Govindarajulu et al., 2011). Here, the whole group, which 481 includes the genus Aeschynomene with its 3 subgenera and its 7 allied genera, is shown to 482 have experienced cladogenesis leading to five main lineages, including the Nod-independent 483 clade, with diploid species found in all these lineages. The multigene data analysis provided 484 robust evidence that two of them, represented by the two diploid species A. patula and S. 485 semperflorens, are involved in an ancient allotetraploidization process that gave rise to the 486 different polyploid lineages clustering in a polytomy. Separate allopolyploidization events 487 from the same diploid parents or a single allopolyploid origin are plausible explanations for 488 the formation of these lineages. However, the consistent resolution of the phylogenetic tree 489 obtained with the combined gene data, where A. patula and S. semperflorens are sisters to the 490 A and B subgenomic sequences, favours the hypothesis of a single allopolyploid origin, as 491 also argued for other ancient plant allopolyploid events in Asimitellaria (Saxifragaceae) and 492 Leucaena (Leguminosae) (Govindarajulu et al., 2011; Okuyama et al., 2012). The 493 phylogenetic network analysis also supports the one-allopolyploidisation hypothesis. 494 However, additional nuclear genes will be needed to conclusively confirm that no additional 495 hybridization event occurred. Although not the focus of the present study, it is worth noting 496 that most diploid species are found in the Neotropics, the two modern representatives of the A 497 and B genome donors that gave rise to the 4x lineages are located on different continents (S. 498 semperflorens in South America and A. patula in Madagascar) and that all the 4x lineages are 499 located in the Palaeotropics (Lewis et al., 2005). This raises questions about the evolution of 500 the whole group and the origin of the 4x lineages. In addition, the presence of a polytomy 501 suggests that this allopolyploid event preceded a rapid and major diversification of 4x groups 502 that have been ascribed to different Aeschynomene subgenera or totally distinct genera that 503 altogether represent more than 80% of the total species of the whole group (LPWG, 2017; 504 Whitefiel & Lockhart, 2007). Diversification by allopolyploidy occurred repeatedly in the 505 genus Aeschynomene since several neopolyploid species are found in both the A. evenia clade 506 and the A. afraspera clade as exemplified by A. indica (4x, 6x) and A. afraspera (8x) (Arrighi 507 et al., 2014; Chaintreuil et al., 2016b). Dense sampling for several Aeschynomene taxa or 508 clades also a more precise delimitation of species boundaries (for morphologically similar 509 taxa but which are genetically differentiated or correspond to different cytotypes) and 510 revealed intraspecific genetic diversity that is often geographically-based as showed for the pantropical species A. americana (this study), A. evenia, A. indica and A. sensitiva 511 512 (Chaintreuil et al., 2018). All these Aeschynomene species share the presence of adventitious 513 root primordia on the stem that correspond to the infection sites for nodulation. The consistent 514 presence of adventitious root primordia in all taxa of the whole group, together with an 515 ancestral state reconstruction, substantiates the two-step model proposed earlier for the 516 evolution of stem nodulation in Aeschynomene, with a common genetic predisposition at the 517 base of the whole group to produce adventitious root primordia on the stem, as an adaptation 518 to flooding, and subsequent mutations occurring independently in various clades to enable 519 stem nodulation (Chaintreuil et al., 2013). The ability to interact with photosynthetic 520 bradyrhizobia that are present in aquatic environments also appears to have evolved several 521 times. This photosynthetic activity is important for the bacterial symbiotic lifestyle as it 522 provides energy usable for infection and subsequently for nitrogenase activity inside the stem 523 nodules (Giraud et al., 2000). To date, natural occurrence of nodulation by photosynthetic 524 bradyrhizobia has been reported only for the A. evenia and A. afraspera clades, and for A. 525 fluminensis (Loureiro et al., 1995; Miché et al., 2010; Molouba et al., 1997). Nevertheless, we 526 could not test the photosynthetic strains isolated from A. fluminensis nodules and the nature of 527 the strains present in those of the newly studied species A. patula has not been investigated 528 yet. They would allow the comparison of their nodulation efficiency with the reference 529 photosynthetic Bradyrhizobium ORS278 and ORS285 strains. In addition, we can ask if the 530 semi-aquatic lifestyle and/or nodulation with photosynthetic bradyrhizobia may have 531 facilitated the emergence of the Nod-independent symbiosis in the A. evenia clade.

532

533 *Aeschynomene* species for a comparative analysis of nodulation with *A. evenia*

534 To uncover whether the absence of detection for several key symbiotic genes in the root and 535 nodule transcriptomic data of A. evenia are due to gene loss or extinction, and to identify the 536 specific symbiotic determinants of the Nod-independent symbiosis, a genome sequencing 537 combined with a mutagenesis approach is presently being undertaken for A. evenia in our 538 laboratory. A comparative analysis with Nod-dependent Aeschynomene species is expected to 539 consolidate this genomic and genetic analysis performed in A. evenia by contributing to 540 elucidate the genetic changes that enabled the emergence of the Nod-independent process. 541 Phylogenomics and comparative transcriptomics, coupled with functional analysis, are

542 undergoing increased development in the study of symbiosis. They enabled unravelling gene 543 loss linked to the lack of developing a symbiosis in certain plant lineages but also to identify 544 new symbiosis genes (for arbuscular mycorrhizal symbiosis (Delaux et al., 2014; Bravo et al., 545 2016); for the nodulating symbiosis (Delaux et al., 2015; Griesmann et al., 2018). 546 Comparative work on symbiotic plants is often hindered, however, either by the absence of 547 closely related species which display gain or loss of symbiotic function or, when these are 548 present, by the lack of a well-understood genetic framework, as outlined in Behm et al. 549 (2014), Delaux et al. (2015), Geurts et al. (2016), Sprent (2017). Nevertheless, in the case of 550 the nodulating *Parasponia*/non-nodulating *Trema* system, a fine comparative analysis was 551 very powerful to demonstrate a parallel loss of the key symbiotic genes NFR5, NIN and RGP, 552 in the non-nodulating species, challenging the long-standing assumption that *Parasponia* specifically acquired the potential to nodulate (Behm et al., 2014; Geurts et al., 2016; van 553 554 Velzen et al., 2018). In this respect, the uncovering of the genetic evolution of the genus 555 Aeschynomene and related genera along with the identification of diploid species outside of 556 the Nod-independent clade, provided a robust phylogenetic framework that can now be 557 exploited to guide the choice of Nod-dependent diploid species for comparative genetic 558 research. Among these, some species are discarded because of the lack of nodulation with 559 reference Bradyrhizobium strains or the inability to produce seeds under greenhouse 560 conditions. Based on efficient nodulation, short flowering time and ease of seed production, A. americana (2n=20, 600 Mb) and A. patula (2n=20, 270 Mb) appear to be the most 561 562 promising Nod-dependent diploid species to develop a comparative genetic system with A. evenia (2n=20, 400 Mb). In contrast to A. evenia, A. americana is nodulated only by non-563 564 photosynthetic bradyrhizobia, and in this respect, it behaves in a similar way to other legumes. 565 This species is widespread in the tropics, so that adequate quantities of germplasm are 566 available, and it has already been subject to detailed studies, notably to isolate its nodulating 567 Bradyrhizobium strains, among which is the DOA9 strain (Noisangiam et al., 2012; Teamtisong et al., 2014). As A. americana belongs to the most basal lineage in the 568 569 Aeschynomene phylogeny, it may be representative of the ancestral symbiotic mechanisms 570 found in the genus. On the other hand, A. patula has a restricted Madagascan distribution with 571 only one accession being available, but it is of interest due to its relatively smaller plant size 572 and genome size (actually the smallest diploid genome in the group) making this species the 573 "Arabidopsis" of the genus Aeschynomene. As for A. americana, this species is efficiently nodulated by non-photosynthetic bradyrhizobia, but it is also compatible with the 574 575 photosynthetic nod gene-containing ORS285 strain. This property makes A. patula 576 particularly interesting as it allows direct comparisons of mechanisms and pathways between 577 it and A. evenia without the problem of potential strain effects on symbiotic responses. In 578 addition, when considering the Aeschynomene phylogeny, A. patula is more closely related to 579 A. evenia than is A. americana, and so it may be more suitable to demonstrate the changes 580 necessary to switch a Nod-dependent to a Nod-independent process or vice-versa. Developing 581 sequence resources and functional tools for A. americana and/or A. patula is now necessary to 582 set up a fully workable comparative Aeschynomene system. In the long run, handling such a 583 genetic system will be instrumental in understanding how photosynthetic *Bradyrhizobium* and 584 some Aeschynomene species co-evolved and in unravelling the molecular mechanisms of the 585 Nod-independent symbiosis.

586 587

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599 AUTHOR CONTRIBUTIONS

J.F.A. designed the experiments. L.B., C.C., R.R., J.F., M.G.M, S.C.B., C.H., M.D. and J.F.A.
performed the experiments and obtained the data. P.S., C.S., L.M. undertook the phylogenetic
analyses. P.M., J.Q., G.P.L., X.P., A.D'H., E.G. and J.F.A. analysed the data. M.G., R.D., H.
Randriambanona, H. Ramanankierna, H.V. and M.Z. contributed to the acquisition and
analysis of accessions. J.F.A. wrote the paper. L.B. and C.C contributed equally. All authors
read and approved the final manuscript.

- 606 607
- 608 **REFERENCES**
- 609

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- 772
- 773

774 FIGURE LEGENDS

775

Figure 1: Phylogeny of the genus *Aeschynomene* and allied genera.

The Bayesian phylogenetic reconstruction was obtained using the concatenated *ITS* (Internal Transcribed Spacer) + *matK* sequences. Numbers at branches indicate posterior probability above 0.5. The five main lineages are identified with a circled number and the two previously studied *Aeschynomene* groups are framed in a red box bordered with a dashed line. On the right are listed *Aeschynomene* subgenus *Aeschynomene* (in green), other *Aeschynomene* subgenera or species groups (in blue) and related genera (in orange).

783

784 Figure 2: Genomic characteristics and phylogenetic relationships.

785 (a) Simplified Bayesian ITS+matK phylogeny with representative species of different lineages 786 and groups. The A. evenia, A. afraspera and BRH (Bakerophyton-Rueppelia-Humularia) 787 clades are represented by black triangles and the polytomy is depicted in bold. Chromosome 788 numbers are indicated in brackets. (b) Phylogenetic relationships based on the combination of 789 4 concatenated nuclear low-copy genes (CYP1, eifla, SuSy and TIP1;1 genes detailed in 790 Figure S5). Diploid species (2n=20) are in blue, polyploid species $(2n\geq28)$ in black. The A 791 and B subgenomes of the polyploid taxa are delineated by red and green boxes in dashed 792 lines, respectively. Nodes with a posterior probability inferior to 0.5 were collapsed into 793 polytomies. Posterior probability above 0.5 are indicated at every node. (c) The one-794 allopolyploidisation hypothesis (N1-best) obtained with the phylogenetic network analysis 795 based on the T2 tree with reticulations in blue (detailed in Fig. S7).

796

797 Figure 3: Occurrence of adventitious root primordia and of stem nodulation.

798 (a) Simplified Bayesian ITS+matK phylogeny of the whole group with the A. evenia, A. 799 afraspera and BRH (Bakerophyton-Rueppelia-Humularia) clades represented by black 800 triangles. The polytomy is depicted in bold. The shared presence of adventitious root 801 primordia is depicted on the stem by a blue circle. Dashed red boxes indicate groups 802 comprising aerial stem-nodulating species. Asterisks refer to illustrated species in (b) for 803 aerial stem-nodulation. (b) Stems of representatives for the different lineages and groups. 804 Small spots on the stem correspond to dormant adventitious root primordia and stem nodules 805 are visible on the species marked by an asterisk. Bars: 1cm.

806

807 Figure 4: Comparison of the root nodulation properties.

(a) Species of different lineages and groups that were tested for nodulation are listed in the
 simplified Bayesian phylogeny on the left. Root nodulation tests were performed using the

B10 DOA9, ORS285, ORS285 Δ nod and ORS278 strains. E, effective nodulation; e, partially B11 effective nodulation; i, ineffective nodulation, -, no nodulation; blank, not tested. (b) Number B12 of nodules per plant, (c) relative acetylene-reducing activity (ARA) and (d) aspect of the B13 inoculated roots developing nodules or not (some nodules were cut to observe the B14 leghemoglobin color inside) after inoculation with *Bradyrhizobium* DOA9, ORS285 and B15 ORS278 on *A. americana*, *A. patula*, *A. afraspera* and *A. evenia*. Error bars in (b) and (c) B16 represent s.d. (n=6). Scale bar in (d): 1 mm.

817

818 Figure 5: Characteristics of diploid species.

819 (a) Development and germplasm data for species that are listed in the simplified phylogeny on 820 the left. A. evenia from the Nod-independent clade (NI) is also included for comparison. 821 Germplasm numbers correspond to the sum of accessions available at CIAT, USDA, Royal 822 Botanic Gardens (Kew), AusPGRIS, IRRI and at LSTM. (b) Multi-dimensional scaling 823 (MSD) plots of the genetic diversity among A. americana (left) and A. villosa (right) 824 accessions according to coordinates 1 and 2 (C1, C2). Identified groups are delimited by 825 circles and labeled with numbers. (c) Geographical distribution of the of the A. americana and 826 A. villosa accessions. Taxon colours and group numbers are the same as in (b). Details of the 827 accessions are provided in Table S4.

828

Figure S1: *matK* phylogeny of the genus *Aeschynomene* and allied genera.

Bayesian phylogenetic reconstruction obtained using the chloroplast *matK* gene. Numbers atbranches are posterior probability.

832

Figure S2: *ITS* phylogeny of the genus *Aeschynomene* and allied genera.

834 Bayesian phylogenetic reconstruction obtained using the Internal Transcribed Spacer (*ITS*)

- sequence. Numbers at branches are posterior probability.
- 836

837 Figure S3: Chromosome numbers in *Aeschynomene* species.

- 838 Root tip metaphase chromosomes stained in blue with DAPI (4',6-diamidino-2-phenylindole).
- 839 Chromosome numbers are indicated in brackets. Scale bars: $5 \,\mu m$.
- 840

841 Figure S4: Chromosome numbers in species of genera related to *Aeschynomene*

- Root tip metaphase chromosomes stained in blue with DAPI (4',6-diamidino-2-phenylindole).
- 843 Chromosome counts are indicated in brackets. Scale bars: $5 \mu m$.

Figure S5: Phylogenetic trees based on nuclear low-copy genes.

Bayesian phylogenetic reconstructions obtained for the *CYP1*, *eif1a*, *SuSy* and *TIP1;1* genes. Diploid species (2n=20) are in blue, polyploid species $(2n\geq28)$ in black excepted *A. afraspera* for which the A and B gene copies are distinguished in red and green respectively. -A, -A1, -A2, -B, -B1 and -B2 indicated the different copies found. Putative A and B subgenomes of the polyploid taxa are delineated by red and green boxes in dashed lines, respectively. Numbers at branches represent posterior probability.

852

Figure S6: Ancestral state reconstruction of ploidy levels in the genus *Aeschynomene* and allied genera.

Ancestral state reconstruction was estimated in SIMMAP software using the 50% majorityrule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Ploidy levels are indicated by different colors. Unknown ploidy levels are denoted by a dash.

858

Figure S7: Phylogenetic networks based on the four nuclear *CYP1*, *eif1a*, *SuSy* and *TIP1;1* genes.

(a) No-allopolyploidisation hypothesis (T1) based on the concatenated gene tree obtained
taking into account the group A (Fig. 2b). (b) No-allopolyploidisation hypothesis (T2) based
on the concatenated gene tree obtained taking into account the group B (Fig. 2b). (c) Oneallopolyploidisation hypothesis (N1-best). (d) Two-allopolyploidisation hypothesis (N2-best).
Blue lines indicate reticulations while other nodes of the network are associated to speciation

866 events. Reconciliation scores obtained for each phylogenetic network are indicated.

867

Figure S8: Ancestral state reconstruction of adventive root primordia in the genus *Aeschynomene* and allied genera.

Ancestral state reconstruction was estimated in SIMMAP software using the 50% majorityrule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Data on the adventitious root primordia come from the present analysis and pertinent previously published data. Presence or not of adventitious root primordia is indicated by different colors.

874

Figure S9: Ancestral state reconstruction of ecological habit in the genus *Aeschynomene*and allied genera.

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877 Ancestral state reconstruction was estimated in SIMMAP software using the 50% majority-

rule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Data on

the species ecology come from pertinent previously published data. Ecological habits are

- 880 indicated by different colors.
- 881

Figure S10: Ancestral state reconstruction of the aerial stem nodulation ability in the genus *Aeschynomene* and allied genera.

- Ancestral state reconstruction was estimated in SIMMAP software using the 50% majorityrule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Data on the occurrence of stem nodulation come from pertinent previously published data. Occurrence or not of stem nodulation is indicated by different colors.
- 888

Figure S11: Ancestral state reconstruction of the ability to nodulate with the photosynthetic *Bradyrhizobium* strains in the genus *Aeschynomene* and allied genera.

- Ancestral state reconstruction was estimated in SIMMAP software using the 50% majorityrule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Data on nodulation with photosynthetic *Bradyrhizobium* strains come from the present analysis and pertinent previously published data. Nodulation with photosynthetic *Bradyrhizobium* strains is considered positive only if reported as occurring naturally or being efficient *in vitro*.
- 896

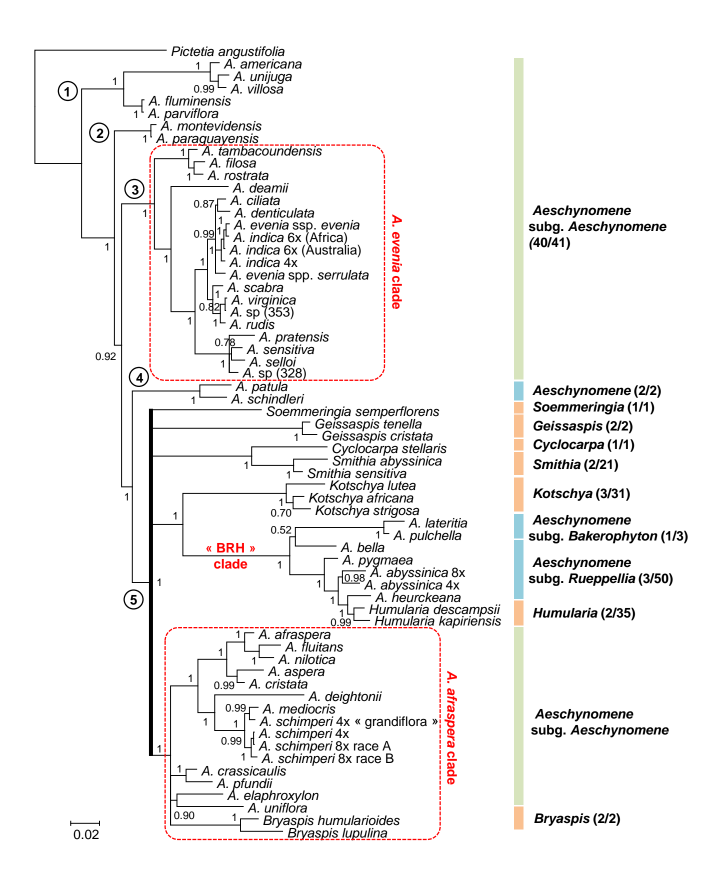
Figure S12: Ancestral state reconstruction of the ability to nodulate with the photosynthetic *Bradyrhizobium* strain ORS278 in the genus *Aeschynomene* and allied genera.

900 Ancestral state reconstruction was estimated in SIMMAP software using the 50% majority-

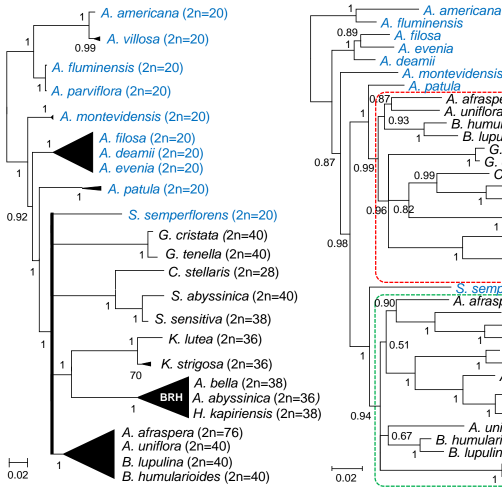
⁹⁰¹ rule topology obtained by Bayesian analysis of the combined *ITS+matK* sequences. Data on

nodulation with ORS278 come from the present analysis and pertinent previously published

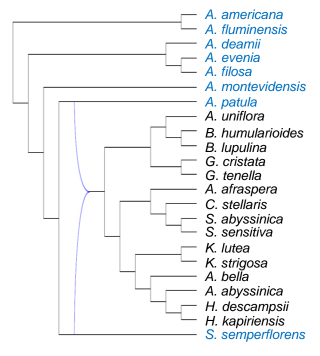
data. Ability or not to nodulate with ORS278 is indicated by different colors.



a ITS + matK



C Phylogenetic network



b Concatenated nuclear genes

