Stem nodulation: diversity and occurrence in Aeschynomene and Sesbania legumes from wetlands of Madagascar Faustin F. Manantsoa¹, Marrino F. Rakotoarisoa², Clémence Chaintreuil³, Adamson T.E. Razakatiana¹, Frédéric Gressent⁴, Marjorie Pervent⁴, Mickaël Bourge⁵, Martial D. Andrianandrasana¹, Nico Nouwen³, Herizo Randriambanona¹, Heriniaina Ramanankierana¹, Jean-François Arrighi^{3*} ¹Laboratoire de Microbiologie de l'Environnement-Centre National de Recherches sur l'Environnement BP 1739 Fiadanana Antananarivo Madagascar ²Department of Ethnobotany and Botany, National Center for Applied Pharmaceutical Research, Antananarivo 101, Madagascar; ³IRD, Plant Health Institute of Montpellier (PHIM), UMR IRD/SupAgro/INRAE/UM/CIRAD, TA-A82/J -Campus de Baillarguet, 34398 Montpellier, France. ⁴INRAE, Plant Health Institute of Montpellier (PHIM), UMR IRD/SupAgro/INRAE/UM/CIRAD, TA-A82/J - Campus de Baillarguet, 34398 Montpellier, France. ⁵Cytometry Facility, Imagerie-Gif, Université Paris-Saclay, CEA, CNRS, Institute for Integrative Biology of the Cell (I2BC), 91198, Gif-sur-Yvette, France. (*Author for correspondence: email jean-francois.arrighi@ird.fr)

34 Abstract

35

As an adaptation to flooding, few legume species have the original ability to develop nitrogen-fixing nodules on the stem. By surveying wetlands of Madagascar, we found a large occurrence and diversity of stem nodulation in Aeschynomene and Sesbania legumes. They represent opportunities to investigate different modalities of the nitrogen-fixing symbiosis in legumes.

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42 Introduction

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44 The symbiosis between legume plants and soil rhizobia results in the formation of nitrogen-fixing 45 nodules, generally exclusively appearing on the roots. However, in a handful of tropical legumes growing in wetlands, nodulation with rhizobia can also occur at stem-located dormant root 46 47 primordia, a process that is referred as stem nodulation. Seen the semi-aquatic lifestyle of these 48 legumes, it has been hypothesized that the stem nodulation trait is an evolutionary adaptation to 49 flooding (Ladha et al., 1992). Stem nodulation was first described for the African Aeschynomene 50 afraspera (not to be confounded with Aeschynomene aspera found in Asia) (Hagerup, 1928). Stem 51 nodulation gained agricultural interest after the discovery that the profuse stem nodulation as found 52 in A. afraspera and Sesbania rostrata results in a high nitrogen fixation activity (Dreyfus and 53 Dommergues, 1981; Alazard and Becker, 1987). To date, stem nodulation has been reported for 54 species belonging to four legume genera: Aeschynomene, Discolobium, Neptunia and Sesbania 55 (Boivin et al., 1997). While, in the three latter genera, stem-nodulation has been described for one or 56 very few species, more than 20 Aeschynomene species have been shown to form stem nodules 57 (Chaintreuil et al., 2013; 2016). These legume species can differ in their stem nodulation ability and 58 intensity, and actually S. rostrata and A. afraspera are the ones for which stem-nodulation is the 59 most profuse (Boivin et al. 1997a).

60 Strikingly, S. rostrata is stem-nodulated only by Azorhizobium caulinodans while other 61 Sesbania species root-nodulate with rhizobia of the Ensifer genus (Boivin et al., 1997a,b). Similarly, 62 different types of Bradyrhizobium strains have been identified as nodulating Aeschynomene species, 63 (Boivin et al., 1997a; Alazard, 1985). The most important difference between the strains were the 64 presence or absence of photosynthetic activity and nod genes to produce Nod factors (Molouba et 65 al., 1999; Giraud et al., 2007; Miché et al., 2011). So far, photosynthetic Bradyrhizobium strains have 66 been exclusively found in nodules of stem-nodulating Aeschynomene species (Boivin et al., 1997a) 67 and strains lacking nod genes have been isolated from nodules of Aeschynomene species that cluster

in a single clade (Chaintreuil *et al.*, 2013, 2016). The *Bradyhrizobium* ORS278-*A. evenia* interaction
serves as model for the deciphering of the latter very specific interaction called Nod-independent
symbiosis (Quilbé *et al.*, 2021). In contrast, strains having *nod* genes, such as *Bradyrhizobium*ORS285, have been isolated from nodules of *A. afraspera* that is one of the *Aeschynomene* species
using a Nod-dependent interaction (Arrighi *et al.*, 2014; Brottier *et al.*, 2018). *S. rostrata* stemnodulating *A. caulinodans* is not photosynthetic and has *nod* genes, but in both cases, the symbiotic
interaction is very specific.

75 Although research has shed some light on the genetics of both partners in the 76 Bradyrhizobium-Aeschynomene and A. caulinodans-S. rostrata symbiotic systems, our understanding 77 of the molecular mechanism that causes stem nodulation is still in its infancy. Furthermore, our 78 knowledge of the diversity and occurrence of stem nodulation in natura is relatively limited as it has 79 been investigated in only a few geographical regions (e.g., James et al., 2001; Molouba et al., 1999; 80 Miché et al., 2011). To fill this gap, we conducted a field study in Madagascar that contains a variety 81 of wetlands with an important plant biodiversity. A series of expeditions were organized to explore 82 wetland-rich regions in the Central (RN1-Itasy Lake), Northern (Nosy Be), Western (RN4-Majunga) 83 and Eastern (RN2-Alaotra Lake) parts of Madagascar (Fig. 1a). In these regions, we found stem-84 nodulated Aeschynomene, Sesbania and Neptunia spp. Ommiting these latters from this analysis, as 85 nodules were only found on floating stems forming adventitious roots, a total of 69 Aeschynomene 86 and Sesbania samples was collected (Table S1). Field observations were completed with molecular 87 and flow cytometry analyses for accurate specimen identification (Tables S2 and S3). Here, we report 88 on the stem-nodulated Aeschynomene and Sesbania species from Malagasy wetlands and discuss the 89 opportunity of these resources to fuel research on the nitrogen-fixing symbiosis in legumes.

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91 Stem nodulation in Nod-independent Aeschynomene species

92 Three stem-nodulated Aeschynomene species belonging to the Nod-independent clade were found in 93 Malagasy wetlands (Fig. 1b). A. evenia was by far the most widespread one, being seen in all parts of 94 Madagascar and in all wetland types: river banks, marshes and ricefields. A. evenia has a transatlantic 95 distribution and a well-defined geographically-structured genetic diversity (Chaintreuil et al., 2018). 96 The form present in Madagascar was previously classified as the Eastern African genotype. As a 97 result, it is closely related to the reference line from Malawi that was selected for the genetic 98 dissection of the Nod-independent symbiosis (Quilbé et al., 2021). Stem-nodulated A. evenia plants 99 were observed in 24 out of the 25 sampling sites. Nodules were green, indicative of the presence of 100 chloroplasts, and often located in the lower part of the stem, but their distribution could extend to

the upper branch-containing part and they were usually present in a profuse fashion. These stem
 nodules were hemispherical and with a broad attachment to the stem (Fig. 2a).

103 The second typical species found in different wetlands is A. sensitiva. Similarly to A. evenia, 104 this species has also a transatlantic distribution and the genotype occurring in Madagascar is 105 concommitantly present in Africa and Eastern Brazil (Chaintreuil et al., 2018). A. sensitiva is well-106 known because the model strain ORS278, a photosynthetic nod gene-lacking Bradyrhizobium, was 107 isolated from stem nodules in Senegal (Giraud et al., 2000, 2007). The photosynthetic activity of this 108 strain was demonstrated to be important for the efficiency of stem nodulation. In addition, A. 109 sensitiva has the particularity to develop unique 'collar' nodules on the stem (Fig. 2b). These were 110 green and readily observed in all 14 sampling sites. Although A. sensitiva was globally less frequently 111 found as compared to A.evenia, at the collection sites the two species were frequently growing 112 adjacent to each other (Fig. 2c). The two species are known to have overlapping distributions in 113 Madagascar but A. sensitiva is absent in the drier regions whereas A. evenia does (Dupuy et al., 114 2002).

115 Unexpected was the discovery of *A. indica* in marshes from Majunga and in ricefields in Nosy 116 Be because this species is not native of Madagascar. A. evenia and A. indica are morphologically 117 similar but they can be distinguished from each other via the flowers that on A. indica plants are in 118 general larger than those on A. evenia (Fig. 2d). This high resemblance is due to their belonging to a 119 same polyploid species complex where A. evenia 2x, A. indica 4x and A. indica 6x forms are present 120 (Arrighi et al., 2014; Chaintreuil et al., 2018). To confirm the visual species identification, we 121 sequenced the nuclear ITS and chloroplastic matk gene and determined the genome size of the 122 specimens JFA29 and JFA109. The obtained data were typical for *A. indica* 4x plants, confirming the 123 visual identification (Fig. S1, Table S2) (Arrighi et al., 2014). In the four collection sites, A. indica 124 plants were stem-nodulated, the nodules varying in shape and size but always with an enlarged base 125 and with a green color (Fig. 2d). Similar to other Aeschynomene species, pink root nodules could be 126 observed in unflooded conditions, (Fig. 2d).

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128 Stem nodulation in Nod-dependent Aeschynomene species

Among *Aeschynomene* species falling outside of the Nod-independent clade, the pantropical *A. americana* and the endemic *A. patula* were frequently present in explored sites. Both species are of interest because they have been proposed as model plants complementary to *A. evenia* to develop a comparative genetic system to study the Nod-independent and Nod-dependent symbioses in *Aeschynomene* (Brottier *et al.*, 2018). However, for both species no stem-nodulated plants were found at the collection sites (Fig. 1b).

135 Two Aeschynomene species, A. elaphroxylon and A. schimperi, which are believed to be 136 introduced in Madagascar, were found in the Central Plateaux (Fig. 1b) (Du Puy et al., 2002). A. 137 elaphroxylon was specially present around the Aloatra lake. It is a distinctive Aeschynomene species 138 as it can form large shrubs to small trees with very showy yellow flowers and spiny stems (Fig. 3a). In 139 the two sampled sites, plants were scarely nodulated and, if so, only on the submerged parts of the 140 stems. In that case, nodules were green and had a flattened hemispherical shape with a broad 141 attachment to the stem (Fig. 3a). In contrast, A. schimperi was frequently found in ricefields. For this 142 species, previous genetic analysis uncovered the presence of 4x and 8x cytotypes, and Malagasy 143 specimens correspond to the 8x cytotype (Chaintreuil et al., 2016). In non-waterlogged conditions, 144 numerous pink nodules could be observed on the main root, while under flooded conditions (9 of the 145 12 sampled sites) green nodules were present in the lower part of the stem (Fig. 3b). These nodules 146 were spherical with a narrow attachment to the stem.

147 In regions at lower altitude, in the North and West of Madgascar, two other Nod-dependent 148 Aeschynomene species were found. A. uniflora was present in the region of Majunga and in Nosy Be. 149 Here it grew either at the edge of marshes or in ricefields where often also A. evenia or A. sensitiva 150 species were present. Strikingly, two plant morphotypes were observed: one corresponding to 151 erected plants with well-visible flowers (3 sampled sites), and a second represented by shrubby 152 plants bearing small flowers (two sampled sites) (Fig. 3c). It is likely that the two morphotypes 153 correspond to the botanical varieties A. uniflora var. grandiflora and var. uniflora (Gillet et al., 1971). 154 ITS and matk sequencing for the samples JFA_51 and JFA_71 (examples of plants with well-visible 155 and small flowers, respectively) revealed consistent differences while flow cytometry measurements 156 were relatively homogenious (Fig. 1b, Fig. S2, Table S2) (Chaintreuil et al., 2016). In both 157 morphotypes, numerous green nodules with a spherical shape and a narrow neck, running on the 158 stem were visible (Fig. 3c).

159 A. cristata was found in pristine marshes and slow-flowing streams between Majunga and 160 Mitsinjo where they often formed important stands of flooded plants. At sites where A. cristata 161 plants were found A.evenia plants could be also present (Fig. 3d). A. cristata has retained little 162 attention until it was shown to be one of the genome donors of A. afraspera and being a sister 163 species of the Asian A. aspera, both species being profusely stem nodulated (Chaintreuil et al., 2013, 164 2016; Devi., 2013a,b). Whereas the A. cristata specimen characterized by Chaintreuil et al. (2016) 165 had densely-hairy stems, those found and collected in Madagascar were glabrous. Based on this 166 characteristic, they may be tentatively associated to A. cristata var. cristata (hairy stems) and var. 167 pubescens (glabrous stems) as decribed by Gillet et al. (1971). However, sequence and flow 168 cytometry data additional to those obtained for sample JFA_34 are required to assess any genetic 169 differentiation (Fig. 1b, Fig. S3, Table S2). In all sampling sites, A. cristata specimens cought attention

very quickly due to the profuse nodulation all over the stem (Fig. 3d). These stem nodules were green and hemisperical with a broad base as described for the related *A. afraspera* species (Alazard and Duhoux, 1987).

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174 Stem nodulation in Sesbania rostrata

175 S. rostrata was observed in only two sampling sites in the region of Majunga. S. rostrata has been 176 used a research model for nodulation due to its profuse stem nodulation and its ability to switch 177 from classical nodulation to Lateral Root Base nodulation in flooded conditions (Capoen et al., 2009; 178 Dreyfus and Dommergues, 1981). S. rostrata is the single species of its lineage in the Sesbania 179 phylogeny (Furraggia et al., 2020). However, S. rostrata specimens of Senegal and Madagascar were 180 shown to be morphologically different and both hybridization and grafting experiments were less 181 successful when interspecific (Ndiaye, 2005). For this reason, it has been proposed they could 182 represent different subspecies of S. rostrata. ITS and matk sequencing of S. rostrata samples JFA 21 183 and JFA 45 also point to genetic differentiation when compared to an accession from Senegal, while 184 flow cytometry analysis revealed similar genome sizes for the same accessions (Fig. 1c, Fig. S4, Table 185 S2). In plants of both sampling sites, nodulation all along the stem was remarkable for its profusion 186 and nodulation sites were typically distributed in vertical rows (Fig. 3e). These stem nodules were 187 green, prominent, and had a constricted base.

188

189 **Conclusions and perspectives**

190 In 2001, James et al. identified several stem-nodulated legumes in the Brazilian Pantanal wetlands. To broaden our knowledge on stem nodulating legumes, this research inspired us to explore 191 192 wetlands in a geographically distinct tropical region, Madagascar. In Malagasy wetlands, a large 193 occurrence and diversity of stem nodulation in Aeschynomene and Sesbania legumes was detected. 194 While some of the species found are already well-known to form stem nodules (A. elaphroxylon, A. 195 evenia, A. indica 4x, A. schimperi, A. sensitiva and S. rostrata), for two other identified species, A. 196 cristata and A. uniflora, this has been once furtively mentioned in a review article and subject to very 197 limited experimentations (Ladha et al., 1992; Chaintreuil et al., 2016). Here, we report on their stem 198 nodulation in the field. Following the traditional classification of stem-nodulating legumes, A. cristata 199 was convincingly a bona fide profusely stem-nodulated species, equaling the stem nodulation level 200 found for A. afraspera and S. rostrata (Boivin et al., 1997a; Ladha et al., 1992). A. uniflora stem 201 nodulation did not reach such level in the field. However, in greenhouse conditions A. uniflora stem 202 nodulation has been shown to be exceptionally dense and to occur all along the stem (Chaintreuil et al., 2016), indicating a profuse stem-nodulation capacity. The discovery of all these species and the
 demonstration of genetic diversity in several of them (*A. cristata, A. uniflora,* and *S. rostrata*) point
 out that more systematic studies including the collection of plants to evaluate species and ecotypes
 endowed with stem-nodulation are required.

207 What lessons could we learn from these studies? On the plant side, profuse stem nodulation 208 may be indicative of high nitrogen fixation activity, the presence of two stem-nodule morphologies 209 (either hemispherical with a broad attachment to the stem or spherical with a narrow base) supports 210 the existence of two developmental programs, and in both cases the photosynthetic activity in stem 211 nodules (inferred from their green color) points to a physiology that likely differs from those of root 212 nodules (discussed in Ladha et al., 1992; Legocki and Szalay, 1984). On the bacterial side, while A. 213 caulinodans has been isolated from S. rostrata stem nodules in both Senegal and Madagascar (Boivin 214 et al., 1997b), a great variety of Aeschynomene-nodulating Bradyrhizobium strains do exist but their 215 genetics are insufficiently understood (Molouba et al., 1999; Miché et al., 2011; Okazaki et al., 2016). 216 New photosynthetic strains having or lacking *nod* genes are expected to be present in stem nodules 217 of A. cristata (because it is a parent species of A. afraspera) and of the Nod-independent 218 Aeschynomene species (e.g., A. evenia, A. indica 4x, and A. sensitiva) respectively. Intringingly, 219 previously only non-photosynthetic strains were isolated from A. uniflora root nodules (Molouba et 220 al., 1999). It would thus be very interesting to investigate the nature of those present in its stem 221 nodules. Given all the valuable information that can be gained from stem-nodulating legumes, we 222 advocate reviving their study as this would significantly increase our understanding on the diversity 223 of mechanisms underlying the nitrogen-fixing symbiosis in legumes.

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233

234 Competing interests

235 None declared.

236

237 Author contributions

JFA and NN wrote the manuscript. FFM, MFR, CC, ATER, MDA, H Randriambanona, H
Ramanankierana and JFA carried out specimen collection, photography, identification and herbarium
conservation. FF, MP and MB performed sequence and flow cytometry analyses.

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242 Data availability

- 243 Data obtained in this study are listed in Tables S1 to S3, and the methods are given in Methods S1.
- 244 The DNA sequences generated in this study were deposited in GenBank under accession numbers

245 OR448903-OR448909 (nuclear ITS) and OR463925-OR463932 (chloroplastic matK).

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247 **References**

- Alazard D. 1985. Stem and root nodulation in *Aeschynomene* spp. *Appl. And Environ. Microbiol.*50:732-734.
- Alazard, D., and M. Becker. 1987. *Aeschynomene* as green manure for rice. *Plant Soil* 101:141-143.
- Alazard, D., Duhoux, E. 1987. Nitrogen-fixing stem nodules on *Aeschynomene afraspera*. *Biological fertility Soils* 4: 61-66.

Arrighi JF, Cartieaux F, Brown SC, Rodier-Goud M, Boursot M, Fardoux J, Patrel D, Gully D, Fabre S,
 Chaintreuil C, Giraud E. 2012. *Aeschynomene evenia*, a model plant for studying the molecular
 genetics of the Nod-independent rhizobium-legume symbiosis. *Molecular Plant-Microbe Interactions* 25(7): 851-861.

- 257 Arrighi JF, Chaintreuil C, Cartieaux F, Cardi C, Rodier-Goud M, Brown SC, Boursot M, d'Hont A,
- 258 **Dreyfus B, Giraud E. 2014.** Radiation of the Nod-independent *Aeschynomene* relies on multiple 259 allopolyploid speciation events. *New Phytologist* **201**: 1457-68.
- Becker M, JK Ladha, Je Ottow. 1988. Stem-nodulating legumes as green manure for lowland rice.
 Phil. J. Crop Sci. 13(3):121-127.
- Boivin C, Ndoye I, Molouba F, De Lajudie P, Dupuy N, Dreyfus B. 1997a. Stem nodulation in
 legumes: diversity, mechanisms, and unusual characteristics. *Critical Reviews in Plant Sciences* 16(1): 1-30.
- Boivin C, Ndoye I, Lortet G, Ndiaye A, De Lajudie P, Dreyfus B. 1997b. The Sesbania root symbionts
 Sinorhizobium saheli and S. teranga bv. sesbaniae can form stem nodules on Sesbania rostrata,
 although they are less adapted to stem nodulation than Azorhizobium caulinodans. Appl Environ
 Microbiol. 63(3):1040-7.

269	Brottier L, Chaintreuil C, Simion P, Scornavacca C, Rivallan R, Mournet P, Moulin L, Lewis GP,
270	Fardoux J, Brown SC, et al. 2018. A phylogenetic framework of the legume genus Aeschynomene
271	for comparative genetic analysis of the Nod-dependent and Nod-independent symbioses. BMC
272	Plant Biol. 18(1) :333.
273	Capoen, W., Oldroyd, G., Goormachtig, S., and Holsters, M. (2010). Sesbania rostrata: a case study
274	of natural variation in legume nodulation. New Phytol 186: 340-345.
275	Chaintreuil C, Arrighi JF, Giraud E, Miche L, Moulin L, Dreyfus B, Munive-Hernadez J, Villegas-
276	Hernadez M, Béna G. 2013. Evolution of symbiosis in the legume genus Aeschynomene. New
277	Phytologist 200 :1247-59.
278	Chaintreuil C, Perrier X, Martin G, Fardoux J, Lewis GP, Brottier L, Rivallan R, Gomez-Pacheco M,
279	Bourges M, Lamy L, et al. 2018. Naturally occurring variations in the nod-independent model
280	legume Aeschynomene evenia and relatives: a resource for nodulation genetics. BMC Plant Biol.
281	18(1) :54.
282	Chaintreuil C, Gully D, Hervouet C, Tittabutr P, Randriambanona H, Brown SC, Lewis GP, Bourge M,
283	Cartieaux F, Boursot, et al. 2018. The evolutionary dynamics of ancient and recent polyploidy in
284	the African semiaquatic species of the legume genus Aeschynomene. New Phytologist.
285	211(3) :1077-91.
286	Devi WR. 2013a. Aeschynomene aspera L., a nitrogen fixing stem nodulation plant from Manipur.
287	International Journal of Scientific & technology Research 2(6) : 232-235.
288	Devi WR. 2013b. Aeschynomene aspera L., a potential stem nodulated plant as green manure for rice
289	cultivation in Manipur. Journal of Medicinal Plants 1(4) :28-31.
290	Dreyfus BL and YR Dommergues. 1981. Nitrogen-fixing nodules induced by Rhizobium on the stem of
291	the tropical legume Sesbania rostrata. FEMS Microbiol. Letter. 10:313-317.
292	Du Puy DJ, Labat J-N, Rabevohitra R, Villiers J-F, Bosser J, Moat J. 2002. The Leguminosae of
293	Madagascar. Royal Botanic Gardens, Kew.
294	Farruggia FT, Lavin M, Wojciechowski MF. 2018. Phylogenetic Systematics and Biogeography of the
295	Pantropical Genus Sesbania (Leguminosae). Systematic Botany 43(2):414-429.
296	Gillett JB, Polhill RM, Verdcourt B. 1971. Leguminosae (part 3): subfamily Papilionoideae (part 1). In
297	Milne-Redhead E & Polhill RM (eds). Flora of Tropical East Africa. <i>Royal Botanic Gardens, Kew.</i>
298	Giraud E, Hannibal L, Fardoux J, Vermeglio A, Dreyfus B. 2000. Effect of Bradyrhizobium
299	photosynthesis on stem nodulation of <i>Aeschynomene sensitiva</i> . PNAS 97(26) : 14795-14800.
300	Giraud E, Moulin L, Vallenet D, Barbe V, Cytryn E, Avarre JC, Jaubert M, Simon D, Cartieaux F, Prin
301	Y, et al. 2007. Legumes symbioses: absence of Nod genes in photosynthetic bradyrhizobia.
302	Science. 316(5829): 1307-12.

- Hagerup O. 1928. En hygrofil baelgplante (*Aeschynomene aspera* L.) med bakterieknolde paa
 staenglen. *Dansk Bot. Arkiv* 14: 1-9.
- 305 James EK, Loureiro FM, Pott A, Pott VJ, Martins CM, Franco AA, Sprent JI. 2001. Flooding-tolerant
- legume symbioses from the Brazilian Pantanal. *New Phytologist* **150**: 723-738.
- Ladha JK, Pareek RP, Becker M. 1992. Stem-nodulating Legume-Rhizobium symbiosis and its
 agronomic use in lowland rice. Advances in Soil Science. pp 147–192.
- 309 Legocki RP and Szalay AA. 1983. Molecular biology of stem nodulation. In DPS Verma et al. (eds).
- 310 Genes involved in Microbe-Plant Interactions. Springer Verlag, Berlin, Heidelberg. pp. 210-219.
- Loureiro MF, Defaria SM, James EK, Pott A, Franco AA. 1994. Nitrogen-fixing stem nodules of the
 legume Discolobium pulchellum Benth. New Phytologist 128(2): 283-295.
- Loureiro MF, James EK, Sprent JI, Franco AA. 1995. Stem and root nodules on the tropical wetland
 legume Aeschynomene fluminensis. New Phytologist 130: 531-544.
- 315 Miché L, Moulin L, Chaintreuil C, Contreras-Jimenez JL, Munive-Hernández JA, Del Carmen Villegas-
- 316 Hernandez M, Crozier F, Béna G. 2011. Diversity analyses of Aeschynomene symbionts in Tropical
- 317 Africa and Central America reveal that nod-independent stem nodulation is not restricted to
- 318 photosynthetic bradyrhizobia. *Environ Microbiol.* **12(8)**:2152-64.
- 319 Molouba F, Lorquin J, Willems A, Hoste B, Giraud E, Dreyfus B, Gillis M, de Lajudie P, Masson-
- Boivin C. 1999. Photosynthetic bradyrhizobia from *Aeschynomene* spp. are specific to stemnodulated species and form a separate 16S ribosomal DNA restriction fragment length polymorphism group. *Appl Environ Microbiol.* 65(7):3084-94.
- 323 Ndiaye M. 2005. Caractérisation taxonomique de deux provenances de Sesbania rostrata Brem :
- 324 Sesbania rostrata du Sénégal, Sesbania rostrata de Madagascar. PhD thesis. Université Cheikh
 325 Anta Diop (UCAD), Dakar, Senegal.
- 326 Okazaki S, Tittabur P, Teulet A, Thouin J, Fardoux J, Chaintreuil C, Gully D, Arrighi JF, Furuta N,
- Miwa H et al. 2016. Rhizobium-legume symbiosis in the absence of Nod factors: two possible scenarios with or without the T3SS. *ISME J.* 10(1): 64-74.
- 329 Quilbé J, Lamy L, Brottier L, Leleux P, Fardoux J, Rivallan R, Benichou T, Guyonnet R, Becana M,
- Villar I, et al. 2021. Genetics of nodulation in *Aeschynomene evenia* uncovers mechanisms of the
 rhizobium-legume symbiosis. *Nat Commun.* 12(1):829.
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334 Supporting Information

- Fig. S1 Alignments of the nuclear *ITS* and chloroplastic *matK* gene sequences for *A. indica* 4x.
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337	Fig. S2 Alignments of the (a) nuclear ITS and (b) chloroplastic matK gene sequences for A. uniflora.
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339	Fig. S3 Alignments of the (a) nuclear ITS and (b) chloroplastic matK gene sequences for A. cristata.
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341	Fig. S4 Alignments of the (a) nuclear ITS and (b) chloroplastic matK gene sequences for S. rostrata.
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343	Methods S1 Material and methods.
344	
345	Table S1 Accessions characterized in this study, origin and nodulation status.
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347	Table S2 Genetic and morphological characteristics of species and samples included in this study.
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349	Table S3 GenBank numbers for the sequences used in the phylogenetic analyses.
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352	Key words: Aeschynomene, Sesbania, stem nodulation, nitrogen-fixation, symbiosis
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355	Methods S1 Material and methods.
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	Methods S1 Material and methods. Description of the collecting areas
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356 357	Description of the collecting areas
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356 357 358 359	Description of the collecting areas To select collection sites of <i>Aeschynomene</i> and <i>Sesbania</i> species in Madagascar, we made use of general information about their distribution as described in the compendium "The Leguminosae of
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356 357 358 359 360 361 362 363 364 365 366 366	Description of the collecting areas To select collection sites of <i>Aeschynomene</i> and <i>Sesbania</i> species in Madagascar, we made use of general information about their distribution as described in the compendium "The Leguminosae of Madagascar" (Dupuy, 2002) and utilized precise location data of previous isolated specimens present in collections through the Global Biodiversity Information Facility (GBIF - https://www.gbif.org/) and the Tropicos database (https://www.tropicos.org/). This resulted in the definition of four collecting areas: 1) the region of Majunga including the vicinity of the Kinkony lake where many temporary to permanent marshes locally named « matsabory » are present. This area also comprises the plain of Marovoay where numerous ricefields form the main rice granary of Madagascar, 2) the Itasy region in the Center of Madagascar where the Itasy Lake and ricefields are present, 3) the Alaotra region in the East side of Madagascar that corresponds to a large basin containing the Alaotra Lake. The
356 357 358 359 360 361 362 363 364 365 366 367 368	Description of the collecting areas To select collection sites of <i>Aeschynomene</i> and <i>Sesbania</i> species in Madagascar, we made use of general information about their distribution as described in the compendium "The Leguminosae of Madagascar" (Dupuy, 2002) and utilized precise location data of previous isolated specimens present in collections through the Global Biodiversity Information Facility (GBIF - https://www.gbif.org/) and the Tropicos database (https://www.tropicos.org/). This resulted in the definition of four collecting areas: 1) the region of Majunga including the vicinity of the Kinkony lake where many temporary to permanent marshes locally named « matsabory » are present. This area also comprises the plain of Marovoay where numerous ricefields form the main rice granary of Madagascar, 2) the Itasy region in the Center of Madagascar that corresponds to a large basin containing the Alaotra Lake. The presence of extensive ricefields make this region the second rice granary of Madagascar, and 4) Nosy

corresponds to the flowering period for most *Aeschynomene* and *Sesbania* species. Plants were
collected in the above mentionned areas but also « en route » from ricefields, rivers and marshes
present along the RN1, RN2 and RN4 national roads and secondary roads leading to the Itasy lake,

the Alaotra lake and the region of Majunga, respectively.

375

376 Plant and data collection

At each sampling location, the presence or receding of water in the aquatic ecosystem was recorded. The stem nodulation status of individuals for each species present in the population was examined and correlated to their positions relative to the flooding area. Whenever possible, three individuals of each species were chosen at random to determine their root nodulation status. Both stems and roots of these individuals were photographed *in situ*. Plant material was frequently collected for germplasm conservation and production of voucher specimens. The latters were deposited at the Herbarium of the CNARP Institute in Antananarivo (Madagascar).

384

385 Plant culture

Seeds collected in the field were dried at 34°C for one week and used for plant cultivation when fresh
material production was required. Seed scarification and plant growth in the greenhouse were
performed as indicated in Arrighi *et al.* (2012).

389

390 Gene sequencing and sequence analysis

391 Genomic DNA was isolated from fresh leaves using the CTAB extraction method. The nuclear 392 ribosomal internal transcribed spacer region (ITS: ITS1-5.8S rDNA gene-ITS2) and the chloroplast 393 matk gene were amplified and sequenced as published in Chaintreuil et al. (2016). Additionnal ITS 394 and *matK* sequences were retrieved from Chaintreuil *et a*l. (2016, 2018) and Brottier *et al.* (2018) for 395 Aeschynomene species and from Farrugia et al. (2018) for Sesbania species. To analyse sequence 396 aligned variations. sequences were using Multalin 397 (http://multalin.toulouse.inra.fr/multalin/multalin.html). ML phylogenetic tree reconstructions were 398 obtained by aligning nucleotide sequences with the MUSCLE program that is incorporated in the 399 MEGA X (v10.1.8) software. Aligned sequences were further processed in MEGA X using the 400 maximum likelihood approach and the Kimura 2-parameter model with a 1000x bootstrap (BS).

401

402 Genome size estimation

Flow cytometry measurements were performed using fresh leaf material as described in Arrighi *et al.*, 2012. Genome size estimations were based on the measurements of three plants per accession
using *Lycopersicum esculentum* (Solanaceae) cv « Roma » (2c = 1.99 pg) as the internal standard.

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407

408 **Figure legends**

409 Fig. 1 Aeschynomene and Sesbania species collected in wetlands of Madagascar. (a) Map of 410 Madagascar with a zoom on the four collecting sites, including RN1-Itsay Lake, RN2-Alaotra Lake, 411 RN4-Majunga and Nosy Be. (b) Phylogeny of Aeschynomene species. (c) Phylogeny of Sesbania 412 species. In (b) and (c), Maximum likelihood phylogenetic reconstructions were obtained using the 413 concatenated ITS + matk sequences. Numbers at nodes represent bootstrap values (% of 1000 414 replicates). Dashed boxes delineate clades where stem nodulation has been reported to occur. Taxa 415 collected in Madagascar are in bold and numbers on their right correspond to the occurrence of stem 416 nodulation in the different collection sites. In orange: no stem nodulation observed, in blue: stem 417 nodulation observed in the present study.

418

419 Fig. 2 Stem nodulation in Aeschynomene species of the Nod-independent clade. (a) A. evenia. From 420 left to right: 1- plant growing in a ricefield near Camp Bandro (Alaotra lake), 2- axillary axis bearing a 421 yellow flower and a mature pod, 3-4- stem nodules located at the base or middle portion of the 422 stem, 5- morphology of stem nodules. (b) A. sensitiva. From left to right: 1- plant growing in a 423 ricefield at Manakambahiny, in the direction of Alaotra lake, 2- axillary axis bearing yellow flowers 424 and developping pods, 3-4- more or less flattened stem nodules, 5- morphology of collar stem 425 nodules. (c) Co-occurrence of A. evenia and A. sensitiva. From left to right: 1- Stand of a mixed 426 population in a shallow sandy river at Marofotroboka (next to RN4), insets show stem nodules of A. 427 sensitiva and A. evenia, respectively, 2-4- permanent marsh at Belobaka (Majunga) where A. sensitiva 428 and A. evenia where found to grow side-by-side with entangled roots, both plants displaying stem 429 nodules. Ae: A. evenia, As: A. sensitiva. (d) A. indica 4x. From left to right: 1- plant growing in a 430 ricefield in Nosy Be, 2- axillary axis bearing a yellow flower and developping pods, 3-4- stem nodules 431 located near the base or at the middle portion of the stem, 5- morphology of stem nodules. Note the 432 presence of numerous pink root nodules in 3.

433

Fig. 3 Stem nodulation in Nod-dependent *Aeschynomene* and *Sesbania* species. (a) *A. elaphroxylon*. From left to right: 1- stand of shrubby plants growing at the edge of a marsh near the Alaotra lake, 2-« showy » yellow flower with visible anthers, 3- woody stem with spines, 4- nodules developping at the base of the stem, 5- morphology of stem nodules. (b) *A. schimperi*. From left to right: 1- young plants in a ricefield at Maharefo, South of the Alaotra lake, 2- yellow flower, 3- pink root nodules, 4stem nodules, 5- morphology of stem nodules. *A. uniflora*. From left to right: 1- plants growing in a

- ricefield at Bongomena next to the RN4, 2- yellow flowers with the smaller variant form shown in the
- inset, 3 and 4- stem nodules of the of the normal and morphological variant, respectively, 5-
- 442 morphology of stem nodules. (d) A. cristata. From left to right: 1- Plant growing in a swampy palm
- grove, 2- showy yellow flower, 3- Stand of A. cristata (Ac) in the center of a marsh lined with A.
- 444 evenia (Ae), 4- nodules present all along the stem, 5- morphology of stem nodules. (e) S. rostrata.
- 445 From left to right: 1- plant growing on the edge of a marsh at Amboromalandy next to the RN4, 2-
- 446 Yellow flower, 3 and 4- arrows of nodules running all along on the stem, 5- morphology of stem
- 447 nodules.

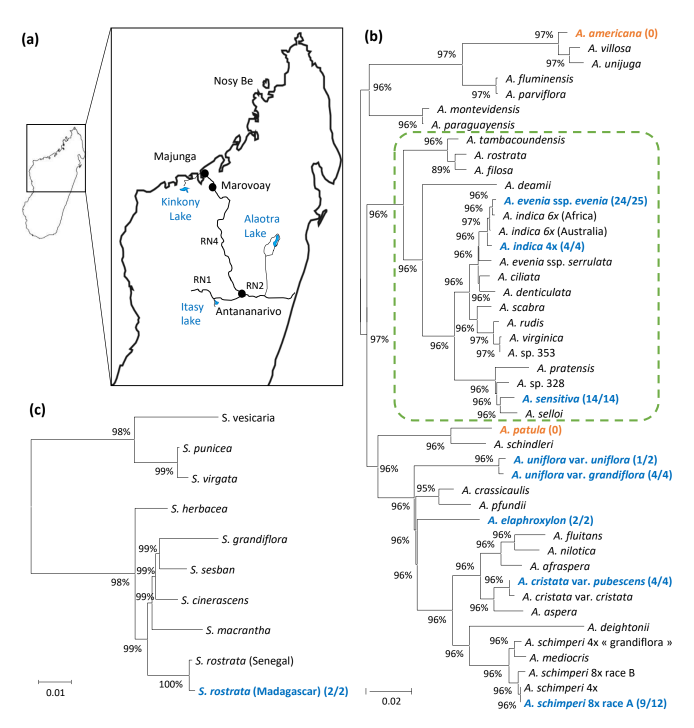


Fig. 1 *Aeschynomene* and *Sesbania* species sampled in wetlands of Madagascar. (a) Map of Madagascar with a zoom on the four collecting sites, including RN1-Itsay Lake, RN2-Alaotra Lake, RN4-Majunga and Nosy Be. (b) Phylogeny of *Aeschynomene*. (c) Phylogeny of *Sesbania*. In (b) and (c), Maximum likelihood phylogenetic reconstructions were obtained using the concatenated the ITS + *matK* sequences. Numbers at nodes represent bootstrap values (% of 1000 replicates). Dashed box delineates the Nod-independent clade. Taxa collected in Madagascar are in bold and numbers on their right correspond to the occurrence of stem nodulation in the different sampling sites. In orange: no stem nodulation observed, in blue: stem nodulation observed in the present study.

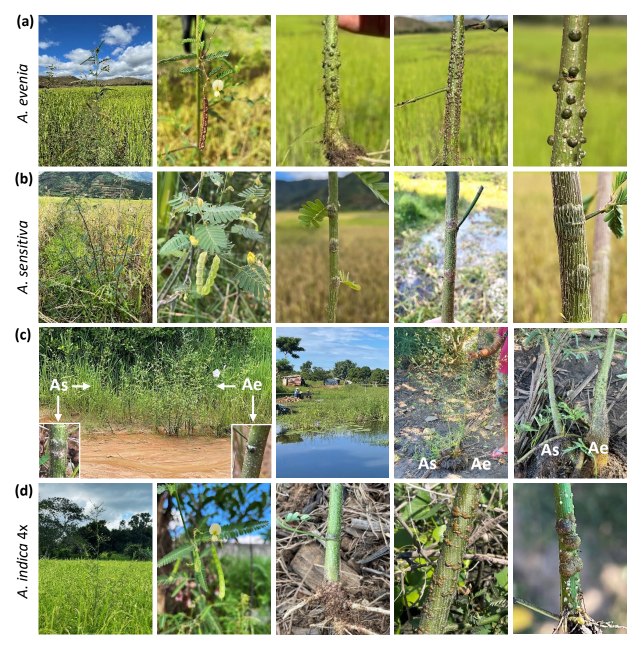


Fig. 2 Stem nodulation in *Aeschynomene* species of the Nod-independent clade. (a) *A. evenia*. From left to right: 1- plant growing in a ricefield near Camp Bandro (Alaotra lake), 2- axillary axis bearing a yellow flower and a mature pod, 3-4- stem nodules located at the base or middle portion of the stem, 5- morphology of stem nodules. (b) *A. sensitiva*. From left to right: 1- plant growing in a ricefield at Manakambahiny, in the direction of Alaotra lake, 2- axillary axis bearing yellow flowers and developping pods, 3-4- more or less flattened stem nodules, 5- morphology of collar stem nodules. (c) Co-occurrence of *A. evenia* and *A. sensitiva*. From left to right: 1- Stand of mixed population in a shallow sandy river at Marofotroboka (on RN4), insets show stem nodules of *A. sensitiva* and *A. evenia*, respectively, 2-4- permanent marsh at Belobaka (Majunga) where *A. sensitiva* and *A. evenia*, As: *A. sensitiva*. (d) *A. indica* 4x. From left to right: 1- plant growing in a ricefield in Nosy Be, 2- axillary axis bearing a yellow flower and a developping pods, 3-4- stem nodules located near the base or at the middle portion of the stem, 5- morphology of stem nodules. Note the presence of numerous pink root nodules in 3.

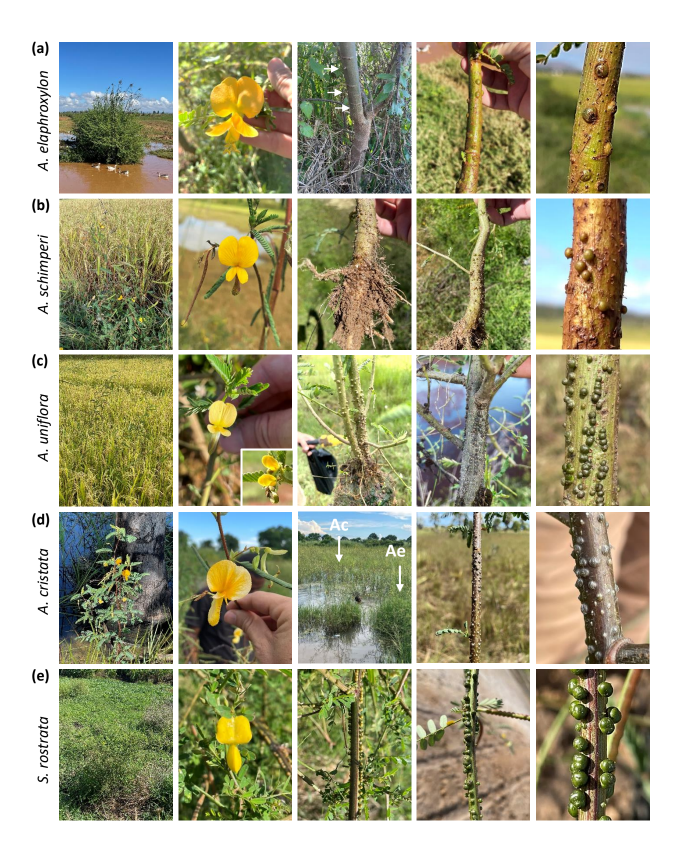


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