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Citation: Takahashi Y, Somta P, Muto C, Iseki K, Naito K, Pandiyan M, et al. (2016) Novel Genetic Resources in the Genus *Vigna* Unveiled from Gene Bank Accessions. PLoS ONE 11(1): e0147568. doi:10.1371/journal.pone.0147568

Editor: Vijai Gupta, National University of Ireland - Galway, IRELAND

Received: September 24, 2015

Accepted: January 5, 2016

Published: January 22, 2016

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Data Availability Statement: All relevant data are within the manuscript and uploaded to DNA Databank of Japan (DDBJ). Accession numbers are available in Table 1.

Funding: This work was supported by JSPS KAKENHI Grant Number 13J09808 and 26850006 to YT. Role of funders: data collection and analysis.

Competing Interests: The authors have declared that no competing interests exist.

RESEARCH ARTICLE

Novel Genetic Resources in the Genus *Vigna* Unveiled from Gene Bank Accessions

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Abstract

The genus *Vigna* (Fabaceae) consists of five subgenera, and includes more than 100 wild species. In *Vigna*, 10 crops have been domesticated from three subgenera, *Vigna*, *Plectro-tropis*, and *Ceratotropis*. The habitats of wild *Vigna* species are so diverse that their genomes could harbor various genes responsible for environmental stress adaptation, which could lead to innovations in agriculture. Since some of the gene bank *Vigna* accessions were unidentified and they seemed to be novel genetic resources, these accessions were identified based on morphological traits. The phylogenetic positions were estimated based on the DNA sequences of nuclear rDNA-ITS and chloroplast *atpB-rbcL* spacer regions. Based on the results, the potential usefulness of the recently described species *V. indica* and *V. sahyadriana*, and some wild *Vigna* species, i.e., *V. aconitifolia*, *V. dalzelliana*, *V. khandalensis*, *V. marina* var. *oblonga*, and *V. vexillata*, was discussed.

Introduction

The genus *Vigna*, in the family Fabaceae, comprises more than 100 wild species [1]. It is an agriculturally important taxon, which includes 10 domesticated species (crops) such as cowpea (*Vigna unguiculata* (L.) Walpers), mung bean (*Vigna radiata* (L.) Wilczek) and azuki bean (*Vigna angularis* (Willd.) Ohwi & Ohashi). Since some of their wild relatives inhabit extreme environments such as arid land, sandy beaches, and limestone karsts [2], they are expected to harbor adaptive genes, which could be used for developing stress-resistant crops for agriculturally unsuitable lands. Moreover, since they have evolved a symbiotic relationship with root-nodulating bacteria, which is also adapted to these extreme environments and contributes toward nitrogen fixation, these legumes have a high potential to contribute toward low-input sustainable agriculture [3, 4].

To introduce useful traits of wild relatives to related crops, interspecific hybridization is the most efficient and reliable strategy. Sequence-based phylogenetic relationships among species play a fundamental role as indicators to predict interspecific cross-compatibility. To increase the genetic diversity of a wild *Vigna* collection for environmental stress screening, *Vigna*

accessions were introduced from several gene banks. Since some of the gene bank accessions were unidentified and seemed to be novel genetic resources that have not been analyzed at the molecular level, these accessions were identified based on morphological traits, and were included in the phylogenetic analysis.

Although Maréchal et al. [5] described seven subgenera in the genus *Vigna*, two of them, *Macrorhynchus* and *Sigmoidotropis*, have been proposed to be distinct genera, i.e., *Wajira* and *Sigmoidotropis*, respectively, based on morphological and molecular phylogenetic analyses [6, 7]. Among the five subgenera presently recognized (*Ceratotropis*, *Haydonia*, *Lasiospron*, *Plectrotropis*, and *Vigna*), crop species have been developed only from three subgenera (*Ceratotropis*, *Plectrotropis*, and *Vigna*). Therefore, we have focused on the species belonging to these subgenera in the present study.

The subgenus *Ceratotropis*, also known as the Asian *Vigna*, is agronomically the most important taxonomic group, from which seven crops have been domesticated, i.e., moth bean (*Vigna aconitifolia* (Jacq.) Maréchal), minni payaru (*Vigna stipulacea* Kuntze), mung bean, black gram (*Vigna mungo* (L.) Hepper), creole bean (*Vigna reflexo-pilosa* Hayata), rice bean (*Vigna umbellata* (Thunb.) Ohwi & Ohashi), and adzuki bean (*Vigna angularis* (Willd.) Ohwi & Ohashi). Tomooka et al. [8] described 21 species, which were divided into three sections: five species in section *Aconitifoliae* N. Tomooka & Maxted, 12 species in section *Angulares* N. Tomooka & Maxted, and four species in section *Ceratotropis* N. Tomooka & Maxted. Although four new species were recently described in the subgenus *Ceratotropis* [9–12], their molecular phylogenetic positions have not been studied. In the present study, two newly described species (*V. indica* and *V. sahyadriana*) and four wild species (wild *V. aconitifolia* (Jacq.) Maréchal, *Vigna dalzelliana* (O. Kuntze) Verdcourt, *Vigna khandalensis* (Santapau) Raghavan & Wadhwa, *V. subramaniana* (Babu ex Raizada) Raizada) of the subgenus *Ceratotropis*, which had not been used in previous molecular phylogenetic studies, were analyzed.

Maréchal et al. [5] described seven species, consisting of two sections in the subgenus Plectrotropis (four species in section *Plectrotropis* and three species in section *Pseudoliebrechtsia*). The subgenus *Plectrotropis* contains a lesser known but potentially important food legume called 'tuber cowpea' (Vigna vexillata (L.) A. Rich.) [13]. This fully domesticated form is still cultivated in Bali and Timor, Indonesia. Maréchal et al. [5] recognized six botanical varieties (var. vexillata, angustifolia, doichonema, macrosperma, pluriflora, and yunnanensis) in V. vexillata. Among these varieties, var. macrosperma was reported as a domesticated taxa but its origin was unknown. Later, considering some proposals for new treatments [14, 15], Maxted et al. [16] accepted seven taxonomic varieties in V. vexillata (var. vexillata, angustifolia, davyi, dolichonema, lobatifolia, macrosperma, and ovata). V. vexillata var. davyi and V. vexillata var. lobatifolia were treated as distinct species (Vigna davyi H. Bol., Vigna lobatifolia Baker) in the subgenus Plectrotropis in Maréchal et al. [5] V. vexillata var. ovata was formerly treated as Strophostyles capensis (Thunb.) E. Mey. As such, the taxonomic treatments of the species in the subgenus *Plectrotropis* are still considered immature, and phylogenetic discussions based on molecular sequence information are necessary. In the present study, accessions of V. vexillata var. vexillata, var. angustifolia, var. lobatifolia, var. macrosperma, var. ovata, and Bali domesticated accessions were analyzed.

In the subgenus *Vigna*, from which cowpea (*Vigna unguiculata*) and bambara groundnut (*Vigna subterranea*) have been domesticated, Maréchal et al. [5] described 36 species in six sections (two species in section *Catiang*, two in *Comosae*, one in *Liebrechtsia*, two in *Macrodontae*, nine in *Reticulatae*, and 20 in *Vigna*). Cowpea is classified under *Catiang*, and bambara groundnut is in the section *Vigna*. For *Vigna*, we are currently focusing on *Vigna marina* (Burm.) Merrill, which inhabits sandy beaches, as a genetic resource for salinity tolerance, and *Vigna luteola* (Jacq.) Bentham, which inhabits riversides, as a flood-tolerant genetic resource

[17, 18]. These two species are closely related, and Padulosi and Ng [19] described *V. marina* ssp. *oblonga* Padulosi as being distributed in coastal areas of West Africa. Sonnante et al. [20] confirmed the genetic independence of *V. luteola*, *V. marina* ssp. *marina*, and *V. marina* ssp. *oblonga* based on isozymes and RAPD. In addition, they showed that *V. marina* ssp. *oblonga* was more closely related to *V. luteola* than to *V. marina* ssp. *marina*. However, *V. marina* ssp. *oblonga* was not included in subsequent phylogenetic analysis based on DNA sequences, although Pasquet et al. [15] described *V. marina* ssp. *oblonga* as being a synonym of *V. luteola*.

We therefore performed a phylogenetic characterization of the aforementioned taxa. To our knowledge, a phylogenetic study using DNA sequences had not been conducted on these taxa based on the DNA sequences of the internal transcribed spacer region of the ribosomal DNA on the nuclear genome (hereafter rDNA-ITS), and the *atpB-rbcL* intergenic spacer on the chloroplast genome (hereafter *atpB-rbcL*).

Materials and Methods

Plant materials

Seventy-one accessions of the genus *Vigna*, consisting of 28 species and three subgenera (*Ceratotropis*, *Plectrotropis*, and *Vigna*) conserved at the National Institute of Agrobiological Sciences, Japan, were used (<u>Table 1</u>). Originally, nine accessions were either unidentified, or seemed to be misidentified, as shown by the bold texts in <u>Table 1</u>. For the morphological analysis and DNA extraction, all the accessions were planted in six 0.3-L plastic pots (one seed/pot), and a 5-L plastic pot (six seeds/pot), and kept in a greenhouse where the temperature was maintained above 20°C with 12 hours of day length. The morphology of each plant was evaluated. For *V. aconitifolia*, weight of a hundred grains, pod shattering, and water absorbency of the seed were evaluated as domesticated traits. When evaluating pod shattering, 20 pods were dried overnight in a circulating incubator at 40°C. Twenty seeds were submerged in a Petri dish at 25°C for two days, and the number of seeds that absorbed water was recorded. We used common bean (*Phaseolus vulgaris* cv. Taisho-kintoki) as an outgroup for molecular phylogenetic analysis.

DNA Sequencing

We sequenced the rDNA-ITS and *atpB-rbcL* of 72 accessions. DNA was extracted from young leaves using a modified CTAB method [21]. PCR primers were designed according to the previous study [22]; C2 (5' -TCCTCCGCTTATTGATATGC-3') and G1 (5' -GGAAGGAGAAGT CGTAACAAGG-3') for rDNA-ITS, and AT1 (5' -AGAACCAGAAGTAGTAGGAT-3') and RB (5' -ACACCAGCTTTGAATCCAAC-3') for atpB-rbcL. The PCR mixture, containing KOD-Plus-Neo one unit (TOYOBO), 1 x PCR Buffer supplied by the manufacturer, 200 µM dNTPs, 1.5 mM MgSO₄, 10 ng of the DNA template, and 0.2 µM of each primer pair, was prepared in a total volume of 50 μL. The PCR cycle was as follows: 94°C for 2 min, 35 cycles of 98°C for 10 sec and 68°C for 1 min. The amplified PCR product was mixed with 2 μ L of ExoSAP-IT, which had been diluted 20-fold, and incubated at 37°C for 30 min, and 80°C for 15 min. The sequencing reaction was conducted according to the protocol of BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems). The reactant was precipitated using ethanol, dried, and dissolved in 10 µL Hi-Di Formamid. The mixture was treated at 95°C for 5 min, and the DNA sequence was determined using a ABI PRISM 3130xl DNA Analyzer (Applied Biosystems). Sequencing was repeated until the depth of each base was greater than five, and the nucleotide sequence was determined according to majority rule in cases where a single nucleotide polymorphism was present. The accession numbers of the sequence information deposited in the DNA Data Bank of Japan (<u>www.ddbj.nig.ac.jp/</u>) are shown in <u>Table 1</u>.

ID Section	Species Name	Status	Origin	JP No.	Original Conservation Site	Original ID and Species Identification	rDNA-ITS Sequence Length (bp)	rDNA-ITS DDBJ Accession No.	atpB-rbcL Sequence Length (bp)	atpB-rbcL DDBJ Accession No.
Subgenus Ceratotropis										
Aconitifoliae	V. aconitifolia	Domesticated	India	245857 T	TNAU GB	2009TN58	562	LC082015	200	LC082267
2 Aconitifoliae	V. aconitifolia	Domesticated India	India	245897 T	TNAU GB	2009TN99	562	LC082017	669	L C082269
3 Aconitifoliae		Domesticated	Pakistan	104332 N	NIAS GB	2752(5)	562	LC082016	669	L C082268
4 Aconitifoliae		Wild	India	235416 A	Australian GB	AUSTRCF106324, Vigna sp.	562	LC082014	699	LC082266
5 Aconitifoliae	V. aconitifolia	Wild	India	245864 T	TNAU GB	2009TN66, Vigna sp.	562	LC082012	699	LC082264
		Wild	India	245865 T	TNAU GB	2009TN67, Vigna sp.	562	LC082013	200	LC082265
Aconitifoliae		Wild	Sri Lanka	205894 N	NIAS GB	2000S-11	561	LC082018	689	LC082270
8 Aconitifoliae		Wild	Sri Lanka	205896 N	NIAS GB	2000S-2	561	LC082019	689	LC082271
9 Aconitifoliae	V. aridicola	Wild	Sri Lanka	207977 N	NIAS GB	2001SL-28	561	LC082020	069	LC082272
10 Aconitifoliae	V. indica	Wild	India	235417	ILRI GB	IL-25019, V. trilobata	562	LC082011	697	LC082263
11 Aconitifoliae	V. khandalensis	Wild	India	253828 T	TNAU GB	VC76	561	LC082005	687	LC082257
12 Aconitifoliae	V. stipulacea	Domesticated India	India	245503 T	TNAU GB	2008TN29	561	LC082007	069	LC082259
13 Aconitifoliae		Wild	Sri Lanka	205892 N	NIAS GB	2000S-6	562	LC082006	069	LC082258
14 Aconitifoliae	V. subramaniana	Wild	India	229278 A	Australian GB	AUSTRCF106193, Vigna sp.	562	LC064351	696	LC064361
15 Aconitifoliae	V. subramaniana	Wild	India	229284 A	Australian GB	AUSTRCF85155, V. radiata var. sublobata	562	LC064350	697	LC064360
16 Aconitifoliae	V. trilobata	Wild	India	245881 T	TNAU GB	2009TN83	562	LC082010	069	LC082262
17 Aconitifoliae	V. trilobata	Wild	Sri Lanka	210605 N	NIAS GB	2000S-5-1	562	LC082009	690	LC082261
18 Aconitifoliae	V. trilobata	Wild	Sri Lanka	205895 N	NIAS GB	2000S-13	562	LC082008	069	LC082260
19 Angulares	V. angularis var. angularis	Domesticated	Japan	37752 N	NIAS GB	ERIMOSHOUZU	557	LC081992	688	LC082244
20 Angulares	V. angularis var. nipponensis	Wild	Japan	87910 N	NIAS GB	CED96101602	557	LC081993	688	LC082245
21 Angulares	V. angularis var. nipponensis	Wild	Laos	226665 N	NIAS GB	2005L34	557	LC081995	688	LC082247
22 Angulares	V. dalzelliana	Wild	India	235419 A	Australian GB	AUSTRCF85146	557	LC081997	689	LC082249
23 Angulares	V. dalzelliana	Wild	Myanmar	210811 N	NIAS GB	2001M24, Vigna sp.	557	LC081996	696	LC082248
24 Angulares	V. exilis	Wild	Thailand	205884 N	NIAS GB	99T-10-1	557	LC081985	069	LC082237
25 Angulares	V. hirtella	Wild	Sri Lanka	218935 N	NIAS GB	9902-48	557	LC081984	690	LC082236
26 Angulares	V. hirtella	Wild	Thailand	109681 N	NIAS GB	CED891122-(9)	557	LC081983	691	LC082235
27 Angulares	V. hirtella	Wild	Laos	220137 N	NIAS GB	2003L-14	558	LC081988	689	LC082240
28 Angulares	V. hirtella	Wild	Thailand	108562 N	NIAS GB	96120305	563	LC081989	689	LC082241
29 Angulares	V. minima	Wild	Thailand	107869 N	NIAS GB	CED891125-(10)	556	LC081998	069	LC082250
30 Angulares	V. minima	Wild	Indonesia	218938 E	Belgian GB	NI1363	556	LC082000	069	LC082252
31 Angulares	V. minima	Wild	Papua N.G.	226877 N	NIAS GB	2005PNG15	556	LC081999	692	LC082251
32 Angulares	V nakachimaa	Wild	Janan	107879 N	NIAS GB	Ukushima	556	1 0.0820.02	69.3	I C082254

ID Section	Species Name	Status	Origin	JP No.	Original Conservation Site	Original ID and Species Identification	rDNA-ITS Sequence Length (bp)	rDNA-ITS DDBJ Accession No.	atpB-rbcL Sequence Length (bp)	atpB-rbcL DDBJ Accession No
33 Angulares	V. nepalensis	Wild	Nepal	107881	NIAS GB	Nepalen	557	LC081994	689	LC082246
34 Angulares	V. reflexo-pilosa var. glabra	Domesticated	Philippines	109684	AVRDC GB	V1160	557	LC081986	698	LC082238
35 Angulares	V. reflexo-pilosa var. reflexo- pilosa	Wild	Malaysia	108867	NIAS GB	510–1	557	LC081987	698	L C082239
36 Angulares	V. riukiuensis	Wild	Japan	108810	NIAS GB	Y-4-1	556	LC082001	692	LC082253
37 Angulares	V. tenuicaulis	Wild	Myanmar	227438	NIAS GB	KYONKADON	557	LC081991	688	LC082243
38 Angulares	V. tenuicaulis	Wild	Thailand	109682	NIAS GB	CED891122-(8)	557	LC081990	688	LC082242
39 Angulares	V. trinervia	Wild	Malaysia	108840	NIAS GB	503-4	561	LC064352	698	LC064362
40 Angulares	V. umbellata	Domesticated Japan	Japan	99485	NIAS GB	Menaga	557	LC081982	689	LC082234
41 Angulares	V. umbellata	Wild	Thailand	210639	NIAS GB	99T-2	557	LC064307	689	LC064328
42 Angulares	V. umbellata	Wild	Thailand	109675	NIAS GB	(6)-1-1	557	LC081981	689	LC082233
43 Angulares	Vigna sp.	Wild	Thailand	210644	NIAS GB	6 - 166	557	LC064303	689	LC064324
44 Ceratotropis	V. grandiflora	Wild	Thailand	107862	NIAS GB	CED891119-(1)	562	LC064345	694	LC064355
45 Ceratotropis	V. mungo var. mungo	Domesticated Th	Thailand	109668	NIAS GB	Subsomotod	562	LC064346	689	LC064356
46 Ceratotropis	V. mungo var. silvestris	Wild	India	107874	NBPGR	TC2211	562	LC064347	690	LC064357
47 Ceratotropis	V. radiata var. radiata	Domesticated	Thailand	110830	NIAS GB	CN60	595	LC064348	688	LC064358
48 Ceratotropis	V. radiata var. sublobata	Wild	Madagascar	107877	AVRDC GB	TC1966	587	LC064349	688	LC064359
49 Ceratotropis	V. radiata var. sublobata	Wild	Papua N.G.	226874	NIAS GB	2005PNG08	597	LC082004	688	LC082256
50 Ceratotropis	V. sahyadriana	Wild	India	235420	Australian GB	AusTRCF104896, Vigna sp.	568	LC082003	689	LC082255
51 Ceratotropis	Vigna sp.	Wild	India	110836	Belgian GB	NI 1135, V. radiata var. setulosa	564	LC064353	688	LC064363
52 Ceratotropis	Vigna sp.	Wild	India	245506	TNAU GB	2008TN32, V. hainiana	559	LC064354	688	LC064364
Subgenus Plectrotropis										
53 Plectrotropis	V. vexillata	Domesticated Indonesia	Indonesia	235863	Belgian GB	NI 1858	560	LC082032	683	LC082284
54 Plectrotropis	V. vexillata	Wild	Brazil	202337	USDA GB	PI 406391	562	LC082035	684	LC082287
55 Plectrotropis	V. vexillata	Wild	Papua N.G.	230747	NIAS GB	2006PNG-37	563	LC082037	683	LC082289
56 Plectrotropis	V. vexillata	Wild	Suriname	202334	USDA GB	PI 406383	563	LC082036	684	LC082288
57 Plectrotropis	V. vexillata var. angustifolia	Wild	Columbia	235869	Belgian GB	NI 936	563	LC082038	684	LC082290
58 Plectrotropis	V. vexillata var. Iobatifolia	Wild	Namibia	235903	Belgian GB	NI 546	557	LC082031	686	LC082283
59 Plectrotropis	V. vexillata var.	Domesticated	Sudan	235905	235905 Belgian GB	NI 111	559	LC082034	684	LC082286

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60	60 Plectrotropis	V. vexillata var. ovata	Wild	South Africa	235908	235908 Belgian GB	NI 1869	562	LC082033	684	LC082285
61	61 Plectrotropis	V. vexillata var. vexillata	Wild	Congo	235912	235912 Belgian GB	NI 245	563	LC082039	684	LC082291
	Subgenus Vigna										
62	2 Catiang	V. unguiculata	Domesticated Nigeria	Nigeria	86801	IITA GB	IT 84S 2246	581	LC082027	686	LC082279
63	63 Catiang	V. unguiculata	Domesticated Sudan	Sudan	86877	IITA GB	TVU 11979	581	LC082026	686	LC082278
64	64 Catiang	V. unguiculata	Domesticated Sudan	Sudan	86879	IITA GB	TVU 11986	581	LC082028	686	LC082280
65	65 Catiang	V. unguiculata ssp. dekindtiana	Wild	Mali	89083	IITA GB	TVNU 457	575	LC082030	684	LC082282
99	66 Catiang	V. unguiculata ssp. sesquipedalis	Domesticated Sri Lanka	Sri Lanka	81610	NIAS GB	MA	581	LC082029	686	LC082281
67	67 Vigna	V. Iuteola	Wild	Australia	236246	236246 Australian GB	AUSTRCF 320527	566	LC082021	689	LC082273
89	3 Vigna	V. Iuteola	Wild	Brazil	235855	Belgian GB	NI 858	566	LC082023	689	LC082275
69) Vigna	V. marina ssp. marina	Wild	Japan	235813	235813 NIAS GB	2009IRIO-1	569	LC082022	069	LC082274
20	70 Vigna	V. marina ssp. oblonga	Wild	Benin	233389	233389 NIAS GB	2006BENIN29	567	LC082024	690	LC082276
71	71 Vigna	V. subterranea	Domesticated unknown	unknown	79992	NIAS GB	L15-20-2	575	LC082025	690	LC082277
72		Phaseolus vulgaris	Domesticated Japan	Japan	219310	219310 NIAS GB	TAISHOU KINTOKI	554	LC082303	679	LC082302
Ī	ne accessions v	vhich were original	Ily either uniden	ntified, or seem	to be	misidentified are	Nine accessions which were originally either unidentified, or seemed to be misidentified are shown by bold texts.				

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Multiple alignment was conducted for each rDNA-ITS and *atpB-rbcL* using Clustal W [23]. The sequence frame was determined according to the previous study [22], and the trimmed sequence was used to construct a phylogenetic tree by the maximum likelihood estimation using MEGA6 [24]. Bootstrap analysis was conducted with 1000 replications.

Results

Morphology-based species identification

Among the nine unidentified or misidentified accessions, six accessions were identified as the following four species (*V. aconitifolia*, *V. dalzelliana*, *V. indica*, and *V. sahyadriana*) based on morphological observation.

Accessions ID-4, ID-5, and ID-6, which were collected in India, were identified as the wild forms of moth bean (*V. aconitifolia*). Seedling, stipule, and seed morphologies of the domesticated and newly identified wild forms of *V. aconitifolia* are shown in Fig 1. Both domesticated and wild forms showed similar variations in leaflet shape, ranging from entire to deeply lobed. Only seeds of the wild forms were covered with a semi-transparent seed coat covering. While the domesticated forms were characterized by larger seeds with water-permeable seed coat and non-shattering pods, the wild forms were found to have smaller seeds, with a water-proof seed coat and high shattering pods (Table 2).

Morphologies of the seedling, style beak, and seed of the remaining accessions newly identified as *V. dalzelliana*, *V. indica*, and *V. sahyadriana* are shown in Fig 2. Accession ID-23, collected in southern Myanmar, showed hypogeal germination with petiolate primary leaves, glabrous pods, seeds without seed coat coverings (smooth seed coat), small yellow flowers, left curved keel petal with protuberance on left keel (keel pocket), indicating that this accession belonged to the section *Angulares* in the subgenus *Ceratotropis*. Additionally, it had a flat style beak (spoon-like shape), which is a key characteristic of *V. dalzelliana*. Therefore, we have identified this accession as *V. dalzelliana*.

Accession ID-50, collected in India, was introduced from the gene bank of Australia (AusTRCF104896), where it was treated as *Vigna* sp. (<u>Table 1</u>). It showed epigeal germination with sessile primary leaves, seeds with seed coat covering, hairy pods, yellow flower, and left curved keel petal with a prominent protuberance on the left keel petal (keel pocket), indicating that this accession belongs to the section *Ceratotropis* in the subgenus *Ceratotropis*. Seed morphology and very long style beak matched the characteristics of *V. mungo*, whereas the direction of laterally attaching pods to the peduncle matched that of *V. radiata*. These characteristics matched the key characters of *V. sahyadriana* well, which was described as a new species by Aitawade et al. [10].

Accession ID-10, collected in India, was introduced from the ILRI (International Livestock Research Institute) gene bank (IL-25019), where it was conserved as *V. trilobata*. It showed epigeal germination with sessile primary leaves, seeds with seed coat covering, hairy pods, small yellow flowers, left curved keel petal with a small protuberance on left keel petal (keel pocket), and a protruding growth habit with deeply lobed leaflets, indicating this accession belongs to the section *Aconitifoliae* in the subgenus *Ceratotropis*. At a glance, it had a very similar overall morphology to *V. trilobata*. However, its stipule was lanceolate, and its seed was rectangular with a very short, non-protruding hilum, which did not match the key characters of *V. trilobata*. These characteristics matched those of *V. indica*, which was described as a new species by Dixit et al. [9].

Accession ID-43, collected in Thailand, was originally identified as *V. umbellata*. However, it showed some features that did not match the key characteristics of *V. umbellata*. Accession ID-51, collected in northern India, was introduced from a Belgian gene bank (NI 1135) as *V*.

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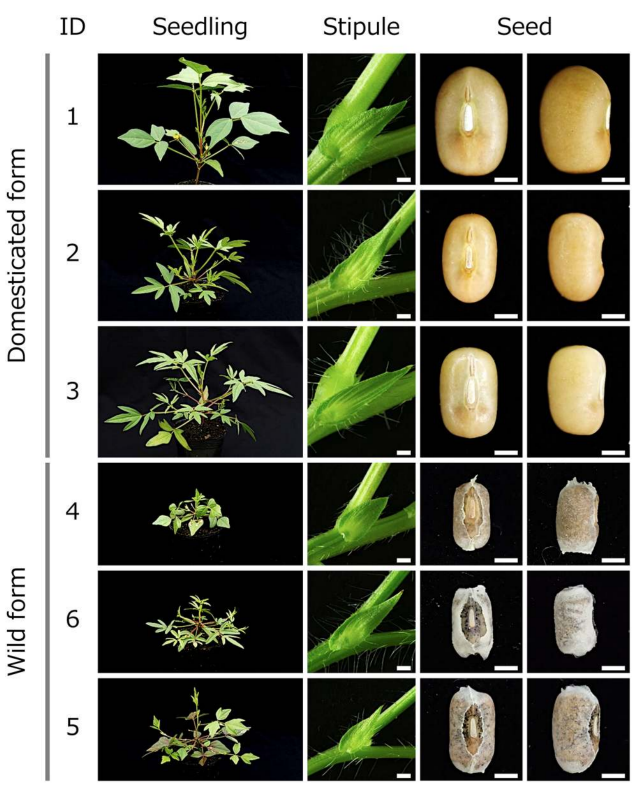


Fig 1. Domesticated form and wild ancestral form of moth bean (V. aconitifolia). Scale bars are 1 mm.

ID	Status	Seed weight ± SD (g/100 grains) ¹	Shattering pods (%)	Germination (%)
1	Domesticated	3.39 ± 0.42 a	0	100
2	Domesticated	2.03 ± 0.38 b	0	100
3	Domesticated	2.20 ± 0.11 b	0	100
4	Wild	0.86 ± 0.14 c	100	0
5	Wild	1.15 ± 0.08 c	100	0
6	Wild	1.26 ± 0.11 c	100	0

Table 2. Comparison of domestication related traits in domesticated and wild V. aconitifolia.

¹ Averages of 3 replications. Different letters indicate that seed weights are significantly different, by Tukey —Kramer's HSD test (P < 0.05).

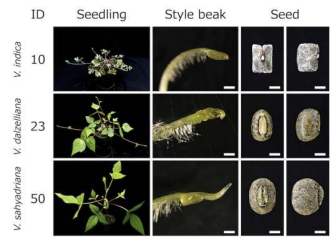
doi:10.1371/journal.pone.0147568.t002

radiata var. *setulosa*. Accession ID-52, collected in southern India, was introduced from the Tamil Nadu Agricultural University (TN32) as *V. hainiana*. Both of these accessions had a similar morphology to that of *V. radiata* in general. However, they showed some features that did not match the key characteristics of *V. radiata*. Therefore, we could not determine the taxonomic identification for these three accessions based on the morphological analysis in the present study.

Molecular phylogenetic analysis

DNA sequences of rDNA-ITS and *atpB-rbcL* were determined for 71 accessions of the genus *Vigna*. For rDNA-ITS, the total length ranged from 556–597 bp; *V. minima*, *V. riukiuensis*, and *V. nakashimae* had the shortest (556 bp), and *V. radiata* had the longest rDNA-ITS (587–597 bp). The total lengths of *atpB-rbcL* ranged from 683 to 700 bp; *V. unguiculata* and *V. vexillata* had the shortest (683–686 bp), whereas *V. aconitifolia* had the longest *atpB-rbcL* (699–700 bp) (Table 1). The numbers of polymorphic sites in rDNA-ITS and *atpB-rbcL* were 211 and 80, respectively.

Based on these sequences of rDNA-ITS and *atpB-rbcL*, phylogenetic trees for respective regions were constructed (Figs <u>3</u> and <u>4</u>). In both phylogenetic trees, the subgenus *Ceratotropis*





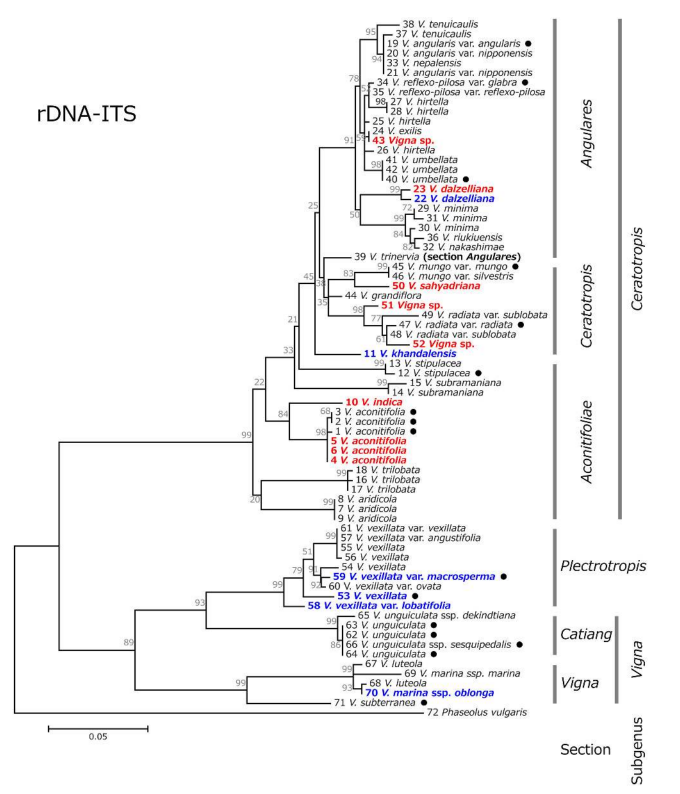


Fig 3. Maximum likelihood tree based on nuclear rDNA-ITS region for the genus *Vigna*, with *Phaseolus vulgaris* as an outgroup. Numbers beside branches represent bootstrap values (%) based on 1000 replications. Scale indicates genetic distance. Domesticated accessions are indicated with black circles, accessions which have been introduced as unidentified or misidentified accessions are indicated with red text, and taxa in which phylogenetic discussion using DNA sequences had not been conducted are indicated with blue text.

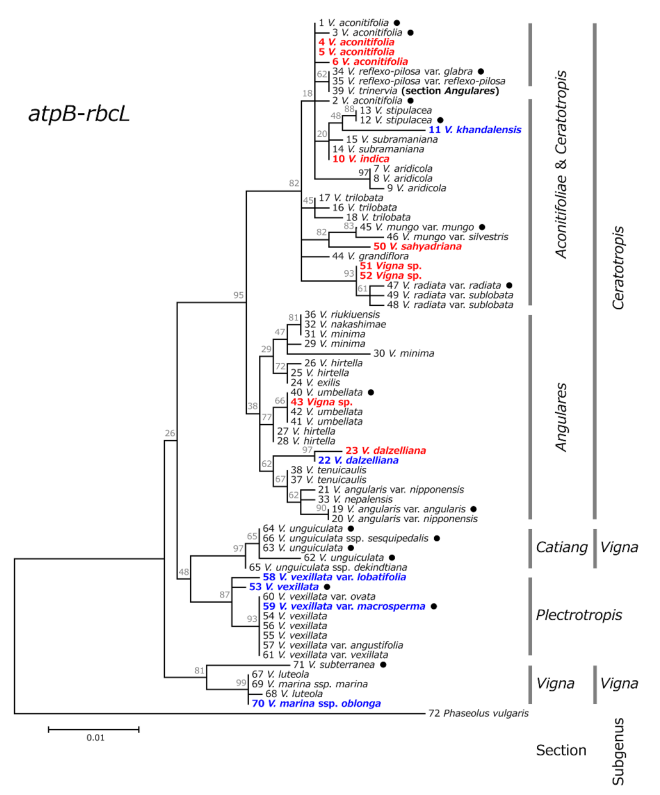


Fig 4. Maximum likelihood tree based on chloroplast *atpB-rbcL* spacer region for the genus *Vigna*, with *Phaseolus vulgaris* as an outgroup. Numbers beside branches represent bootstrap values (%) based on 1000 replications. Scale indicates genetic distance. Domesticated accessions are indicated with black circles, accessions which have been introduced as unidentified or misidentified accessions are indicated with red text, and taxa of which phylogenetic discussion using DNA sequences had not been conducted are indicated with blue text.

formed a single cluster, distinct from the subgenera *Vigna* and *Plectrotropis*. The section *Catiang* in the subgenus *Vigna* allied with the subgenus *Plectrotropis* forming a single cluster, while the section *Vigna* in the subgenus *Vigna* was distantly allied.

The phylogenetic tree based on rDNA-ITS divided the section *Aconitifoliae* into multiple branches, and clustered the section *Ceratotropis* and *Angulares* independently (Fig.3). Alternatively, the phylogenetic tree based on *atpB-rbcL* divided the subgenus *Ceratotropis* into two groups, i.e., a blended group comprising the sections *Aconitifoliae* and *Ceratotropis*, and the section *Angulares* (Fig.4). While the section *Angulares* clustered distinctly from other groups, the interspecific genetic distances within the *Angulares* cluster were small.

The phylogenetic analysis revealed the species most closely related to the accessions that were newly identified in this study. Accession ID-4, ID-5, and ID-6, identified as a wild form of moth bean, were most closely related to moth bean (*V. aconitifolia*). Accession ID-23 (Myanmar), identified as *V. dalzelliana*, was most closely related to the *V. dalzelliana* collected in India. Accession ID-50, identified as *V. sahyadriana*, was most closely related to *V. mungo*. Accession ID-10, identified as *V. indica*, was most closely related to *V. aconitifolia* in the phylogenetic tree based on rDNA-ITS, and to *V. subramaniana* in the phylogenetic tree based on *atpB-rbcL*. Accession ID-43 (*Vigna* sp.) was closely related to *V. exilis* in the rDNA-ITS tree, whereas it was allied with *V. umbellata* in the *atpB-rbcL* tree. Accessions ID-51 and ID-52 were most closely related to *V. radiata* in both trees.

V. khandalensis (accession ID-11) was differentiated substantially from other species, but was relatively close to *V. stipulacea*. Accessions within *V. vexillata* showed considerable levels of genetic variation. The accession ID-58 (*V. vexillata* var. *lobatifolia*), and the Indonesian domesticated form (accession ID-53) noticeably differentiated from other *V. vexillata* accessions. *V. marina* ssp. *oblonga* (accession ID-70), which was found on the coast of West Africa, was more closely related to *V. luteola* than to *V. marina* ssp. *marina*.

Discussion

Genetic differentiation within the genus Vigna

The subgenus Ceratotropis is thought to have emerged from the subgenus Vigna via the subgenus Plectrotropis [16, 25, 26]. The theoretical basis of this hypothesis is that, while the subgenus Vigna has a symmetric keel without pocket, the subgenus Plectrotropis has a curved keel with a pocket, and the subgenus Ceratotropis has a more prominently twisted keel with a more prolonged pocket. However, the phylogenetic tree using rDNA-ITS in this study suggested the following genetic differentiation patterns. The common ancestor of the genus Vigna first diverged into the common ancestor of the subgenera Vigna plus Plectrotropis, and the common ancestor of the subgenus Ceratotropis. Then, the common ancestor of the subgenera Vigna plus Plectrotropis diverged into the common ancestor of the section Vigna (subgenus Vigna) and the common ancestor of the section Catiang (subgenus Vigna) plus subgenus Plectrotropis. This is supported by the fact that the species in the section Catiang (subgenus Vigna) and the subgenus Plectrotropis have purple flowers, while those in the section Vigna (subgenus Vigna) have yellow flowers. Similar species relationships to our phylogenetic tree were obtained in previous studies using other molecular markers [7, 20, 27]. Therefore, it seems more appropriate to raise the rank of the section *Catiang* as a subgenus level. However, we leave this taxonomic revision for future work, since we used the limited number of species in the section Catiang, Vigna, and the subgenus *Plectrotropis*.

"*Plectrotropis*", which represents the subgenus, and the section including *V. vexillata*, has been misspelled as "*Plectotropis*" in Maréchal et al. [5], and in many subsequent publications such as Tomooka et al. [8] and Maxted et al. [16], but the former should be the correct spelling,

as it appeared in Schumach [28] and Baker [29] as a genus name and a subgenus name, respectively.

After cowpea and *V. vexillata* were shown to be relatively close to each other by molecular analysis [30], an interspecific hybrid between the two species was obtained [31]. Moreover, an interspecific hybrid was obtained between cowpea and *V. luteola*, which are more distantly related species [32]. In the present study, we propose that *V. marina* is worth trying for producing interspecific hybrids with bambara groundnut (*V. subterranea*), based on their relatively close phylogenetic positions. *V. marina* is highly tolerant to salinity and alkaline soil [17, 33], while bambara groundnut is a crop that is adapted to arid lands [34]. Drought, saline, and alkaline soils are the most important environmental stresses to be addressed in agriculture.

Novel genetic resources in the genus Vigna

Vigna indica T.M. Dixit, K.V. Bhat & S.R. Yadav. Accession ID-10 is revealed to be the only germplasm of *V. indica* currently available at the gene bank. Although a holotype (*Rothe 6229a*) of this species was described as *V. trilobata* (L.) Verdcourt var. *pusilla* Naik et Pokle [35], results of the phylogenetic analysis supported Dixit et al. [9], in that this taxon is an independent species in the section *Aconitifoliae*. Whereas *V. indica* was reported to be morphologically most similar to *V. aridicola* by Dixit et al. [9], it was also similar to the wild form of *V. aconitifolia* in its stipule and flower morphology.

In this study, *V. indica* showed the closest relationship with *V. aconitifolia* in the rDNA-ITS tree. Conversely, it showed almost the same *atpB-rbcL* sequence as that of *V. subramaniana*. These facts suggest the possibility that *V. indica* is derived from an interspecific hybrid between *V. subramaniana* and *V. aconitifolia*. Further studies are necessary to confirm the origin of this species. Additionally, useful traits screening and interspecific cross-compatibility of *V. indica* should be conducted to determine its usefulness as a genetic resource, especially for moth bean (*V. aconitifolia*), the most closely related crop.

Vigna sahyadriana Aitawade, K.V. Bhat et S.R. Yadav. Accession ID-50 is the only germplasm of *V. sahyadriana* available from the gene bank at present. This species was recently described as a new species distributed in Maharashtra, India [10]. Since accession ID-50 was collected in Madhya Pradesh, India, the distribution range of this species seems to have expanded toward the inland of India.

Accession ID-50 was most closely related to, but clearly distinguishable from, black gram (*V. mungo*) in both phylogenetic trees (Figs $\underline{3}$ and $\underline{4}$). This suggests that the useful traits and interspecific cross-compatibility of *V. sahyadriana* should be investigated to determine if it can be used as genetic resources for black gram.

Vigna aconitifolia (Jacq.) Maréchal: Wild ancestor of moth bean. Although the wild form of moth bean was documented to be distributed in India [36], living samples have not been identified in the gene bank [27], and therefore its identity and useful traits have not been studied. In this study, we found the wild ancestor of moth bean in a gene bank collection. Accessions ID-5 and ID-6 were collected in Tamil Nadu, and accession ID-4 was collected in Andhra Pradesh, India. The collection sites of these three accessions suggest that the primary habitat of the wild form of moth bean is southeastern India.

Moth beans have been cultivated mainly in arid lands from India to Pakistan, and also in some other counties including Bangladesh, Myanmar, and China [<u>37</u>]. Since moth bean is reported as a crop most tolerant to drought and heat in the subgenus *Ceratotropis* [<u>38</u>, <u>39</u>], it is generally thought to be suitable as a crop in tropical arid lands.

Recently, we have found that the wild ancestor of moth bean showed higher drought tolerance than the domesticated forms, and we successfully obtained the F_2 lines among the two forms (data not shown). Moreover, since the interspecific hybrid between mung bean and moth bean has been reported [40], wild moth bean would be useful to develop moth bean and mung bean varieties with higher drought tolerance.

Vigna dalzelliana (O. Kuntze) Verdcourt. The geographical distribution of this species was thought to be limited to India and Sri Lanka [8]. Although Thuan [41] reported *V. dalzelliata* in the Indo-China region (Vietnam, Laos, and Cambodia), it was the result of a misidentification of *V. minima* specimens [39]. More recently, John et al. [42] reported that they found *V. dalzelliana* in the Andaman Islands. Identification of accession ID-23 as *V. dalzelliana* in this study revealed an additional range of geographical distribution for this species, southern Myanmar.

The dissemination pathway of *V. dalzelliana* from India to southern Myanmar is unknown. Further explorations in the broad areas along the Bengal Gulf (Bangladesh and Myanmar) are necessary. However, since *V. dalzelliana* also inhabits Sri Lanka and the Andaman Islands [8, <u>42</u>], researchers must consider the possibility that the distribution range expanded from India to Myanmar via these Islands.

Based on the rDNA-ITS tree, *V. dalzelliana* is located at the basal position with a *V. minima* species complex (*V. minima*, *V. nakashimae*, *V. riukiuensis*) [43], and both of these species are well differentiated within the section *Angulares* (Fig 3). Since *V. dalzelliana* is the only species known to be distributed in south India, where species of the other two sections are rich, it could be the ancestral species of the section *Angulares*. Investigating the process of the species emergence and expansion will provide important insights to understand the evolution of this section.

Vigna khandalensis (Santapau) Raghavan & Wadhwa. *Vigna khandalensis* was reported to inhabit a rainforest climate area in the Western Ghats and the Deccan Plateau in India [44]. It is the only wild species to have an erect plant type in the subgenus *Ceratotropis* in *Vigna*. Its seeds were collected as a food during famines [45]. While Tomooka et al. [8] classified this species in the section *Aconitifoliae* based on the short keel pocket and style beak; Bisht et al. [46] reported that this species is morphologically similar to species in the section *Ceratotoropis*. The phylogenetic trees in this study suggested that *V. khandalensis* is a species in the section *Aconitifoliae*, and located at the basal position to the species in the section *Ceratotoropis*. *V. khandalensis* was most closely related to *V. stipulacea* in the section *Aconitifoliae*, and the two species were similar in that they had large stipules. Since *V. stipulacea* is a creeping plant cultivated as food, fodder, and green manure in Tamil Nadu, India [2], *V. khandalensis* might be used to improve *V. stipulacea* growth. *V. khandalensis* may also be useful as a genetic resource for other section *Ceratotoropis* crops, since the interspecific hybrid between this species and mung bean was obtained [47].

Vigna marina (Burm.) Merrill ssp. *oblonga* Padulosi. *V. marina* ssp. *oblonga* was proposed for the plants growing on the coastal zones of West Africa [19]. The phylogenetic tree using rDNA-ITS in this study confirmed that *V. marina* ssp. *oblonga* was more closely related to *V. luteola* than to *V. marina* ssp. *marina* (Fig 3), which was suggested by isozyme and RAPD analyses [20]. Additionally, phylogenetic trees suggest that there is a large intraspecific variation in *V. luteola*.

To address the evolution of *V. luteola* and *V. marina*, we need to consider *V. oblongifolia* A. Rich., a species closely related to these, although it was not included in this study. In *V. oblongifolia*, two botanical varieties have been described [25]. Phylogenetic trees in the previous studies have shown that *V. oblongifolia* var. *parviflora* is more closely related to *V. luteola* than to *V. marina*, and *V. oblongifolia* var. *oblongifolia* is more closely related to *V. oblongifolia* var. *parviflora* is more distant from these [48, 49]. This suggests that *V. marina* ssp. *oblonga* may be more closely related to *V. oblongifolia* var. *parviflora* than to *V. marina* ssp. *marina*. Therefore, the taxonomic treatment of *V. marina* ssp. *oblonga*,

and *V. oblongifolia* var. *parviflora* should be reconsidered based on intra and inter-specific variations in *V. marina*, *V. luteola*, and *V. oblongifolia*.

Since there are no interspecific crossing barriers among *V. marina* ssp. *marina*, *V. marina* ssp. *oblonga*, and *V. luteola* [17, 50], and interspecific hybrid plants between *V. oblongifolia* and *V. luteola* were obtained [51], these are thought to form a primary gene pool. Therefore, to introduce the salinity and alkaline tolerance of *V. marina* into bambara groundnut, interspecific cross-compatibility should be investigated, taking into consideration the use of bridging species in the section *Vigna*. In Maxted et al. [16], there are 18 species listed in the section *Vigna*.

Vigna vexillata (L.) A. Rich. The wild forms of this species are widely distributed in pantropical regions, including Africa, Asia, Oceania, and America, and its swollen roots have been collected as food [52–54]. This species includes two domesticated forms that are morphologically distinct from each other. One is a twining plant without any taxonomic rank at an intraspecific level, which is cultivated in Bali, Indonesia [13]. Another is an erect plant named *V. vexillata* var. *macrosperma*, which is collected in Africa, Central America, and Australia. For both, the domestication origins are unknown.

In this study, the Indonesian domesticated form (accession ID-53) was found to be genetically differentiated from other species. This suggests that the Indonesian domesticated form, and *V. vexillata* var. *macrosperma* (accession ID-59), have been domesticated independently from different wild forms. This notion was also supported by the fact that a hybrid among the two domesticated forms was not obtained [55]. Moreover, there is an intraspecific crossing barrier between the Indonesian domesticated form and some wild forms [55]. Therefore, the ancestor of the Indonesian domesticated form is unknown.

Similarly, *V. vexillata* var. *lobatifolia* was found to be genetically differentiated from other species. This taxon was described originally as *V. lobatifolia* Baker [56], then classified as an independent species in the section *Pseudoliebrechtsia* [25], or the section *Plectrotropis* [5] in the subgenus *Plectrotropis*, and then given the current rank as botanical variety of *V. vexillata* based on isozyme polymorphisms [15, 16, 57]. However, since *lobatifolia* has a unique habitat (Namib Desert), and is morphologically distinct, we do not reject the taxonomic systems of Verdcourt [25] and Maréchal et al. [5], in which it was treated as an independent species. However, only nine accessions in five varieties of *V. vexillata* were analyzed for the subgenus *Plectrotropis* in this study, and thus further studies are required to systematize the taxonomy of this subgenus, and clarify the rank of the Indonesian domesticated forms and *V. vexillata* var. *lobatifolia*.

The natural habitat of *V. vexillata* was very diverse, including arid lands, coastal areas, acidic soil, and alkaline soil [16, 58, 59]. Some accessions have been reported to harbor flood resistance and pest resistance [60-63]. It is therefore believed that this species contains highly useful genetic resources to breed crops for agriculturally unfavorable lands.

Future perspectives

In recent years, research on the use of wild relatives has been actively pursued. In addition to interspecific cross-breeding, new concepts have been proposed such as 'Reverse Breeding' [64], which involves regaining the crop stress tolerance, which has been lost in the breeding or domestication process, by backcrossing with the wild form. Another strategy is 'Neo-Domestication' [18], or the domestication of the stress-tolerant wild species that cannot be crossed with crop species. This process could be achieved by using mutation breeding, and mutant screening could be accelerated by TILLING, a screening method using the sequence information of domesticated genes. To advance these wild species breeding strategies, more information

concerning the correct taxonomic placement, and genetic relationships among species, should be acquired to predict interspecific cross-compatibility, and to select an appropriate breeding strategy.

Acknowledgments

This work was partly supported by JSPS KAKENHI Grant Number 13J09808 and 26850006.

Author Contributions

Conceived and designed the experiments: YT PS NT. Performed the experiments: YT CM KI KN. Analyzed the data: YT CM. Contributed reagents/materials/analysis tools: YT PS MP NS NT. Wrote the paper: YT NT.

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