

Adapting clonally propagated crops to climatic changes: a global approach for taro (*Colocasia esculenta* (L.) Schott)

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Abstract Clonally propagated crop species are less adaptable to environmental changes than those propagating sexually. DNA studies have shown that in all countries where taro (*Colocasia esculenta* (L.) Schott) has been introduced clonally its genetic base is narrow. As genetic variation is the most important source of adaptive potential, it appears interesting to attempt to increase genetic and phenotypic diversity to strengthen smallholders' capacity to adapt to climatic changes. A global experiment, involving 14 countries

from America, Africa, Asia and the Pacific was conducted to test this approach. Every country received a set of 50 indexed genotypes in vitro assembling significant genetic diversity. After on-station agronomic evaluation trials, the best genotypes were distributed to farmers for participatory on-farm evaluation. Results indicated that hybrids tolerant to taro leaf blight (TLB, *Phytophthora colocasiae* Raciborski), developed by Hawaii, Papua New Guinea and Samoa breeding programmes outperformed local

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cultivars in most locations. However, several elite cultivars from SE Asia, also tolerant to TLB, outperformed improved hybrids in four countries and in one country none of the introductions performed better than the local cultivars. Introduced genotypes were successfully crossed (controlled crossing) with local cultivars and new hybrids were produced. For the first time in the history of Aroids research, seeds were exchanged internationally injecting tremendous allelic diversity in different countries. If climatic changes are going to cause the problems envisaged, then breeding crops with wide genetic diversity appears to be an appropriate approach to overcome the disasters that will otherwise ensue.

Keywords Allelic diversity · *Colocasia esculenta* · Crossing · In vitro distribution · On-farm evaluation · Selection

Introduction

There are uncertainties regarding regional climate change prediction models (Ramirez-Villegas et al. 2013). Hence, the development of strategies aiming at strengthening smallholders' adaptation to climatic changes is problematical and the identification of breeding objectives for the creation of new varieties is challenging. The situation for clonally propagated crop species is quite complex as they are less adaptable to changes in environmental conditions than those propagating sexually (Dodd and Duhovnikoff 2016; Mercer and Perales 2010). As far as tropical root and tuber crops (cassava, sweet potato, yams and aroids) are concerned, smallholders in

developing countries cannot rely on organized breeding programmes or national seed supply systems to renew their germplasm. The vast majority rely on their own cultivars (landraces) obtained through local selection and traditional exchanges. As the development of a new variety takes about 10 years without guarantee of wide adoption, there is a need to find alternative ways of accelerating smallholders' adaptation capacity (Ceballos et al. 2015).

Studies conducted with DNA markers revealed that for most of these roots and tuber crop species, genetic distances correspond to geographic distances and there are different gene pools. Markers also revealed that new variants spontaneously appearing in farmers' plots are hybrids (Scarcelli et al. 2006). In practice, farmers intercrop different genotypes which may flower, genetically recombine and produce viable seeds that germinate and the most attractive volunteers are cloned (Roullier et al. 2013). These new variants are then tested and if they satisfy farmers' needs, they are exchanged with others (Sardos et al. 2012). This suggests that farmers can manipulate germplasm if they have access to sufficient allelic diversity. Traditional networks between communities can then distribute clones efficiently through family ties (VandenBroucke et al. 2015).

Taro (*Colocasia esculenta* (L.) Schott) is a neglected root crop and an "orphan" species with no international research centre with the mandate to assist producers with the development of improved varieties (unlike cassava, sweet potato and yams). It has, however, a recognized potential to strengthen food security. It has been grown in tropical Asia for more than 10,000 years (Fullagar et al. 2006) and is now cultivated throughout the wet tropics (Matthews et al. 2017). According to FAO databases, taro produce the

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lowest yields of all root crops, with an average of only 6.5 tons of fresh corms per ha. In most countries, taro is either a backyard or home garden crop, or is cultivated by smallholders within a shifting agroforestry system with very limited inputs. The world production in 2014 was approximately 11 million tons of fresh corms and cormels from 1.5 million ha but many countries do not maintain or supply statistics for taro (e.g., India, Bangladesh, Burma, Indonesia, Vietnam, Cuba and others) although they are significant producers. The highest yields (>20 t/ha) are obtained in subtropical zones (Egypt, China) (www.fao.org 2014).

The genetic diversity of taro has been well documented. The study of the isozyme variation in more than 1400 cultivars (diploids and triploids) and wild forms from Asia and Oceania revealed greater variation in Asia (Lebot and Aradhya 1991). Isozymes and ribosomal DNA were useful to study the variation in cultivars from China, Taiwan and Japan (Matsuda 2002). AFLP markers have confirmed that diversity is greater in Asia but that the genetic diversity within most countries is low (Kreike et al. 2004). More recently, SSRs were used to investigate diversity in 19 countries of Asia, the Pacific, Africa and America. The highest diversity was again observed in Asia, mainly in India, and clonal reproduction appeared predominant in African and American countries. In West Africa, cultivars were found to have originated from India while in South Africa cultivars shared lineages with Japan. Surprisingly, cultivars from the West Indies were found to have originated from the Pacific, while in Costa Rica they were from India or Asia. To sum up, in all countries where taro has been introduced clonally, its genetic base is narrow (Chair et al. 2016).

Producing crops ready for change—changes to climate, pests and diseases or a need for processing—is particularly difficult for those that are vegetatively propagated and where seed propagation requires special procedures. As genetic variation is the most important source of adaptive variation, it appears interesting to attempt to increase taro genetic variation within smallholders' portfolios. We can expect farmers to use it to adapt to climatic changes. This could be achieved through the geographical distribution of allelic diversity and through the distribution of selected genotypes. A core sample representing the useful diversity of the species has been assembled with the best cultivars from South East Asia (Lebot et al.

2004). It would therefore be interesting to distribute these elite genotypes for direct use or for breeding to countries with narrow genetic bases. Breeding programmes in Papua New Guinea (PNG), Samoa and Hawaii have produced taro hybrids tolerant to the taro leaf blight (TLB, *Phytophthora colocasiae* Raciborski) and the distribution of these could also contribute to strengthen smallholders' adaptation. Finally, segregating progenies obtained from true botanical seeds resulting from controlled crosses between selected parents could be distributed directly to farmers to allow farmers' selection of hybrids corresponding to local needs. Consequently, taro appears as a good model species to study how allelic diversity could be injected in farmers' fields through the propagation, distribution and on-farm evaluation of selected genotypes introduced following international guidelines for the safe movement of germplasm (Zettler et al. 1989).

We conducted a global experiment to compare the performances of taro cultivars and improved hybrids distributed to farmers in 14 different countries of Asia, Africa, America and the Pacific. We present here the results obtained for the research stations propagation process and agronomic evaluation, the on-farm evaluation of introduced germplasm and the first attempts to cross introduced genotypes with local cultivars. We also discuss the practicalities of this approach for taro and its potential for adapting other root and tuber crops to climatic changes.

Materials and methods

Selected genotypes

The geographical origins and major characteristics of cultivars and hybrids distributed and evaluated in this study are presented in Table 1. Fifty-nine cultivars originated from: Indonesia (21), Japan (6), Malaysia (9), the Philippines (2), Thailand (20) and Vietnam (1) were considered. They correspond to some of the best farmers' varieties selected in each country by the Taro Network for South East Asia and Oceania (TANSOA) (Kreike et al. 2004). Forty-one hybrids originated from: the University of Hawaii breeding programme (6), the NARI (National Agricultural Research Institute) breeding programme in PNG (8) and the Secretariat of the Pacific Community breeding

Table 1 List of cultivars (cv) and improved hybrids (hb) distributed to country partners (identified by their international two letters codes)

Code	Name	Origin	Type	TLB	Taste	BF	CR	CU	GH	IN	ID	MG	NG	NI	PG	PH	SA	TT	VU
ID06	ID155	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID18	ID237	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID14	Lamputara	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID31	Manokwari	Indonesia	cv.	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID08	ID178	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID32	ID512	Indonesia	cv.	Susceptible	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID10	ID225	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID12	Kidal	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID16	Lebak	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID19	Hejo	Indonesia	cv.	Tolerant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID20	Apu	Indonesia	cv.	Tolerant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID26	Enrekang	Indonesia	cv.	Tolerant	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID09	ID218	Indonesia	cv.	Tolerant	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID17	Gelang	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID07	ID167	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID23	Lampbar	Indonesia	cv.	Susceptible	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID24	ID392	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID01	ID472	Indonesia	cv.	Tolerant	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID35	Kudo	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
ID22	Semarang	Indonesia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP01	Tsuronoko	Japan	cv.	n.a.	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP03	Miyako	Japan	cv.	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP08	Takenoko	Japan	cv.	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP04	Shogatsu	Japan	cv.	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP06	Akame	Japan	cv.	n.a.	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
JP02	Wasehasuba	Japan	cv.	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY12	Klang	Malaysia	cv.	Resistant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY14	Kluang	Malaysia	cv.	Resistant	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY06	Hitam	Malaysia	cv.	Immune	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY07	Segamat	Malaysia	cv.	Resistant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY11	Banting	Malaysia	cv.	Resistant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY02	Cina	Malaysia	cv.	Immune	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY13	Jenjarum	Malaysia	cv.	Immune	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY03	MY35	Malaysia	cv.	Resistant	Very good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
MY09	Sekinchah	Malaysia	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x

Table 1 continued

Code	Name	Origin	Type	TLB	Taste	BF	CR	CU	GH	IN	ID	MG	NG	NI	PG	PH	SA	TT	VU
PH05	<i>Kalpao</i>	Philippines	cv.	Tolerant	Very good							x							
PH14	<i>PH157</i>	Philippines	cv.	Tolerant	Good													x	
TH24	<i>Boklua</i>	Thailand	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH05	<i>Srisamrong</i>	Thailand	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH08	<i>Ta Daeng</i>	Thailand	cv.	Susceptible	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH09	<i>Klonglan</i>	Thailand	cv.	Susceptible	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH04	<i>Hom</i>	Thailand	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH10	<i>Khamin</i>	Thailand	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH01	<i>Surin</i>	Thailand	cv.	Susceptible	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
TH02	<i>Banmao</i>	Thailand	cv.	Susceptible	Good							x	x						
TH07	<i>Looch Lom</i>	Thailand	cv.	Susceptible	Good							x	x						
TH25	<i>Phayao</i>	Thailand	cv.	Susceptible	Acceptable			x		x	x		x						x
TH30	<i>Sangkom</i>	Thailand	cv.	n.a.	n.a.	x				x	x						x		
TH17	<i>Tha-u-then</i>	Thailand	cv.	Susceptible	Acceptable					x	x								
TH14	<i>TH48</i>	Thailand	cv.	Susceptible	Acceptable				x										x
TH03	<i>Maechan</i>	Thailand	cv.	Susceptible	Acceptable														x
TH19	<i>Phan</i>	Thailand	cv.	Susceptible	Very good		x												
TH12	<i>Hom dam</i>	Thailand	cv.	Susceptible	Acceptable	x													
TH16	<i>Lorncak</i>	Thailand	cv.	Susceptible	Acceptable			x											
SM97	<i>Phuek</i>	Thailand	cv.	Susceptible	Good														x
TH13	<i>TH47</i>	Thailand	cv.	n.a.	Acceptable								x						
TH15	<i>Wangsaiphon</i>	Thailand	cv.	Susceptible	Acceptable														
VN01	<i>Chom tim</i>	Vietnam	cv.	Tolerant	Good										x	x			
HW26	<i>BC99-11</i>	Hawaii	hb	Tolerant	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
HW37	<i>Pa 'akala</i>	Hawaii	hb	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
HW08	<i>PEXPH15-6</i>	Hawaii	hb	n.a.	n.a.	x	x	x	x	x	x	x	x	x	x	x	x	x	x
HW05	<i>2000-21</i>	Hawaii	hb	Tolerant	Acrid					x									
HW12	<i>BC99-6</i>	Hawaii	hb	Tolerant	Good	x	x								x	x			
HW03	<i>19-2001-70</i>	Hawaii	hb	Tolerant	Acrid										x				
PG10	<i>C3-12</i>	PNG	hb	Resistant	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
PG03	<i>C2-E3</i>	PNG	hb	Resistant	Good	x	x		x										
PG11	<i>C3-22</i>	PNG	hb	Resistant	Acceptable	x	x		x										
PG13	<i>C3-46</i>	PNG	hb	Resistant	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x
PG08	<i>C2-E11</i>	PNG	hb	Resistant	Good	x	x	x	x	x	x	x	x	x	x	x	x	x	x
PG09	<i>C3-10</i>	PNG	hb	Resistant	Acceptable	x	x	x	x	x	x	x	x	x	x	x	x	x	x

Table 1 continued

Code	Name	Origin	Type	TLB	Taste	BF	CR	CU	GH	IN	ID	MG	NG	NI	PG	PH	SA	TT	VU
PG12	C3-44	PNG	hb	Resistant	Acceptable					x	x						x		
PG46	C4-70	PNG	hb	Tolerant	Very good					x	x								
SM80	Alafua	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM13	Fanuatapu	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM158	Lalomanu	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM149	Lepa	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM151	Letogo	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM116	Manono	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM115	Manu	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM120	Nu 'utele 2	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM104	Pauli	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM152	Saleapaga	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM43	Samoa43	Samoa	hb	Tolerant	Good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM138	Vaimauga	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM157	Myaeo-le-la	Samoa	hb	Tolerant	Very good					x	x	x	x	x	x	x	x	x	x
SM132	Sapapalii	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM147	Myaela 2	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM135	Matautu	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM136	Moataa 2	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM134	Apia	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM143	Asau	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM111	Samoa13	Samoa	hb	Tolerant	Good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM114	Myaela	Samoa	hb	Tolerant	Very good	x	x		x	x	x	x	x	x	x	x	x	x	x
SM83	Samoaana	Samoa	hb	Tolerant	Very good					x	x	x	x	x	x	x	x	x	x
SM97	Gagasavea	Samoa	hb	Tolerant	Very good	x	x			x	x	x	x	x	x	x	x	x	x
SM46	Samoa46	Samoa	hb	Tolerant	Good	x	x			x	x	x	x	x	x	x	x	x	x
SM152	Samoa10	Samoa	hb	Tolerant	Very good					x	x	x	x	x	x	x	x	x	x
SM10	Tolo-gataua	Samoa	hb	Tolerant	Very good					x	x	x	x	x	x	x	x	x	x
SM111	Salelologa	Samoa	hb	Tolerant	Very good					x	x	x	x	x	x	x	x	x	x

BF Burkina Faso, CR Costa Rica, CV Cuba, GH Ghana, IN India, ID Indonesia, MG Madagascar, NG Nicaragua, NI Nigeria, PH Papua New Guinea, PG Papua New Guinea, SA South Africa, TT Trinidad and Tobago (St Vincent), VU Vanuatu

programme in Samoa (27). All hybrids were obtained through controlled crosses and successive recurrent selection cycles. Overall, 100 genotypes were selected for their geographical distances and genetic diversity, their corm shape and quality, and their tolerance or resistance to TLB (Table 1).

Indexation of germplasm

All taro accessions were maintained *in vitro* in the Pacific Community (SPC) Centre for Pacific Crops and Trees, Suva, Fiji. Tissue cultured accessions were grown in 100 mL glass jars (Cospak, Australia), containing 20 mL of Murashige and Skoog basal medium, supplemented with 3% sucrose, benzylaminopurine (1.0 mg/L) and naphthalene-acetic acid (0.3 mg/L). Cultures were maintained at a temperature of 20 °C under a day length of 16 h (Taylor 2002). *In vitro* plantlets were raised from parent plants which were inspected, screened and found to be free from *P. colocasiae*. Further, the taro plantlets were derived from mother plants which have been indexed negative at three, and again at 6 months of growth, for each of the four known taro viruses (*Dasheen mosaic virus*, *Taro bacilliform virus*, *Taro vein chlorosis virus*, and *Colocasia bobone disease virus*) using highly sensitive polymerase chain reaction, with both negative and positive controls included in all tests. Only suckers derived *in vitro* from an original meristem, which was tested negative for viruses, were considered as negative for those viruses. These suckers were used to provide the source from which all clones of selected genotypes were obtained. The virus indexing protocol used was developed by Queensland University of Technology in Australia under the AusAID-funded Taro Genetic Resources: Conservation and Utilization project (1998–2003) (Harding et al. 2004).

International distribution

Between June and November 2011, 50 genotypes (three *in vitro* clones per genotype) were sent to 14 different country partners: Costa Rica (University of Costa Rica, San José), Nicaragua (University of Nicaragua, Managua), Cuba (INIVIT, Instituto Nacional de Investigaciones de Viandas Tropicales, Santa Clara), St Vincent (CARDI, Caribbean Agricultural Research and Development Institute, Trinidad and Tobago), Burkina Faso (Université de Ouagadougou),

Ghana (CSIR, Plant Genetic Resources Research Institute), Nigeria (NRCRI, National RootCrops Research Institute, Umudike), South Africa (ARC, Agricultural Research Council, Pretoria), Madagascar (FOFIFA, Centre National de la Recherche Appliquée au Développement Rural, Antananarivo), India (CTCRI, Central Tuber Crops Research Institute, Trivandrum), Indonesia (LIPI, Indonesian Centre for Research and Development in Biotechnologies, Bogor), the Philippines (PhilRootCrops, Baybay, Leyte), Papua New Guinea (NARI, National Agricultural Research Institute, Lae), Vanuatu (VARTC, Vanuatu Agricultural Research Training Centre, Santo). Expeditions were done as soon as an import permit was signed by the authorized official institution in the importing country partners. Distributions from SPC to the partners were made under the Standard Material Transfer Agreement of the International Treaty on Plant Genetic Resources for Food and Agriculture.

On-station propagation and evaluation

Upon receipt, all country partners transferred the introduced *in vitro* plantlets into nurseries. The plastic pots were filled with a compost/soil mix sterilized with good drainage and aeration. The plantlets roots were washed thoroughly to completely remove the culture medium containing nutrients and sugar which attract fungi and bacteria. The plantlets were then covered with clear plastic bag without holes for the first 4 weeks to maintain high humidity and help the plants to adjust to their new environment. After 4 weeks, the plastic bags were gradually removed. Developing plants were watered three or four times a week on alternative days depending on the weather and screen house conditions. Field propagation was initiated in 2012. During 2 years, plants of selected varieties were intensively propagated on-station (through headsets corresponding to the summit of the corm and/or suckers). During the propagation and on-station evaluation process, local cultivars were used for comparison with introduced genotypes. Local and introduced genotypes were planted at 1 × 1 m spacing. The local checks were the best local cultivars identified in each participating country during the previous years of characterization of national germplasm collections. A range of qualitative and quantitative morphological and agronomic data were collected as part of the

evaluations; these included, stolon and sucker production, flowering ability, vigour, plant height, TLB tolerance and corm yield. Five plants per genotype were scored during two successive years before selecting the most interesting genotypes for further propagation.

Production of true taro seeds

Partners produced controlled crosses following protocols described in details elsewhere (Ivancic and Lebot 2000). Briefly, emasculation was conducted 2 days before the inflorescence opened by cutting the upper part of the spathe and the whole male portion of the spadix. After pollination, female flowers were protected with the lower green part of the spathe which was removed during emasculation. Fruit heads were usually ready to be harvested 1 month after pollination. Seeds can be germinated 1 week after sowing and/or can be kept for several years in a glass flask or ziplock plastic bag with silica gel in deep freeze. In each country, the best parents were selected after on-station evaluation, crosses were made and seeds were collected and germinated. In Vanuatu and PNG, seeds obtained from open pollination between selected hybrids, were collected and bulked. In June–July 2015, these seeds were distributed to all partners and germinated upon reception. Seedlings were raised in nurseries and hybrid plants in their first clonal generation were distributed to farmers for screening, each farmer receiving only one clone per hybrid in batches of 30 hybrids (Fig. 1).

On-farm evaluation

Genotypes for on-farm trials were selected primarily on yield performance. All introduced genotypes producing 10 and more t/ha (more than 1 kg/plant) were included in the list for on-farm trials. In order to increase genetic diversity in farmer fields, the best and popular local varieties were not included even though they produced good yield because they were already quite widespread in farmer's fields. Although, several local cultivars produced yields between 4 and 9 t/ha, preference was always given to exotic genotypes that showed high or moderate levels of resistance to TLB and had good eating qualities. A total of 30 accessions were selected for on-farm trials. In mid-2014, headsets of selected genotypes were distributed to farmers for

evaluation. Depending on the number of headsets available and on the number of volunteer farmers, the number of sites (villages) varied per country partner (Table 2). At harvest, corms were weighted and farmers were invited to taste their organoleptic characteristics following the most common cooking preparation: peeling, washing, cutting into pieces of approximately 5 × 5 cm and boiling in water. Farmers were invited to score each genotype (cultivar or hybrid) for taste, texture, aroma and acidity on a scale from 1 (poor) to 5 (excellent) without being replicated.

Results

Due to insufficient availability of indexed *in vitro* plantlets, partners received different sets of cultivars and hybrids. However, 18 genotypes including cultivars from Japan (JP01), Malaysia (MY12, MY14), Thailand (TH24, TH08) and hybrids from Hawaii (1), and Samoa (12) were distributed to 14 country partners and 33 genotypes were common to 10 countries (Table 1). Each of the 14 participating countries received cultivars and hybrids from different geographical origins. Cultivars originating from Indonesia, Japan, Malaysia and Thailand had already been fingerprinted with isozymes and DNA markers (Lebot and Aradhya 1991; Kreike et al. 2004) to confirm their genetic distances and were distributed to all countries. Hybrids from the breeding programmes in Hawaii, PNG and Samoa were also distributed to all partners (of course, hybrids from PNG were not sent to PNG). These hybrids have been produced through crosses between TANSO elite genotypes and local cultivars and were therefore highly heterozygous (Cho 2004; Iosefa et al. 2012). Overall, significant allelic diversity was therefore introduced in each country with 50 selected genotypes (Table 1).

The regional germplasm centre of SPC in Fiji has developed an efficient system for distribution of plants in plastic pouches (Taylor et al. 2004) and contamination rates were extremely low, and accordingly transfers to soil were highly successful. In most countries, the number of introduced genotypes evaluated on-station was high with rates of 100% in Costa Rica, the Philippines and Papua New Guinea. However, in three countries (Burkina Faso, Nigeria, Indonesia), the number of genotypes successfully evaluated on-station was lower and this was reported

Fig. 1 **A** In vitro plantlets received by partners, **B** transplanting young plants into plastic pots in nurseries, **C** on-station propagation plots after cyclone Haiyan in November 2013 in Baybay, Leyte, the Philippines, **D** field propagation of selected genotypes in Nicaragua, **E** preparing batches of headsets for distribution to farmers in the East Coast of Madagascar, **F** distribution of batches to farmers in Costa Rica, **G** discussing vegetative growth with farmers on the Highlands of Madagascar, **H** harvesting on-farm trials in Morobe Province, Papua New Guinea, **I** cross-pollinated fruit head in Santo, Vanuatu, **J** true botanical seeds ready for international exchange



Table 2 Selected genotypes distributed to farmers for participatory evaluation

Country	Local cultivars maintained on-station	Introduced genotypes evaluated on-station	Local cultivars used for comparison	Genotypes (cvs + hbs) selected on-station	Genotypes distributed to farmers	On-farm trials sites (villages)	Farmers involved
Costa Rica	2	50	2	30	30	9	52
Nicaragua	6	43	8	25	31	8	16
Cuba	102	48	5	18	0	0	0
St Vincent	2	45	2	9	30	8	25
Burkina Faso	18	28	18	22	22	4	66
Ghana	81	37	81	30	30	8	60
Nigeria	5	25	5	9	9	4	50
South Africa	81	49	29	15	20	4	10
Madagascar	8	37	8	10	27	5	143
India	424	37	3	37	35	12	52
Indonesia	252	26	4	30	30	4	50
Philippines	240	50	5	16	16	6	100
Papua New Guinea	279	50	10	30	34	11	21
Vanuatu	302	47	50	30	24	8	16

to be due to poor adaptation observed at the nursery stage (Table 2). In 2012, several partners provided photographs of their nurseries and/or propagation plots and these have been posted on the International Network for Edible Aroids (INEA) website (www.ediblearoids.org). Overall, the in vitro introduction process was considered by all partners as fairly simple and technically easy to implement.

There were noticeable differences in the performance of the introductions, and the results from the evaluations, both on-station and in farmers' fields (Table 2). For instance, at the nursery stage, it soon appeared that Japanese genotypes were not adapted to the tropical conditions of most countries and their growth and yield measured on-station were too low to allow further field propagation. However, some of these Japanese cultivars presented good eating quality (observed in PNG) and good pollen fertility (observed in Cuba) and were therefore kept cautiously in germplasm collections and were used as parents in crosses. The number of local checks used during the on-station evaluation process varied from only two local cultivars (Costa Rica), whereas there were 50 in Vanuatu and 81 in Ghana. In Vanuatu and Ghana, introduced genotypes were planted and harvested together with the local germplasm collections, so that

the introductions could be compared with a large number of local cultivars. In most countries, the on-station evaluation trials confirmed the superior performance of the introductions in terms of yield, vigour, TLB tolerance of the breeding lines from the Hawaii, PNG and Samoa breeding programmes. TLB has been introduced to western Africa and is now spreading very rapidly throughout this region (Omame et al. 2012). However, *P. colocasiae* is not a threat yet in Costa Rica, Nicaragua, Cuba, St Vincent, South Africa and Vanuatu but could be introduced anytime.

After 2 years of on-station evaluation, the number of selected genotypes (local or introduced cultivars and hybrids) ranged from nine in St Vincent and Nigeria to 37 in India. Taro farmers in St Vincent are focusing on the export market to the US. In this case, it was thought more appropriate to select on-station only those genotypes with traits corresponding to the local cultivar used to satisfy market demands; accordingly, only nine genotypes were selected for on-farm evaluation. The number of genotypes distributed to farmers ranged from 35 genotypes in India to zero in Cuba, where on-farm evaluation continues before distribution of selected genotypes through the national scheme. Hence, the number of distinct evaluation sites and farmers involved in on-farm trials varied greatly

between countries. In Madagascar, for instance 143 farmers received selected genotypes for on-farm evaluation (Table 2).

After two successive years of on-farm evaluation trials, partners ranked their best five genotypes based on on-station and on-farm evaluation trials (Table 3). Hybrids from Hawaii breeding programme (HW nos) performed well in Costa Rica, Nicaragua, Cuba and Ghana. Hybrids from Samoa (SM nos) performed well in Burkina Faso, Madagascar, the Philippines and Vanuatu. One hybrid from PNG (PG03) performed well in Nigeria. Interestingly, introduced cultivars performed better than hybrids in St Vincent, South Africa, India and Indonesia. In Papua New Guinea, an improved hybrid from the fifth recurrent selection cycle (C5-353) outperformed introduced genotypes. However, in South Africa, none of the 50 introduced genotypes performed better than the local cultivars. Overall, in the 14 countries, 64 introduced genotypes (54 hybrids and 10 cultivars) were ranked in the five best genotypes and performed better than local taro cultivars.

Except for PNG, India and Vanuatu, in most countries, taro breeding was at a very early stage before this global experiment was initiated. Therefore, the introduction of genotypes aimed at introducing parents to produce local hybrids tolerant to TLB with good quality corms, and to distribute them to farmers to make adaptation to climatic changes possible. The results of the crosses conducted in participating countries between introduced genotypes and local

cultivars are presented in Table 4. In Costa Rica, nine crosses were conducted and after evaluation of 90 seedlings, there are nine selected hybrids under further test in 2017. These are from crosses between local cultivars and hybrids from Samoa and Hawaii. Selections were made for low numbers of suckers and a large round main corm. In Cuba, considerable seedling variation has been produced from 23 successful hybridisations, using female parents from Thailand, Malaysia, Samoa and Japan. The cultivar *Miyako* (Japan) and the Samoa hybrid *Manu* (SM135) were found to be the best pollen donors.

In Burkina Faso, the best exotic genotypes used as parents were PG10, ID14, ID06, ID24, SM13, SM138 and TH12. More than 200 crosses were made, of which 96 were successful and the germination rate was 34%. There are now selected 70 hybrids and these are being multiplied. In South Africa, flower induction of the local cultivars was successful but pollinations failed. Most of the local cultivars were found to be triploids (Chair et al. 2016). In future, the introduced genotypes will be used as female parents and the local ones as pollen donors. In Madagascar, hybridisations were attempted but the local cultivars did not flower, so crosses were made between introductions and between them and wild taro genotypes flowering naturally. As taro, has been introduced clonally with the first Austronesian migrations ca. 2000 years ago, these wild types are in fact escaped from cultivation and naturalised. Of the 60 crosses made, 37 came to maturity, and more than 20,000 seeds were sown. The

Table 3 Best five selected genotypes after two years of on-station evaluation (2012–14) and two years of on-farm evaluation (2015–16) (cultivars are in italics)

Country	1st	2nd	3rd	4th	5th
Costa Rica	HW37	SM46	SM116	SM128	PG13
Nicaragua	HW26	PG03	SM151	SM158	<i>ID24</i>
Cuba*	HW37	SM143	C3-12	SM128	SM80
St Vincent	<i>ID24</i>	SM80	SM149	SM83	PG09
Burkina Faso	SM80	SM135	PG11	HW05	SM120
Ghana	HW37	SM151	SM10	<i>ID24</i>	SM134
Nigeria	PG03	<i>TH05</i>	SM158	<i>ID12</i>	SM80
South Africa	<i>Thandizwe</i>	<i>Amzam</i>	<i>Mabhida</i>	<i>Ngubane</i>	<i>Gumede</i>
Madagascar	SM80	SM115	SM157	SM43	SM128
India	<i>ID06</i>	<i>TH10</i>	<i>TH07</i>	SM116	SM151
Indonesia	<i>TH05</i>	<i>Mentega</i>	SM135	SM157	SM134
Philippines	SM115	SM151	SM132	SM80	<i>MY112</i>
Papua New Guinea	C5-353	SM143	SM148	SM43	C5-245
Vanuatu	SM13	SM138	PG11	SM149	SM120

* On-station evaluation only

Table 4 Summary results of crosses conducted between introduced and local genotypes

Country	Parents selected	Crosses conducted	Seeds obtained	Seedlings transplanted	Selected local hybrids
Costa Rica	6	9	286	90	9
Nicaragua	6	3	0	0	0
Cuba	19	23	2037	n.a.	n.a.
St Vincent	–	–	–	–	–
Burkina Faso	38	205	22,365	7604	70
Ghana	12	12	n.a.	1680	12
Nigeria	14	22	7316	259	180
South Africa	–	–	–	–	–
Madagascar	19	60	n.a.	>20,000	1053
India	12	10	n.a.	234	32
Indonesia	30	–	n.a.	–	–
Philippines	16	12	n.a.	1111	n.a.
Papua New Guinea	60	200	n.a.	>4000	931
Vanuatu	22	18	2319	2040	242

seedlings (7604) are still under evaluation on the East coast lowlands (Ilaka research station). In addition, seeds were obtained from Vanuatu in 2015. More than 5000 seedlings were raised and 1790 given to ten farmers and 800 planted on-station on the East coast. In Ghana, twelve successful hybridizations resulted from crosses between ten introduced genotype and two local cultivars. A total of 1680 F1 seedlings were distributed to 28 farmers for on-farm evaluation and twelve were selected and are under propagation. In Nigeria, 180 seedlings have been cloned and included in clonal evaluation trials. The seeds received have also been germinated and F1 hybrids are being evaluated.

In Indonesia, profuse natural seeds set occurred naturally among introduced genotypes and local cultivars. As various *Diptera* spp. are active pollinators, open pollinated seeds were produced. In the Philippines, the introduced genotypes have increased the number of potential parents for hybridization. Natural flowering occurred in SM115 and SM151 and they produced open pollinated seeds. Artificial induction of flowering had to be done as synchronization of flowering with local cultivars was a problem. In India, breeding was also hindered due to asynchrony in flowering and an attempt was made to cryostore pollen at different time intervals ranging from 1 week to 2 months and to use the cryostored pollen for hybridization with introduced genotypes. Fruit setting was observed within a week of crossing introduced

genotypes with cryopreserved pollens (Mukherjee et al. 2016). Introduced genotypes were also crossed in Bhubaneswar (Orissa) and 31 crosses recombining TANSO cultivars and Samoa hybrids were successful. The seeds of these 31 full-sib families were sent to VARTC in Vanuatu where they were germinated and 5214 hybrids were obtained and are now under selection. In Papua New Guinea, 57 genotypes were selected as parents for hybridizations, including 15 cultivars from the NARI core collection, 28 introduced genotypes, and 14 hybrids from the NARI breeding programme. Two hundred crosses were made of which 93% developed seeds and of these 60% germinated. Subsequently, over 4000 seedlings are now under selection in 46 progeny trials in Madang and Morobe provinces (northern coast of PNG).

In Vanuatu, controlled crosses were made to produce full sib families aimed at evaluating broad sense heritability and genetic gain for traits related to vegetative growth, yield, and corm quality. A fully randomized-block trial consisting of 13 full-sib families (2040 F1 hybrids) was established and measured in F1 (seminal generation) and first clonal generation (C1). Statistical tests revealed the stability of the presence or absence of stolons, suckers, and inflorescences between F1 and C1, suggesting strong genetic control for such traits. The number of stolons, the number of suckers, fresh corm weight, and dry matter content were found to be the most heritable traits, indicating that breeders should focus on those for

eliminating undesirable hybrids in early clonal generations (Soulard et al. 2016).

Discussion

If climatic changes are going to cause the problems envisaged (severe droughts, heavy rains and/or development of pests and diseases), then breeding crops with wide genetic diversity is an appropriate approach to overcome the disasters that will otherwise ensue. The work described here has successfully broadened the genetic bases of taro in 14 countries through the introduction, on-station and on-farm evaluation of exotic germplasm. Country partners have conducted the first crosses and generated thousands of seedlings. These new genotypes have been screened, evaluated, and distributed to producers, representing an exceptional resource of genetic diversity. Also, partners started to exchange internationally seeds of taro and this is a major breakthrough as it is the first time in the history of Aroids research that it is done on such a large scale. This process has injected tremendous allelic diversity in different countries.

During this 6 years global experiment, we have shown that when introduced genotypes of taro satisfy farmers' needs, they are readily adopted, although more time is obviously needed to confirm their final adoption. For subsistence farmers, quality taste criteria are always the most important for adopting new clones but farmers are also interested in variation and in new morphological traits (Camus and Lebot 2010). If the introduced genotypes continue to correspond to local agronomic expectations, and if their cooking properties, organoleptic qualities and texture remain acceptable over the next clonal generations, they will be propagated and shared with other communities. The more they are appreciated, the more they are propagated and exchanged with other farmers (Sardos et al. 2012). Several countries reported on-station theft of introduced genotypes before they could be distributed because of their attractive morphological appearance (e.g., red petioles or waxiness of leaf blades). Local adaptation and yield are, of course, also appreciated but their evaluation takes several clonal generations before it can be confirmed. In the present study, only 2 years of on-farm evaluation trials were conducted and it is too soon to conclude but it is possible to say that introductions had a significant impact on taro

diversity as in all countries, except South Africa, cultivars or hybrids were selected.

Once parents are selected, the controlled crosses aim at producing the maximum number of seedlings for field evaluation. Unfortunately, around the world, the most interesting cultivars of *C. esculenta* do not flower naturally and it is necessary to induce flowering with gibberellic acid for breeding purposes. Detailed crossing protocols to assist taro breeders are now available (www.ediblearoids.org) but in some countries GA₃ is difficult to find. Also, success rates in controlled crosses of taro are frequently very low (approx. 30% of hand crosses produce seeds) and depending on the parents, the number of seedlings per cross may vary from 0 to 400 (Ivancic and Lebot 2000). In the present global experiment, all countries followed the same approach: identification of the best parents and induction of flowering. When successful controlled crosses were made with a low success rate but with sufficient number of hybrids (often several hundreds) (Table 4), the screening of hybrids was done using visual tools (e.g., presence or absence of suckers and stolons) to allow the rapid elimination of undesirable genotypes in the seminal (F1) and first clonal generation (C1).

More advanced breeding programmes in Samoa (Iosefa et al. 2012), Hawaii (Cho 2004) and PNG (Ivancic and Lebot 2000) have proceeded similarly for the last three decades and lessons have been learned for taro breeding. Selection is always made on the individual value of the parents. In Papua New Guinea, the breeding programme attempted to use a wild genotype from Thailand (called *Bangkok*) as a source of resistance to TLB (Singh and Okpul 2000). Unfortunately, it was soon evident that the incorporation of resistance brought along several deleterious traits, such as stolons and acidity. Now this programme and others are using tolerant parents to produce horizontal resistance. The present study has confirmed the validity of this approach with several PNG hybrids successfully selected in Costa Rica, Nicaragua, Cuba, St Vincent, Burkina Faso, Nigeria, and Vanuatu (Table 3). In Samoa, the breeding programme was initiated with the introduction of good (non-acrid) foreign cultivars resistant to TLB, first from Micronesia and then from SE Asia (TANSO) and presenting significant genetic diversity (Kreike et al. 2004). Several selected hybrids were distributed and accepted by farmers (Fonoti 2005).

The results obtained in the present global experiment indicate that the Samoan breeding programme has also been successful as many hybrids (SM genotypes) are now accepted by farmers in 14 countries. A similar approach was used in Hawaii and the introduction of TANSO cultivars allowed the production of hybrids resistant to TLB. In this case again, taro breeding has been successful as demonstrated by the number of hybrids (HW genotypes) selected by farmers (Table 3).

However, in most countries there are still serious technical constraints needing solutions before taro improvement programmes can progress efficiently. The somewhat limited number of successful controlled crosses (Table 4) is impairing the creation of diversity needed for further selection but asynchrony, pollen viability and the receptiveness of female flowers are severely impacting the success rates of these crosses. Protocols for the cryopreservation of pollen, either freeze dried or cryostored in liquid nitrogen have been developed (Mukherjee et al. 2016) but more work is needed, especially with freeze dried pollen in order to ease and speed up the control crosses.

When several full-sib families are obtained there is then a need for robust field comparison and there is also a need for a field design able to control the environmental variance. If there are between 100 and 300 hybrids per family, the field experiment layout becomes fairly expensive to maintain. As taro breeders need efficient visual tools to screen thousands of seedlings successfully at the F1 and C1 stages (Ivancic et al. 2003), a Vegetative Growth Index (VGI) was developed to predict the final yield of hybrids during their vegetative phase (before 5 months). This multi criteria index combines the measurements of the number of suckers, stolons, the plant height, leaf length and width to predict the yield of the plant. The correlation analysis between individual plants in F1 and C1, has confirmed the suitability of VGI as a corm yield prediction tool for screening hundreds of hybrids (Soulard et al. 2016). In several countries (Madagascar, PNG, Vanuatu), the VGI has been shown to be a practical and efficient tool.

Field evaluation of hybrids is then constrained by the production of homogenised propagules: headsetsets of taro corms produce vigorous plants while young suckers and stolons do not. When evaluating new hybrids, it is observed that intraclonal variation is often very high between different propagules of the

same clone (e.g., headset, suckers or stolons or corms originating from the same clone). It is therefore difficult to obtain sufficiently homogenised planting materials to establish reliable field trials with the needed replications for accurate phenotyping. Hence, this physiological heterogeneity gives significantly different results during the first evaluation trials and is often confusing. The availability of sufficient homogenised planting materials (number of clones per genotype) impacts directly the reliability of the evaluation data because of the insufficient number of replications, the small plot sizes, and the small number of sites where these genotypes are evaluated. Also, the relatively long growth cycle for taro (8–10 months) is a potential source of errors when evaluating genotypes for yield and dry matter content as these two major traits can vary with the month of harvest.

Finally, G x E interactions necessitate the establishment of multi-locations trials but the constraints imposed by accurate evaluation are rendering this approach very complex (Ivancic and Lebot 2000). A way of coping with these practical and financial constraints is to work with farmers for the evaluation, as shown in the present study (Table 3). As diversity exists because of genetic resources distribution and adaptation by localized populations (Namkoong et al. 2004), the introduction of allelic diversity represents an attractive approach for strengthening farmers' adaptation if this diversity is adopted and used (Lebot et al. 2005). Farmers are often pushing crop populations through a new evolution, adding favourable alleles to the gene pool while maintaining diversity (Birnbaum 2006). Farmers often maintain landraces while adopting modern varieties (Brush 2000). They know how to manage their varietal portfolios to optimise risk management. If allelic diversity is introduced, farmers' capacity to adapt to forthcoming changes will be strengthened by the access to new diversity.

Conclusions

This global experiment was set out to develop a simple system for speeding up the improvement of neglected clonally propagated crops using taro as no one has the international mandate to work on this crop. To test the approach, this experiment shared germplasm of wide allelic diversity among 14 partners, evaluated and

compared 50 introduced genotypes with local cultivars, gave the best selections to farmers for evaluation and initiated local breeding programmes. The approach was participatory, effectively distributing genotypes to farmers in the shortest time possible due to the time needed for bulking clonally the planting materials. In most participating countries it worked well, although there were obvious differences depending on the local means available. In order to strengthen smallholders' capacity to adapt to climatic changes, the global approach tested for taro could be used for other species, especially neglected ones such as minor yams (*Dioscorea* spp.) and aroids (*Alocasia macrorrhiza*, *Cyrtosperma chamissonis*, *Xanthosoma sagittifolium*). Plant breeding offers a solution to climate change adaptation but financial costs and the lack of research capacity are major constraints. On-farm evaluation and selection might represent a cost efficient approach.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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