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Evolutionary Processes Involved in the Emergence and Expansion of an Atypical *O. sativa* Group in Madagascar



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Abstract

Understanding crops genetic diversity and the evolutionary processes that accompanied their worldwide spread is useful for designing effective breeding strategies. Madagascar Island was one of the last major Old World areas where human settlement brought the introduction of Oryza sativa. Early studies in the island had reported the presence of a rice group specific to Madagascar. Using 24 K SNP, we compared diversity patterns at the whole genome and at haplotype (30 SNP-long segments along the genome) levels, between 620 Malagasy and 1929 Asian rice accessions. The haplotype level analysis aimed at identifying local genotypic variations, relative to the whole genome level, using a group assignment method that relies on kernel density estimation in a Principal Component Analysis feature space. Migration bottleneck had resulted in 10–25% reduction of diversity among the Malagasy representatives of indica and japonica populations. Compared to their Asian counterpart, they showed slightly lower indica and japonica introgressions, suggesting the two populations had undergone less recombination when migration to the island occurred. The origins of the Malagasy indica and japonica groups were delineated to indica subpopulation from the Indian subcontinent and to tropical japonica from the Malay Archipelago, respectively. The Malagasy-specific group (Gm) had a rather high gene diversity and an original haplotype pattern: much lower share of indica haplotypes, and much higher share of Aus and japonica haplotypes than indica. Its emergence and expansion are most probably due to inter-group recombination facilitated by sympatry between indica-Aus admixes and "Bulu" type landraces of japonica in the central high plateaux of Madagascar, and to human selection for adaptation to the lowland rice cultivation. Pattern of rice genetic diversity was also tightly associated with the history of human settlement in the island. Emergence of the Gm group is associated with the latest arrivals of Austronesians, who founded the Merina kingdom in the high plateaux and developed lowland rice cultivation. As an intermediary form between Aus, indica and japonica, the three pillars of O. sativa domestication, Gm represents a very valuable genetic resource in breeding for adaptation to cold tolerance in tropical highlands. We proposed the name Rojo for this new rice group.

Keywords: Rice, *Oryza sativa*, Madagascar, Diversity, Evolutionary process

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Introduction

Rice, Oryza sativa L., displays very large morphological and physiological variations contributing to its cultivation as a food crop in a very large range of environments (Maclean et al., 2002). This phenotypic diversity is associated with differentiation into two subspecies, indica and japonica (Oka, 1983), and at least two minor ecotypes, Aus and Basmati, distinguishable by isozyme (Glaszmann, 1987) and DNA markers (Garris et al., 2005). Recently, analysis of population structure of O. sativa, from the 3000 Rice Genomes Project (3 K-RG, 2014), identified nine subpopulations, of which four belonged to the indica subspecies, three to the japonica subspecies, and the remaining two encompassed the Aus and Basmati ecotypes. The differentiation of the subpopulations was associated with some eco-geographical specialisation. The *indica* subpopulations named XI-1A, XI-2 and XI-3 mainly originated from East Asia, South Asia and Southeast Asia, respectively. The subpopulation XI-1B assembled modern indica varieties of diverse origins. The japonica subpopulations named GJ-trop, GJsbtrop and GJ-temp mainly originated from tropical Southeast Asia, subtropical Southeast Asia, and temperate East Asia, respectively. The subpopulations encompassing Aus and Basmati ecotypes were named cAus and cBas, respectively (Wang et al., 2018).

While the domestication of O. sativa goes back to an estimated 10,000 years (Higham and Lu 1998; Sweeney and McCouch, 2007), the seniority of *indica-japonica* differentiation within its wild ancestor O. rufipogon has been estimated at more than 100,000 years (Wang et al., 1992; Ma and Bennetzen, 2004). Based on this early differentiation, it was often concluded that O. sativa had undergone two independent domestications from the divergent pools of O. rufipogon (Second, 1982; Cheng et al., 2003; Vitte et al., 2004). However, Molina et al. (2011), analysing the polymorphism of 630 gene fragments, using a demographic modelling approach, concluded to a single origin of Asian domesticated rice. Likewise, Huang et al. (2012), analysing the domestication sweeps and genome-wide patterns in genome sequences of 446 accessions of O. rufipogon and 1083 accessions of O. sativa, concluded that japonica rice was first domesticated from a specific population of O. rufipogon in southern China. Indica rice was subsequently developed from crosses between japonica rice and local wild rice as the initial cultivars spread into South East and South Asia. On the other hand, considering the domestication not as an event but as an evolutionary process promoted by interactions between plant and man, Oka (1988) suggested multiple and diffuse domestications of O. sativa, in a large area stretching from the Himalayan footsteps of India to China. Glaszmann (1988), reporting marked isozymic differentiation among minor ecotypes, such as Aus and Aromatic, in the Indian subcontinent, reinforced the assumption of diffuse domestications and the major contribution of local wild forms. The fact that these ecotypes contained unique alleles not found in indica and japonica was later confirmed by Jain et al. (2004) and Garris et al. (2005) using SSR markers. Likewise, a recent reinterpretation of Huang et al. (2012) data concluded on three sources of domestication, distributed across the Indica, Japonica and Aus, and the Basmati-like cluster to be a hybrid between Aus and Japonica (Civan et al., 2015). However, the presence of some weedy rice among the plant material somewhat compromises this interpretation. The hypothesis of multiple independent domestication events was further reinforced by the analysis of patterns of introgressions in nine important domestication genes in the 3 K-RG accessions (Wang et al., 2018).

Understanding the evolutionary processes that accompanied rice domestication and spread has been complicated in Asia (i) by large-scale migration of cultivated rice accompanying human movements across the continent in areas where wild ancestor populations were already present, (ii) by recombination between different O. sativa subpopulations (Kovach et al., 2009, Santos et al., 2019) despite some level of reproductive barrier, and (iii) more recently, by the trading of cultivated rice varieties. Analysis of rice diversity pattern in areas where O. rufipogon is absent may provide insights into O. sativa expansion processes. The Madagascar Island represents a good case of such an expansion area. According to human genetic data, the island's human settlement of Indonesian origin goes back most probably between 1400 and 1200 years ago (Tofanelli et al., 2009; Cox et al., 2013). Rice is the country's staple food and is grown wherever possible with very ingenious developments in inland valleys and terraces on hillsides. Among the wild relatives of O. sativa, so far only the presence of O. longistaminata and O. punctata has been reported (FAO, 1996); and the African cultivated rice, O. glaberrima, is absent from the island.

Studying the variability of morpho-physiological traits in the national collection of rice varieties established in the 1940s and conserved ex situ by the national agronomic research institute (FoFiFa), Ahmadi et al. (1988) identified, in addition to representatives of *indica* and *japonica* subspecies, an atypical group specific to Madagascar and preferentially present in the central high plateaux of the country. Compared to the *indica* landraces, the atypical landraces were taller, had a larger stem diameter, longer and larger leaves, longer panicles, shorter but wider grains and a shorter growth cycle. The finding of an atypical group was confirmed later using isozymic data (Ahmadi et al., 1991). More recently, Mather et al. (2010), using sequence data from 53 gene

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fragments, reported a clear differentiation of Madagascar *indica*-type landraces from Asian *indica*, as well as evidence of *indica/japonica* recombinations. Likewise, Radanielina et al. (2012), analysing the in situ ecogeographical distribution of *O. sativa* in the highlands of Madagascar, using SSR markers, confirmed the presence of the atypical group and reported a high level of withinvariety diversity. Understanding the evolutionary processes (e.g. founder events, hybridisation and selection) that accompanied the introduction and spread of rice in Madagascar will also affect the conservation and utilisation strategies for the valuable genetic resources it represents.

The availability of the 3 K-RG project data and the feasibility of producing dense genotypic data for a large set of Malagasy rice accessions provided us with the opportunity to analyse the original diversity of the Malagasy rice gene pools on a fine scale. Here, using a 24 K SNP dataset, we describe the pattern of rice diversity in Madagascar relative to the reference pattern observed in Asia. The observed pattern confirmed the presence in Madagascar of a group not identifiable as such in Asia. The evolutionary processes most probably involved in the emergence and expansion of this original ecotype are (i) multiple hybridisations and recombinations between indica and cAus subpopulations in the context of their sympatry in south Asia; (ii) multiple hybridisations and recombinations between the indica-cAus intermediary form that reached Madagascar with lowland-grown tropical japonica subpopulation in the context of their sympatry in the highlands of the country; (iii) favourable selection pressure for the survival of the recombinant forms in the cold climate of the island's high plateaux.

Materials and Methods

Plant Material

The study included a Malagasy diversity panel and an Asian reference panel. The Malagasy panel was composed of 620 Malagasy rice landraces belonging to the Malagasy national collection of rice varieties. Among those accessions, 410 (111 collected before 1980, and 299 collected in 2012) represented the countrywide rice diversity. The remaining 210 accessions were collected in 2007, in the high-plateau (Vakinankaratra) region of the country (Supplementary Table 1). Each accession of the Malagasy panel was georeferenced. The Asian reference panel was composed of 1929 Asian rice landraces. They were extracted from the list of over 3000 accessions that were re-sequenced in the framework of the 3000 rice genomes project (3 K RGP, 2014). The extraction was based on two selection criteria applied successively. The first criterion was the geographic origin. Only accession of Asian origin was retained. The second criterion was the status, landrace versus improved, of the accessions. Only accessions with a high likelihood of being a landrace (based on the structure of the name) were retained. According to the Wang et al. (2018) classification of the 3 K RGP accessions, among the 1929 accessions of our Asian panel, 1139 belong to *indica*, 495 to *japonica*, 169 to cAus, and 62 to cBas subpopulations. The remaining 64 were considered as Admix (Supplementary Table 2). Hereafter we will refer to the Malagasy panel as M-panel, to the Asian panel as A-panel, to Asian ensembles as subpopulations and to Malagasy ensembles as groups.

Genotypic Data

The M-panel was genotyped using the genotyping by sequencing (GBS) methodology. DNA libraries were prepared at the Regional Genotyping Technology Platform (http://www.gptr-lr-genotypage.com) hosted by Cirad, Montpellier, France. For each accession, genomic DNA was extracted from the leaf tissues of a single plant, using the MATAB method. Each DNA sample was diluted to 100 ng/µl and digested separately with the restriction enzyme ApekI. DNA libraries were then singleend sequenced in a single-flow cell channel (i.e. 96-plex sequencing) using an Illumina HiSeq[™]2000 (Illumina, Inc.) at the Regional Genotyping Platform (http://get. genotoul.fr/) hosted by INRA, Toulouse, France. The fastq sequences were aligned to the rice reference genome, Os-Nipponbare-Reference-IRGSP-1.0 with Bowtie2 (default parameters). Non-aligning sequences and sequences with multiple positions were discarded. Single nucleotide polymorphism (SNP) calling was performed using the Tassel GBS pipeline v5.2.29. (Bradbury et al., 2007). The initial filters applied were the quality score (> 20), the count of minor alleles (> 1), and the bi-allelic status of SNPs. In the second step, loci with minor allele frequency (MAF) below 2.5%, with heterozygosity rate > 5% and with more than 20% missing data were discarded. The missing data were imputed using Beagle v4.0. The process yielded 34,614 SNP, which represents an average marker density of one SNP every 13.2 kb.

This working dataset can be downloaded in HapMap format from http://tropgenedb.cirad.fr/tropgene/JSP/interface.jsp?module=RICE study Genotypes, study type M-panel_GBS_data.

Genotypic data for the A-panel was extracted, during December 2018, from the SNP-Seek database (http://snp-seek.irri.org/; Mansueto et al. 2017), which contained over 29 million bi-allelic markers from 3 K-RG project. The selection criteria were: (i) matching with one of the 34,614 SNP obtained through GBS for the M-panel, (ii) rate of missing data < 20%, (iii) heterozygoty < 5% and (iv) MAF > 1%. The selection process yielded 23, 981 SNPs, which represents an average marker density of one SNP every 18.8 kb. The missing data were

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imputed using Beagle v4.0. The heat map of distribution of the two genotypic datasets along the chromosomes is presented in Supplementary figure 1.

Analysis of the Genotypic Data

Analyses of genotypic data aiming at characterisation of the genetic diversity of the M-panel, per se, were performed with the 34,614 SNPs data set. Analyses of genotypic data aiming at characterisation of the genetic diversity of the M-panel, relative to the genetic diversity of the A-panel were performed with the 23,981 common SNPs, hereafter referred to as 24 k SNPs.

Whole-Genome Level Analyses

The structure of the two panels were first analysed separately, then jointly. In each case, estimates of ancestry coefficients were obtained, using the inference algorithm "sparse nonnegative matrix factorization" (sNMF) (Frichot et al., 2014). The sNMF algorithm produces estimates of ancestry proportions similar to STRUCTURE (Pritchard et al., 2000) or ADMIXTURE (Alexander and Lange, 2011) with much shorter runtimes. The algorithm was implemented under the R package LEA (Frichot and Francois, 2015).

The distances between individual accessions were investigated using a simple-matching dissimilarity index, with DARwin software (Perrier and Jacquemoud-Collet, 2006). An unweighted neighbour-joining tree was constructed based on this dissimilarity matrix.

Differentiation between populations was estimated using pairwise F_{ST} which estimates genetic differentiation based on allele frequency (Wright, 1931). The F_{ST} statistics were calculated over the 23,981 SNPs common to M-panel and A-panel, using Arlequin 3.5.2.2 software (Schneider et al., 2000). The same software was used to estimate gene diversity within each population. Gene diversity was defined as the probability that two randomly chosen haplotypes are different in the sample and was estimated as $\hat{H} = \frac{n}{n-1} \left(1 - \sum_{i=1}^k P_i^2\right)$, where n is the number of gene copies in the sample, k is the number of haplotypes, and P_i is the sample frequency of the i-th haplotype (Nei, 1987).

The speed of decay of linkage disequilibrium (LD) in the two panels was estimated by computing $\rm r^2$ between pairs of markers on a chromosome basis, using the "full matrix" option of Tassel 5.2.63 software (Bradbury et al., 2007), and then by averaging the results by classes of distance using XLSTAT.

Characterisation of Local Haplotypes

This characterisation aimed at identifying local, haplotype level, genotypic variations among the accessions, relative to the global, whole genome level, classification resulting from the above-described whole genome analyses. We implemented the characterisation and assignment method developed by Santos et al. (2019). It relies on kernel density estimation (KDE) in a principal component analysis (PCA) feature space. KDE is a non-parametric method that estimates the density of probability of random variables. It produces an approximation of the distribution of data in the form of combination of kernels (McKinney, 2012). Briefly, first a PCA is performed using the genotypic data of all accessions under study (in our case Mpanel and A-panel) at a given number of adjacent SNP loci of a given chromosome. Then, for each accession, the first five coordinates of the PCA are submitted to the KDE algorithm to compute the likelihood of membership of the accession to each of the reference populations predefined at the whole genome level. Finally, the confrontation of the membership likelihoods computed by the KDE algorithm to a predefined threshold leads to assignment of the local haplotype to one of the reference populations or to one of the haplotypes intermediate between the reference populations, or to an unknown, outlier, haplotype. The process is repeated systematically along each chromosome. In our case, the number of reference populations retained was three, i.e., the accessions belonged to either indica, or japonica or cAus populations, or were classified as admixed. In other words, accessions identified as cBas in Wang et al. (2018) classification were considered as admix. This choice was in line with the three-pillar view of rice domestication (Civan et al., 2015) and offered the advantage of limiting the number of intermediary classes of haplotype to four. Thus, the final number of classes of haplotypes was equal to eight: ind, jap, cAus, ind-jap, indcAus, *jap-*cAus, indi*-jap-*cAus, and outlier. After several tests, the size of the beans retained was of 30 consecutive SNP loci. The overlap between two adjacent beans was of 15 SNPs loci. The local classification was implemented with the function KernelDensity of the python package sklearn.neighbors (Pedregosa et al. 2011) that was used with a Gaussian Kernel. The p-values ratio threshold for classification of the local haplotypes into the three reference classes and the four intermediate classes was set to 3. All local haplotypes, with p-values of assignment to each of the three reference classes below 0.0005 were classified as outliers.

Results of the local assignments were summarised in ideograms. For each accession and each chromosome, in order of increasing first SNP, windows of the same classification were merged into a single block. A new block was created with each change in class.

Results

Population Structure

The structure analysis performed on the 1929 accessions of the A-panel indicated K = 3 as the most likely number

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of populations, i.e., the lowest rate (10%) of accessions with ancestry coefficients below the threshold of 0.80. The three populations corresponded to the indica, japonica and cAus. Assignment of individual accessions to one of the three populations was almost completely in line with the Wang et al. (2018) classification of the 3 K RGP accessions, based on ~4.8 million SNPs. The rate of identical classification was of 96% for the indica population (1090 out of 1139), 100% for the japonica population (495 out of 495), and 92% for cAus (156 out of 169). The 187 admix accessions, with estimated ancestry coefficients below 0.80, included 90 accessions classified as Admix, XI-adm or GJ-adm by Wang et al. (2018), and 16, 62 and 19 accessions assigned by the same authors to cAus, cBas and indica populations, respectively. The 62 cBas accessions derived a large share (0.68%) of their ancestry coefficient from the japonica population (Supplementary figure 2). The hypotheses of K = 4 and K = 5 subdivided the *indica* population into its components XI-1, XI-3 and XI-2, and at K = 6, the cBas accessions formed a separate population. However, under these hypotheses, the rate of accessions with ancestry coefficients below 0.80 increased drastically, reaching 39%, 46% and 43%, respectively. Thus, one can consider that our 24 K SNP genotypic dataset provided a reasonably accurate description of the global structure of the A-panel.

When a similar clustering analysis was performed on the A-panel plus M-panel of 620 accessions, the hypothesis of K = 2 separated a population including the *indica* accessions from another including the japonica accession (Fig. 1a). At K = 3, the population including the *indica* accessions split into 2, and at K = 4 it split into 3 subpopulations. At K = 5, the population including the japonica accessions split into 2 subpopulations. K values above five did not improve population discrimination and did not reduce the number of Admix. Among the populations identified at K = 5, four corresponded to indica, japonica, cAus and cBas from the corresponding A-panel, plus 21%, 7%, 0.2% and 0.3% of accessions from the M-panel, respectively. The fifth population, exclusively composed of Malagasy accessions (38% of the accessions of the M-panel), corresponded to the group specific to Madagascar first described by Ahmadi et al. (1988). Thirteen percent of accessions of A-panel and 33% of accessions of M-panel with less than 80% of their estimated ancestry deriving from one of the identified populations were considered as admixed.

Lastly, when structure analysis was performed on the 620 accessions of the M-panel alone, the most likely number of populations was K = 3, among which two corresponded to *indica* and *japonica* and the third to the group specific to Madagascar (Fig. 2a). The M-panel representatives of *indica*, hereafter called *G1* group,

accounted for 25% of the accessions, the japonica accessions, hereafter called G6 group, for 7%, and the group specific to Madagascar hereafter called *Gm* for 36%. The remaining 32% accessions were mainly admixtures of the G1 and Gm groups, except two Malagasy aromatic varieties with more than 70% of their estimated ancestry deriving from G6, and one cAus accessions. The G1 group consisted of lowland rice varieties cultivated in low elevation areas (altitudes < 800 m). The G6 was composed of upland rice varieties cultivated in the low elevation forest areas of the east coast (altitude < 750 m) on the one hand, and a few lowland rice varieties cultivated in high elevation areas (altitude > 1250 m) on the other hand. Lastly, the Gm assembled lowland rice varieties mainly cultivated at altitudes between 800 m and 1600 m (Supplementary figure 3).

The distance-based neighbour-joining tree supported the results of the structure analyses for the joint analysis of the A-panel and M-panel (Fig. 1b) as well as for the analysis of the M-panel alone (Fig. 2b). It also allowed to separate rather clearly the component subpopulations of the indica and japonica, as subdivided by Wang et al. (2018). The indica XI-1A subpopulation mainly originated from China, the XI-1B subpopulation of diverse origins, the XI-2 subpopulation mainly originated from the Indian subcontinent, and the XI-3 subpopulation mainly originated from South-East Asia. The GJ-temp component mainly originated from Japan, China and Korea, the GJ-trop component, mainly originated from Indonesia, Philippines, Malaysia and Sri Lanka, and the GJ-sbtrop component mainly originated from Bhutan, India, Laos, Myanmar and Thailand (Fig. 1b). The Malagasy G1 accessions split into two ensembles, one clustering with the Asian XI-1B subpopulation of very diverse origins, and the other, together with the Malagasy admix accessions, with the Asian XI-2 subpopulation of South-Asian origin. The Malagasy G6 accessions formed a distinct cluster among the Asian GJ-trop subpopulation, mainly of Indonesian origin. The Malagasy Gm group formed a distinct cluster at the vicinity of the Asian XI-2 subpopulation.

Genetic Characteristics of the Asian Subpopulations and Malagasy Groups

In the A-panel, average gene diversity over the 24 k common SNPs ranged from 0.030 ± 0.0142 for GJ-temp subpopulation to 0.177 ± 0.084 for XI-2 (Table 1). Average gene diversity in Malagasy G1 (0.138 ± 0.089) and G6 (0.100 ± 0.048) was smaller compared to their Asian counterpart *indica* (0.183 ± 0.086) and *japonica* (0.111 ± 0.037). The Gm group had a medium level gene diversity, 0.094 ± 0.044 .

Important differences were observed between subpopulations for r^2 estimate of the initial LD, i.e. pairwise

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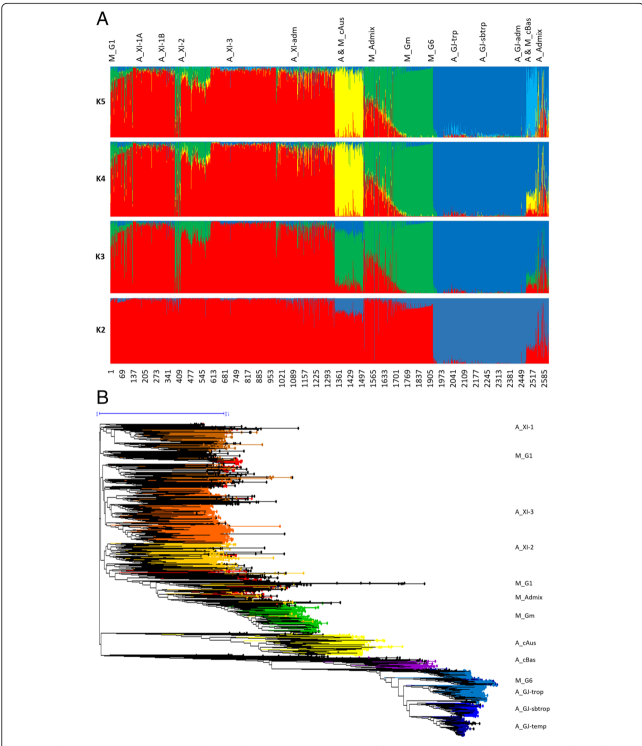


Fig. 1 Population structure in 1929 Asian and 620 Malagasy rice accessions estimated from 23,981 genome wide SNPs. **a**: output of sNMF for K = 2 to K = 5. From top to bottom, the Asian accessions (prefix A) are organised first by their classification in Wang et al. 2018 groups and subgroups, then by alphabetic order of country (not shown). Malagasy accessions (prefix M) are organised first by groups identified by sNMF with K = 3, then by increasing values of ancestry coefficient. **b** Unweighted neighbour-joining tree of simple matching distances. Tree branches are coloured according to membership in sNMF groups at K = 3 and ancestry coefficient 0.8, for Malagasy accessions, according to Wang et al. (2018) subpopulations of Asian accessions

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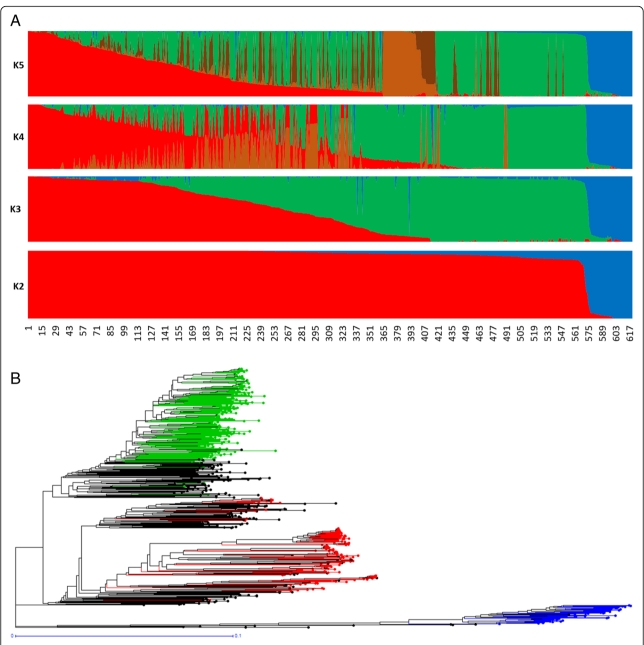


Fig. 2 Population structure in 620 Malagasy rice accessions estimated from 32,614 SNPs. **a:** Output of sNMF for K = 2 to K = 5; **b:** Unweighted neighbour-joining tree of simple matching distances. Tree branches are coloured according to membership of sNMF groups at K = 3 and ancestry coefficient threshold of 0.8

distance between markers below 25 kb (from 0.405 for XI-2 subpopulation to 0.724 for GJ-sbtrop) and for its subsequent decay (Supplementary Table 3). While the distance at which the LD went below 0.2 was between 125 and 150 kb for XI-2, it was above 700 kb for GJ-sbtrop and GJ-temp. Pattern of LD decay in G1 was similar to the one of the *indica* population but the absolute values were slightly higher (0.05 point) whatever the pairwise distance between markers (Fig. 3). The G6

group had a high initial LD ($r^2 = 0.594$) and rather low rate of LD decay ($r^2 < 0.2$ at pairwise distances between markers above 1.5 Mb). The Gm group had the highest initial LD ($r^2 = 0.648$ for distances below 25 kb) but the LD decay was rather fast ($r^2 < 0.2$ at distances above 600 kb) and similar to the one of G1 for distance above 2 Mb. Compared to G1 group and Asian subpopulations, G6 and Gm showed larger inter-chromosome variations of LD pattern. For instance, in Gm, the average r^2 values

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Table 1 Gene diversity, linkage disequilibrium and genetic differentiation between populations estimated by F_{ST} statistic

	N	Gene diversity	$\begin{array}{l} \text{Distance}^{a} \\ \text{(kb) for } r^{2} \leq 0.2 \end{array}$		Asian	panel									Mala	agasy el	
Population				Genetic differentiation between subpopulations estimated by F_{ST} statistic										stic	_		
				indica	XI-1A	XI-1B	XI-2	XI-3	cAus	japonica	GJ- trop	GJ- sbtrop	GJ- temp	cBas	G1	Gm	G6
Indica (XI)	1139	0.19 ± 0.09	75–100														
- XI-1A	178	0.13 ± 0.06	250–300		0.00												
- XI-1B	49	0.15 ± 0.07	250–300		0.25	0.00											
- XI-2	179	0.18 ± 0.08	125–150		0.23	0.17	0.00										
- XI-3	392	0.16 ± 0.06	125–150		0.25	0.18	0.15	0.00									
cAus	169	0.16 ± 0.08	200–250	0.42	0.53	0.49	0.42	0.49	0.00	0.73							
japonica (GJ)	495	0.11 ± 0.04	350–400	0.66													
- GJ-trop	210	0.09 ± 0.04	500-600		0.74	0.74	0.68	0.68	0.70		0.00						
- GJ-sbtrop	105	0.05 ± 0.02	700–800		0.76	0.78	0.69	0.69	0.71		0.25	0.00					
- GJ-temp	150	0.03 ± 0.01	700–800		0.79	0.83	0.72	0.71	0.75		0.30	0.43	0.00				
cBas	62	0.12 ± 0.06	500-600	0.54	0.65	0.62	0.56	0.59	0.56	0.51	0.48	0.56	0.64	0.00			
G1	156	0.14 ± 0.09	125–150	0.07	0.14	0.14	0.11	0.16	0.44	0.73	0.69	0.70	0.74	0.56	0.00		
Gm	220	0.09 ± 0.05	600–700	0.42	0.57	0.58	0.41	0.52	0.55	0.78	0.77	0.80	0.82	0.70	0.46	0.00	
G6	45	0.10 ± 0.05	1250–1500	0.66	0.70	0.69	0.63	0.65	0.64	0.17	0.13	0.34	0.43	0.46	0.62	0.75	0.00

N number of accessions; in the cases of *indica* (XI) and *japonica* (GJ), N includes 342 XI-adm and 30 GJ-adm accessions, respectively. r^2 : Linkage disequilibrium (LD); r^2 : threshold of pairwise distance between markers above which the mean r^2 reaches values below 0.2. The 64 Admix accessions of the A-panel, the 196 Admix, 1 cAus and 2 cBas accessions of the M-Panel were excluded from LD, gene diversity and genetic differentiation analyses

(over 28 levels of pairwise distance between markers) of chromosomes 6 and 11 were +42% and +33% higher than in other chromosomes, respectively.

Genetic differentiation between the three Malagasy groups was significant (p < 0.05 for 1023 permutations), ranging from 0.46, for G1-Gm, to 0.75, for G6-Gm (Table 1). However, "among population" variance, calculated by AMOVA, represented only 40% of the total molecular variance (p < 0.0001), indicating large within-group differentiation. Differentiations between the Malagasy G1 and G6 groups and their Asian counterparts were rather low but highly significant suggesting a specific history of the Malagasy rice varieties. The G1 - cAus F_{ST} and the G1 - cBas F_{ST} were of the same magnitude as the indica - cAus and the indica - cBas F_{ST} , respectively. Similar pattern was observed for G6 - cAus and G6 - cBas G- cBas

the Asian subpopulations were also high, ranging from 0.41 to 0.82 (Table 1), confirming the fact that Gm could not be directly affiliated to one of those subpopulations. The F_{ST} between Gm and the Asian subpopulations was lowest with XI-2.

Haplotype Level Genetic Variations

Results of the KDE-based local classification are represented for each chromosome in Fig. 4 and are summarised over the whole genome in Table 2. As expected, a large percentage of haplotypes of accessions of the Apanel that were considered as members of the *indica*, *japonica* or cAus subpopulations at the whole genome level (Wang et al., 2018), was attributed by the KDE-based classification to their homologous classes of haplotypes, *ind*, *jap* and cAus. On average, the percentage was of 68.9, 58.4 and 53.1 for *indica*, *japonica* and cAus accessions, respectively. However, variations of up to 35%

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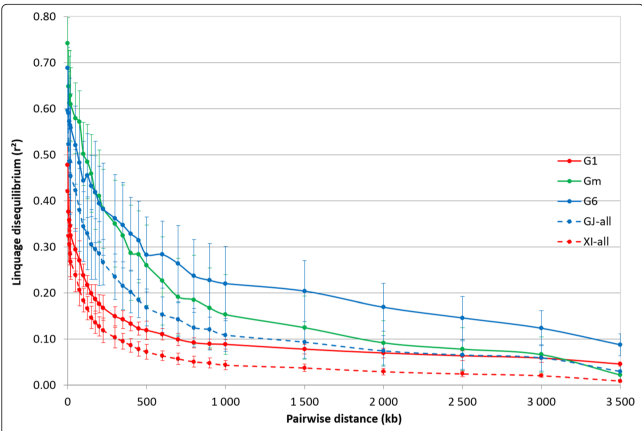


Fig. 3 Patterns of decay in linkage disequilibrium in the *indica* and *japonica* population of the Asian panel and the three groups of the Malagasy rice panel, genotyped with 23,981 SNP. The curve represents the average r² according to pairwise distance between markers among the 12 chromosomes and the bars represent the associated standard deviation

of these shares existed between chromosomes. The percentages of cross-classification of haplotypes of accessions among the three subpopulations (i.e. indica to jap and vice-versa, indica to cAus and vice-versa, and japonica to cAus and vice-versa) were low, rarely exceeding 2%. The percentage of haplotypes attributed to each of the four intermediate classes (ind-jap, ind-cAus, japcAus, and ind-jap-cAus) depended on the classification of accessions at the global level. In the case of indica accessions, on average 12.3% of the haplotypes were classified as ind-cAus and 9% as ind-jap-cAus, while the average share of the ind-jap and jap-cAus haplotypes were of 4.8% and 2.9%, respectively. In the case of the *ja*ponica accessions, on average 13.8% of the haplotypes were classified as ind-jap and 17.7% as ind-jap-cAus, while the average share of ind-cAus and jap-cAus haplotypes were of 3.8% and 4.0%, respectively. Accessions belonging to the cAus subpopulation were characterised by a large share of haplotypes (on average 26.4%) classified as ind-cAus. The proportions of haplotypes of indica, japonica and cAus accessions classified as outliers were very low, less than 1%, on average. Accessions of the Apanel belonging to the cBas subpopulation of Wang et al. (2018) were characterised by large shares of *jap* haplotypes, 31.2% on average, and outlier haplotypes, 11.0% on average.

Among accessions of the M-Panel, the shares of the eight classes of haplotypes in the *G1* group were similar to the ones observed for its Asian counterpart, *indica*. In the case of *G6* the shares differed slightly from its Asian counterpart, *japonica*, with higher percentages of haplotypes classified as *ind* and cAus. Patterns of distribution of the eight classes of haplotypes among the accessions of *Gm* group, significantly diverged from the ones observed for all other groups. It displayed a significantly lower share of *ind* haplotypes (50.6% on average) and significantly higher shares of cAus, *jap* and outlier haplotypes than the Asian *indica* and the Malagasy *G1* group, 8.2%, 5.4% and 2.6% on average, respectively. The Malagasy Admix had a haplotype pattern similar to *Gm*, with slightly lower shares of cAus and *jap* haplotypes.

To further investigate the relationship between the *Gm* and the Asian *indica*, we compared the patterns of distribution of haplotypes of *Gm* accessions with the ones of each of the four *indica* subpopulations, XI-1, XI-2, XI-3 and XI-adm (Wang et al., 2018). Patterns of

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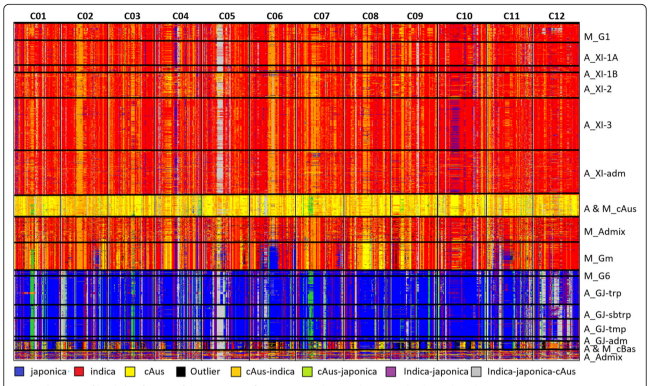


Fig. 4 Ideogram of local classification of the genomes of 1929 Asian and 620 Malagasy rice landraces, based on 23,982 SNP. Patterns are organised per chromosome from left to right and by accessions from top to bottom. The Asian accessions (prefix A) are organised first by their classification in Wang et al. (2018) groups and subgroups, then by alphabetic order of country (not shown). Malagasy accessions (prefix M) are organised first by groups identified by structure analysis with K = 3, then by increasing values of ancestry coefficient

distribution of eight classes of haplotypes in the *Gm* group diverged from the ones of the four *indica* subpopulations by lower share of *ind* haplotypes and higher shares of cAus and *jap* haplotypes. The divergence was the lowest with the XI-2 subpopulation.

The *Gm* accessions also displayed larger variations of the shares of each class of haplotype across the 12 chromosomes. For instance, chromosome 6 displayed an exceptionally high share of *jap* haplotypes (24% on average), chromosome 8 a large share of cAus haplotypes (19.8% on average) and chromosome 11 a low share of *ind* and high shares of *ind-jap* and *ind-jap-c*Aus haplotypes, 34.9%, 13.2% and 19.2%, respectively. In some cases, these haplotypes were contiguous and covered a large segment of the chromosome. For instance, on chromosome 6, one of the *jap* haplotype segments spread over 4.7 Mb (13.0 to 17.7 Mb) and on chromosome 8 one of the cAus haplotype segments spread over 5.4 Mb (10.9–16.3 Mb).

Dissymmetry based neighbour-joining tree constructed with the genotypic data of individual chromosomes (Supplementary figure 4) confirmed the heterogeneity of shares of different classes of haplotype among the 12 chromosomes. It also showed the discriminatory power of genetic diversity at the individual chromosome level.

Accessions belonging to Gm were always clustering in a well-individualised branch of the tree, often near the cAus branch. M-Admix accessions almost systematically clustered with the *Gm* accessions. The chromosome level neighbour-joining trees also allowed, in some cases, to identify accessions of the M-panel and A-panel that displayed similar patterns of haplotypes shares. In the case of the M-panel, the three accessions most often clustering with Gm belonged to G1 originated from the Mahajanga region on the west coast of Madagascar. In the case of the A-panel, more than 95% of accessions clustering near Gm were of Indian subcontinental origin and belonged to XI-2 subgroup (66%), XI-adm (21%) and cAus (13%). Three accessions of the A-panel most frequently present in the vicinity of the *Gm* cluster were of Indian origin: Larha Mugad (IRGC 52339-1) from Karnataka, Adukkan (IRGC 81783-1) from Kerala and Cuttack 29 (IRGC 49573-1) from Odisha. Larha Mugad and Adukkan belonged to XI-2, Cuttack 29 was an XIadm.

Characteristics of the Large cAus and *Jap* Chromosomic Segments in the *Gm* Group

The two largest *jap* and the largest cAus chromosomic segments in the *Gm* accessions were investigated for

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Table 2 Mean shares (%) of the eight categories of haplotypes identified by the KDE-based local classification, in each population and subpopulation of the Asian and Malagasy diversity panels

Diversity panel Asian Panel	Global	KDE-based local classification										
	classification	ind	cAus	Jap	ind-jap	ind-cAus	jap-cAus	ind-jap-cAus	Outlier			
	indica	68.9	1.2	0.9	4.8	12.3	2.9	8.9	0.1			
		(59.8-84.8)	(0.9-1.9)	(0.4-2.7)	(2.1-7.4)	(5.8–19.2)	(1.6-4.5)	(3.8–15.7)	(0-0.4)			
	- XI-1	67.6	0.5	1.0	5.2	12.4	3.4	9.8	0.1			
		(57.6–85.6)	(0.2-1.6)	(0.3-2.9)	(2-7.9)	(4.3-17.9)	(1.7-9)	(4.1–18.9)	(0-0.2)			
	- XI-2	65.6	2.9	0.5	4.4	14.5	2.8	8.5	0.8			
		(51.4-80)	(1.9-4.8)	(0.2-1.6)	(1.7-6.6)	(8.4-23.2)	(-0.4-4.7)	(1.2–15.1)	(0-8.6)			
	- XI-3	72.7	0.3	0.5	4.6	10.9	2.7	8.3	0.1			
		(60.8–88.5)	(0.1-0.7)	(0.1-2.4)	(2-9.3)	(4.1–19.9)	(1.5-4.5)	(3.3–13.9)	(0-0.3)			
	- X-adm	67.2	1.5	1.4	5.0	12.6	3.0	9.2	0.1			
		(57.5-83.5)	(1.2-2.3)	(0.7-3.3)	(2.2-7.2)	(6.2-18.9)	(1.7-4.6)	(4.1–16.1)	(0-0.5)			
	japonica	1.0	0.4	58.4	13.8	3.8	4.0	17.7	0.8			
		(0.4-1.8)	(0.3-0.7)	(36.4–71.6)	(3-23.2)	(2.7-6.5)	(2.7-6.5)	(9.7–27.3)	(0.1-7.2)			
	- GJ-trop	1.1	0.5	58.2	14.7	3.7	3.9	16.0	2.0			
		(0.3-2.0)	(0.3-1.1)	(36.3-71.9)	(3.1-23.2)	(1.9-6.4)	(1.8-6.3)	(8.4–27.4)	(0.1-9.8)			
	- GJ-sbtrop	0.8	0.4	58.4	14.5	4.1	4.2	16.5	1.3			
		(0.1–1.7)	(0.1-0.9)	(40.9-73.4)	(2.9-23.9)	(2.8-6.9)	(2.8-6.4)	(9.1–25.5)	(0.1-6.6)			
	- GJ-temp	1.0	0.2	55.7	14.6	4.5	4.6	18.2	0.8			
		(0.4-2.4)	(0.0-0.6)	(33.7–71.4)	(3.2-23.0)	(2.8-6.9)	(2.9-7.1)	(11.1–28.1)	(0.0-3.9)			
	- GJ-adm	2.7	0.7	54.8	14.8	4.4	4.4	17.1	1.5			
		(1.4-5.0)	(0.3-1.1)	(35.3-70.3)	(5.1-22.7)	(3.0-6.5)	(2.8-6.6)	(9.8–27.9)	(0.0-6.6)			
	cAus	1.8	53.1	0.8	3.3	26.4	3.5	10.8	0.2			
		(1.3-2.3)	(37.6–63.8)	(0.5-2)	(1.7-6.7)	(14.4-42.2)	(1.8-6.7)	(4.8-18.4)	(0-0.4)			
	cBas	16.9	13.2	31.2	6.9	7.1	2.6	11.1	11.0			
		(10.6-27.4)	(4.9-22.4)	(20.7-38.7)	(2.2-12.9)	(2.9-10.9)	(-0.9-5.4)	(4.4–21.2)	(6.1–23.7)			
	Admix	24.4	12.4	24.3	7.4	10.6	4.3	10.3	6.2			
		(20.1-27.2)	(8.7-16.4)	(16.4–30.9)	(2.3-11.3)	(7.5–14.5)	(2-13.3)	(1.8–17.2)	(4.9–9.7)			
Malagasy panel	G1	69.7	1.3	0.6	5.0	11.9	2.8	8.4	0.4			
		(59.9-84.2)	(0.6-2.3)	(0.1-2.1)	(2.0-7.6)	(4.5-17)	(1.6-4.2)	(3–16.2)	(0.1-1.2)			
	G6	2.7	2.0	57.4	12.5	3.9	3.8	15.0	3.1			
		(1.5-4.8)	(0.8-4.1)	(38.3-69.0)	(3.0-18.9)	(2-7)	(1.6-6.8)	(7.1–23.1)	(1.2–10.8			
	Gm	50.6	8.2	5.4	5.7	14.3	3.3	9.8	2.6			
		(34.9–79.6)	(2.4–19.8)	(1.2-24)	(2.1-13.2)	(6-21.5)	(1.8-6.3)	(3.7–19.2)	(1.3-3.8)			
	Admix	58.9	6.2	3.4	4.6	12.5	2.8	7.4	4.1			
		(49.2-75.0)	(2.4-8.3)	(2.6-5.1)	(1.9-8.8)	(5.0-20.5)	(1.0-5.2)	(1.6-12.9)	(2.3-6.2)			

kinship with their homologous segments in other Malagasy groups and in Asian subpopulations, and for their functional proprieties. Dissimilarity based phylogenetic neighbour-joining trees were constructed using the genotypic data from each of the three segments (240, 221 and 225 SNP loci for chromosome 6, 8 and 11, respectively) for kinship analysis. The functional properties of the three large introgression segments was investigated through gene ontology analysis (Mi et al. 2013)

and compilation of results of ten recent genome-wide association studies (GWAS) for response to low temperature stress, between 2015 and 2020.

In the neighbour-joining trees constructed with the genotypic data from the 13.0 to 17.7 Mb segment of chromosome 6, the *Gm* accessions and a significant share of the Malagasy Admix accessions formed a rather compact branch in the vicinity of the *japonica* cluster (Supplementary figure 5). *Japonica* accessions most

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closely clustering with Gm accessions were members of the Malagasy G6 group, (Fig. 5). The non-Malagasy accessions clustering near Gm were GJ-trop from Indonesia, Malaysia and the Philippines, plus one Admix accession from India.

In the neighbour-joining trees constructed with the genotypic data from the 10.6 to 15.7 Mb segment of chromosome 11, *Gm* accessions formed again a rather compact branch near the *japonica* cluster. *Japonica* accessions most closely clustering with *Gm* belonged to *G6*, to GJ-trop from Indonesia and, to a lesser extent, to GJ-sbtrop from India and GJ-tem from China (Supplementary figure 5).

In the neighbour-joining trees constructed with the genotypic data from the 10.9–16.3 Mb segment of chromosome 8, *Gm* accessions formed a compact cluster near the cAus subpopulation. Two cAus accessions from Bangladesh and India were embedded in the *Gm* cluster. The closest neighbours to the *Gm* cluster were 30 cAus, 4 XI-2 and one XI-adm accessions of Bangladeshi and Indian origin (Supplementary figure 6).

Gene ontology analysis (Supplementary Table 4) detected highly significant (78.5 fold) enrichment in "cold

acclimation" genes in the japonica introgression of chromosome 11. It also detected small but significant enrichment among the unclassified genes of the three introgression segments. Compilation of results GWAS for response to low temperatures led to 610 quantitative trait loci (QTL) of which 22, 11 and 8 were located in the introgression segment of chromosomes 6, 8 and 11 respectively (Supplementary Table 5). These numbers were on average of 1.35 QTL per Mb of genome length outside the introgression segments and on average 4.63, 2.02 and 1.56 QTL per Mb, in the introgression segment of chromosomes 6, 8 and 11 respectively. Lastly, a literature review for gene families involved in cold tolerance and response to abiotic stresses, identified 29, 25 and 14 genes with possible involvement in response to low temperatures, in the introgression segment of chromosomes 6, 8 and 11 respectively (Supplementary Table 6).

Discussion

The objective of this work was to use the power of highresolution genotypic data to gain new insights into the

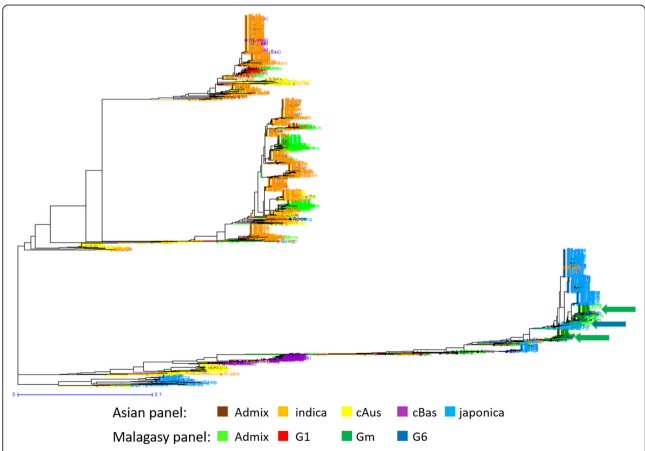


Fig. 5 Unweighted neighbour-joining tree of simple matching distances, constructed with the genotypic data at 13.0–17.7 Mb segment of chromosome 6 (240 SNP). Accessions of the Asian panel are coloured according to their membership to major populations defined by Wang et al. (2018). Accessions of the Malagasy panel are coloured according to their membership of sNMF groups at K = 3 and ancestry coefficient threshold of 0.8. Positions of G6 and Gm accessions on the trees are highlighted with coloured arrows corresponding to their group membership

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organisation of rice genetic diversity in Madagascar and, beyond, into the evolutionary processes accompanying *O. sativa*'s geographical expansion.

The population structure of rice in Madagascar, with its three major groups G1, G6 and Gm, revealed in this study corresponded to the large scale Asian differentiation of O. sativa, but with two significant differences: (i) only the two major Asian subpopulations, *indica* and japonica were significantly represented; the cBas and cAus subpopulations were each represented by two accessions only. (ii) Madagascar hosted an additional group (Gm) that could not be assigned to any of the four major Asian subpopulations. The existence of an atypical rice group was first reported by Ahmadi et al. (1988), using morpho-physiological data, and then confirmed using isozyme (Ahmadi et al., 1991) and SSR markers (Radanielina et al. 2012). However, so far, the Gm group has not been identified as such in studies directly comparing Malagasy varieties with a diversity panel originating from Asia. Indeed, Mather et al. (2010), who compared 45 Malagasy landraces with 39 Asian accessions (21 indica and 18 GJ-trop), identified only one group among the Malagasy lowland accessions and called it Malagasy-indica. Zhao et al. (2011), who genotyped a rice diversity panel of 415 accessions with 44 K SNP, classified the 6 Malagasy accessions present as admixed and not as belonging to one of the four major Asian subpopulations. Thus, our study is the first one comparing a Malagasy and an Asian diversity panel on a large enough scale to distinguish the *Gm* group from both Malagasy and Asian originating indica subpopulations. The Malagasy-indica of Mather et al. (2010) most likely corresponded to our *Gm* group.

Gene diversity in *G1* and *G6* represented only 75% and 66% of the diversity in the Asian *indica* and *japonica*, respectively. This reduced diversity represents the bottleneck associated with the introduction of rice to Madagascar from different Asian sources. Consistent with the relative diversity of Asian and Malagasy-*indica* reported by Mather et al. (2010), the gene diversity of *Gm* represents 51% of that of *indica*.

The three Malagasy groups are not evenly distributed across the country. One major factor shaping this uneven geographic distribution is the climatic conditions associated with altitude, leading to a preferential habitat for each group sketched in Fig. 6. Similar to the area of cultivation of *indica* in tropical Asia and elsewhere in the world, *G1* accessions are mainly cultivated in the low-elevation coastal wetlands. The preferential habitat of *Gm* is the wetlands of 1200-1700 m elevation, while in tropical Asia such areas are mainly occupied by GJ-temp. The eco-geographic distribution of *G6* is more complex, some accessions being cultivated in the upland ecosystem of the Eastern-forests areas and some others in the lowland ecosystem of the central high plateaux.

The structuring of rice genetic diversity in three major groups and their uneven eco-geographical distribution in Madagascar is also related to the history of the introduction of rice in the island summarised in Fig. 6. The tight clustering of G6 accessions with the GJ-trop accessions originating from Indonesia on the neighbour-joining tree corroborates historical and linguistic data of the Austronesian (present Malaysia and Indonesia) origin of the human population that first settled the Madagascar island 1400-1200 years ago (Dahl, 1951; Tofanelli et al., 2009; Cox et al., 2013). The swidden cultivation of upland rice along the east coast of Madagascar started with these first colonisations of the island (Beaujard, 2011). The Malagasy lowland japonica varieties ("Vary Lava" and "Latsika" vernacular families, well known for very long and thick grains, and for cold tolerance, respectively) also belong to the GJ-trop subpopulation. Varieties belonging to the Bulu ecotype of Indonesia are the archetypes of lowland-grown GJ-trop (Glaszmann, 1987; Thomson et al. 2007). Some examples have also been reported in the Philippines, with varieties such as Azucena. The presence of lowland grown GJ-trop in Madagascar is thus another clue backtracking the origin of the Malagasy japonica varieties to Indonesia. Interestingly almost all Indonesian lowland-grown landraces of GJ-trop genotype originated from Central Java, Kalimantan and Sumatra islands (Thomson et al., 2007) where more than 500,000 ha of rice are grown in terraced fields at an altitude above 700 m (Harahap, 1979). In these areas, typical indica varieties such as IR8 yield very poorly and coldtolerant varieties are needed (Harahap, 1979). As the cold tolerance of the GJ-temp is generally associated with round grain type, the Malagasy long grain "Vary Lava" varieties constitute a useful source for the diversification of grain quality in breeding for temperate regions and cold-prone tropical areas.

The characteristics of the Malagasy G6 group also provide insight into the evolutionary process that accompanied japonica subspecies' expansion. First domesticated in the Yangzi region of China (Liu et al., 2007; Molina et al., 2011), the japonica subspecies has accompanied the southward migrations of human populations reaching Taiwan by 3000 BC and Central Indonesia by 2000 BC (Bellwood, 2011). When the japonica subspecies began its migration south-westward to Madagascar more than 2500 years later (in the middle of the first millennium AD), it had already reached its current balance of recent and ancestral alleles, as shown by the almost equal mean rate of jap haplotypes in G6 compared to the japonica subpopulation. On the other hand, compared to japonica, G6 showed a slightly higher mean rate of ind and cAus haplotypes, suggesting some genetic expansion during the south-westward waves of migration or after Ahmadi et al. Rice (2021) 14:44 Page 14 of 18

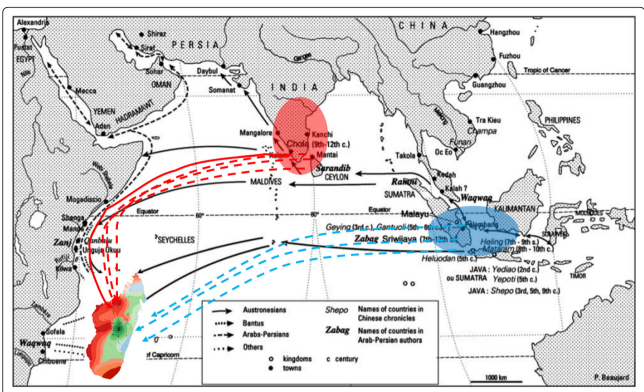


Fig. 6 History of human migrations to Madagascar, the associated introductions of different rice genetic groups, and the current ecogeographical distribution of rice groups in the country. The map of human migrations (black arrows) was borrowed from Beaujard (2011). The red and blue circles indicate the most probable area of origin of the Malagasy *indica* and *japonica* rice groups, respectively. The blue dotted lines indicate the arrival in Madagascar of the *japonica* rice varieties, associated with the South-westward migration of population of Malay origin (1200–1400 AD). The red dotted lines indicate the arrival of the *indica* rice varieties associated with the waves of Austronesians reaching Madagascar up to the fifteenth century AD. The red bold line represents the latest wave of Austronesians migrants who established the Merina kingdom in the central highlands of the country and developed rice cultivation in those areas. The red, blue and green colours covering the Malagasy island indicate du current eco-geographical distribution of rice groups in the country

insulation in Madagascar. Given the migration route of the early Austronesian settlers, the latter option seems the most probable. Whatever the origin of the genetic expansion, it has endowed the *G6* group with enough diversity to allow adaptation to both the warm upland ecosystem of East coast-forest areas and the cold lowland ecosystem of the high plateaux.

According to historical and linguistic data (Boiteau, 1977; Domenichini-Ramiaramanana 1988; Allibert, 2007; Beaujard, 2011), waves of Austronesians reaching Madagascar went on until the fifteenth century AD, using more complex routes and borrowing Dravidian and Arabo-Persian linguistic, cultural and rice cropping practices on their way. While the lowland rice farming vocabulary of Malagasy originates in the Malay Archipelago, yet terms from Indian, Arabic and Swahili origins are also frequent, especially on the west coast and in the central high plateaux (Beaujard, 2003, 2011). These data are consistent with our finding of preferential clustering of a large share of *G1* accessions with accessions of the XI-2 subpopulation of *indica* originating from

Bangladesh, India and Sri-Lanka. A tight relationship between Madagascar and Indian subcontinent *indica* accessions was also reported by Mather et al. (2010), based on the clustering of some of their Indian *indica* accessions with the Malagasy-*indica* group.

The latest arrivals of Austronesians, who disembarked on the west coast and later founded the Merina kingdom in the central high plateaux, are believed to have played an important role in the development of lowland rice cultivation in Madagascar (Boiteau, 1977; Beaujard, 2011). The establishment of the *Gm* group in the high plateaux is probably associated with this last major immigration wave. Indeed, rice cultivation was absent from the high plateaux before the establishment of the Merina ethnic group (Raison, 1972; Abé, 1984) and the contemporary preferential area of distribution of the *Gm* group coincides with the area of establishment of the Merina and their Betsileo cousins.

Accessions of the *Gm* group are characterised by a low mean share of *ind* haplotypes, down to 35% in chromosome 11, and high mean shares of cAus and *jap*

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haplotypes, up to 20% and 24%, in chromosome 8 and 6, respectively. Within the A-panel, apart from cAus per se, such high share of cAus haplotypes is observed only in some cBas accessions. Given the almost absence of representatives of cAus subpopulation in Madagascar, the most parsimonious hypothesis regarding the origin of the cAus haplotypes in Gm group is a recombination between indica and cAus in the Asian area of origin. Analysis of the origin of the large segments of chromosomes 1 and 8 composed of cAus haplotypes in Gm group indicates a high proximity with two cAus, four XI-2 and one XI-adm accessions of Bangladeshi and Indian origin. Among these accessions at least three [Mugad (IRGC 52339-1), Adukkan (IRGC 81783-1), and Cuttack 29 (IRGC 49573-1)] are from the East (Odisha) and South-West (Kerala, Karnataka) coastal states of India, situated along the maritime roads of the latest Austronesians' migrations to Madagascar, described by Beaujard (2011). Interestingly, (i) western Odisha is recognized as part of the center of origin of Aus ecotypes (Sharma et al. 2000). (ii) Glaszmann (1988) reported the presence of indica, Aus, Aromatic and intermediate accessions among the primitive cultivars of the Jeypore tract of Odisha, thought to have been isolated from alien rice and thus representative of local domestication processes. (iii) The Odisha State includes rice growing areas at elevation above 750 m on the one hand and dry season rice cultivation practices with temperature below 18 °C (Das, 2012), on the other hand.

Regarding the origin of the high mean share of jap haplotypes in Gm accessions, its occurrence in all chromosomes (with larger proportions in chromosomes 6, 7 and 11) with complex patterns, sometimes corresponding to very unlikely double or triple crossing-overs, argues for multiple and rather ancient hybridisation events resulting from sympatry between GJ-trop accessions and indica-cAus intermediate forms. The rarity of representatives of the GJ-trop subpopulation in the rims of the Indian subcontinent, and the rarity of representatives of indica-cAus intermediate forms in the Malo-Indonesian archipelago, excludes the possibility of their sympatry in these places. Quite the reverse, the high plateaux of Madagascar have been hosting representatives of the two subpopulations for some five centuries. Interestingly, a large majority of G6 accessions most closely clustering with the Gm group in the neighbour-joining tree constructed with genotypic data from the large jap haplotypes of chromosomes 6 and 11, originates from the high plateaux of Madagascar (collection mean altitude of 1240 m) and is cultivated in the lowland ecosystem. The presence in Madagascar of a large number of Admix accessions, that cluster with Gm when cAus haplotypes are considered and differ from *Gm* for the share of jap haplotypes (mean share of 3.4% per chromosome against 5.4% for Gm), reinforces the hypothesis of a local, Malagasy, origin of Gm group. It emerged from multiple hybridisation and recombination between indica-cAus intermediary forms and japonica (G6) groups. The footprint of the most recent such hybridisations is still visible in chromosomes 6 and 11 of G6, with much slower LD decay. The hypothesis of Gm emergence from local hybridisation is also in agreement with the high rate of fertility (reaching 70%) of F1 hybrids of indica x Bulu crosses (Morinaga and Kuriyama, 1958). However, as the usual fate of the recombinant progenies of such crosses is a replacement by parental forms (Oka, 1983), their survival and expansion is most likely due to some forms of positive selection. The preferential ecogeographic distribution of Gm accessions in the high plateaux of Madagascar (1200-1600 m), argues for the selection for cold tolerance.

Mather et al. (2010) also reported evidence of several separate indica x japonica hybridisations and recombinations in Madagascar. The extent of the *Gm* group's gene diversity revealed in the present study (equivalent to the one of GJtrop subpopulation and much larger than the ones of GJsbtrop and GJ-temp), and the pattern of its LD decay (faster than the ones of the three subpopulations of *japonica*) argue for its recognition as an original group. The recognition as an original group, instead of "Admix", would be helpful for its future utilisation as a valuable genetic resource for adaptation to high elevation areas. Indeed, "Admix" accessions are seldom included in genetic studies or used in breeding programs. We propose the name "Rojo" for this original rice group. This prefix is shared by almost 25% of names of the accessions belonging to *Gm* and refers to their grain shape associating a medium-long length with a large width (Peltier 1970; Ahmadi et al. 1988).

Despite the complex underlying evolutionary and demographic events in the making of the Malagasy rice gene pool, we were able to backtrack different components of the O. sativa diversity present in Madagascar to their Asian geographic area of origin and genetic subpopulations and to confirm the presence of the original Gm group. Thanks to the multiplicity of introduction events borrowing several routes, the bottleneck undergone by G1 and G6 groups has not been too severe. The evolutionary process involved in the emergence and expansion of the Gm ecotype, are most probably (i) multiple hybridisations and recombinations between indica and cAus subpopulations in the context of their sympatry in south Asia. (ii) Several hybridisations and recombinations between representatives of indica-cAus admixes and G6 group in the context of sympatry in the high plateaux of Madagascar. (iii) Human selection for adaptation to the lowland rice cultivation in the high plateaux. The rather large genetic diversity of the Gm group, the existence of intermediaries or admixed forms Ahmadi et al. Rice (2021) 14:44 Page 16 of 18

between G1 and Gm, and the complementary preferential habitats of G1 and Gm in terms of altitude and associated climatic conditions, provide a unique opportunity to investigate rice adaptation to thermal aspects of climate changes, using landscape genomic approach.

Abbreviations

3 K RGP: 3000: rice genomes project; cAus: Genetic group including the Aus ecotype; cBas: Genetic group including the Basmati ecotype; Fst: Genetic differentiation; G1: Genetic group representative of the indica subspecies in Madagascar; G6: Genetic group representative of the japonica subspecies in Madagascar; GJ: Japonica subspecies; GJ-adm: Admixed japonica; GJ-sbtrop: Subtropical japonica; GJ-temp: Temperate japonica; GJ-trop: Tropical japonica; Gm: Genetic group specific to Madagascar; KDE: Kernel density estimation; LD: Linkage disequilibrium; MAF: Minor allele frequency; PCA: Principal component analysis; sNMF: Sparse nonnegative matrix factorization; SNP: Single nucleotide polymorphism; XI: Indica subspecies; XI-1A: Indica subpopulation mainly originated from East Asia; XI-1B: Indica subpopulation of divers origine; XI-2: Indica subpopulation mainly originated from South Asia; XI-3: Indica subpopulation mainly originated from Southeast Asia: XI-adm: Admixed indica

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12284-021-00479-8.

Additional file 1: Supplementary Figure S1: Heat map of distribution of the two set SNP loci (32,614 and 23,981) along the genome.

Additional file 2: Supplementary Figure S2: Population structure in 1929 accessions of the Asian panel estimated from 23,981 genome wide SNPs. The subpopulations are coloured according to their membership to groups defined by Wang et al. (2018).

Additional file 3: Supplementary Figure S3: Altitudinal distribution of the Malagasy panel 620 rice accessions according to their membership to groups defined by the sNMF-based analysis of population structure.

Additional file 4: Supplementary Figure 54: Unweighted neighbour-joining tree of simple matching distances, constructed with the genotypic data of individual chromosomes (C1 to C12). Accessions of the Asian panel are coloured according to their membership to subpopulations defined by Wang et al. (2018). Accessions of the Malagasy panel are coloured according to their membership to sNMF groups at K=3 and ancestry coefficient threshold of 0.8. Positions of the Malagasy accessions on the trees are highlighted with coloured arrows corresponding to their group membership. Accessions of the Asian panel are coloured according to their membership to major populations defined by Wang et al. (2018).

Additional file 5: Supplementary Figure S5: Unweighted neighbour-joining tree of simple matching distances, constructed with the genotypic data at 11.6 – 15.7 Mb segment of chromosome 11 (225 SNP). Accessions of the Asian panel are coloured according to their membership to major populations defined by Wang et al. (2018). Accessions of the Malagasy panel are coloured according to their membership to sNMF groups at K = 3 and ancestry coefficient threshold of 0.8.

Additional file 6: Supplementary Figure S6: Unweighted neighbour-joining tree of simple matching distances, constructed with the genotypic data at 10.9 – 16.3 Mb segment of chromosome 8 (221 SNP). Accessions of the Asian panel are coloured according to their membership to major populations defined by Wang et al. (2018). Accessions of the Malagasy panel are coloured according to their membership to sNMF groups at K = 3 and ancestry coefficient threshold of 0.8.

Additional file 7: Supplementary Table S1: List and main characteristics of the 620 Malagasy rice accessions used in the present study.

Additional file 8: Supplementary Table S2: List of the 3K-RG project accessions used in the present study.

Additional file 9: Supplementary Table S3: Decay of pairwise linkage disequilibrium with distance between markers, among the three groups of the Malagasy panel and in seven subpopulations of the Asian panel.

Additional file 10: Supplementary Table S4: GO enrichment analysis in the three in introgression segments of Gm group, using tool proposed at http://geneontology.org/.

Additional file 11: Supplementary Table S5: Compilation of results of ten genome wide association studies for response to low temperature stress in rice.

Additional file 12: Supplementary Table S6: list of genes in the three in introgression segments of Gm group, with possible connection with response to low temperature.

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Authors' Contributions

NA and AR conceived the study and wrote the manuscript. NA, AR and TR assembled the Malagasy rice accessions. JF produced the genotypic data. JDS: Contributed to haplotype level analysis.. NA, analyzed the data. All authors read and approved the final manuscript.

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Availability of Data and Materials

The genotypic datasets generated and analysed during the current study are available in hapmap format at the following address http://tropgenedb.cirad.fr/tropgene/JSP/interface.jsp?module=RICE, in the tab "STUDIES" then selecting for "Genotypes" in *Study type* and for "GS-Ruse_IRRI-Reference-Population.

Declarations

Ethics Approval and Consent to Participate

Doesn't apply.

Consent for Publication

Doesn't apply.

Competing Interests

The authors declare that they have no conflict of interest.

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