

African rice (*Oryza glaberrima*) genomic introgressions impacting upon panicle architecture in Asian rice (*O. sativa*) lead to the identification of key QTLs

Hélène Adam (✉ helene.adam@ird.fr)

UMR DIADE, University of Montpellier, IRD

Andrés Gutierrez

Alliance Bioversity-CIAT

Marie Couderc

UMR DIADE, University of Montpellier, IRD

François Sabot

UMR DIADE, University of Montpellier, IRD

Fabrice Ntakirutimana

UMR DIADE, University of Montpellier, IRD

Julien Serret

UMR DIADE, University of Montpellier, IRD

Julie Orjuela

UMR DIADE, University of Montpellier, IRD

James Tregear

UMR DIADE, University of Montpellier, IRD

Stefan Jouannic

UMR DIADE, University of Montpellier, IRD

Mathias Lorieux

UMR DIADE, University of Montpellier, IRD

Research Article

Keywords: rice, chromosome segment substitution line, QTL, panicle architecture, *Oryza glaberrima*, *Oryza sativa*.

Posted Date: May 11th, 2023

DOI: <https://doi.org/10.21203/rs.3.rs-2884677/v1>

License: © ⓘ This work is licensed under a Creative Commons Attribution 4.0 International License. [Read Full License](#)

Abstract

Background: Developing high yielding varieties is a major challenge for breeders tackling the challenges of climate change in agriculture. The panicle (inflorescence) architecture of rice is one of the key components of yield potential and displays high inter- and intra-specific variability. The genus *Oryza* features two different crop species: Asian rice (*Oryza sativa* L.) and the African rice (*O. glaberrima* Steud). One of the main morphological differences between the two independently domesticated species is the structure (or complexity) of the panicle, with *O. sativa* displaying a highly branched panicle, which in turn produces a larger number of grains than that of *O. glaberrima*. The genetic interactions that govern the diversity of panicle complexity within and between the two species are still poorly understood.

Results: To identify genetic factors linked to panicle architecture diversity in the two species, we used a set of 60 Chromosome Segment Substitution Lines (CSSLs) issued from third generation backcross (BC₃DH) and carrying genomic segments from *O. glaberrima* cv. MG12 in the genetic background of *O. sativa* Tropical Japonica cv. Caiapó. Phenotypic data were collected for rachis and primary branch length, primary, secondary and tertiary branch number and spikelet number. A total of 15 QTLs were localized on chromosomes 1, 2, 3, 7, 11 and 12 and QTLs associated with enhanced secondary and tertiary branch numbers were detected in two CSSLs. Furthermore, BC₄F_{3,5} lines carrying different combinations of substituted segments were produced to decipher the effects of the identified QTL regions on variations in panicle architecture. A detailed analysis of phenotypes versus genotypes was carried out between the two parental genomes within these regions in order to understand how *O. glaberrima* introgression events may lead to alterations in panicle traits.

Conclusion: Our analysis led to the detection of genomic variations between *O. sativa* cv. Caiapó and *O. glaberrima* cv. MG12 in regions associated with enhanced panicle traits in specific CSSLs. These regions contain a number of key genes that regulate panicle development in *O. sativa* and their interspecific genomic variations may explain the phenotypic effects observed.

Background

Improving or enhancing the sustainability of rice yields continues to be crucial challenge in the breeding of this crop, especially in the context of a growing world population and with regard to climate change. Inflorescence architecture in rice directly affects yield potential through the regulation of grain number and harvesting ability. Grain number per panicle depends on orders and numbers of branches (i.e., primary, secondary and potentially tertiary branches as well as both lateral and terminal spikelets) and on the length of each axis. Rice panicle architecture is based on the production of a series of lateral meristems with distinct identities (Itoh et al., 2005). After the floral transition, the Shoot Apical Meristem (SAM) is converted into an indeterminate Rachis Meristem (RM) which produces several axillary meristems, the primary branch meristems (PBMs) that subsequently give rise to primary branches (PBs). Once the RM becomes degenerate (i.e., it has lost its activity), the newly formed PBM elongates and initiates a variable number of axillary meristems forming a new branch order. These acquire the identity of secondary branch meristems (SBMs) and develop into secondary branches (SBs), which in turn may produce tertiary branch meristems (TBMs), or alternatively differentiate directly into lateral spikelet meristem (SpMs) and then florets. The basic architecture of the panicle is thus determined by patterns of axillary meristem formation and the specification of their identities. Overall, the complexity and diversity of panicle branching can be considered as governed by two key elements: the number of axillary meristems produced during the indeterminate phase; and the rate of meristem fate transition, which determines whether an axillary meristem grows into a higher-order branch or differentiates into a spikelet.

Over recent decades, a number of genes and quantitative trait loci (QTLs) associated with panicle development and affecting its architecture have been identified and functionally described in rice (Xing and Zhang 2010; Chongloi et al., 2019; Yin et al., 2021; Wang et al., 2021). Typically, these genes and QTLs are implicated in axillary meristem formation or in the conversion of indeterminate meristems to spikelets, thus influencing the branching complexity of the panicle (Komatsu et al., 2003a; Komatsu et al., 2003b; Kobayashi et al., 2009; Yoshida et al., 2013; Bai et al., 2016a).

Most studies of rice panicle development have focused on the Asian crop plant *Oryza sativa* L. However, a second species of cultivated rice, *O. glaberrima* Steud, was domesticated independently of Asian rice, in the inner delta of Niger river (Cubry et al., 2018) from the wild relative species *O. barthii* around 2,500 years ago. Cultivation of *O. sativa* was introduced into West Africa around 400 years ago and this crop has since largely replaced *O. glaberrima* – although the latter is still grown, it represents only 1–2% of the total cultivated area. *O. glaberrima* is more adapted to a variety of ecologically and climatically diverse regions, its agronomically useful characters including higher tolerance to drought, high temperatures and infertile soils as well as a greater resistance to various biotic stresses (Jones et al., 1997; Linares, 2002; Thiémélé et al., 2010; Wambugu et al., 2013; Li et al., 2015; Wambugu et al., 2021; Lorieux et al., 2013). While *O. sativa* is less adapted to the African environment, this species has a higher yield potential than *O. glaberrima*, the difference being partly explained by the higher panicle complexity of Asian rice (Harrop et al., 2019). Even if molecular pathways associated with the regulation of axillary meristem identity seem to be conserved between the two species, it was possible to identify a set of genes displaying differential expression in relation to panicle phenotypic variability between African and Asian rice (Harrop et al., 2019). Moreover, a Genome Wide Association Study (GWAS) carried out on a panel of African rice accessions allowed the identification of several new genomic regions associated with panicle branching diversity and climatic variables (Cubry et al., 2020).

In the context of climate change, achieving high tolerance to environmental stresses is of paramount importance to rice agriculture while knowledge relating to the genetic control of traits governing yield potential in African rice remains very limited. Agronomic traits such as heading date, yield, plant height and grain size are controlled by many genes exerting major and/or minor effects. Identifying the genes and genomic regions associated with interspecific variation in the number of grains produced per panicle is challenging due to the polygenic nature and environmental sensitivity of panicle branching traits.

Libraries of introgression lines (ILs) provide a useful means to investigate genetic phenomena from an interspecific perspective. An introgression line library is a collection of lines with a common genetic background (the recipient genome), each carrying one or a few genomic regions originating from a donor genome. The ILs are generally obtained through recurrent backcrossing onto the recipient parent, followed by several rounds of self-fertilization (BC_nF_n) or double

haploidization (BC_nDH). The genome of the donor genotype is then represented in the library by discrete homozygous chromosomal fragments. Such ILs are also called Chromosome Segment Substitution Lines (CSSLs). In contrast to segregating populations, such as Recombinant Inbred Lines (RILs), Doubled Haploid (DH), Backcross (BC) or F_{2:3} populations, QTL analyses using CSSLs are suitable to evaluate minor allelic differences conferred by additive QTLs in a uniform genetic background in a reduced population size (Eshed and Zamir, 1995; Koumproglou et al., 2002). Genetic dissection of complex traits can thus be achieved by combining genetic variation with introgressed genomic fragments so as to reduce interference effects between QTLs. Hence, CSSLs provide a powerful means to identify QTLs for complex traits with minor and/or additive effects (Balakrishnan et al., 2019; Zhang et al., 2022). This approach may provide access to novel and potentially beneficial genes "hidden" in the genetic background of a related species that can be discovered when placed in the genetic background of a cultivated species.

In the present study, we evaluated several panicle morphological traits in a BC₃DH CSSL population developed from an interspecific cross between the recipient parent *O. sativa ssp. japonica* (cv. Caiapó) and the donor parent *O. glaberrima* (cv. MG12) (Table 1) (Gutiérrez et al., 2010). QTLs relating to modifications in panicle morphology were mapped on the rice genome, and 4th generation backcross (BC₄F_{3.5}) generations were produced to evaluate the effect of each region on panicle phenotype variation. Finally, DNA polymorphisms between the two parental genomes were investigated in detail for some key genes that were previous implicated in panicle branching diversity in *O. sativa*. The results obtained will help allow the identification of functionally significant polymorphisms and, more generally, provide an insight into the genetic bases of panicle architecture diversity between the two species.

Table 1
Panicle Traits analyzed in the CSSLs

Variable	Name	Measurement unit
PBN	Primary branch number	Number/panicle
SBN	Secondary branch number	Number/panicle
SBN/PBN	Secondary branch number per Primary branch number	Number/panicle
TBN	Tertiary branch number	Number/panicle
SpN	Spikelet number	Number/panicle
PBL	Primary Branch length	cm/panicle
RL	Rachis Length	cm/panicle

Results

Panicle Traits in the CSSL population

In order to identify new genetic factors governing the panicle branching diversity observed between *O. sativa* cv. Caiapó (referred to as *Os_Caiapó* hereafter) and *O. glaberrima* cv. MG12 (referred to as *Og_MG12* hereafter), phenotypic measurements were performed on a population of 60 BC₃DH CSSLs and their parents (Additional file1: Table S1) (Gutiérrez et al. 2010).

Six quantitative traits were evaluated per panicle: rachis length (RL); primary, secondary and tertiary branch numbers (PBN, SBN, TBN respectively); primary branch length average (PBL); and spikelet number (SpN) (Table 1, Additional file1: Table S2). The two parents showed contrasting panicle phenotypes with a higher panicle complexity observed in *Os_Caiapó* compared to *Og_MG12*. More specifically, the *Os_Caiapó* parent displayed panicles with more PBN and SBN leading to a higher SpN compared to *Og_MG12* (Fig. 1A). In contrast, the *Og_MG12* parent produced a panicle with longer PBs compared to *Os_Caiapó*. Since the trials were conducted as two repeated experiments, we computed broad sense heritability for each experiment (Additional file1: Table S3). All the measured variables showed broad sense heritability values higher than 0.8 except for TBN. For all traits, the mean of the CSSL population was similar to the mean of the recurrent *Os_Caiapó* parent (Additional file1: Table S4). The coefficient of variation for the SBN/PBN ratio was higher in the CSSL population than in *Os_Caiapó* (26.10% versus 18.26%). Histograms of the CSSLs for panicle traits showed a continuous distribution for all traits except TBN (Fig. 1B). The observed distributions were similar between the two repeated experiments. The abnormal distribution of the TBN trait was associated with a high value of CV (Coefficient Variation) and a low heritability value. This observation probably results from the rare and unstable nature of this trait which is dependent on environmental conditions and does not appear frequently in the *Os_Caiapó* parent. Moreover, the formation of tertiary branches has not been described in *O. glaberrima* populations (Cubry et al., 2020), nor was it observed in our earlier experiments.

By analyzing relationships between the different panicle traits, we observed a strong correlation (0.95) between SpN and SBN (Fig. 2A). Other traits were moderately correlated with each other, notably SBN and PBN. We also observed a low correlation between PBL and SBN. Principal component analysis showed a separate distribution of *Os_Caiapó* and *Og_MG12* parents on the PC1 and PC2 axes. In contrast, we observed an overlapping of the BC₃DH lines with the *Os_Caiapó* recurrent parent (Fig. 2B). Analysis of variable contributions showed that SpN is the main trait contributing to the diversity observed in this population.

Detection of QTLs associated with panicle architecture traits

The Dunnett's test revealed a total of 37 lines with significant differences compared to the *Os_Caiapó* recurrent parent for all traits combined (Additional file1: Table S5). In general, these lines bear at least two *Og_MG12* introgression fragments. For this reason, assigning the underlying QTL to a single chromosome segment was not straightforward. The CSSL Finder software was therefore used to detect genomic regions associated with panicle phenotype variation with a

F-test value at each marker (Gutiérrez et al., 2010). Graphical genotype representations were also produced in which the CSSLs were ordered by trait value for each evaluated trait (Additional file 2: Figure S1). We considered the F-test as significant when its value was higher than 10.0. QTLs were assigned to the *Og_MG12* introgressed regions of these CSSLs if the F-test was significant and confirmed by the graphical genotype logical analysis.

In total, 15 QTLs were detected for all combined traits analyzed with the exception of rachis length (Fig. 3). The F-test value of the markers allowing the detection of each QTL are reported in Table 2 along with their positions in the *O. sativa* and *O. glaberrima* reference genomes (cv. Nipponbare IRGSP-1.0 and cv. CG14 OglRS2 respectively).

Table 2
QTLs for morphological panicle traits detected in the CSSL population

Traits	ID_QTL	Chromosome	SSR Marker	F-Test value	significance F-Test	Position (bp) in <i>O. sativa</i> Caiapo	Position (bp) in <i>O. glaberrima</i> MG12	Position (bp) in <i>O. sativa</i> MSU 7	Position (bp) in <i>O. glaberrima</i> OMAPv2	Representative CSSL	Positi allele
PBN	qPBN1	1	RM472	3.48	NS	38 918 631	35 483 127	37 890 275	34 691 456	L_10	<i>Og</i>
			RM165	12.56	*	41 165 563	37 610 087	40 107 107	36 797 582	L_46	<i>Og</i>
			RM104	7.92	NS	41 226 405	37 667 912	40 167 993	36 855 255		
	qPBN2	2	RM191	0.01	NS	13 357 683	13 118 242	13 367 915	13 091 268	L_10	<i>Og</i>
			RM341	15.49	*	19 822 576	19 710 485	19 342 082	19 060 676	L_46	<i>Og</i>
			RM262	7.87	NS	21 260 857	21 173 898	20 800 911	20 552 457		
	qPBN7	7	RM10	8.88	NS	22 227 559	20 124 022	22 189 175	19 801 671	L_10	<i>Og</i>
			RM351	16.44	*	23 991 049	21 742 091	23 926 002	21 416 312	L_46	<i>Og</i>
			RM234	7.08	NS	25 545 695	23 016 334	25 473 749	22 717 218		
PbL	qPBL1	1	RM315	3.91	NS	37 748 125	34 409 891	36 735 198	33 646 637	L_10	<i>Og</i>
			RM472	12.23	*	38 918 631	35 483 127	37 890 275	34 691 456	L_46	<i>Og</i>
			RM165	22.55	*	41 165 563	37 610 087	40 107 107	36 797 582		
			RM104	14.08	*	41 226 406	37 667 913	40 167 994	36 855 255		
			RM3362	11.51	*	44 150 041	40 246 380	43 044 090	39 436 568		
	qPBL3	3	RM55	1.08	NS	30 677 384	28 677 802	29 052 298	28 518 740	L_10	<i>Og</i>
			RM3525	14.51	*	32 028 705	30 028 933	30 393 958	29 873 318	L_46	<i>Og</i>
			RM7000	12.89	*	35 524 810	33 277 548	33 802 689	33 128 938		
			RM227	1.08	NS	36 679 424	34 296 048	34 932 380	34 146 855		
	qPBL7	7	RM10	8.14	NS	22 227 559	20 124 022	22 189 175	19 801 671	L_10	<i>Og</i>
			RM351	15.78	*	23 991 049	21 742 091	23 926 002	21 416 312	L_46	<i>Og</i>

Traits	ID_QTL	Chromosome	SSR Marker	F-Test value	significance F-Test	Position (bp) in <i>O. sativa</i> Caiapo	Position (bp) in <i>O. glaberrima</i> MG12	Position (bp) in <i>O. sativa</i> MSU 7	Position (bp) in <i>O. glaberrima</i> OMAPv2	Representative CSSL	Positi allele
			RM234	8.8	NS	25 545 695	23 016 334	25 473 749	22 717 218		
SBN	qSBN/PBN11	11	RM21	1.14	NS	20 161 293	19 279 722	19 639 289	18 598 655	L_42	<i>Os</i>
			RM206	13.74	*	22 961 364	21 790 549	22 480 895	21 092 821	L_55	<i>Og</i>
			RM254	15.09	*	24 651 269	23 182 998	24 230 511	22 477 748	L_56	<i>Og</i>
			RM224	9.2	NS	29 246 818	25 784 047	27 673 312	25 499 087		
	qSBN/PBN12	12	RM28130	9.09	NS	16 565 178	15 215 344	16 706 323	15 275 476	L_42	<i>Og</i>
			RM277	14.84	*	21 403 653	20 209 650	22 364 793	20 270 371	L_55	-
			RM235	2.69	NS	25 324 296	23 473 458	26 141 561	23 604 232	L_56	-
SpN	qSpN3	3	RM60	10.8	*	109 164	86 928	106 972	77 908	L_55	<i>Og</i>
			RM22	9.45	NS	1 561 917	1 493 255	1 520 630	1 485 562	L_11	<i>Og</i>
										L_26	<i>Os</i>
	qSpN11	11	RM21	1.14	NS	20 161 293	19 279 722	19 639 289	18 598 655	L_55	<i>Og</i>
			RM206	13.74	*	22 961 364	21 790 549	22 480 895	21 092 821	L_11	<i>Os</i>
			RM254	15.09	*	24 651 269	23 182 998	24 230 511	22 477 748	L_26	<i>Og</i>
			RM224	9.2	NS	29 246 818	25 784 047	27 673 312	25 499 087		
TBN	qTBN1	1	RM226	0.31	NS	35 105 017	31 957 160	34 034 101	31 200 058	L_10	<i>Og</i>
			RM265	22.67	*	36 243 983	33 029 536	35 197 671	32 275 011	L_4	<i>Og</i>
			RM315	6.09	NS	37 748 125	34 409 891	36 735 198	33 646 637	L_46	
			RM472	31.11	*	38 918 631	35 483 127	37 890 275	34 691 456		
			RM165	4.62	NS	41 165 563	37 610 087	40 107 107	36 797 582		
	qTBN3-1	3	RM175	1.93	NS	3 911 091	3 633 683	3 866 833	3 625 881	L_10	<i>Og</i>
			RM7576	21.91	*	6 107 599	5 772 156	6 079 319	5 767 071	L_4	<i>Og</i>
			RM7	0.27	NS	9 810 142	9 300 611	9 829 641	9 280 539	L_46	
	qTBN3-2	3	RM227	0.72	NS	36 679 424	34 296 048	34 932 380	34 146 855	L_10	<i>Og</i>

Traits	ID_QTL	Chromosome	SSR Marker	F-Test value	significance F-Test	Position (bp) in <i>O. sativa</i> Caiapo	Position (bp) in <i>O. glaberrima</i> MG12	Position (bp) in <i>O. sativa</i> MSU 7	Position (bp) in <i>O. glaberrima</i> OMAPv2	Representative CSSL	Positi allele
			RM148	15.58	*	37 552 877	35 129 500	35 843 050	34 980 430	L_4	<i>Og</i>
			RM85	0.72	NS	38 025 225	35 613 693	36 348 204	35 464 999		
	qTBN5	5	RM159	6.76	NS	496 697	433 400	488 155	404 817	L_4	<i>Og</i>
			RM267	25.12	*	2 871 237	2 517 535	2 881 410	2 485 558		<i>Os</i>
			RM194	25.12	*	5 179 705	4 653 443	5 329 987	4 472 981		
			RM169	0.96	NS	7 361 060	6 805 643	7 498 060	6 637 671		
	qTBN7	7	RM11	0.1	NS	19 315 836	17 468 875	19 257 970	17 138 872	L_10	<i>Og</i>
			RM10	12.61	*	22 227 559	20 124 022	22 189 175	19 801 671	L_4	<i>Og</i>
			RM351	17.42	*	23 991 049	21 742 091	23 926 002	21 416 312	L_46	
			RM234	66.31	**	25 545 695	23 016 334	25 473 749	22 717 218		
			RM18	62.09	**	25 725 478	23 182 832	25 653 583	22 884 015		
			RM134	26.55	*	26 708 250	24 069 776	26 637 574	23 770 479		
			RM118	33.95	*	26 708 384	24 069 876	26 637 674	23 770 550		
			RM420	0.09	NS	29 538 599	26 687 759	29 432 340	26 361 122		

RL trait: 13 CSSLs showed significant changes in RL value in comparison to the *Os_Caiapo* recurrent parent (Additional file1: Table S5). However, no QTLs were detected for this trait (Additional file2: Figure S1).

SBN/PBN trait: a total of 16 CSSLs showed a significant difference in secondary branch number per primary branch (SBN/PBN ratio) compared to the *Og_MG12* parent (Additional file 1: Table S5). Two QTLs associated with a decreased SBN/PBN ratio were detected (qSBN/PBN11 and qSBN/PBN12, maximum F-test scores of 15.09 and of 14.94 respectively) (Table 2; Additional file2: Figure S1). The average SBN/PBN ratio values were 3.0 for *Os_Caiapo* and of 1.7 for *Og_MG12*, corresponding to a decrease of 42% in *Og_MG12* relative to *Os_Caiapo*. The CSSLs L_42, L_55 and L_56 showed a decrease of 44.7%, 42.4% and 36.3% respectively relative to the *Os_Caiapo* parent, suggesting a high effect of the *Og_MG12* introgression(s) in these lines (Additional file1: TableS5). The phenotyping of these lines was repeated for a second year to confirm panicle trait variation (Additional file3: FigureS2). The L_55 and L_56 lines contain an *Og_MG12* segment in chromosome 11, but with missing marker information at the position of qSBN/PBN12. In contrast, the CSSL L_42 displays only an *Og_MG12* segment in chromosome 12 (Additional file3: Figure S2).

SpN trait: a total of 12 CSSLs showed a significant difference compared to the *Os_Caiapo* recurrent parent (Additional file 1: Table S5). Two QTLs (qSpN3 and qSpN11) were associated with a decreased SpN with maximum F-test scores of 10.8 and 15.1 respectively (Table 2; Additional file2: Figure S1). CSSLs L_55, L_11 and L_26 showed decreases of 37.9%, 32.3% and 30.8% respectively compared to the *Os_Caiapo* parent (Additional file1: Table S5). Lines L_55 and L_26 contain *Og* segments in both chromosomes 3 and 11 in the positions of the detected QTLs (Additional file3: Figure S2). Line L_11 contains only one *Og_MG12* introgression in chromosome 11, related to qSpN11 (Additional file 3: Figure S2).

PBN trait: a total of 13 CSSLs had significantly different PBN values compared to the *Os_Caiapo* recurrent parent (Additional file1: Table S5). Three QTLs corresponding to a decreased PBN (qPBN1, qPBN2 and qPBN7) with maximum F-test scores of 12.56, 15.49 and 16.44 were detected on chromosomes 1, 2 and 7 respectively (Table 2; Additional file2: Figure S1). Average PBN values were 11.9 for *Os_Caiapo* and 8.2 for *Og_MG12* (Additional file1: Table S5). The PBN values of two CSSLs (L_10 and L_46) differed significantly from the *Os_Caiapo* recurrent parent with a reduction of 24.1% and 20.9% respectively (Additional file 1: Table S5). These 2 CSSLs harbor similar *Og_MG12* segments in chromosomes 1, 2 and 7 (Additional file3: Figure S2).

PBL trait: Fourteen CSSLs exhibited significant differences with the *Os_Caiapo* recurrent parent for the PBL trait. Three QTLs corresponding to an increased PBL (qPBL1, qPBL3 and qPBL7) with maximum F-test scores of 22.55, 14.51 and 15.78 were detected on chromosomes 1, 3 and 7 respectively (Table 2; Additional file2: Figure S1). Average PBL values were 11.1 for *Os_Caiapo* and 12.3 for *Og_MG12* (Additional file1: Table S4). Two CSSLs (L_10 and L_46)

showed increased PBL values of 19% and 12.8% respectively in comparison to the *Os_Caiapó* recurrent parent and harbored similar *Og_MG12* segments in chromosomes 1, 3 and 7 (Additional file1: Table S5; Additional file3: Figure S2).

TBN trait: Surprisingly, several CSSLs showed a significantly increased TBN value in our field conditions (Additional file1: Table S5) and five QTLs were detected: qTBN1, qTBN3-1, qTBN3-2, qTBN5 and qTBN7 with maximum F-test scores of 31.11, 21.91, 15.98, 25.12 and 66.09 respectively (Table 2; Additional file2: Figure S1). The L₁₀ and L₄ lines have similar *Og_MG12* segments in chromosomes 1, 3 and 7. Within chromosome 5, only the L₄ line contains a segment from the *Og_MG12* parent (Additional file3: Figure S2). As the L₄₆ line contains otherwise similar *Og_MG12* introgressions in chromosome 1, 3 and 7, this CSSL was included in the second phenotyping campaign, in which it revealed the presence of tertiary branches on its panicles in contrast to L₄ (Additional file3: Figure S2). Overall, these results support an association of *Og_MG12* introgressions with the presence of tertiary branches.

For the PBN, PBL and TBN panicle traits, the same BC₃DH CSSLs (L₁₀ and L₄₆) led to the detection of several QTLs. The 2 lines in question display a similar panicle phenotype with a higher number of tertiary branches associated with a decreased PBN and increased PBL and SBN/PBN ratio values compared to the *Os_Caiapó* recurrent parent (Fig. 4A, B). The L₁₀ and L₄₆ BC₃DH lines contain a complex association of *Og_MG12* introgressions in their genomes (Fig. 4C). For clarification, the regions corresponding to colocalized QTLs, associated with different traits, were renamed without specifying the associated traits: q₁ (for qPBL1, qPBN1 and qTBN1); q₂ (for qPBN2), q₃ - 1 (for qTBN3-1); q₃ - 2 (for qPBL3 and qTBN3-2); q₅ (for qTBN5); and q₇ (for qTBN7, qPBN7 and qPBL7).

Substitutions of Os_Caiapó genomic regions by corresponding Og_MG12 segments can result in an added value panicle phenotype

As the altered phenotype observed in lines L₁₀ and L₄₆ was associated with multiple substitution segments (Fig. 4C), it was critical to determine whether this phenotype, and notably the formation of tertiary branches, involved one or several QTLs. Thus, different BC₄F_{3:5} lines were obtained from the BC₃DH lines by backcrossing and self-pollination (Fig. 4C), after which homozygosity was checked using the SSR markers carried by the introgressed *Og_MG12* segments present in the BC₃DHs. Panicle traits were phenotyped for five BC₄F_{3:5} lines containing the different individual *O. glaberrima* introgressed segments affecting TBN, together with *Os_Caiapó* and *Og_MG12* (Fig. 4B; Additional file1: Table S1). A second phenotypic campaign was carried out for validation of the BC₄ lines L_B, L_D and L_E (Additional file4: Figure S3).

Divergent panicle traits compared to the *Os_Caiapó* parent were observed in all BC₄ lines except for L_C (Fig. 5B). The latter contains an introgression at the beginning of q₇ (RM11 to RM10), suggesting that this region does not influence panicle architecture. All BC₄ lines except L_C showed longer primary branches compared to the *Os_Caiapó* parent, suggesting that introgressions in chromosomes 1, 3 and 7 could independently influence primary branch length (Fig. 5A). The L_A line, containing *Og_MG12* regions corresponding to q₁, q₂ and q₃ - 2, also showed a decreased PBN value associated with a higher SBN/PBN ratio. However, no tertiary branches were observed. This suggests that the association of the q₁, q₂ and q₃ - 2 regions could induce a reduction in PBN associated with an increased SBN.

Line L_B containing only one *Og_MG12* introgression (from RM60 to RM7) in place of q₃ - 1 showed increased values for four panicle traits (PBN, PBL, SBN and SpN) but not for TBN. However, these results were not observed during the second phenotyping of this line, except for an increase in PBL. Thus, it can be deduced that the q₃ - 1 region positively influences primary branch length in a manner that is independent of the other detected QTLs.

Finally, the BC₄ lines L_D and L_E were the only ones observed to develop tertiary branches (Fig. 5A). In both lines, this phenotype is associated with an increased SBN/PBN and a longer PBL. The higher SBN/PBN ratio is associated with a decreased PBN in L_D, meaning that in this line more secondary branch meristems are established during panicle development. Similar results for SBN/PBN and TBN were observed in the second round of phenotyping, with a comparable decrease in PBN in the L_E line (Additional file4: FigureS3).

The two aforementioned lines contain contiguous introgression fragments in chromosome 7, from RM10 to RM18 for L_D and from RM134 to RM118 for L_E. Since the exact recombination positions of the introgressed segments for each BC₄ is not known, the regions between SSRs with different genotypes are included in the QTL intervals. In Fig. 5A, it can be observed that lines L_D and L_E, which share the same phenotypes except for PBN, may harbor introgressions that overlap over only a small region between RM18 and RM134. Based on these observations, two different hypotheses can be proposed with regard to QTL position(s) in the q₇ region (Fig. 5A). The first one postulates the existence of a common QTL located between RM18 and RM134 that controls PBL, SBN/PBN and TBN. The second hypothesis is that two different QTLs, q₇ - 1 between RM10 and RM134, and q₇ - 2 between RM18 and RM420, exert a similar effect on PBL, SBN/PBN and TBN but act differently on PBN.

Taken together, the above results suggest that panicle architecture is a complex trait controlled by various different genomic regions which positively or negatively influence branching. An association of q₁, q₂ and q₃ - 2 produces opposing effects on PBN and SBN while only the q₃ - 1 region, comprised between RM175 and RM7, may impact positively upon PB length. Finally, the region in q₇, which negatively influences the number of primary branches and may be associated with the formation of tertiary branches and an increase in SBN per primary branch, may be defined as being either between the RM18 and RM134 marker positions or associated with two QTLs positioned between RM10 and RM134 for q₇ - 1 and from RM18 to RM420 for q₇ - 2.

Co-location of QTLs with known genes and QTLs detected with other populations.

Many studies have reported QTLs associated with panicle architecture and/or yield traits or both, using bi-parental (QTL mapping) or diversity panels GWAS (Crowell et al., 2014; Ya-fang et al., 2015; Bai et al., 2016b; Crowell et al., 2016; Rebolledo et al., 2016; Ta et al., 2018; Zhang et al., 2019; Cubry et al., 2020; Bai et al., 2021; Zhong et al., 2021). We found 70 common sites between the QTLs detected in this study and those reported earlier (Additional file 1: TableS6). For instance, co-located sites relating to similar traits (SBN and SpN) were found on chromosomes 11 and 12. Nevertheless, despite the significant number of

common sites identified in the different studies, the traits that they have been reported to affect are not always strictly comparable or even similar, even if they relate in some way to panicle architecture or yield.

To evaluate the synteny of the QTL regions between the genomes of the two CSSL parents, sequences from the assemblies of the *Os_Caiapó* and *Og_MG12* genomes obtained by ONT sequencing (<https://doi.org/10.23708/QMM2WH>; <https://doi.org/10.23708/1JST4X>; Additional file1: TableS7) were used to carry out local alignment for each QTL region. The assembled genomes are of high continuity and completeness, with BUSCO score of 98.4% allowing precise detection of structural variations. For the majority of the QTL regions, no major structural variation was observed between the two genomes (Additional file5: Figure S4). However, some notable differences were detected at the qPBN2, qTBN5, qTBN7, qSpN3 and qSBN12/PBN12 sites. Within the QTLs qPBN2 and qTBN7, inverted segments of 1,296,899 and 230,762 bp were observed between *Os_Caiapó* and *Og_MG12* respectively (Additional file1: Table S8). The genomic regions for QTLs qTBN5, qSpN3 and qSBN/PBN12 displayed InDels between *Os_Caiapó* and *Og_MG12* (Additional file1: Table S8). With the exception of qSBN11, we observed a high sequence similarity in the QTL regions between the two genomes, suggesting similar content in terms of coding sequences. Synteny analysis was extended by the alignment of the *Os_Caiapó* and *Og_MG12* QTL regions with those of the *O. sativa* cv. Nipponbare reference genome (Additional file5: Figure S4). Based on the high conservation of the corresponding regions between *Os_Caiapó* and *Os_Nipponbare*, we then used as a reference, for subsequent analyses, the *O. sativa* cv. Nipponbare (IRGSPv1.0) functional gene annotation databases (i.e., MSU7 and RAP_db) for candidate gene prioritization in relation to panicle architecture and flowering.

For all QTLs identified in the present study, we found several known genes with relevant functions that related to flowering and/or panicle development (Table 2; Additional file6: Figure S5). Genes such as *LAX PANICLE1* (*LAX1*), *OsMADS1/LEAFY HULL STERILE1* (*LHS1*), *OsMADS14*, *OsMADS34/PANICLE PHYTOMER2* (*PAP2*), *OsINDETERMINATE SPIKELET 1* (*OsIDS1*), *OsMADS18*, *DENSE AND ERECT PANICLE2* (*DEP2*) and *FRIZZY PANICLE* (*FZP*) are known to be involved in the control of the panicle development and were suggested as candidates that might contribute to the panicle branching diversity observed between the two parents *Os_Caiapó* and *Og_MG12* (Chongloi et al., 2019; Wang et al., 2021; Yin et al., 2021).

q₇ genetic variations between *Os_Caiapó* and *Og_MG12*

We paid particular attention to the q₇ region, as it was found to be associated with several panicle morphological traits and to span a genomic region that included several panicle-associated genes. To understand if specific genetic modifications present in the q₇ QTL region could be associated with panicle architecture variation, we further compared the constituent genes in the region between the RM10 and RM420 markers in the *Os_Caiapó* and *Og_MG12* genomes on chromosome 7 (22.23–29.59 Mbp in *Os_Caiapó* vs 20.12–26.69 Mbp in *Og_MG12*). For this purpose, gene annotation comparison and BLAST analyses were performed to explore in detail gene synteny and presence/absence of genes between the *Os_Caiapó* and *Og_MG12* genomes within this region (Fig. 5B; Additional file1: Table S9). Between the RM10 and RM420 marker positions, we observed a variation in the number of genes between the two genomes (Additional file1: Table S9). This variation is due to several factors: the differential presence of corresponding orthologs between the two genomes; differential gene duplication within the region; and different locations of a given gene within the two genomes (i.e., genomic rearrangement) (Fig. 5B; Additional file1: Table S9). Among the genes that differ between the two genomes in this region, many are related to transposable elements (TEs) or are hypothetical protein-encoding genes. None of them has a known function relating to panicle development or flowering control, nor to any other developmental processes.

Special attention was paid to common genes present in the q₇ region that were known to be related to inflorescence development and/or meristem activity and maintenance. Based on a bibliographic survey, no candidate genes of special interest were identified within the region between the RM18 and RM134 markers. In contrast, several interesting candidate genes were identified in the q₇ – 1 and q₇ – 2 regions: the *OsHOX14*, *OsMADS18* and *DEP2* genes in q₇ – 1 and the *FZP* and *WOX11* genes in q₇ – 2 (Fig. 5C). SNP and InDel sites associated with the aforementioned genes that were polymorphic between the *Os_Caiapó* and *Og_MG12* genomes were analyzed in order to detect amino acid modifications, open reading frame alterations or transcription factor binding site (TFBS) variations within promoter regions.

OsHOX14 (LOC_Os07g39320/Os07g0581700) encodes a protein of the Homeodomain-leucine zipper (HD-Zip) TF family and is involved in the regulation of panicle development (Beretta et al., 2023; Shao et al., 2018). Two of the identified SNPs cause amino acid variations in the coding sequence, including one in the homeobox domain (Fig. 5C; Additional file1: Table S10). Comparison of the *OsHOX14* and *OgHOX14* promoter regions revealed InDel and SNP variations leading to the loss of 9 TFBSs and a gain of three new TFBSs in the CSSLs bearing *Og_MG12* q₇ – 1 introgression (Fig. 5C; Additional file1: Table S11).

OsMADS18 (LOC_Os07g41370/Os07g0605200) encodes a protein of the AP1/FUL-like MADS-box TF subfamily (Masiero et al., 2002). In addition to its role in flowering time promotion (Fornara et al., 2004), *OsMADS18* is known to specify inflorescence meristem identity (Kobayashi et al., 2012). SNPs were detected in the coding sequence; one of the SNPs leads to a non-synonymous change in the *OgMADS18* protein outside the known binding or functional domains (Fig. 5C; Additional file 1: Table S10). Scanning of the promoter regions of *OsMADS18* and *OgMADS18* in the *Os_Caiapó* and *Og_MG12* genomes respectively revealed several SNPs and InDels that lead to variations between the two genomes (presence/absence) of TFBSs recognized by AP2, NAC or HB TFs (Fig. 5C, Additional file1: Table S11).

DEP2 (LOC_Os07g42410/Os07g061600) encodes a plant-specific protein of unknown function associated with panicle size (Li et al., 2010). Various SNPs and one InDel are observed between the coding sequences of *OsDEP2* and *OgDEP2* which result respectively in changed amino acid identities between the two protein orthologs (Fig. 5C; Additional file 1: Table S10) and an insertion of two amino acids in the *OgDEP2* protein. Sequence variations were also observed between the promoters of *OsDEP2* and *OgDEP2*; these variations lead to the loss of bZIP, B3, AT-Hook and AP2 TFBSs in the CSSLs carrying q₇. On the other hand, in comparison to *O. sativa*, the CSSLs carrying q₇ include NAC and bZIP TFBSs specific to the promoter of *OgDEP2*.

Moreover, among the genes contained in the q₇ – 2 region, *FZP* (LOC_Os07g47330/Os07g0669500) encodes an AP2/ERF TF known to play a role in the regulation of floral meristem determinacy in rice (Komatsu et al., 2003; Zhu et al., 2003). Several SNP and InDel variations observed in the coding region of

FZP lead to coding sequence amino acid changes between the two genomes (Fig. 5C; Additional file 1: Table S10). An InDel of 9 bp in *OgFZP* results in the insertion of three histidine amino acids outside the single AP2 domain in the *OgFZP* protein. Several recent studies have shown that variations in the promoter region and 3' UTR (Untranslated Transcribed Region) of the *OsFZP* gene affect panicle architecture by through changes in the number of branches produced (Bai et al., 2016, 2017; Huang et al., 2018; Wang et al., 2020; Chen et al., 2022). Here, we observed several SNPs and InDels between the promoter of the two orthologs, which result in a differential presence of TFBSs (Fig. 5C, Additional file1: Table S11). No variation was observed in the copy number variant (CNV) motif of 18 bp described by Bai et al. (2017). Huang et al. (2018) revealed a deletion of 4 bp in the *OsFZP* 5' regulatory region in comparison to the sequence of *O. rufipogon*. This deletion is observed in *Os_Caiapó* and not in the genome of *Og_MG12* (Fig. 5C). Recently, it has been shown that CU-rich elements (CUREs) present in the 3' UTR of the *OsFZP* mRNA are crucial for efficient *OsFZP* translational repression (Chen et al., 2022). Three CUREs are detected in the 3' UTR of *OsFZP* in the *Os_Caiapó* genome. In contrast, a deletion of the third CURE sequence is observed in the *Og_MG12* genome (Fig. 5C).

Although the WUSCHEL-related homeobox *WOX11* gene (LOC_Os07g48560/Os07g0684900) has been mainly described as playing an important role in root development (Zhao et al., 2009; Zhang et al., 2018), this gene is also required to regulate rice shoot development and panicle architecture (Cheng et al., 2018). An InDel in the coding region of *WOX11* leads to an Asn amino acid insertion in the *OgWOX11* protein in *Og_MG12* without affecting the homeobox domain of the protein (Fig. 5C, Additional file1: Table S10). In the promoter region, variations between *Os_Caiapó* and *Og_MG12* lead to the presence/absence of several TFBSs in the promoter of *OgWOX11* (Fig. 5C; Additional file 1: Table S11). We also observed a large insertion of 3,760 bp, containing about 31 AP2 TFBSs, in *Og_MG12* compared to the *Os_Caiapó* genome.

Overall, our analysis revealed variations in synteny that demonstrate an absence of certain *Os_Caiapó* genes and the addition of other loci as a consequence of *Og_MG12* genomic introgressions within the CSSLs: some of these changes can be hypothesized to play a role in determining panicle trait diversity. Moreover, our analysis of candidate genes revealed TFBS variations and protein coding sequence polymorphisms that may lead to variations in transcript levels and/or protein activity in CSSLs harboring the q₇ introgression.

Discussion

Added value of the *O. sativa* x *O. glaberrima* CSSL population

Inter-specific *O. sativa* x *O. glaberrima* CSSL genetic resources are of great interest, as *O. glaberrima* can provide a gene pool with potential for rice improvement in terms of resistance to biotic and abiotic stresses and ecological adaptability (Gutiérrez et al. 2010). CSSL libraries are also good pre-breeding materials for the simultaneous identification, transfer and pyramiding of key genes in crop improvement programs. They are also useful for the study of traits lost or retained during the process of evolution, domestication and selection (Ali et al., 2010; Balakrishnan et al., 2019). Genetic strategies based on CSSL populations have the advantage of using a relatively small number of lines for experiments, leading to the possibility of replicating evaluations, thereby enhancing statistical strength for complex and time-consuming phenotypic assessments.

In this study, backcross introgression lines (BC₃DH) of *O. sativa_Caiapó* x *O. glaberrima_MG12* were phenotyped for panicle morphological traits, which are key components of yield potential. Our analyses led to the detection of 15 QTLs, localized on chromosomes 1, 2, 3, 5 and 7 and comprising several genes known to be involved in panicle architecture determination (Table 1). Colocalization was observed with previous QTLs and GWAS sites associated with panicle branching traits, supporting the implication of the QTLs detected in this study in the regulation of panicle morphology. Moreover, since the genomic sites described here were detected using various different sets of populations, we propose that the genomic regions in question play a major role in determination of panicle architecture diversity.

Some of the BC₃DH lines used in this analysis contain several *Og_MG12* introgression regions, thus potentially allowing the detection of several QTLs with epistatic interactions. For most of the traits studied, we detected several QTLs within the same line, as observed in the lines L₁₀ and L₄₆ that allowed the localization of QTLs governing TBN, PBN- and PBL-related traits. This is not surprising in the case of a complex trait such as panicle architecture, which is controlled by a large number of genes and QTLs with small effects that could be influenced by environmental and epistatic interactions (Xing et al., 2002). An additional advantage of the CSSL-based approach is that it provides novel interspecific combinations of parental alleles which may significantly affect panicle branching traits, thereby providing an insight into how inflorescence architecture is governed in the two species and how their respective genetic factors interact. The fact that we observed a colocalization of QTLs for different panicle traits suggests a likely genetic interdependence between these traits; this hypothesis is supported by correlations observed between phenotypic values for morphological characters such as SpN, SBN, PBN, PBL and RL that share common QTLs.

Impact of *O. glaberrima* introgressions on panicle traits in the *O. sativa* background

Compared to the *Os_Caiapó* recurrent parent, the *Og_MG12* donor parent displays a global reduction in panicle constituent numbers (i.e., numbers of primary branches, secondary branches and spikelets) and produces longer primary branches. Tertiary branches were not observed in the *Og_MG12* donor parent. On the basis of these observations, the QTLs detected in the CSSLs were expected to be associated with a global decrease in panicle constituent numbers (PBN, SBN, SpN) and an increase of PB length in comparison to the *Os_Caiapó* parent. Indeed, this is mainly what was observed. In most lines studied, introgressions of *Og_MG12* alleles of panicle-regulating genes produced a negative effect on the regulatory network controlling panicle morphological traits, indicating that some of the genomic variations observed between *Og_MG12* and *Os_Caiapó* in these regions are functionally important.

Surprisingly, we also detected QTLs associated with an increase in panicle trait values in comparison to the *Os_Caiapó* recurrent parent. Two of the BC₃DH CSSLs (L₁₀ and L₄₆) carried four substitution segments from the *Og_MG12* parent donor and displayed panicle branching modifications that included not only a reduction of PBN and an increase of PBL as expected, but also an increase in secondary branch number per primary branch and a formation and development of tertiary branches (TB). As confirmed with the corresponding BC₄ lines, both of the CSSL lines displayed a phenotype with an added value for

one or several panicle traits. For example, the *Og_MG12* introgression localized on chromosome 3 in place of q_3 – 1 induces a globally higher complexity of panicle branching with an increase in PBN, SBN/PBN and SpN, associated with an increased PBL. This region includes the gene *OsMADS1/LHS1* belonging to the *SEPALLATA MADS-Box* TF family which is known to play an important role in determining floral meristem identity and in floral organ development (Jeon et al., 2000; Prasad et al., 2005; Cui et al., 2010). In *O. sativa*, *OsMADS1/LHS1* directly regulates several other transcription factor genes (MADS box and Homeodomain family members) and hormone signaling pathways implicated in floral meristem specification, maintenance and determinacy (Khanday et al., 2013).

The unexpected results obtained for the aforementioned lines indicate that introgression of certain *Og_MG12* regions into the *Os_Caiapó* genetic background can lead to an enhanced panicle phenotype and suggest a perturbation of the pathways regulating inflorescence development in *Os_Caiapó*. This perturbation could be explained by several mechanisms. Firstly, it is possible to postulate the presence of a negative regulator of branching in *Os_Caiapó* that is absent in the *Og_MG12* introgression, leading to an upregulation of branching in the CSSL. A second hypothesis could be proposed whereby one or more genes specific to *O. glaberrima* are present in the *Og_MG12* introgression compared to the corresponding *Os_Caiapó* region. One or more of these genes could be positive regulators of panicle architecture, with this regulatory potential being either latent or only very weakly expressed in the *Og_MG12* background. A third possible explanation could be proposed involving *cis*-variations associated with orthologous genes conserved between the two genomes. In this case, the addition of new *Og_MG12* genes, or their variant allelic forms into the *Os_Caiapó* genomic background could be postulated to exert an additive effect on panicle branching in the CSSLs, by affecting regulatory interactions within the *Os_Caiapó* network governing panicle branching.

In a similar way, our observations on the effects of QTL q_7, which favored increased SB and TB numbers, might have more than one possible explanation. Comparative analysis of the BC₄ lines led us to suggest two hypotheses (above) regarding the position of QTL(s) on q_7 region (Fig. 5A). However, based on quantitative and qualitative differences between the BC₄ line phenotypes, we favor the hypothesis whereby the presence of two distinct regions could influence panicle trait variation differentially (i.e., the q_7 – 1 and q_7 – 2 regions). In this scenario, both would be associated with increased PBL, SBN and TBN traits values whereas q_7 – 1 would have an additional negative effect on PBN. Moreover, the overlapping region of the introgressed segments in lines L_D and L_E, which spans a region of ~ 984 Kbp, is recombined in both lines. This implies that Lines L_D and L_E can bear the *Os* allele at any locus in this region so the probability that both lines bear the *Og* allele at a same putative QTL position is low. In addition, this region only contains a few genes whose predicted function is unknown.

In parallel, we observed a good synteny between the two species within the q_7 – 1 and q_7 – 2 regions, ruling out the possibility that large-scale genomic rearrangements might explain the phenotypic variations observed. Indeed, the strong global synteny between these regions suggests rather that CSSL phenotypes might be accounted for by modifications within the genomic regions of key regulatory genes that give rise to variations in their expression or in their protein functions and/or interactions through gene regulatory networks (GRNs). Mutations affecting transcription factor proteins and/or DNA sequences or both, such as enhancers and promoters, fall into this category (Hill et al., 2021). Moreover, traits associated with dynamic processes are more readily modified through their *cis*- and/or *trans*-regulation rather than through coding sequence mutations (Wray, 2007). The q_7 – 1 and q_7 – 2 regions comprise several genes known for their role in panicle development and numerous SNPs and InDels have been detected in the promoter regions and coding sequences of these candidate genes.

Within the q_7 – 1 region, three genes have been reported to have a direct function in panicle architecture establishment. The homeodomain-leucine zipper transcription factor gene *OsHOX14* has been shown to be involved in the regulation of panicle development and its loss of function leads to a reduction in PBN compared to wild type (Shao et al., 2018; Beretta et al., 2023). The second candidate gene *OsMADS18*, in addition to its role in flowering time promotion (Fornara et al., 2004), is known to specify inflorescence meristem identity through interaction with PANICLE PHYTOMER2 (PAP2/*OsMADS34*) and the two other members of the AP1/FUL subfamily, *OsMADS14* and *OsMADS15* (Kobayashi et al., 2012). *DEP2* encodes a plant-specific protein without a known functional domain, mutations of which affect elongation of the rachis and both primary and secondary branches, caused by a defect in cell proliferation (Li et al., 2010; Mao et al., 2020; Wang et al., 2021b). For each of these three genes, the *Og_MG12* allelic version shows variations in comparison to *Os_Caiapó* in both protein coding sequences and in putative TFBSs within the promoter region. The presence of these binding sites may reveal a divergence in the regulation of the expression of these genes between the two parents which could lead to the morphological variations observed in the CSSLs (i.e., decrease in PBN).

Concerning the q_7 – 2 region associated with the formation of higher order panicle branches (i.e., secondary and tertiary branches), the candidate genes *WOX11* and *FZP* were detected in both genomes. The WUSCHEL-related homeobox gene *OsWOX11* is necessary and sufficient to promote crown root emergence and growth in rice by responding to or regulating auxin and cytokinin signaling (Zhao et al., 2009, 2015; Cheng et al., 2014, 2015; Zhang et al., 2018). *OsWOX11* can recruit a histone acetyltransferase complex to activate downstream target genes to stimulate cell proliferation in crown root development (Zhou et al., 2017). Cheng et al. (2018) showed that *OsWOX11* and JM705 cooperatively control shoot growth and commonly regulate the expression of a set of genes involved in meristem identity (Cheng et al., 2018) indicating that the role of this gene is not confined to roots. A comparison of the promoter regions of *WOX11* orthologs revealed numerous SNP and InDel differences between the two genomes and an insertion of 3,760 bp in *Og_MG12*, which includes numerous AP2 TFBS potentially recognized by the *MULTI-FLORET SPIKELET 1 (MFS1)* and *OsFZP* genes, known to play an important role in the regulation of spikelet meristem determinacy (Komatsu et al., 2001, 2003; Zhu et al., 2003; Ren et al., 2013). In an *O. sativa* genomic environment, the promoter of *OgWOX11* containing new AP2 TFBSs could potentially be programmed differently in its expression during the reproductive phase of the plant due to altered promoter activity, with possibly effects on cell proliferation in the axillary meristem leading to higher order branching in the CSSL lines.

The *OsFZP* gene plays an important role for the transition from branch to spikelet primordium during panicle development (Komatsu et al., 2001, 2003; Zhu et al., 2003). Reduced expression of *OsFZP* at the reproductive stage increases the extent of higher order branching of the panicle, resulting in increased grain number (Bai et al., 2016). Moreover, recent studies revealed *OsFZP* as the causal gene of QTLs for grain number per panicle (Bai et al., 2017; Fujishiro et al., 2018; Huang et al., 2018; Wang et al., 2020) and showed that the fine-tuning of *OsFZP* expression at the transcriptional and post-translational levels could

determine panicle architecture (Bai et al., 2017; Huang et al., 2018; Chen et al., 2022). Some SNPs and InDel variations between the *Os_Caiapó* and *Og_MG12* genomes were detected within the promoter region of the two *FZP* orthologs. Notably, the *OgFZP* promoter does not possess a 4bp deletion observed in the *Os_Caiapó* genome adjacent to an auxin response element (AuxRE) 2.7 Kbp upstream *OsFZP*. This deletion is known to affect the binding activity of OsARF6 to the promoter of *OsFZP*, leading to decreased accumulation of *OsFZP* mRNAs, in comparison to its *O. rufipogon* Griff. wild-relative ortholog. This reduced amount is associated with an increase in secondary branch number in *O. sativa*, in comparison to *O. rufipogon* (Huang et al., 2018). Consequently, *OgFZP* is more similar to its *O. rufipogon* ortholog in this part of the regulatory region. We also observed a deletion in the CU-rich elements (CUREs) localized in the 3'UTR of *OgFZP* in comparison to its *Og_Caiapó* ortholog. This element is known to be crucial for efficient OsFZP translation repression, via the binding with OsPTB1 and OsPTB2 polypyrimidine trans-binding proteins acting as repressors (Chen et al., 2022). Accordingly, these two variations observed in regulatory regions of *OgFZP* could result in an increase in transcript and/or protein amounts, leading to a precocious transition from branch to spikelet primordium. Such a scenario would be corroborated by the less-branched panicle phenotype of *Og_MG12* compared to *Os_Caiapó* but not by the phenotype of the CSSLs containing *OgFZP*. The analysis of the combinatorial effect of variations in these regulatory elements would be useful to explain the observed phenotypes.

Conclusion

This study was carried out with the broad aim of identifying discrete genetic elements that govern rice panicle architectural diversity using a population of interspecific introgression lines. We detected several QTLs associated with rice yield and panicle branching diversity observed between the two cultivated rice species *O. sativa* and *O. glaberrima*, with decreased values – as expected – in the CSSLs but also unexpectedly with added values for some QTLs compared to the *O. sativa* recurrent parent. This was the case for QTL q_7 on chromosome 7, which causes a global modification of panicle morphology due to two distinct genomic regions: q_7 – 1 and q_7 – 2. For both regions, the *Og_MG12* introgression causes an increase in higher orders of branches and q_7 – 1 is additionally associated with a reduction in PBN. A detailed comparative genomic analysis revealed variations in gene content, promoter region sequence and protein coding sequences for a number of different key genes that act in the control of panicle architecture. When an *Og_MG12* allelic variant is placed in an *Os_Caiapó* genomic background context, it might have an additive effect on panicle branching in the CSSLs, through its addition to the *Os_Caiapó* gene regulatory network (GRN) regulating panicle branching. Moreover, it cannot be ruled out that the higher complexity panicle architecture observed in corresponding CSSLs could result from the combination of these key genes in the *Os_Caiapó* GRN. However, it should be borne in mind that the biological functions and regulation of the genes present in the q_7 – 1 and q_7 – 2 regions have been described in *O. sativa*. The functional analysis of their homologs in *O. glaberrima* has yet to be investigated to confirm similar roles and/or to assess their regulation. Further analysis of the regulation of these genes will be beneficial in improving rice breeding for the exploitation of favorable alleles present in this CSSL population. Some genomic regions associated with multiple traits point to the possibility of enhancing yield potential. The genetic material created here will provide knowledge and resources to facilitate the retention of favorable alleles in breeding programs and to develop new improved rice cultivars using *O. glaberrima* as a genetic resource for enhancing morphological traits in *O. sativa*.

Materials and Methods

Plant Materials

For QTL detection, a set of 60 introgression lines, or Chromosome Segment Substitution Lines (CSSL) was used. The CSSLs were derived from a cross between *O. sativa* subgroup tropical japonica (cv. Caiapó) as the recurrent parent and *O. glaberrima* (cv. MG12; acc. IRGC103544) as the donor parent (Gutiérrez et al., 2010). The CSSLs used were BC₃DH, that is, obtained after three rounds of backcrossing to the recurrent parent followed by double haploidization of BC₃F₁ male gametes (Additional file 1: Table S1). The selection of this set of 60 lines covering the entire donor genome from an initial panel of 312 BC₃DH lines genotyped using 200 simple-sequence repeat (SSR) markers was carried out with the help of the program CSSL Finder v0.9 (<http://mapdisto.free.fr>; Gutiérrez et al., 2010). The effects of *O. glaberrima* introgressions on panicle trait variation were evaluated by choosing five BC₄F_{3:5} lines, each containing the target region and as few introgressed *O. glaberrima* genomic segments in the rest of the genome as possible (Additional file 1: Table S1).

Phenotypic evaluation

The 60 BC₃DH lines, along with the parents (*Os_Caiapó* and *Og_MG12*), were grown together in November 2012-January 2013 (Year 1) in the International Center for Tropical Agriculture (CIAT, now Alliance Bioversity-CIAT) headquarters experimental fields (Palmira, Colombia) (76° 21'W, 3° 30'N and 967masl) under irrigated conditions. The experimental design was a randomized complete block design with two replications and 62 plots. Fives plants per line per plot were grown. Panicle traits were evaluated for subsequent QTL analysis and validation. For each line, the three main panicles from three randomly chosen plants (9 panicles total) per line per replicate were collected. Each panicle was spread out on a white background and held in place with metal pins for photography shooting. A total of 1,443 panicles from the 60 BC₃DH lines and the parents were dissected and scored manually. In January-April 2014 (Year 2), a total of eight BC₃DH lines were re-evaluated using P-TRAP software (Al-Tam et al., 2013), which performs automatic scoring of panicle traits. Five BC₄F₃/F₅ lines with parents, plus L_10 and L_46 BC₃DH, were grown in the greenhouse in Montpellier, France (March 2021) at 28°C / 80% relative humidity under short days conditions (11h light-13h dark), and were scored for panicle traits using P-TRAP. Panicle phenotypes were re-evaluated in the same conditions for three BC₄ lines (June 2021).

For each analysis (2013, 2014 and 2021), 6 morphological panicle traits were scored: rachis length (RL); number of primary branches (PBN); number of secondary branches (SBN); number of tertiary branches (TBN); spikelet number (SpN); and average primary branch length (PBL) per panicle (Additional file 1: Table S2).

Statistical Analysis

Basic statistical analyses were performed for each trait using Analysis of variance (ANOVA) functions in the R software package (version 4.0.4, R Core Team 2022), taking into consideration all factors (replicates, blocks within replicates and lines) as fixed effects. Trait heritability was computed using the line effect based on the variance among phenotypic measurements between the two replicates of the population phenotyping assay. The corplot, ade4 and devtool R packages were used to analyze phenotypic correlation between traits.

The CSSL Finder program was used to display graphical genotypes of the 200 SSRs in the 60 BC₃DH CSSLs together with phenotypic values for each trait. QTLs were detected using the graphical genotyping tool proposed by CSSL Finder, that allows combination of logical genotype-phenotype association with single-marker ANOVA F-test. The F-test was considered significant when its value was higher than 10.0.

Comparison between the CSSLs and the recurrent parent *Os_Caiapó* was performed by a Dunnett's multiple comparison test ($p < 0.05$). The relative effect *RE* of an *O. glaberrima* introgression in a given CSSL was calculated using the least square means (LSMEANS) output of the GLM procedure as follows:

$$RE = 100 \times [LSMean(Lines) - LSMean(Os_Caiapó)] / LSMean(Os_Caiapó)$$

A QTL was taken into consideration when the trait value of a given line was significantly different from that of the parent *Os_Caiapó* ($p < 0.001$) and when at least two lines having overlapping segments shared similar phenotypes. The phenotypic variation of these lines was further evaluated in a second phenotyping experiment in Year 2.

Genome sequencing, assembly, quality assessment and annotation

Samples of young leaves from *O. sativa Caiapó* and *O. glaberrima MG12* were used to extract genomic DNA and for preparation of SKL-LSK109 libraries as described in Serret et al., 2021.

Genome sequencing was performed on a Nanopore MinION Flow Cell R9.4.1 (Oxford technologies, Oxford Science Park, UK). The long-read sequences of *Os_Caiapó* and *Og_MG12* generated by the ONT sequencing were assembled using the following steps. After basecalling and Q > 8 filtering with Guppy V6.1.2, with the SUP model, the preliminary genome was assembled with Flye V2.9 and default assembly parameters for Sup ONT (Kolmogorov et al. 2019). Medaka V1.6.1 was then used to create consensus sequences and to improve the accuracy of the assembly (polishing). The contigs were ordered with RagTag V2.1.0 using *O.sativa cv Nipponbare/IRGSP1.0* as a reference and with default parameters (Alonge et al., 2022). An evaluation of genome assembly completeness was carried out using BUSCO (Benchmark Universal Single Copy Orthologs) v5.2.2 with default parameters and the Poale database (Additional file1: Table S7) (Manni et al., 2021).

The Liftoff tool was used for gene prediction and annotation using the genome reference *O. sativa cv Nipponbare* (IRGSPv1.0, MSU7.0) and associated gff files for both *Os_Caiapó* and *Os_MG12* and by using the *O. glaberrima* genome reference OglRS2 along with its associated gff files (https://ftp.gramene.org/oryza/release-6/gff3/oryza_glaberrima/) in the case of *Og_MG12* (Additional file1: Table S7) (Tranchant-Dubreuil et al., 2023).

Genomic Alignment

The sequences of the QTL regions and the genes of interest were extracted from the three genomes *O. sativa cv. Caiapó*, *O. glaberrima cv. MG12* and *O. sativa cv. Nipponbare* MSU7 using Seqkit version 2.4.0 (Shen et al. 2016). Sequences of QTL regions were aligned and plotted using the minimap2 aligner implemented in the D-GENIES online tool (<https://dgenies.toulouse.inra.fr/run>) using the option Few repeats).

QTL colocalization, identification and bio-analysis of candidate genes

For QTL and gene colocalization, the qTARO database (<http://qtaro.abr.affrc.go.jp>), the funRiceGenes database (<https://funricegenes.github.io/>) (Huang et al. 2022) and various datasets from recently published works were used to identify QTLs and genes that overlapped with the QTL regions detected in this study. To identify genes potentially associated with the QTL regions, we used the *O. sativa cv Nipponbare* MSU7.0 (<https://riceplantbiology.msu.edu>) and the RAP_db (<https://rapdb.dna.affrc.go.jp>) annotation databases. From the annotated gene list, the candidate genes were identified based on their predicted function (biological processes), their referencing in the funRiceGenes database and/or their expression pattern with respect to the trait of interest (Harrop et al., 2016; Harrop et al., 2019). Regions spanning the candidate genes and their associated 5 Kbp upstream region (considered as their promoter regions) were obtained for *O. sativa* (IRGSPv1.0, MSU7.0) and for *O. glaberrima* (OglRS2), then nucleotide and protein alignments were performed using CLUSTALW in order to identify SNPs, InDels and amino acid changes between the two species (Thompson et al. 1994). The Plant Promoter Analysis Navigator web facility (<http://plantpan.itsp.ncku.edu.tw/>) was used to detect transcription factor binding sites (TFBSs) and regulatory elements (CpG islands and tandem repeats), with a cut-off at 0.8, in the promoter region of candidate genes (Chow et al., 2018). Variations between the two genomes, in terms of promoter and gene structures along with SNP and InDel positions and information, were drawn using the R package KaryoploteR (Gel and Serra 2017).

Declarations

Acknowledgments

The authors acknowledge Ndomassi Tando and the ISO 9001 certified IRD itrop HPC (member of the South Green Platform) at IRD Montpellier for providing HPC resources that have contributed to the research results reported within this paper (URL: <https://bioinfo.ird.fr/-http://www.southgreen.fr>). We thank Lady Johanna Arbelaez Rivera (Alliance Bioversity-CIAT), Sophie Chéron (IRD) and Harold Chrestin (IRD) for plant care. We thank Marie-Christine Combes (IRD) and Carole Gauron (IRD) for their help with the panicle phenotyping work. We thank Jean-Francois Rami (Cirad) for his help with R scripts. We also thank Laurence Albar (IRD) for her helpful feedback on the manuscript.

Funding

This research was funded by the Agropolis Foundation through the “Investissements d’avenir” programme (ANR-10-LABX-0001-01), the Fondazione Cariplo (EVOREPRICE 1201-004), and the CGIAR Research program on Rice (RICE CRP).

Availability of data and materials

The CSSLs materials are available from the corresponding authors on reasonable request. Sequencing data generated for this project are available at <https://doi.org/10.23708/QMM2WH> and <https://doi.org/10.23708/1JST4X> for *Os_Caiapó* and *Og_MG12* respectively.

Authors' Contributions: HA, ML designed the project, with input from SJ; HA and ML performed QTL analysis; ML designed the population subset and the markers and supervised field phenotyping; AG performed the SSR analysis on the BC₃DH and BC₄ plants. MC and JS performed the ONT sequencing. MC and FS performed genome assembly and annotation transfer. FN performed the bioinformatic minimap2 alignments. JO developed a Python script for the colocalization analysis. HA, SJ and ML analyzed and interpreted the data. HA wrote the initial draft of the paper. SJ, JT and ML revised the paper. All authors critically reviewed and approved the final manuscript.

Ethics Approval and Consent to Participate

Consent for Publication

Competing Interests

“The authors declare no conflict of interest.”

References

1. Ali ML, Sanchez PL, Yu S, Lorieux M, Eizenga GC (2010) Chromosome Segment Substitution Lines: A Powerful Tool for the Introgression of Valuable Genes from *Oryza* Wild Species into Cultivated Rice (*O. sativa*). *Rice* 3:218–234. <https://doi.org/10.1007/s12284-010-9058-3>
2. Alonge M, Lebeigle L, Kirsche M et al (2022) Automated assembly scaffolding using RagTag elevates a new tomato system for high-throughput genome editing. *Genome Biol* 23:258. <https://doi.org/10.1186/s13059-022-02823-7>
3. Al-Tam FM, Adam H, Anjos AD, Lorieux M, Larmande P, Ghesquière A, Jouannic S, Shahbazkia HR (2013) P-TRAP: a Panicle Trait Phenotyping Tool. *BMC Plant Biol* 13:122. <https://doi.org/10.1186/1471-2229-13-122>
4. Bai S, Hong J, Li L, Su S, Li Z, Wang W, Zhang F, Liang W, Zhang D (2021) Dissection of the Genetic Basis of Rice Panicle Architecture Using a Genome-wide Association Study. *Rice* 14:77. <https://doi.org/10.1186/s12284-021-00520-w>
5. Bai X, Huang Y, Hu Y, Liu H, Zhang B, Smaczniak C, Hu G, Han Z, Xing Y (2017) Duplication of an upstream silencer of FZP increases grain yield in rice. *Nat Plants*. <https://doi.org/10.1038/s41477-017-0042-4>
6. Bai X, Huang Y, Mao D, Wen M, Zhang L, Xing Y (2016a) Regulatory role of FZP in the determination of panicle branching and spikelet formation in rice. *Sci Rep* 6:19022. <https://doi.org/10.1038/srep19022>
7. Bai X, Zhao H, Huang Y, Xie W, Han Z, Zhang B, Guo Z, Yang L, Dong H, Xue W et al (2016b) Genome-Wide Association Analysis Reveals Different Genetic Control in Panicle Architecture Between and Rice. *The Plant Genome* 9:0. <https://doi.org/10.3835/plantgenome2015.11.0115>
8. Balakrishnan D, Surapaneni M, Mesapogu S, Neelamraju S (2019) Development and use of chromosome segment substitution lines as a genetic resource for crop improvement. *Theor Appl Genet* 132:1–25. <https://doi.org/10.1007/s00122-018-3219-y>
9. Beretta VM, Franchini E, Din IU et al (2023) The ALOG family members OsG1L1 and OsG1L2 regulate inflorescence branching in rice. *Plant J*. <https://doi.org/10.1111/tpj.16229>
10. Chen Q, Tian F, Cheng T, Jiang J, Zhu G, Gao Z, Lin H, Hu J, Qian Q, Fang X et al (2022) Translational repression of FZP mediated by CU-rich element/OsPTB interactions modulates panicle development in rice. *Plant J*. <https://doi.org/10.1111/tpj.15737>
11. Cheng S, Huang Y, Zhu N, Zhao Y (2014) The rice WUSCHEL-related homeobox genes are involved in reproductive organ development, hormone signaling and abiotic stress response. *Gene* 549:266–274. <https://doi.org/10.1016/j.gene.2014.08.003>
12. Cheng S, Tan F, Lu Y, Liu X, Li T, Yuan W, Zhao Y, Zhou D-X (2018) WOX11 recruits a histone H3K27me3 demethylase to promote gene expression during shoot development in rice. *Nucleic Acids Res* 46:gky017. <https://doi.org/10.1093/nar/gky017>
13. Cheng S, Zhou D-X, Zhao Y (2015) WUSCHEL-related homeobox gene WOX11 increases rice drought resistance by controlling root hair formation and root system development. *Plant Signal Behav* 11:e1130198. <https://doi.org/10.1080/15592324.2015.1130198>
14. Chongloi GL, Prakash S, Vijayraghavan U (2019) Regulation of meristem maintenance and organ identity during rice reproductive development. *J Exp Bot* 70:1719–1736. <https://doi.org/10.1093/jxb/erz046>
15. Chow C-N, Lee T-Y, Hung Y-C, Li G-Z, Tseng K-C, Liu Y-H, Kuo P-L, Zheng H-Q, Chang W-C (2018) PlantPAN3.0: a new and updated resource for reconstructing transcriptional regulatory networks from ChIP-seq experiments in plants. *Nucleic Acids Res* 47:gky1081. <https://doi.org/10.1093/nar/gky1081>

16. Crowell S, Falcão AX, Shah A, Wilson Z, Greenberg AJ, McCouch SR (2014) High-Resolution Inflorescence Phenotyping Using a Novel Image-Analysis Pipeline, PANorama. *Plant Physiol* 165:479–495. <https://doi.org/10.1104/pp.114.238626>
17. Crowell S, Korniliev P, Falcão A, Ismail A, Gregorio G, Mezey J, McCouch S (2016) Genome-wide association and high-resolution phenotyping link *Oryza sativa* panicle traits to numerous trait-specific QTL clusters. *Nat Commun* 7:10527. <https://doi.org/10.1038/ncomms10527>
18. Cubry P, Pidon H, Ta KN, Tranchant-Dubreuil C, Thuillet A-C, Holzinger M, Adam H, Kam H, Chrestin H, Ghesquière A et al (2020) Genome Wide Association Study Pinpoints Key Agronomic QTLs in African Rice *Oryza glaberrima*. *Rice* 13:66. <https://doi.org/10.1186/s12284-020-00424-1>
19. Cubry P, Tranchant-Dubreuil C, Thuillet A-C, Monat C, Ndjondjop M-N, Labadie K, Cruaud C, Engelen S, Scarcelli N, Rhoné B et al (2018) The Rise and Fall of African Rice Cultivation Revealed by Analysis of 246 New Genomes. *Curr Biol* 28:2274–2282e6. <https://doi.org/10.1016/j.cub.2018.05.066>
20. Cui R, Han J, Zhao S, Su K, Wu F, Du X, Xu Q, Chong K, Theißen G, Meng Z (2010) Functional conservation and diversification of class E floral homeotic genes in rice (*Oryza sativa*). *Plant J* 61:767–781. <https://doi.org/10.1111/j.1365-3113x.2009.04101.x>
21. Eshed Y, Zamir D (1995) An introgression line population of *Lycopersicon pennellii* in the cultivated tomato enables the identification and fine mapping of yield-associated QTL. *Genetics* 141:1147–1162. <https://doi.org/10.1093/genetics/141.3.1147>
22. Fornara F, Parenicová L, Falasca G, Pelucchi N, Masiero S, Ciannanea S, Lopez-Dee Z, Altamura MM, Colombo L, Kater MM (2004) Functional characterization of OsMADS18, a member of the AP1/SQUA subfamily of MADS box genes. *Plant Physiol* 135:2207–2219. <https://doi.org/10.1104/pp.104.045039>
23. Fujishiro Y, Agata A, Ota S et al (2018) Comprehensive panicle phenotyping reveals that qSm7/FZP influences higher-order branching. *Sci Rep-uk* 8:12511. <https://doi.org/10.1038/s41598-018-30395-9>
24. Gel B, Serra E (2017) karyoploteR: an R/Bioconductor package to plot customizable genomes displaying arbitrary data. *Bioinformatics* 33:3088–3090. <https://doi.org/10.1093/bioinformatics/btx346>
25. Gutiérrez AG, Carabalí SJ, Giraldo OX, Martínez CP, Correa F, Prado G, Tohme J, Lorieux M (2010) Identification of a Rice stripe necrosis virus resistance locus and yield component QTLs using *Oryza sativa* x *O. glaberrima* introgression lines. *Bmc Plant Biol* 10:6. <https://doi.org/10.1186/1471-2229-10-6>
26. Harrop TWR, Din IU, Gregis V, Osnato M, Jouannic S, Adam H, Kater MM (2016) Gene expression profiling of reproductive meristem types in early rice inflorescences by laser microdissection. *Plant J* 86:75–88. <https://doi.org/10.1111/tjp.13147>
27. Harrop TWR, Mantegazza O, Luong AM, Béthune K, Lorieux M, Jouannic S, Adam H (2019) A set of AP2-like genes is associated with inflorescence branching and architecture in domesticated rice. *J Exp Bot* 70:5617–5629. <https://doi.org/10.1093/jxb/erz340>
28. Hill MS, Zande PV, Wittkopp PJ (2021) Molecular and evolutionary processes generating variation in gene expression. *Nat Rev Genet* 22:203–215. <https://doi.org/10.1038/s41576-020-00304-w>
29. Huang Y, Zhao S, Fu Y, Sun H, Ma X, Tan L, Liu F, Sun X, Sun H, Gu P et al (2018) Variation in the regulatory region of FZP causes increases in secondary inflorescence branching and grain yield in rice domestication. *Plant J* 96:716–733. <https://doi.org/10.1111/tjp.14062>
30. Huang F, Jiang Y, Chen T et al (2022) New Data and New Features of the FunRiceGenes (Functionally Characterized Rice Genes) Database: 2021 Update. *Rice* 15:23. <https://doi.org/10.1186/s12284-022-00569-1>
31. Itoh JI, Nonomura KI, IKEDA K, Yamaki S, Inukai Y, Yamagishi H, Kitano H, NAGATO Y (2005) Rice plant development: from zygote to spikelet. *Plant Cell Physiol* 46:23–47. <https://doi.org/10.1093/pcp/pci501>
32. Jeon J-S, Jang S, Lee S, Nam J, Kim C, Lee S-H, Chung Y-Y, Kim S-R, Lee YH, Cho Y-G et al (2000) leafy hull sterile1 Is a Homeotic Mutation in a Rice MADS Box Gene Affecting Rice Flower Development. *Plant Cell* 12:871–884. <https://doi.org/10.1105/tpc.12.6.871>
33. Jones MP, Dingkuhn M, Aluko/snm > GK, Semon M (1997) Interspecific *Oryza Sativa* L. X *O. Glaberrima* Steud. progenies in upland rice improvement. *Euphytica* 94:237–246. <https://doi.org/10.1023/A:1002969932224>
34. Kaneko-Suzuki M, Kurihara-Ishikawa R, Okushita-Terakawa C, Kojima C, Nagano-Fujiwara M, Ohki I, Tsuji H, Shimamoto K, Taoka K (2018) TFL1-Like Proteins in Rice Antagonize Rice FT-Like Protein in Inflorescence Development by Competition for Complex Formation with 14-3-3 and FD. *Plant Cell Physiol* 59:458–468. <https://doi.org/10.1093/pcp/pcy021>
35. Khanday I, Yadav SR, Vijayraghavan U (2013) Rice LHS1/OsMADS1 controls floret meristem specification by coordinated regulation of transcription factors and hormone signaling pathways. *Plant Physiol* 161:1970–1983. <https://doi.org/10.1104/pp.112.212423>
36. Kobayashi K, Maekawa M, Miyao A, Hirochika H, Kyoizuka J (2009) PANICLE PHYTOMER2 (PAP2), encoding a SEPALLATA subfamily MADS-box protein, positively controls spikelet meristem identity in rice. *Plant Cell Physiol* 51:47–57. <https://doi.org/10.1093/pcp/pcp166>
37. Kobayashi K, Yasuno N, Sato Y, Yoda M, Yamazaki R, Kimizu M, Yoshida H, Nagamura Y, Kyoizuka J (2012) Inflorescence Meristem Identity in Rice Is Specified by Overlapping Functions of Three AP1/FUL-Like MADS Box Genes and PAP2, a SEPALLATA MADS Box Gene. *THE PLANT CELL ONLINE* 24:1848–1859. <https://doi.org/10.1105/tpc.112.097105>
38. Kolmogorov M, Yuan J, Lin Y, Pevzner PA (2019) Assembly of long, error-prone reads using repeat graphs. *Nat Biotechnol* 37:540–546. <https://doi.org/10.1038/s41587-019-0072-8>
39. Komatsu KK, Maekawa MM, Ujii SS, Satake YY, Furutani II, Okamoto HH, Shimamoto KK, Kyoizuka JJ (2003a) LAX and SPA: major regulators of shoot branching in rice. *Proc Natl Acad Sci USA* 100:11765–11770. <https://doi.org/10.1073/pnas.1932414100>
40. Komatsu M, Chujo A, Nagato Y, Shimamoto K, Kyoizuka J (2003b) FRIZZY PANICLE is required to prevent the formation of axillary meristems and to establish floral meristem identity in rice spikelets. *Development* 130:3841–3850. <https://doi.org/10.1242/dev.00564>
41. Komatsu M, Maekawa M, Shimamoto K, Kyoizuka J (2001) The LAX1 and FRIZZY PANICLE 2 Genes Determine the Inflorescence Architecture of Rice by Controlling Rachis-Branch and Spikelet Development. *Dev Biol* 231:10–10. <https://doi.org/10.1006/dbio.2000.9988>

42. Koumproglou R, Wilkes TM, Townson P, Wang XY, Beynon J, Pooni HS, Newbury HJ, Kearsey MJ (2002) STAIRS: a new genetic resource for functional genomic studies of Arabidopsis. *Plant J* 31:355–364. <https://doi.org/10.1046/j.1365-313x.2002.01353.x>
43. Li F, Liu W, Tang J, Chen J, Tong H, Hu B, Li C, Fang J, Chen M, Chu C (2010) Rice DENSE AND ERECT PANICLE 2 is essential for determining panicle outgrowth and elongation. *Cell Res* 20:838–849. <https://doi.org/10.1038/cr.2010.69>
44. Li X-M, Chao D-Y, Wu Y, Huang X, Chen K, Cui L-G, Su L, Ye W-W, Chen H, Chen H-C et al (2015) Natural alleles of a proteasome $\alpha 2$ subunit gene contribute to thermotolerance and adaptation of African rice. *Nat Genet* 47:827–833. <https://doi.org/10.1038/ng.3305>
45. Linares OF (2002) African rice (*Oryza glaberrima*): history and future potential. *Proc Natl Acad Sci USA* 99:16360–16365. <https://doi.org/10.1073/pnas.252604599>
46. Lorieux M (2012) MapDisto: fast and efficient computation of genetic linkage maps. *Mol Breed* 30:1231–1235. <https://doi.org/10.1007/s11032-012-9706-y>
47. Lorieux M, Garavito A, Bouniol J, Gutiérrez A, Ndjiondjop M-N, Guyo R, Martinez CP, Tohme J, A Ghesquière (2013) Unlocking the *O. glaberrima* treasure for rice breeding in Africa. « Realizing Africa's Rice Promise ». Marco Wopereis and David Johnson, Eds. CABI, London
48. Mao C, He J, Liu L, Deng Q, Yao X, Liu C, Qiao Y, Li P, Ming F (2020) OsNAC2 integrates auxin and cytokinin pathways to modulate rice root development. *Plant Biotechnol J* 18:429–442. <https://doi.org/10.1111/pbi.13209>
49. Manni M, Berkeley MR, Seppey M, Zdobnov EM (2021) BUSCO: Assessing Genomic Data Quality and Beyond. *Curr Protoc* 1:e323. <https://doi.org/10.1002/cpz1.323>
50. Marais GABG, Nicolas MM, Bergero RR, Chambrier PP, Kejnovsky EE, Monéger FF, Hobza RR, Widmer AA, Charlesworth DD (2008) Evidence for Degeneration of the Y Chromosome in the Dioecious Plant *Silene latifolia*. *Curr Biol* 18:5–5. <https://doi.org/10.1016/j.cub.2008.03.023>
51. Masiero S, Imbriano C, Ravasio F, Favaro R, Pelucchi N, Gorla MS, Mantovani R, Colombo L, Kater MM (2002) Ternary Complex Formation between MADS-box Transcription Factors and the Histone Fold Protein NF-YB*. *J Biol Chem* 277:26429–26435. <https://doi.org/10.1074/jbc.m202546200>
52. Nakagawa M, Shimamoto K, Kyozuka J (2002) Overexpression of RCN1 and RCN2, rice TERMINAL FLOWER 1/CENTRORADIALIS homologs, confers delay of phase transition and altered panicle morphology in rice. *Plant J* 29:743–750. <https://doi.org/10.1046/j.1365-313x.2002.01255.x>
53. Neogy A, Garg T, Kumar A, Dwivedi AK, Singh H, Singh U, Singh Z, Prasad K, Jain M, Yadav SR (2019) Genome-Wide Transcript Profiling Reveals an Auxin-Responsive Transcription Factor, OsAP2/ERF-40, Promoting Rice Adventitious Root Development. *Plant Cell Physiol* 60:2343–2355. <https://doi.org/10.1093/pcp/pcz132>
54. Ohmori S, Kimizu M, Sugita M, Miyao A, Hirochika H, Uchida E, Nagato Y, Yoshida H (2009) MOSAIC FLORAL ORGANS1, an AGL6-like MADS box gene, regulates floral organ identity and meristem fate in rice. *Plant Cell* 21:3008–3025. <https://doi.org/10.1105/tpc.109.068742>
55. Prasad K, Parameswaran S, Vijayraghavan U (2005) OsMADS1, a rice MADS-box factor, controls differentiation of specific cell types in the lemma and palea and is an early-acting regulator of inner floral organs. *Plant J* 43:915–928. <https://doi.org/10.1111/j.1365-313x.2005.02504.x>
56. Rebolledo MC, Peña AL, Duitama J, Cruz DF, Dingkuhn M, Grenier C, Tohme J (2016) Combining Image Analysis, Genome Wide Association Studies and Different Field Trials to Reveal Stable Genetic Regions Related to Panicle Architecture and the Number of Spikelets per Panicle in Rice. *Front Plant Sci* 7:1384. <https://doi.org/10.3389/fpls.2016.01384>
57. Ren D, Li Y, Zhao F, Sang X, Shi J, Wang N, Guo S, Ling Y, Zhang C, Yang Z et al (2013) MULTI-FLORET SPIKELET1, Which Encodes an AP2/ERF Protein, Determines Spikelet Meristem Fate and Sterile Lemma Identity in Rice. *Plant Physiol* 162:872–884. <https://doi.org/10.1104/pp.113.216044>
58. Serret J, Mariac C, Albar L, Sabot F (2021) From low cost plant HMW DNA extraction to MinION. <https://doi.org/10.17504/protocols.io.bu3vny6>. sequencing v1
59. Shao J, Haider I, Xiong L, Zhu X, Hussain RMF, Overnäs E, Meijer AH, Zhang G, Wang M, Bouwmeester HJ et al (2018) Functional analysis of the HD-Zip transcription factor genes Oshox12 and Oshox14 in rice. *PLoS ONE* 13:e0199248. <https://doi.org/10.1371/journal.pone.0199248>
60. Shen W, Le S, Li Y, Hu F (2016) SeqKit: A Cross-Platform and Ultrafast Toolkit for FASTA/Q File Manipulation. *PLoS ONE* 11:e0163962. <https://doi.org/10.1371/journal.pone.0163962>
61. Suzuki T, Toriba T, Fujimoto M, Tsutsumi N, Kitano H, Hirano H-Y (2006) Conservation and diversification of meristem maintenance mechanism in *Oryza sativa*: Function of the FLORAL ORGAN NUMBER2 gene. *Plant Cell Physiol* 47:1591–1602. <https://doi.org/10.1093/pcp/pcl025>
62. Ta KN, Khong NG, HA TL, Nguyen DT, Mai DC, Hoang TG, Phung TPN, Bourrié I, Courtois B, Tran TTH, Dinh BY, Tuan N, Do NV, Lebrun M, Gantet P, Jouannic S (2018) A genome-wide association study using a Vietnamese landrace panel of rice (*Oryza sativa*) reveals new QTLs controlling panicle morphological traits. *BMC Plant Biol* 18:282. <https://doi.org/10.1186/s13227-017-0065-y>
63. Tao J, Liang W, An G, Zhang D (2018) OsMADS6 controls flower development by activating rice FACTOR OF DNA METHYLATION LIKE 1. <https://doi.org/10.1104/pp.18.00017>. PLANT PHYSIOLOGY
64. Thiémlé D, Boissard A, Ndjiondjop M-N, Chéron S, Sére Y, Aké S, Ghesquière A, Albar L (2010) Identification of a second major resistance gene to Rice yellow mottle virus, RYMV2, in the African cultivated rice species, *O. glaberrima*. *Theor Appl Genet* 121:169–179. <https://doi.org/10.1007/s00122-010-1300-2>
65. Thompson JD, Higgins DG, Gibson TJ (1994) CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Res* 22:4673–4680. <https://doi.org/10.1093/nar/22.22.4673>
66. Tranchand-Dubreuil C, Blaison M, Chenal C, Albar L, Klein V, Vigouroux Y, Sabot F (2023) FrangiPANE, a tool for creating a panreference using left behind reads. *Nar Genom Bioinform* 5:lqad013. <https://doi.org/10.1093/nargab/lqad013>
67. Wambugu PW, Furtado A, Waters DL, Nyamongo DO, Henry RJ (2013) Conservation and utilization of African *Oryza* genetic resources. *Rice* 6:29. <https://doi.org/10.1186/1939-8433-6-29>

68. Wambugu PW, Ndjiondjop M-N, Henry R (2021) Genetics and Genomics of African Rice (*Oryza glaberrima* Steud) Domestication. *Rice* (N Y) 14:6–14. <https://doi.org/10.1186/s12284-020-00449-6>
69. Wang C, Yang X, Li G (2021a) Molecular Insights into Inflorescence Meristem Specification for Yield Potential in Cereal Crops. *Int J Mol Sci* 22:3508. <https://doi.org/10.3390/ijms22073508>
70. Wang J, Bao J, Zhou B, Li M, Li X, Jin J (2021b) The osa-miR164 target OsCUC1 functions redundantly with OsCUC3 in controlling rice meristem/organ boundary specification. *New Phytol* 229:1566–1581. <https://doi.org/10.1111/nph.16939>
71. Wang S-S, Chung C-L, Chen K-Y, Chen R-K, Genetics (2020) <https://doi.org/10.1534/genetics.119.302862>
72. Wray GA (2007) The evolutionary significance of cis-regulatory mutations. *Nat Rev Genet* 8:206–216. <https://doi.org/10.1038/nrg2063>
73. Xing Y, Tan Y, Hua J, Sun X, Xu C, Zhang Q (2002) Characterization of the main effects, epistatic effects and their environmental interactions of QTLs on the genetic basis of yield traits in rice. *Theor Appl Genet* 105:248–257. <https://doi.org/10.1007/s00122-002-0952-y>
74. Xing Y, Zhang Q (2010) Genetic and molecular bases of rice yield. *Annu Rev Plant Biol* 61:421–442. <https://doi.org/10.1146/annurev-arplant-042809-112209>
75. Ya-fang Z, Yu-yin M, Zong-xiang C, Jie Z, Tian-xiao C, Qian-qian L, Xue-biao P, Shi-min Z (2015) Genome-Wide Association Studies Reveal New Genetic Targets for Five Panicle Traits of International Rice Varieties. *Rice Sci* 22:217–226. <https://doi.org/10.1016/j.rsci.2015.07.001>
76. Yin C, Zhu Y, Li X, Lin Y (2021) Molecular and Genetic Aspects of Grain Number Determination in Rice (*Oryza sativa* L). *Int J Mol Sci* 22:728. <https://doi.org/10.1038/s41467-017-01501-8>
77. Yoshida A, Sasao M, Yasuno N, Takagi K, Daimon Y, Chen R, Yamazaki R, Tokunaga H, Kitaguchi Y, Sato Y et al (2013) TAWAWA1, a regulator of rice inflorescence architecture, functions through the suppression of meristem phase transition. *Proc Natl Acad Sci* 110:767–772. <https://doi.org/10.1073/pnas.1216151110>
78. Zhang B, Shang L, Ruan B, Zhang A, Yang S, Jiang H, Liu C, Hong K, Lin H, Gao Z et al (2019) Development of Three Sets of High-Throughput Genotyped Rice Chromosome Segment Substitution Lines and QTL Mapping for Eleven Traits. *Rice* 12:33. <https://doi.org/10.1186/s12284-019-0293-y>
79. Zhang S, Hu W, Wang L, Lin C, Cong B, Sun C, Luo D (2005) TFL1/CEN-like genes control intercalary meristem activity and phase transition in rice. *Plant Sci* 168:1393–1408. <https://doi.org/10.1016/j.plantsci.2004.10.022>
80. Zhang T, Li R, Xing J, Yan L, Wang R, Zhao Y (2018) The YUCCA-Auxin-WOX11 Module Controls Crown Root Development in Rice. *Front Plant Sci* 9:523. <https://doi.org/10.3389/fpls.2018.00523>
81. Zhang Y, Zhou J, Xu P, Li J, Deng X, Deng W, Yang Y, Yu Y, Pu Q, Tao D (2022) A Genetic Resource for Rice Improvement: Introgression Library of Agronomic Traits for All AA Genome *Oryza* Species. *Front Plant Sci* 13:856514. <https://doi.org/10.3389/fpls.2022.856514>
82. Zhao Y, Cheng S, Song Y, Huang Y, Zhou S, Liu X, Zhou D-X (2015) The Interaction between Rice ERF3 and WOX11 Promotes Crown Root Development by Regulating Gene Expression Involved in Cytokinin Signaling. *THE PLANT CELL ONLINE* 27:2469–2483. <https://doi.org/10.1105/tpc.15.00227>
83. Zhao Y, Hu Y, Dai M, Huang L, Zhou D-X (2009) The WUSCHEL-related homeobox gene WOX11 is required to activate shoot-borne crown root development in rice. *Plant Cell* 21:736–748. <https://doi.org/10.1105/tpc.108.061655>
84. Zhong H, Liu S, Meng X, Sun T, Deng Y, Kong W, Peng Z, Li Y (2021) Uncovering the genetic mechanisms regulating panicle architecture in rice with GPWAS and GWAS. *BMC Genomics* 22:86. <https://doi.org/10.1186/s12864-021-07391-x>
85. Zhou S, Jiang W, Long F, Cheng S, Yang W, Zhao Y, Zhou D-X (2017) Rice Homeodomain Protein WOX11 Recruits a Histone Acetyltransferase Complex to Establish Programs of Cell Proliferation of Crown Root Meristem. *Plant Cell* 29:1088–1104. <https://doi.org/10.1105/tpc.16.00908>
86. Zhu Q-H, Hoque MS, Dennis ES, Upadhyaya NM (2003) Ds tagging of BRANCHED FLORETLESS 1 (BFL1) that mediates the transition from spikelet to floret meristem in rice (*Oryza sativa* L). *BMC Plant Biol* 3:6. <https://doi.org/10.1186/1471-2229-3-6>
87. Zhu W, Yang L, Wu D, Meng Q, Deng X, Huang G, Zhang J, Chen X, Ferrándiz C, Liang W et al (2021) Rice SEPALLATA genes OsMADS5 and OsMADS34 cooperate to limit inflorescence branching by repressing the TERMINAL FLOWER1-like gene RCN4. *New Phytol*. <https://doi.org/10.1111/nph.17855>

Figures

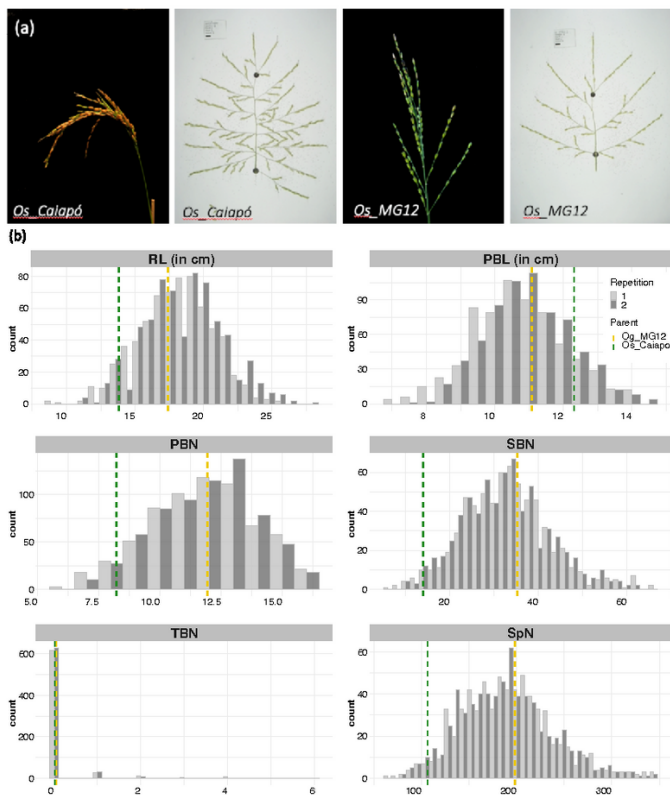


Fig.1 Panicle structure and morphological panicle trait values in the two parental lines and the 60 BC3DH CSSLs. **(a)** Panicle Architecture contrasts between the two parental lines (*Os_Caiapó* and *Os_MG12*). **(b)** Distribution of panicle trait values in the 60 BC3DH CSSLs. Dashed vertical lines correspond to the mean values of each parent (green for *Os_MG12* and yellow for *Os_Caiapó*). Values for repetitions 1 and 2 are shown in light gray and dark gray, respectively. Abbreviations: RL, Rachis Length; PBN, Primary Branch Number; PBL, Primary Branch Length; SBN, Secondary Branch Number; TBN, Tertiary Branch Number; SpN, Spikelet Number.

Figure 1

See image above for figure legend

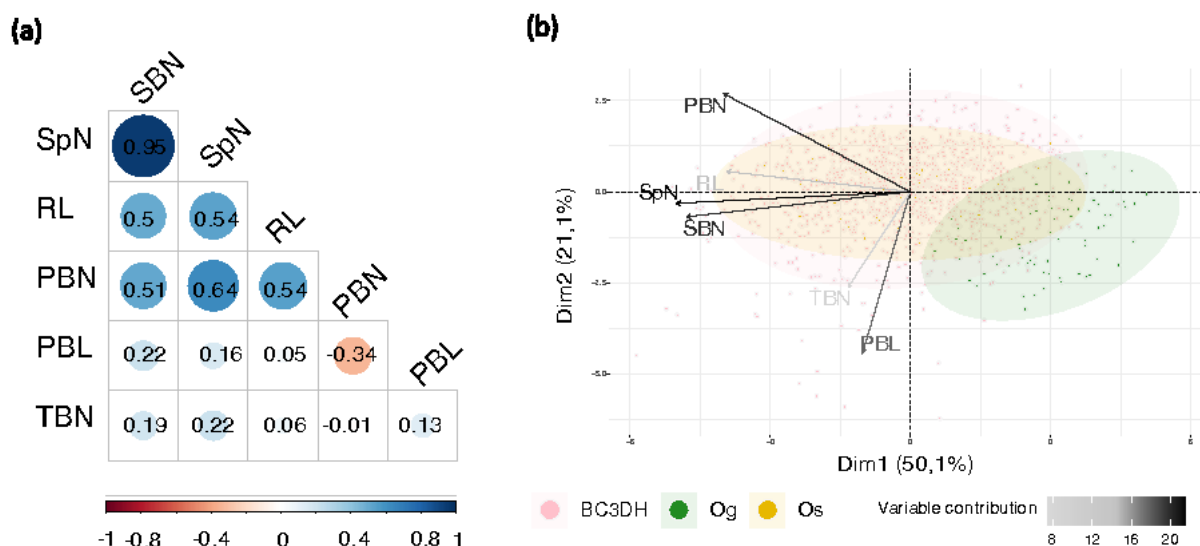


Fig. 2 Description of the morphological panicle traits observed in the BC3DH CSSL lines and parents. **(a)** Correlations between panicle traits. **(b)** Principal component analysis with traits contributions and individual panicle distributions. Abbreviations: RL, Rachis Length; PBN, Primary Branch Number; PBL, Primary Branch Length; SBN, Secondary Branch Number; TBN, Tertiary Branch Number; SpN, Spikelet Number.

Figure 2

See image above for figure legend

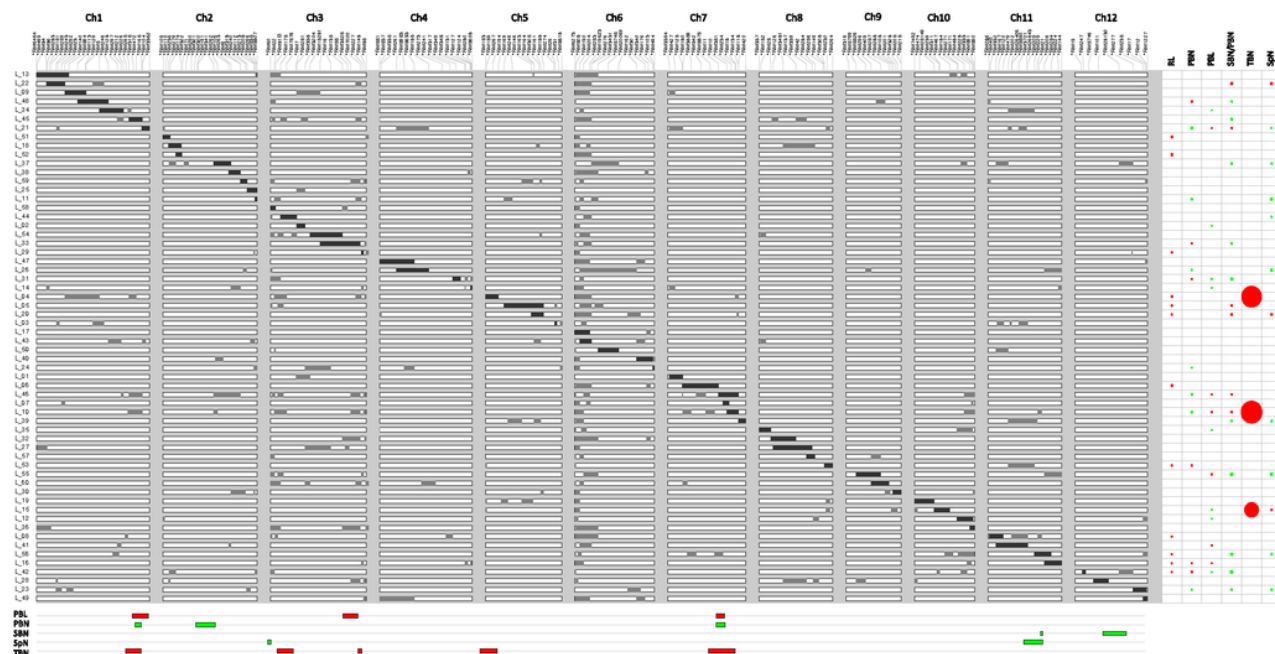


Fig. 3 Graphic representation of the genotypes of the 60 BC3DH CSSLs, showing line x trait significant associations and QTL positions. The 12 chromosomes are covered by 200 evenly dispersed SSR markers. Genotypes are displayed horizontally. Black areas represent the *O. glaberrima* MG12 target chromosome segments, the set of segments broadly covering the entire *O. glaberrima* MG12 genome. White areas represent the *O. sativa* Calapó genetic background and grey areas represent *O. glaberrima* additional chromosomal segments. Phenotypic effects of CSSL lines are represented in relative terms as circles on the right of the figure. The area of each circle is proportional to its relative effect. Horizontal bars at the bottom of the figure indicate QTL positions deduced from analyses performed using CSSL finder software. The colors of the circles and bars represent the effect as follows: red, increasing effect compared to the recurrent parent *O. sativa* Calapó; green, decreasing effect.

Figure 3

See image above for figure legend

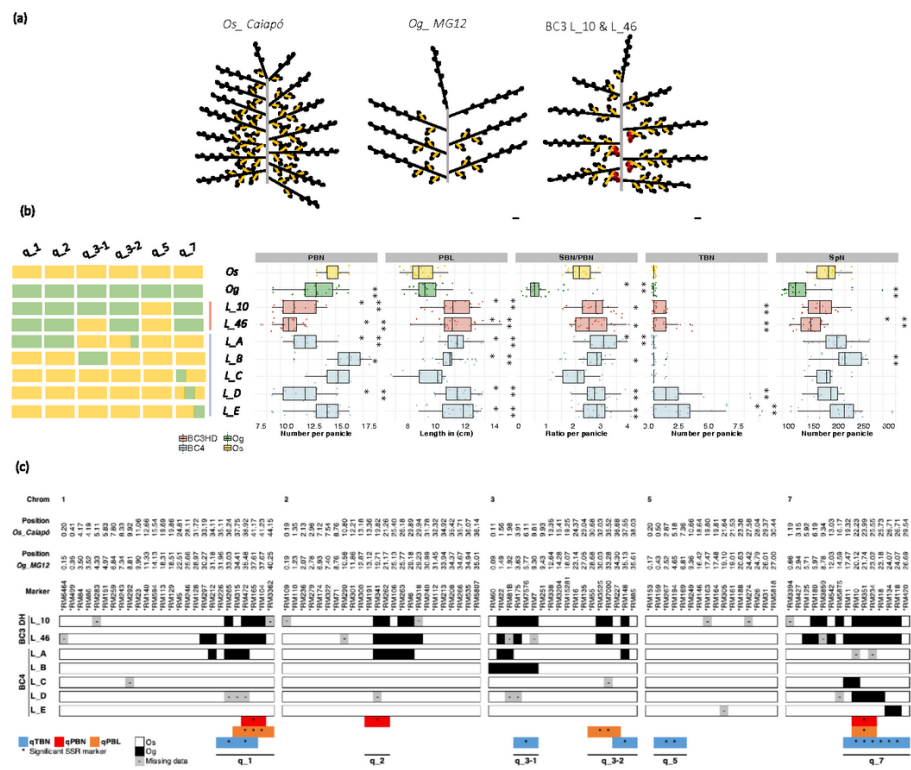


Fig. 4 Phenotypic and genetic description of BC3DH and BC4 lines to dissect the effects of QTLs on panicle traits. **(a)** Schematic representation of the panicle structure observed in *Os_Caiapo*, *Og_MG12* and BC3DH_L10 and L_46, indicating primary branch (PB) in black, secondary branch (SB) in yellow, tertiary branch (TB) in red and spikelet (Sp) in green. Scale bar = 1cm. **(b)** Boxplots of the phenotypic variation observed in *Os_Caiapo* (yellow), *Og_MG12* (green), BC3DH (red) and BC4 (blue) lines. Each point represents the phenotypic value for one panicle. Statistical significance (t-test *p*-values) between *Os_Caiapo* and each line for the panicle morphological traits is indicated as follows: ** *p*-values < 0.01; *** < 0.001. The left hand panel represents the allelic status (*Os_Caiapo* in yellow or *Og_MG12* in green) for each QTL region in each line. **(c)** Visualization of *Og_MG12* segments present in chromosomes 1, 3, 5 and 7 in BC3DH and BC4 lines to dissect the effects of QTLs on panicle traits. The positions of QTLs associated with PBN, PBL and TBN traits are indicated immediately above them (bars shaded in orange, red and blue respectively). Abbreviations: PBN, primary branch number; PBL, primary branch length; TBN, tertiary branch number; SBN/PBN, ratio between secondary branch and primary branch numbers; SpN, spikelet number.

Figure 4

See image above for figure legend

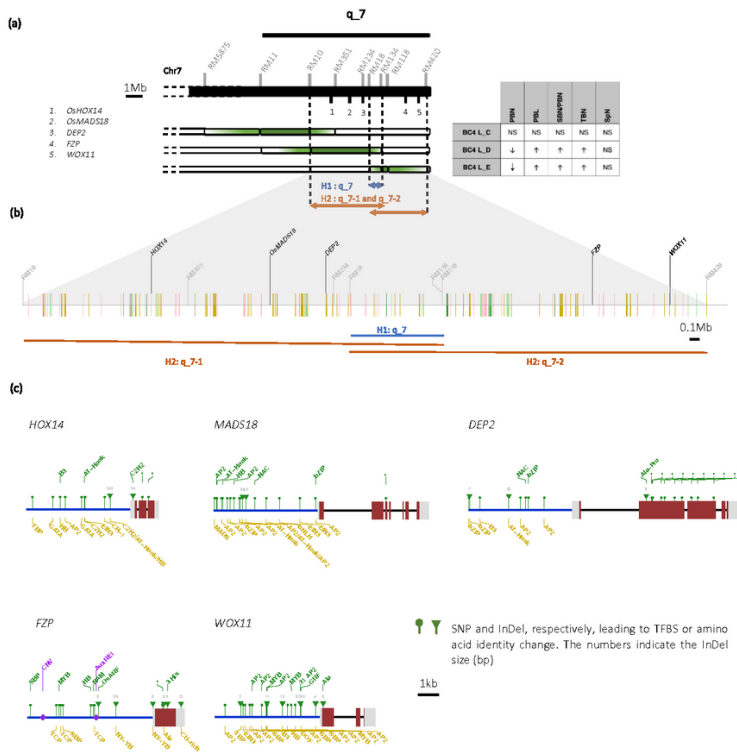


Fig. 5 Description of genetic variations observed in the q₇ region between the *Os_Calipso* and *Oq_MG12* genomes. **(a)** Influence of the q₇ region on panicle variation in the BC₄ lines. Light green and green shading indicate the “extended” and strict introgression positions respectively for each BC₄ line. **(b)** Representation of variation observed in the annotated genes between the *Os_Calipso* and *Oq_MG12* genomes in the region between the RM10 and RM420 SSR markers. Yellow bars show loci only present in *Os_Calipso* genome, green bars indicate loci only present in *Oq_MG12*. Light green bars represent genes that are duplicated in *Oq_MG12* and pink bars represent genes that are annotated in *Oq_MG12* but present in a different genomic region in *Os_Calipso*. **(c)** Schematic representation of sequence divergence in candidate genes between the 2 genomes. Blue lines correspond to promoter regions, red and grey boxes represent exons and introns respectively. SNP and InDel variations leading to TFBS or amino acid changes are represented by dots and triangles respectively. The number above a triangle indicates InDel size. Asterisks indicate amino acid identity modification in the protein sequence. TFBSs present only in *Os_Calipso* or *Oq_MG12* are shown colored in yellow and green respectively. Abbreviations: TFBS, Transcription Factor Binding Site; Ala, alanine; Pro, proline; His, histidine; EIN3, Ethylene-Insensitive-Like3; C2H2, C2H2-Type Zinc finger; AP2, AP2-ERF; bHLH, basic helix-loop-helix; GRF, Growth-regulating factor; TCP, Teosinte-Branched1; SBN, Squamosa Binding Protein; NY-YB, nuclear factor Y; HB, Homeodomain; MADS, MADS-BOX; bZIP, basic leucine zipper; TBP, TATA-box-binding.

Figure 5

See image above for figure legend

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [AdditionalFile1TableS.xlsx](#)
- [AdditionalFile26Figures.pdf](#)