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Developing genome-wide microsatellite markers of bamboo and their applications on molecular marker assisted taxonomy for accessions in the genus *Phyllostachys*

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Morphology-based taxonomy via exiguously reproductive organ has severely limitation on bamboo taxonomy, mainly owing to infrequent and unpredictable flowering events of bamboo. Here, we present the first genome-wide analysis and application of microsatellites based on the genome of moso bamboo (*Phyllostachys edulis*) to assist bamboo taxonomy. Of identified 127,593 microsatellite repeat-motifs, the primers of 1,451 microsatellites were designed and 1,098 markers were physically mapped on the genome of moso bamboo. A total of 917 markers were successfully validated in 9 accessions with ~39.8% polymorphic potential. Retrieved from validated microsatellite markers, 23 markers were selected for polymorphic analysis among 78 accessions and 64 alleles were detected with an average of 2.78 alleles per primers. The cluster result indicated the majority of the accessions were consistent with their current taxonomic classification, confirming the suitability and effectiveness of the developed microsatellite markers. The variations of microsatellite marker in different species were confirmed by sequencing and *in silico* comparative genome mapping were investigated. Lastly, a bamboo microsatellites database (<http://www.bamboogdb.org/ssr>) was implemented to browse and search large information of bamboo microsatellites. Consequently, our results of microsatellite marker development are valuable for assisting bamboo taxonomy and investigating genomic studies in bamboo and related grass species.

As one of the most important non-timber forest resources and fastest-growing plant in the world, bamboos belong to a group of flowering and evergreen plants, classified in the grass family Poaceae, subfamily Bambusoideae, tribe Bambuseae. Because of having a strong carbon fixation capability and a high strength-to-weight ratio, like natural woody, bamboo had received much attention in the ecological and economic aspects. More than 1,250 species, belonging to 75 genera, are reported worldwide¹.

The vegetative features are quite essential for identification of the bamboo members. However, as one of important and special characteristics in bamboo, infrequent and unpredictable flowering events, such as a long flowering cycle of up to 120 years and flowering once before culm death, severely restricts the opportunity to study fresh reproductive materials. Even if their herbarium samples were available, enough morphological resolution was lack, causing confusion in the real field condition². Therefore, traditional morphological taxonomy based on reproductive characteristics has to face great difficulties in bamboo. On the other hand, currently morphology-based taxonomy for bamboo, mainly depending on vegetative characteristics such as culm or culm-sheath, is hard to produce a confirmed taxonomic demarcation and approach for bamboo. Because some variations may be induced in culm, leaf or rhizome of bamboo during different development stages or under different environments. In particular, with the increasing number of identified bamboo species, a few of confused phenomena based on morphological taxonomy were emerged, such as homonym and synonym^{3,4}.

To address this issue properly and assist bamboo taxonomy, considerable progress had been made in the field of modern molecular techniques⁵⁻¹⁰. As one of the most powerful genetic markers in biology, simple sequence



Table 1 | Distribution of microsatellites involved genome size, microsatellite frequency and microsatellite type in six surveyed plant species

Plant species	Genome size (Mb)	SSR/Mb	Identified SSRs			
			Single SSRs (2–6 bp repeat motif)		Compound SSRs	
			Number	%	Number	%
<i>Phyllostachys edulis</i>	2051.7	62.2	106,582	83.53	21,011	16.47
<i>Zea mays</i>	2066.4	52.1	94,683	87.88	13,059	12.12
<i>Oryza sativa</i>	374.5	165.5	49,505	79.88	12,472	20.12
<i>Sorghum bicolor</i>	738.5	91.1	57,016	84.70	10,295	15.30
<i>Brachypodium distachyon</i>	271.9	76.5	17,865	85.93	2,924	14.07
<i>Arabidopsis thaliana</i>	119.7	135.5	12,259	75.59	3,958	24.41

repeats (SSRs), also known as microsatellites, refer to DNA sequences of 1–6 bp in length that are tandem repeated a variable number of times¹¹. Generally, the repeat-motifs with more than mono-nucleotide were selected to develop molecular markers^{12,13}. Compared with other molecular marker, the nature of microsatellites provides a number of advantages such as simple PCR screening, co-dominant, relatively abundant and inherent potential for variation. Therefore, microsatellites were considered to be better suited for construction of high-density linkage maps¹⁴, QTL mapping¹⁵, genetic diversity analysis¹⁶, cultivar identification¹⁷ and marker-assisted selection¹⁸. Meanwhile, microsatellites with a high degree of transferability among species have been widely utilized for comparative genetics and taxonomy research^{19,20}.

Nevertheless, traditional development of microsatellite markers is a time-consuming, money-cost process and limited utilization²¹. The previous results^{22,23} demonstrated that a less number of bamboo microsatellites markers were identified with few polymorphism, due to lacking of a large number of sequence data. Therefore, a genome-wide analysis of bamboo microsatellite on the basis of the draft genome of moso bamboo (*Phyllostachys edulis*) was performed²⁴, to effectively expand our understanding and widely detect polymorphism of microsatellites in bamboo. Thus, we had identified the whole genome-based microsatellite loci and developed some microsatellite markers. Moreover, a selected set of validated and physically mapped markers were used in evaluating polymorphic and diversity potential in 78 accessions belonged to the genus *Phyllostachys*, which is one of the most important genus of bamboo including more than 50 species and accounting for 90% of commercial bamboos in the Bambusoideae. Genetic relationship was revealed among the accessions of *Phyllostachys* by cluster analysis to identify the tallies with morphological taxonomy. Furthermore, the genetic basis of sequence length variation by sequencing and *in silico* comparative genome mapping were investigated as well. Lastly, the module of bamboo microsatellite database was implemented in order to provide worldwide researchers with a diverse and powerful search and browse for bamboo microsatellite information.

Results

Microsatellites frequency analysis. Owing to mono-nucleotide repeat-motif having homo-polymorphism, it was rarely employed as microsatellite markers based on polymorphism analysis^{12,13}. Thus, we identified a total of 127,593 microsatellites with more than mono-nucleotide repeat-motif in moso bamboo genome based on bioinformatics analysis (Table 1). Besides the genome of moso bamboo, a genome-wide investigation of microsatellite based on the 5 important plants (*Zea mays*, *Oryza sativa*, *Sorghum bicolor*, *Brachypodium distachyon* and *Arabidopsis thaliana*) were performed by utilizing their latest genomic sequences and annotations. The result demonstrated the highest microsatellites frequency was found in *O. sativa* (165.5/Mb) which was the minor size of genome, whereas the lower microsatellites frequency was in *Z. mays* (52.1/Mb) which owned the biggest size of genome. Moreover, among the six genomes

analyzed, the genome size of moso bamboo (2,051.7 Mb) was ~17 times of that of *A. thaliana* (119.7 Mb), but the frequency of *A. thaliana* (135.5/Mb) was ~2-fold of that of moso bamboo (62.2/Mb). Compared with these of *Z. mays*, the genome size of moso bamboo was almost identical to that of *Z. mays* (2,066.4 Mb), but the number of microsatellites in moso bamboo was yet increased by 65.2% than that of *Z. mays*.

The microsatellite possessed more than one microsatellite locus contiguously with intervening non-repeat sequence less than 100 bp were identified as compound formation in this study. Compound microsatellite is a special aggregation of microsatellites in which two or more individual microsatellites are found directly adjacent to each other. All identified microsatellites have a minimum length of 15 bp. Whether two or more adjacent microsatellites account as a compound microsatellite depends on the distance separating these microsatellites²⁵. Based on the definition of interrupted length was 1–100 bp in this study, the distribution of interrupted length was analyzed in the selected plants (see Supplementary Table S1 online). The result showed that the mainly interrupted length was concentrated on 3–16 bp. The most of interrupted length was present in 10 bp of moso bamboo, accounted for 6.4%. Moreover, the number of interrupted length was most in moso bamboo, followed by *Z. mays*, while that of *A. thaliana* was slight, revealing the lower complexity of genome than those of moso bamboo and *Z. mays*.

Frequency of motif repeat and length distribution. As a relative frequency based on different microsatellite repeat motifs and length repeat, the major statistical result of moso bamboo was shown in Supplementary Table S2. For the di- and tri- nucleotide repeats, the AG/CT and AT/AT were the dominant repeats compared with AC/GT and CG/CG, and the AAG/CTT was the most abundant, followed by CCG/CGG and AGG/CCT. Moreover, the frequency of ACT/AGT and ACC/CGT repeats were at lower level, accounted for only 2.98% and 3.65%, respectively.

As shown in Figure 1a, besides mono-nucleotide repeat-motif, the length of other microsatellite repeat-motifs was counted in this study and classified into 10 groups. According to the previous study²⁶, the length of microsatellites with more than 20 bp was defined as class I or hypervariable markers, while ones ranged from less than or equal to 20 bp was defined as class II or potentially variable markers. Comparative analysis of the 6 selected plants demonstrated class II occupied a large share, evenly accounted for up to 85%. The number of microsatellites correspondingly decreased with the increasing of microsatellites length.

Microsatellites distribution in different genomic regions. Since microsatellite distribution is the result of selection pressure during evolution²⁷, microsatellites in different genomic regions have different features and may perform varied functions^{28–30}. Here, we analyzed the distribution of microsatellites in genomic regions based on latest genomic sequences and annotations in surveyed plants³. According to the location and annotation of genome, genomic

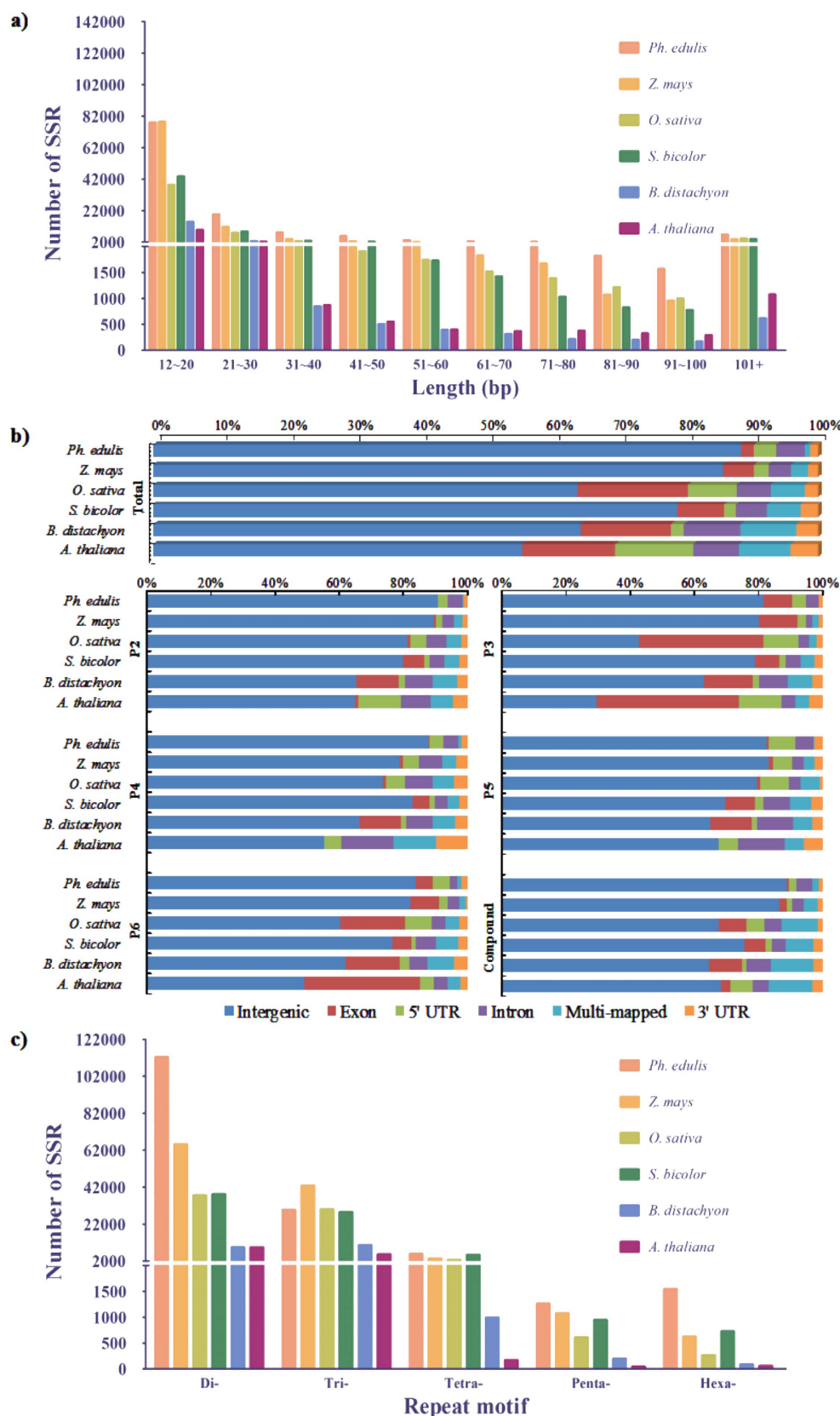


Figure 1 | Distribution of microsatellite repeat motif, length and different genomic regions. (a) Microsatellite length distribution. The x-axis represents the nucleotide length of microsatellites. The y-axis indicates the number of microsatellites with different length in the six selected plant species. (b) The x-axis indicates the proportion of microsatellites with various repeat motifs. The y-axis represents different repeat motif in the six selected plant species. P2: di-nucleotide repeats; P3: tri-nucleotide repeats; P4: tetra-nucleotide repeats; P5: penta-nucleotide repeats; P6: hexa-nucleotide repeats; compound: compound microsatellite. (c) Total number of each repeat motif. The x-axis indicates repeat motif with di- to hexa- nucleotides. The y-axis represents the number of microsatellites with various repeat motifs. The different plant species were marked with different color.



Table 2 | Universal and polymorphic potential of 1,098 microsatellite markers

Type of SSR	Number of SSR	Universality		Polymorphism	
		Number	Probability (%)	Number	Probability (%)
Di-nucleotide	581	474	81.6	193	40.7
Tri-nucleotide	442	377	85.3	130	34.5
Tetra-nucleotide	35	31	88.6	20	64.5
Penta-nucleotide	22	19	86.4	8	42.1
Hexa-nucleotide	9	7	77.8	5	71.4
Compound	9	9	100	9	100
Total	1,098	917	/	365	/

sequences in the 6 surveyed plant species were classified into 6 regions. Besides 5' UTR (Untranslated Regions), intron, exon, 3' UTR and intergenic region, the multi-mapped region was identified via bioinformatics analysis. For instance, one sequence of microsatellite was able to align two or more regions.

As shown in Figure 1b, according to the analysis of sequences, the result illustrated most sequences were commonly mapped onto intergenic regions. The remaining parts were aligned to other regions. For example, the second highest proportion belonged to intron region in moso bamboo, followed by 5' UTR region. Whereas the second one appeared in exon region of *O. sativa*, followed by intron region.

To comprehensively unveil the detailed feature on different type of repeat, distribution based on each type of microsatellites was analyzed. In di-nucleotide repeat, the higher proportion of 5' UTR and exon regions emerged in *A. thaliana* and *B. distachyon*, respectively. In tri-nucleotide repeats, exon region with tri-nucleotide repeat was rather abundant in overall analyzed plants. And the top mapping region was not intergenic regions, but rather exon regions in *A. thaliana*. Excluding tri-nucleotide repeat, exon regions accounted for larger proportion in hexa-nucleotides repeats. The highest proportion of multi-mapped region was found in compound type of motif, because compound microsatellites with longer nucleotides sequences made it possible to align more regions.

Universal and polymorphic potential of microsatellite markers.

Polymorphism of microsatellite marker is the basis to develop molecular markers that are utilized in genetic mapping broadly³¹. Here, we have employed experimental method to detect polymorphism of microsatellite markers in bamboo. Firstly, of all identified microsatellites, 1,451 primer pairs in gene region were designed and 1,098 markers were physically mapped on the genome of moso bamboo. To evaluate universality and polymorphism of developed microsatellite markers, *Ph. edulis*, *Ph. sulphurea* var. *viridis*, *Ph. violascens*, *Ph. nigra*, *Ph. aureosulcata*, *Ph. propinqua*, *Ph. sulphurea*, *Ph. atrovaginata* and *Ph. heteroclada* were amplified using 1,098 primer pairs (see Supplementary Figure S1 online). The result showed 917 of the selected markers produced clear and stable bands with the expected size, among which 365 markers had polymorphism. The universality and polymorphism probabilities of different kind of microsatellites were predicted (Table 2). As effectively molecular marker, microsatellite marker was valuable for identifying relative species. For example, the amplification bands of locus PhEMS-653 were only produced in the *Ph. edulis* and its variant and cultivars (see Supplementary Figure S2a online), therefore the locus was specific for this species and its subordinates. Thus, retrieved from microsatellites validated, 23 primer pairs with high amplification effect were subsequently used for polymorphism analysis among 78 accessions. The 64 alleles ranged from 2 to 5 per loci were detected by 23 SSRs, with an average value of 2.78. The maximum number of alleles was 5 observed in locus PhEMS-561 (see Supplementary Figure S2b

online), and then followed by 4 alleles detected in locus PhEMS-791 (see Supplementary Figure S2c online).

Phylogenetic analysis of *Phyllostachys* accessions. UPGMA (Unweighted Pair Group Method with Arithmetic Mean) analysis of microsatellite data showed that 78 accessions were clustered into 3 classes (Figure 2) based on the genetic distance (see Supplementary Figure S3 online). The accessions in Class I and Class II were belonged to *Phyllostachys* section, while those in Class III were belonged to *Heterocladae* section, in which the cultivars, variants or forma from *Ph. violascens*, *Ph. vivax*, *Ph. edulis*, *Ph. sulphurea*, *Ph. aurea*, *Ph. nigra*, and *Ph. aureosulcata* were together clustered, indicating the majority of the accessions in this clustered result were consistent with their current taxonomic classification³². Class I was the smallest cluster only including 4 accessions (*Ph. propinqua*, *Ph. propinqua* f. *lanugtinosa*, *Ph. nuda* and *Ph. sp.*). The genetic distance between *Ph. nuda* and *Ph. propinqua* was 0.2143 which supported that they were different species with having or not bristles on the back of sheath and coronatines on the internodes of young culm. Class II was the largest group and separated into five subclasses, among which subclass IIa was the group of *Ph. violascens*, including its variant (*prevernalis*) and cultivars ('Notata', 'Viridisulcata', 'Flavistriatus' and 'Anhuiensis'). *Ph. violascens* f. *prevernalis* was the forma of *Ph. violascens* noted in the Flora of China (Chinese), its middle of internode slightly slimmed in morphology and shoot duration was earlier half a month than those of *Ph. violascens*, and the genetic distance between them was 0.111, which supported that they could not be considered as the same one. In addition, *Ph. tianmuensis*, *Ph. primotina*, *Ph. ncarinate*, *Ph. glabrata*, *Ph. dulcis* and *Ph. nigella* were also clustered together within the group of *Ph. violascens*, which their close relationship agreed with the previously morphological classification. The accessions clustered in subclass IIb, IIc and IId were agreed to their current taxonomic classification³².

The genetic analysis based on sequence length variation. On the basis of differences in the copy of microsatellite repeats, allelic length variations were found¹². The mutations, such as insertion, deletion, or translocations, in the regions flanking microsatellite may be considered as polymorphism in the sequence. Therefore, to investigate whether the PCR really amplified the targets, the PCR products were subcloned into T-easy vector and sequenced. For example, the results of sequences alignment showed the locus PhEMS-855 amplified variant alleles from 120 to 147 bp (Figure 3), which revealed mixed type of allelic distribution with the variable number of repeats in the microsatellite motifs (AT)_n accompanied by several point mutations like insertion/deletion or substitution mutations.

In silico comparative genome mapping between moso bamboo and other surveyed species. Based on the experimental results and the previous studies^{12,13}, of 1,098 validated microsatellite, 984 microsatellite markers on the genome of moso bamboo was successfully mapped on four sequenced Poaceae members (Table 3, Figure 4a1-d1). To display syntenic relationships of genome-widely

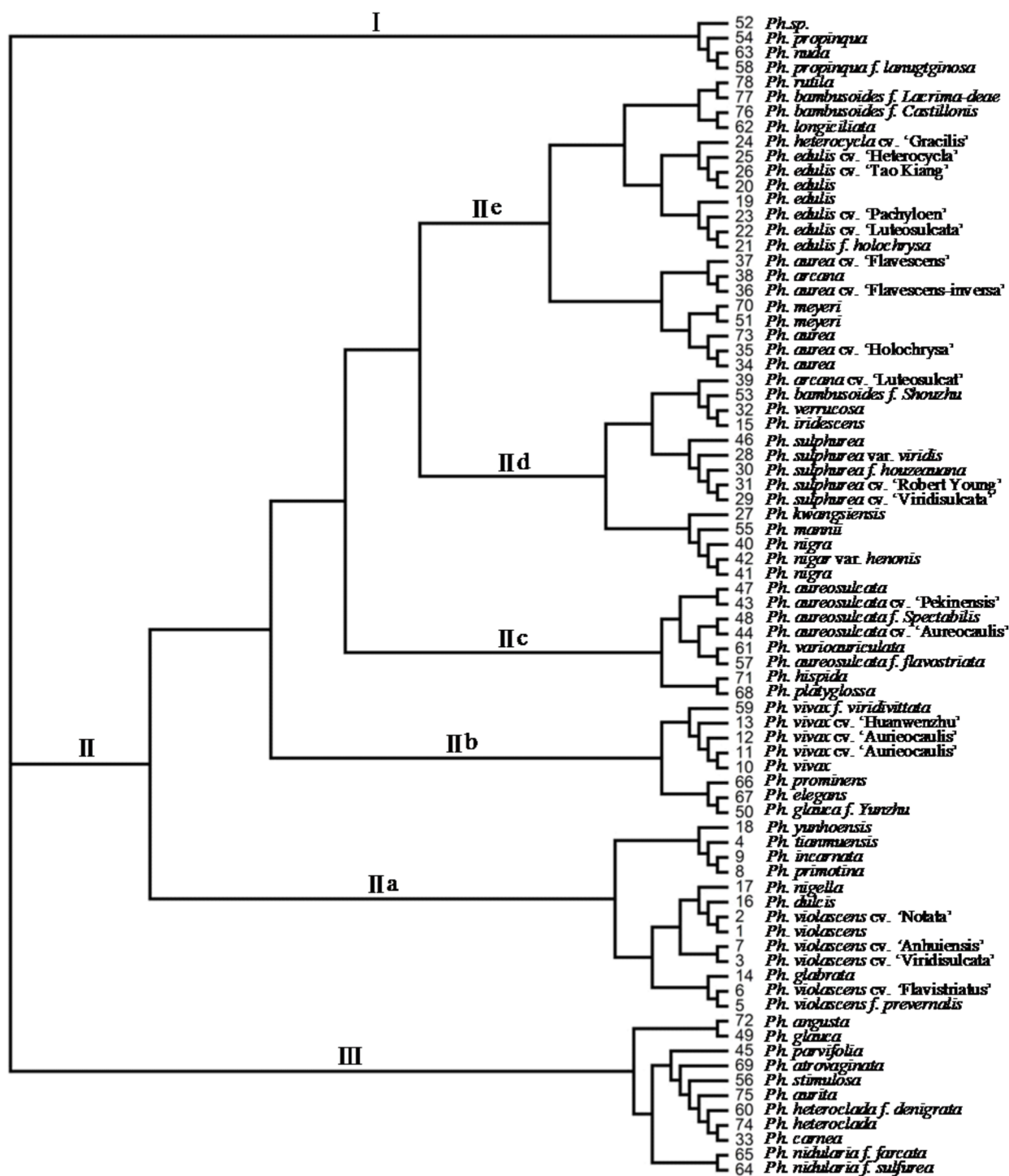


Figure 2 | Phylogenetic analysis of 78 bamboo accessions in genus *Phyllostachys* based on microsatellite data. No. 1–78 represent different bamboo accessions, detailed in Supplementary table S3.

microsatellites marker-based via *in silico* comparative analysis, the scaffolds of moso bamboo were clustering into 6 groups via aligning microsatellites to surveyed genomes, because the draft genome of moso bamboo is scattered and hard to assign scaffolds to chromosomes by now. The results demonstrated the significant proportion of orthology and syntenic relationship of microsatellite marker with *Z. mays* (116, 11.79%), *O. sativa* (185, 18.80%), *S. bicolor* (126, 12.80%), and *B. distachyon* (180, 18.29%). Meanwhile, based on the bioinformatics analysis, of 127,593 identified microsatellites, a total of 101,683 microsatellite markers with available for primers was mapped on four sequenced Poaceae members and their physical

location on genome of other surveyed plants were compared (Table 3, Figure 4a2–d2). Detailed results on microsatellites marker-based comparative mapping were in Supplementary Table S4–S7, which demonstrated a number of moso bamboo microsatellite markers significantly matched with different chromosomes of *Z. mays*, *O. sativa*, *S. bicolor* and *B. distachyon*.

Constructed a database of microsatellites for moso bamboo. To facilitate access and utilization of microsatellite data in moso bamboo, the entire microsatellites had been organized into an online database that was freely accessible via the website (<http://>

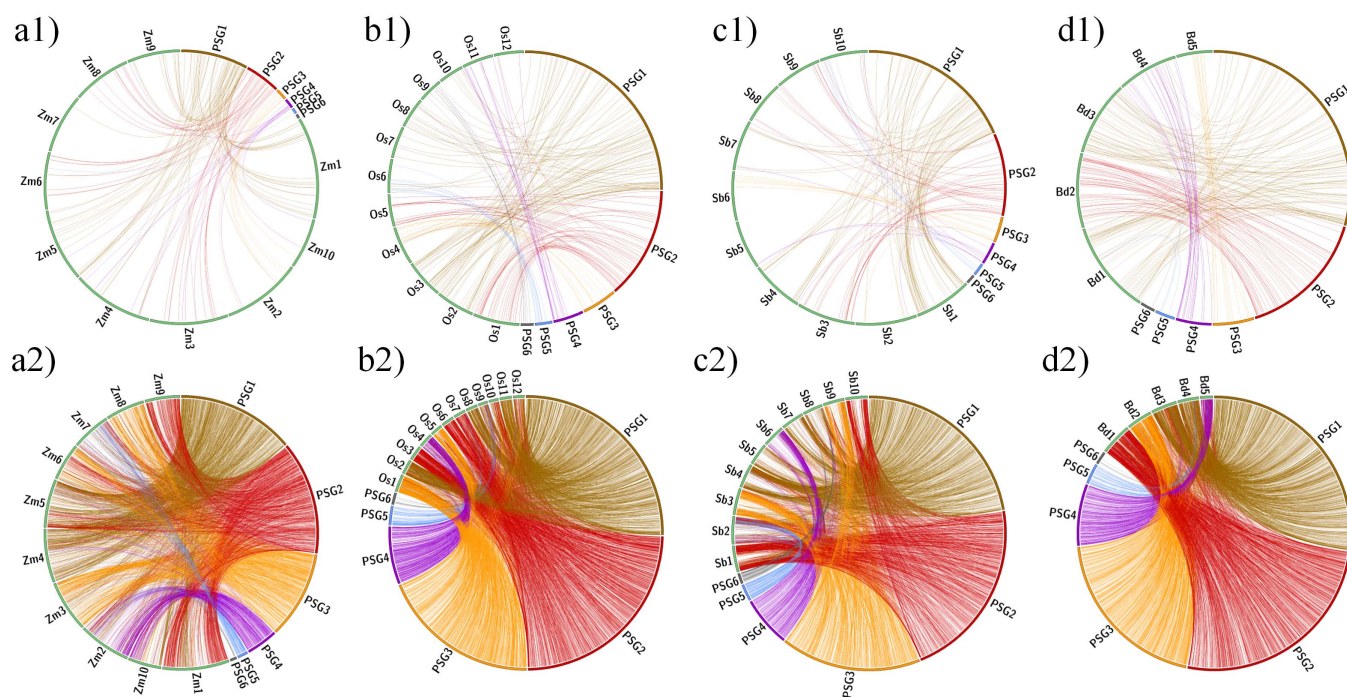


Figure 4 | Genome relationship of moso bamboo with other plant species. Based on the experimentally physical mapping, syntenic relationship of moso bamboo genome with (a1) *Zea mays*, (b1) *Oryza sativa*, (c1) *Sorghum bicolor* and (d1) *Brachypodium distachyon* chromosomes using 984 physically mapped moso bamboo microsatellite markers. Maximum syntenic relationships of the genome of moso bamboo with *Oryza sativa* chromosomes based on experimental microsatellite markers were apparent. Besides, based on the bioinformatics comparative mapping, syntenic relationship of moso bamboo genome with (a2) *Zea mays*, (b2) *Oryza sativa*, (c2) *Sorghum bicolor* and (d2) *Brachypodium distachyon* chromosomes using 101,683 mapped moso bamboo microsatellite markers. Similarly, maximum syntenic relationships of the genome of moso bamboo with *Oryza sativa* chromosomes based on predicted microsatellite markers were apparent as well. Zm1–10 used to be short for the chromosome 1–10 of *Zea mays*; Os1–12 used to be short for the chromosome 1–12 of *Oryza sativa*; Sb1–10 used to be short for the chromosome 1–10 of *Sorghum bicolor*; Bd1–1–5 used to be short for the chromosome 1–5 of *Brachypodium distachyon*; PSG1–6 used to be short for the group of scaffold 1–6 of *Phyllostachys edulis*.

ferent genomic regions have different features and may play various roles^{28–30}. For example, microsatellite frequency in monocot coding regions is 2-fold of that of dicots^{35,36}. Besides, another reason was that the large of scattered and discontinuous sequences may lead to low frequency of microsatellite in the relatively bigger size of genome with high complexity and incomplete assembling.

As a kind of molecular markers, di-nucleotide repeats were considered as the essential resource, because its mutant rate was the highest and the amount was abundant³⁷. For di-nucleotide repeat category, AG/CT repeat was more frequent in *B. distachyon* and *Z. mays* with 50.72% and 47.74%, respectively, whereas AT/AT repeat were more abundant in *S. bicolor* and *A. thaliana* with 54.40% and 52.58%, respectively. Another noting that AG/CT and AT/AT repeat dominated over other di-nucleotide repeats in entire selected plant species. Moreover, the outcome of di-nucleotide indicated that GC-rich motifs were enrichment rather than AT-rich motifs.

Differentiated from di-nucleotide repeat motif, the remaining ones (from tri- to hexa-nucleotides) have a wide diversity in the distribution of repeat motif. For example, in tri-nucleotide repeats category, AAG/CTT repeat of moso bamboo was the abundant, accounted for 20.08%, followed by the CCG/CGG repeat (20.04%), whereas CCG/CGG repeat was plentiful in *O. sativa*, *B. distachyon* and *S. bicolor*. This result may be caused by the high GC content and codon bias in monocotyledons³⁸. Subsequently, the frequency of tetra-, penta and hexa-nucleotide repeats was relatively at low level in all surveyed plants and the number of their motifs was slight.

As the development of microsatellite markers, the main resources derived from the cDNA and EST data owing to lacking of the genome data. cDNA-SSR and EST-SSR were on the basis of transcription region, high conversation and wide versatility. The comparative studies among cDNA-SSR, EST-SSR and genome-SSR depicted that

the most of motif repeats and frequency were similar in moso bamboo^{39,40}. However, there were some significantly differences between the size of examined sequences and the number of identified microsatellites (see Supplementary Table S8 online). For example, tri-nucleotide repeat was the dominant in EST-SSR and cDNA-SSR, while mono- and di-nucleotide repeats were abundant in genome-SSR. Moreover, the analysis indicated that the type and number of microsatellite repeats were more abundant in genome sequence. Taken together, it is necessary that developing microsatellite markers based on the whole genome of moso bamboo to comprehensively and effectively assisting bamboo taxonomy and systematically expand our understanding of microsatellite.

The taxonomy of bamboo has puzzled researchers because of bamboo reproductive characteristics. Especially in the genus *Phyllostachys*, some of same accessions may have different names, while some different accessions may have the same name. With the fast development of molecular biology, the molecular marker of bamboo was developed and used for bamboo genetic diversity and phylogenetic analysis which was helpful for the taxonomy of bamboo. ISSR and AFLP techniques were used to measure genetic diversity and similarity in cultivars of moso bamboo, three taxa including *Ph. kwangsiensis*, *Ph. nigra* var. *henonis* and moso bamboo were separated into three groups while the cultivars of moso bamboo were genetically similar⁴¹. Meanwhile, RAPD analysis was also capable in detecting bamboo species, although it was sometime difficult to distinguish similar species⁶. For example, the previous study on 23 bamboos of *Phyllostachys* based on RAPD method showed the cultivars of *Ph. vivax*, *Ph. violascens* and *Ph. aureosulcata* were not clustered into one clade⁴². There was a difference between this result of cluster and the traditional taxonomy based on classification of biological characteristics. To considerably investigate this problem, we

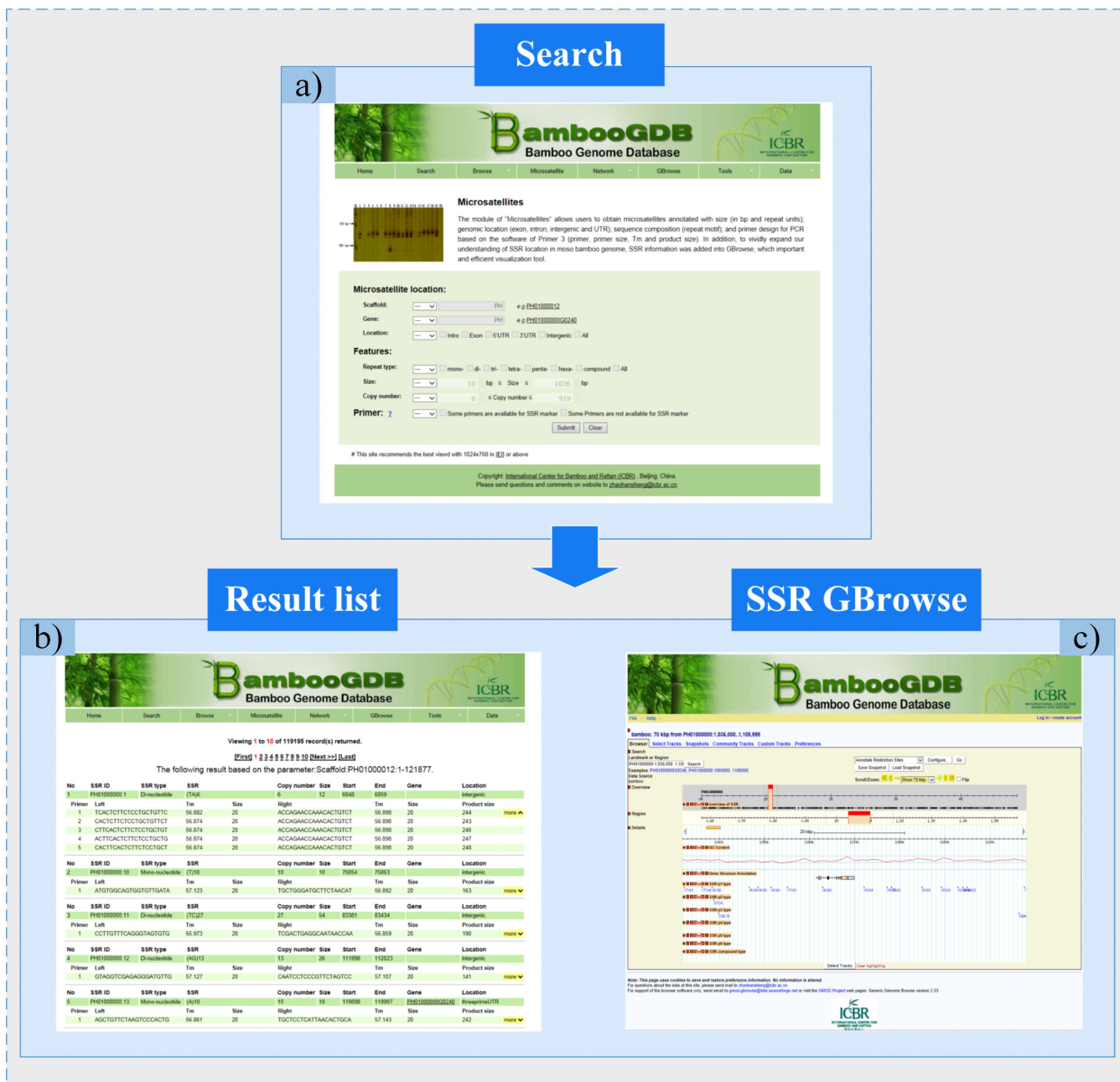


Figure 5 | Screenshot showing the pages of browse and results in bamboo microsatellite database. (a) The Boolean search was provided in the searching page of moso bamboo microsatellites. (b) The results page was included, such as microsatellite ID, microsatellite type, microsatellite position, microsatellite primers and so on. (c) The physical location of microsatellites in moso bamboo genome was displayed in GBrowse tool.

had adopted microsatellite technique based on genome sequence to detect the bamboos of the genus *Phyllostachys*.

Based on the development of microsatellite markers, the clustered result showed that the majority of the accessions were consistent with their current taxonomy. The accessions of synonyms were confirmed by the genetic distance of zero, such as accession 34 'Luo han zhu' and accession 73 'Ren mian zhu' belonged to the same one of *Ph. aurea*, accession 51 'Zhe jiang dan zhu' and accession 70 'Mao huan zhu' were the same one of *Ph. meyeri*. However, there were also few exceptions. For example, in subclass IIe, the cultivars of 'Holochrysa', 'Luteosulcata', 'Pachyloen', 'Gracilis', 'Heterocycla' and 'Tao Kiang' from *Ph. edulis* clustered together demonstrated close genetic relationship among them, the genetic distance was zero among 'Holochrysa', 'Luteosulcata' and 'Pachyloen', in addition to that among 'Gracilis', 'Heterocycla', 'Tao Kiang' and *Ph. edulis* was

also zero, indicating that these cultivars had no genetic variation. The results also indicated that the 23 microsatellite loci selected were hardly used to distinguish the *Ph. edulis* from its cultivars. Another example, the accessions in Class III mainly contain the members of Heterocladae section except accession 49 'Dan zhu' and accession 74 'Shui zhu' which were classified into *Phyllostachys* section according to the reference³². Several of molecular techniques made similar taxonomy but including slight discrepancies. For example, *Ph. nigra* was classified in *Phyllostachys* section using microsatellite, which was consistent with the taxonomy defined by previous research⁴³. However, *Ph. nigra* was grouped internal to Heterocladae section using the RFLP⁴⁴ and AFLP⁴⁵ methods.

These results may be caused by the following reasons, 1) Bias resulted from the cluster owing to the number of microsatellite molecular markers were not enough. 2) The resolution was relatively



lower for the products of amplification using non-denaturing PAGE electrophoresis, which resulted in allelic loci were not easily identified comparing with capillary electrophoresis. 3) Only the easily recognizable bands were recorded by artificial reading, the fuzzy ones were not adopted in data statistics. Therefore, the most abundant of allelic loci was only five in this study, while this number was nine in the population structure and phylogenetic relationships of *Ph. edulis* using capillary electrophoresis in diversity analysis⁴⁶. These reasons for the difference with traditional taxonomy need to be further validated by expanding the number of microsatellite loci with the capillary electrophoresis method.

Due to complex genomic architecture, weak biology research on bamboo, infrequent sexual reproduction and the long periods of time between flowering intervals, the draft genome of moso bamboo is scattered with 277,278 scaffolds by now²⁴. Therefore, it is difficult to assign scaffolds to chromosomes. Here, to vividly demonstrate physical location of microsatellites in genomic regions, information of microsatellite had been added into GBrowse, which is an efficient visualization tool⁴⁷. In our GBrowse, various types of factors, including gene, coding sequence, messenger RNA, RNA-seq, were simultaneously in the context of genomic regions, expanding our understanding of microsatellites (Figure 5c). Users can access to the detailed feature page of corresponding entries from the browser.

Method

Plant materials, genomic DNA isolation and detection. Leaf samples of the accessions belonged to the genus *Phyllostachys* are collected from the base of Anhui Experimental Station of International Center for Bamboo and Rattan, Huangshan City in Anhui Province as well as Lin'an County, Hangzhou City in Zhejiang Province, respectively. The accessions were shown in Supplementary Table S3. These Latin name mainly referred to Flora of China²². Total genomic DNA was extracted with CTAB method described by Doyle and Doyle with minor modifications⁴⁸. The quality and quantity of DNA were detected using Nanodrop 2000.

Identification of microsatellites. The MicroSatellite identification tool (MISA) (<http://pgrc.ipk-gatersleben.de/misa/>) was used to identify microsatellite occurrences in the genome sequence data (moso bamboo data from BambooGDB⁴⁹, others from Phytozome v10³). To identify the presence of microsatellites, the minimum repeat unit was defined as 6 for di-, and 5 for tri-, tetra-, penta-, and hexa-nucleotides. The microsatellite position, number of different repeat types of microsatellite and length (motif bp × number of motif) was analyzed by a bespoke program written in MISA files⁵⁰.

Designing primers and validation of moso bamboo microsatellite markers. The primer pairs for flanking sequences of each unique microsatellite were designed automatically using the Primer3 (See Supplementary Table S9 online). Nine samples of *Ph. edulis*, *Ph. sulphurea* var. *viridis*, *Ph. violascens*, *Ph. nigra*, *Ph. aureosulcata*, *Ph. propinqua*, *Ph. sulphurea*, *Ph. atrovaginata* and *Ph. heteroclada* were selected to validate the suitability of microsatellite loci by PAGE. According to previous studies, PCR amplification was performed and PCR products were separated²⁸. Seventy-eight accessions belonged to the genus *Phyllostachys* (see Supplementary Table S3 online) were used for microsatellite marker validation. UPGMA cluster analysis was performed based on the similarity matrix for 78 bamboo accessions using the PowerMarker program in the statistical analysis⁵¹, then phylogeny was displayed in the TreeView software⁵².

The variation analysis of microsatellite marker. Four primer pairs were selected to detect the microsatellite variation in different bamboo species. The genomic DNA samples of *Ph. edulis*, *Ph. violascens*, *Ph. aureosulcata*, and *Ph. heteroclada* were used as templates for PCR, and the PCR products were subcloned into T-easy vector and sequenced. The sequence alignment analysis was carried out by using DNAMAN version 5.0.

In silico comparative genome mapping between bamboo and other grass species. Based on the previous studies^{12,13}, the flanking sequences of microsatellite marker loci were mapped the genome sequences of moso bamboo were BLAST searched with best hit against genome sequences of *Zea mays*, *Oryza sativa*, *Sorghum bicolor*, *Brachypodium distachyon* to analyze marker-based syntenic relationship between bamboo and other grass species. Before displayed with syntenic relationships in Circos software⁵³, the scattered scaffolds of moso bamboo were clustered into 6 groups via aligning microsatellites to surveyed genomes.

System design and implementation of microsatellite database. For moso bamboo, all data of microsatellite are stored and managed in a MySQL relational database. The website was developed in CentOS Linux 6.4 environment using Java Platform Enterprise Edition, Apache Tomcat, Apache web server and so on. In addition, to

vividly exhibit location of microsatellite in whole genome, microsatellite data was added in GBrowse after being converted into genome feature format (GFF). GBrowse uses MySQL as backend and was built following the configuration files provided by its developer (<http://gmod.org>).

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Author contributions

H.Z. and Z.G. designed the experiments; H.Z. and L.Y. performed experiments; H.Z., L.Y., L.W., L.D. and H.S. analyzed data; Y.L., X.Y. and Z.P. contributed reagents/materials/analysis tools; H.Z. and Z.G. wrote the paper. All authors read and approved the final manuscript.

Additional information

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