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Abstract

Background: Chinese bayberry (Myrica rubra Sieb. & Zucc.) is an important subtropical evergreen fruit tree in southern China. Generally dioecious, the female plants are cultivated for fruit and have been studied extensively, but male plants have received very little attention. Knowledge of males may have a major impact on conservation and genetic improvement as well as on breeding. Using 84 polymorphic SSRs, we genotyped 213 M. rubra individuals (99 male individuals, 113 female varieties and 1 monoecious) and compared the difference in genetic diversity between the female and the male populations.

Results: Neighbour-joining cluster analysis separated M. rubra from three related species, and the male from female populations within M. rubra. By structure analysis, 178 M. rubra accessions were assigned to two subpopulations: Male dominated (98) and Female dominated (80). The well-known cultivars ‘Biqi’ and ‘Dongkui’, and the landraces ‘Fenhong’ are derived from three different gene pools. Female population had a slightly higher values of genetic diversity parameters (such as number of alleles and heterozygosity) than the male population, but not significantly different. The SSR loci ZJU062 and ZJU130 showed an empirical Fst value of 0.455 and 0.333, respectively, which are significantly above the 95% confidence level, indicating that they are outlier loci related to sex separation.

Conclusion: The male and female populations of Chinese bayberry have similar genetic diversity in terms of average number of alleles and level of heterozygosity, but were clearly separated by genetic structure analysis due to two markers associated with sex type, ZJU062 and ZJU130. Zhejiang Province China could be the centre of diversity of M. rubra in China, with wide genetic diversity coverage; and the two representative cultivars ‘Biqi’ and ‘Dongkui’, and one landrace ‘Fenhong’ in three female subpopulations. This research provides genetic information on male and female Chinese bayberry and will act as a reference for breeding programs.

Background

Chinese bayberry (Myrica rubra) is an evergreen fruit tree native to southern China and other Asian countries [1]. The fruit has a delicious flavour and high nutritional value, especially rich in anthocyanins. It can be eaten as fresh fruit and as processed products, and has become popular in China and other countries in recent years [2-4]. Among the six Myrica species in China, only M. rubra is commercially cultivated in Zhejiang, Jiangsu, Fujian and Guangdong, and is also grown in the Yunnan, Guizhou and Hunan provinces [5,6].

Chinese bayberry (2n = 16) belongs to the Myricaceae family, is usually dioecious, is wind pollinated, and only a few individuals are monoecious. Other species in the Myricaceae family such as M. cerifera, M. faya and M. rivas-martinezii are also dioecious plants [7-9], with unclear mechanism of sex determination. Monoecious plants have also been found in M. faya [9]. M. rubra has a symbiotic association with nitrogen-fixing bacterium in the root system, which has also been found in other Myricaceae family members [7,10-12]. The morphology of its inflorescences and flowers varies with the sex of the tree (Fig. 1). Male plants, with a different colour and
shape of staminate catkins, are planted for landscape and pollination purposes [1]. The number of male plants is diminishing year by year because of their low economic benefits. Even though Chinese bayberry was domesticated in southern China more than 2000 years ago, it only has about fifty years of research history. The rich germplasm of Chinese bayberry is reflected by around 300 recorded landraces/cultivars [5], also some landraces including a group rather than a single scion variety. Zhejiang Province has the longest history of Chinese bayberry cultivation, with considerable germplasm resources. The main cultivars ‘Biqi’ (with black fruit and average weight of 11 g) and ‘Dongkui’ (red fruit and average weight of 22 g) are pure scion cultivars, and have been widely cultivated in China. While two landraces ‘Fenhong’ (pink fruit and average weight of 13 g) and ‘Shuijing’ (light yellow fruit and average weight of 15 g) are limited to local regions in Ningbo area of Zhejiang Province. Mutation and natural elite line identification has been the dominant way of breeding new cultivars [6].

Genetic diversity, cultivar identification and the geographic origin of Chinese bayberry have been studied using molecular markers [5,6,13], but based on the dozen available markers [14,15], with the male plants rarely used in the study of cross breeding and genetic diversity. With the development of sequencing techniques, a large number of simple sequence repeat markers (SSRs) have been isolated in recent years [13,16,17]. Using SSR markers, no genetic differentiation has been found between M. rivas-martinezii and M. faya [8]. SSRs are powerful markers which have been widely used in studies on genetic diversity and population structure, and to construct linkage maps [18-22].

Molecular markers linked to sex determination have been reported in a few dioecious fruit crops. A microsatellite (GATA)n revealed sex-specific differences in papaya [23], and SCAR markers were then developed for sex determination [24,25]. Fraser found that sex-linked SCAR markers and the ‘Flower-sex’ phenotype mapped to the same linkage group in the dioecious species Actinidia chinensis (kiwifruit), which revealed putative X/Y sex-determining chromosomes[19]. However, there has been no report on male plant genetic diversity and sex determination in Chinese bayberry or other Myrica species. Here we report, for the first time, the genetic diversity of male plants of Chinese bayberry compared with the female plants, using 84 SSR markers. This information will be useful in the conservation of these diverse individuals and for direct application in the new initiative of the Chinese bayberry cross-breeding program.

Results

Genetic diversity of the M. rubra accessions

The 84 SSRs used in this research were polymorphic in both female and male samples, amplifying a total of 876 alleles, with an average of 10.43 alleles per locus. (Additional file 1: Table S1). The number of alleles per locus (Na) ranged from 3 to 23. The frequency of more than half of the alleles (526, 60.05 %) was less than 5 % and 236 were unique (frequency less than 1 %). Low allele frequencies resulted in a reduced effective number of alleles (Ne, 3.71). The observed heterozygosity (Ho) ranged from 0.11 to 0.90, with an average of 0.49. For most loci (72) the Ho value was lower than the expected (He), which ranged from 0.19 to 0.88, (mean, 0.65). Shannon’s information index (I) for each locus ranged from 0.47 to 2.36, with a mean of 1.48.

The Na, Ne, Ho, and Hs of the male and female populations in 84 SSR loci are shown in Table 1. Across all 84 SSR loci, we found fewer Na (8.71) in the male population than in the female (9.04). The sample of male accessions (99) was smaller than that of the female population (113

| Subset of accessions | Sample size | Na  | Ne   | Ho   | Hs   | F   |
|----------------------|-------------|-----|------|------|------|-----|
| All accessions       | 213         | 10.11 | 3.75 | 0.49 | 0.65 | 0.25 |
| Female population    | 114         | 9.04 | 3.60 | 0.55 | 0.65 | 0.34 |
| Male population      | 99          | 8.75 | 3.34 | 0.40 | 0.62 | 0.15 |

Note: Na: number of observed alleles; Ne: effective number of alleles; Ho: observed heterozygosity; Hs: gene diversity; F: Wright’s fixation index.
accessions), so to correct for this difference in sample size, the means of distinct and private alleles per locus were analysed as a function of the sample size for the two populations (Additional file 2: Figure S1). With increasing sample size, the number of distinct and private alleles also increased, and it was clear that the female population had a slightly higher value than the male population. Moreover, the female population had a slightly higher mean effective number of alleles (3.60) and Hs (0.65) than the male population (3.34 and 0.62, respectively), although not significant (t test, \( P = 0.2530 \)). Apart from the differences between them, these results also illustrate that the female and male of Chinese bayberry have abundant genetic diversity.

**Population Structure**

The 213 *M. rubra* accessions were evaluated for population stratification. The assumed number of populations was set from \( K = 1 \) to 10. According to the method of Evanno [26], the accessions were mainly divided into two subpopulations (\( K = 2 \)). CLUMPP alignment of 10 independent solutions for \( K = 2 \) showed pairwise ‘G’ values around 0.99, indicating that the assignment of genotypes to the subpopulations was well correlated between runs. Setting the membership coefficient to \( Q \geq 0.6 \), 178 genotypes could be generally clustered into two subpopulations (Fig. 2a), one the Male Dominated pop. (green in Fig. 2a) with 82 male and 16 female plants, and the other the Female Dominated pop. (red in Fig. 2a) with 76 female and four male plants. The remaining 35 genotypes were unstructured: about half of them (17) were female cultivars from the Hunan, Jiangsu and Guizhou provinces. The monoecious plants (C2010-4) also clustered within this admixed subpopulation.

STRUCTURE was then run for each of the subpopulations as a first step. The Male Dominated population of plants (Fig. 2a, in green) was further divided into four groups (Fig. 2b). The first, the Mixed male subpopulation

![Fig. 2](image-url) Population stratification based on Bayesian clustering approaches, with a value of \( k \) from 1 to 10. Each individual is shown as a thin vertical line, different subpopulations are in a different colour. The name of each population or subpopulation and the number of individuals included is given at the top of each column. a. The first STRUCTURE step with 213 accessions, \( k = 2 \). The subpopulation was displayed by DISTRUCT, and each subpopulation ordered according to the membership coefficient. b. Nest structure analysis for the Male Dominated pop. mainly divided into four subpopulations. c. Nest structure analysis for the Female Dominated pop. which was further divided into two subpopulations. Note: Unstructured indicates individuals which were not assigned to any population.
(orange in Fig. 2b), included 27 male plants from five provinces of south China. The Mixed sex subpopulation (purple in Fig. 2b) included five female plants from Guangdong and Zhejiang and three male plants from Zhejiang and Jiangsu provinces, and the third subpopulation, Zhejiang male (yellow in Fig. 2b), included 21 male plants and one cultivar from Zhejiang Province. In the fourth subpopulation, the ‘Dongkui’ female series, there were seven landraces from Zhejiang, one cultivar (Heiruilin) from Taiwan and one (Ruiguangmei) from Japan.

The Female-dominated population was further structured into two groups (Fig. 2c). The ‘Biqi’ female series was largely restricted to plants from Zhejiang (41), also with two from Fujian and one from the Jiangxi Provinces. Eight of the ten cultivars from Zhejiang were ‘Biqi’-derived cultivars, with three male plants Y2012-1, Y2012-2 and Y2010-16 also in the ‘Biqi’ female series. The ‘Fenhong’ female series included 31 female and one male (‘Y2012-151’) accessions mainly from Zhejiang Province, with seven and eight of the 31 female accessions belonging to the ‘Shuijing’ and ‘Fenhong’ series respectively.

Based on AMOVA analysis, most variation (81.55 %) was detected within individual accessions, with only 11.00 % attributed to variation among the six subpopulations (Table 2). The overall Fst among the six subpopulations was 0.1100 (p value < 0.05). Pairwise Fst values ranged from 0.0728 (between the ‘Zhejiang male’ and ‘Mixed male’ subpopulations) to 0.2346 (between the ‘Zhejiang male’ and ‘Dongkui female’ series) (Table 3). The genetic diversity of 178 structured accessions was also confirmed by PCoA (Fig. 3). The first three axes together accounted for 63.65 % of the variation. The first and second coordinates accounted for 26.91 % and 19.76 % of the molecular variation, respectively, with the first coordinate separating Male Dominated population from Female Dominated population accessions, and the second coordinate separating the ‘Biqi’ and the ‘Fenhong’ female series.

### Table 2 Analysis of molecular variance (AMOVA) among subpopulations inferred by STRUCTURE analysis, based on the 84 SSR loci of 178 Chinese bayberry accessions

| Source of variation | d.f. | Sum of squares | Variance components | Fixation indices | Percentage of variation |
|---------------------|------|----------------|---------------------|-----------------|-------------------------|
| Among populations   | 5    | 530026         | 1.86626 Va          | Fst = 0.11005   | 11.00                   |
| Among individuals within subpopulations | 146  | 2387945       | 1.26342 Vb          | Fis = 0.00371   | 7.45                    |
| Within individuals  | 152  | 2102000       | 13.82895 Vc         | Fit = 0.18455   | 81.55                   |
| Total               | 303  | 5019970       | 16.95863            |                 |                         |

Fis: inbreeding coefficient of subpopulations, Fit: inbreeding coefficient in the total sample, Fst: genetic differentiation among subpopulations.
0.013 and 0.607 (Fig. 6a), respectively. Moreover, the frequency of 240/240 and 242/242 genotypes in the two populations was distinct (Fig. 6b). Similarly for the ZJU103 locus, ZJU130-160 bp was 0.119 in the female and 0.719 in the male population, while ZJU130-162 bp was 0.487 and 0.097, respectively (Fig. 6c). The main genotype of ZJU130 in the male population was 160/160 bp, while in the female population they were 162/162, 162/164 and 162/168 (Fig. 6d).

When we exclude the two SSR markers associated with sex, the phylogenetic tree (Additional file 6: Figure S3) showed that a small group of the male and female accessions were mixed together, not separated clearly as in Fig. 4.

Discussion

SSR polymorphism and genetic diversity

Here we studied the variability of a heterogeneous collection of 213 Chinese bayberry genotypes. The 99 male individuals were collected from five provinces of south China, with more than two thirds from six cities in Zhejiang Province. In total, the 84 polymorphic SSRs amplified an average of 10.43 alleles per locus, 26.9 % of which were rare alleles, higher than the value observed in previous studies [6,13,15]. These high values can be explained by a larger sample size and high heterogeneity of the sample.

The female and male accessions contributed equally to the variability, with a mean gene diversity value, $H_e$, of 0.65 for the 216 individuals, which suggested the rich genetic diversity of *M. rubra*. This was similar to that (0.66) reported by Xie, where 123 cultivars were analysed with 14 SSRs [6], and slightly lower than the figure of 0.72 estimated by Jiao, who studied 23 landraces and six male accessions of *M. rubra* [13]. Heterozygosity of the females was slightly higher than that of the males (0.65 versus 0.62) but the difference was not significant, in part due to the wide genetic background in the samples and to the cross pollination in dioecious plants. Some of the landraces also clustered within their own gene pools, especially those from Zhejiang Province: ‘Biqi’, ‘Fenhong’ and ‘Dongkui’. Based on our results, it is clear that germplasm from Zhejiang Province, with its rich genetic diversity, should be widely used in breeding programs.

Population structure

Bayesian clustering analysis has been proven to be powerful tools to evaluate the genetic structure of fruit tree populations. A nested clustering strategy has also been used to assign individuals to populations, for example in apple [27] and peach [18]. The primary genetic separation in our data divided the male and female collections into two subpopulations, with a few exceptions, with low genetic structure suggesting that gene flow

### Table 3 Pairwise estimates of Fst among the six subpopulations, based on 84 SSRs

|                  | ‘Biqi’ female series | ‘Fenhong’ female series | Mixed sex | Zhejiang male | ‘Dongkui’ female series | Mixed male |
|------------------|----------------------|-------------------------|-----------|---------------|-------------------------|-----------|
| ‘Biqi’ female series | 0.0000               |                         |           |               |                         |           |
| ‘Fenhong’ female series | 0.0945               | 0.0000                  |           |               |                         |           |
| Mixed sex         | 0.1118               | 0.0756                  | 0.0000    |               |                         |           |
| Zhejiang male     | 0.1621               | 0.1534                  | 0.1335    | 0.0000        |                         |           |
| ‘Dongkui’ female series | 0.2066               | 0.2085                  | 0.1910    | 0.2346        | 0.0000                  |           |
| Mixed male        | 0.0876               | 0.1040                  | 0.0849    | 0.0728        | 0.1919                  | 0.0000    |
Fig. 4 (See legend on next page.)
between them is high. In the STRUCTURE analysis, 82 males and 16 females were assigned to the Male-dominated population, which can be further clustered into four subpopulations (Fig. 2b), suggesting a mixed genetic background of both male and female groups. The population differentiation value between the ‘Dongkui’ female series (close to the male subpopulation) and the two female subpopulations (‘Biqi’ and ‘Fenhong’) indicates early differentiation of the ‘Dongkui’ female from the males. This can explain why, applying growth regulator to control vegetative vigour, the ‘Dongkui’ cultivar is prone to bear male flowers [28], as well as provide evidence that male and female of Chinese bayberry may share common ancestors and co-evolved. Dioecious plants may have evolved from hermaphrodite ancestors [29]. The lower population differentiation value found between the ‘Zhejiang male’ and ‘Mixed male’ subpopulations suggests that sex differentiation occurred earlier than genetic differentiation.

Also using STRUCTURE as the first step, most Zhejiang landraces clustered together (Female Dominated pop., Fig. 2a). However, further analysis separated this cluster into two subpopulations, the ‘Biqi’ female series and the ‘Fenhong’ female series (Fig. 2c), with a low population differentiation value between Zhejiang landraces, revealing different germplasm sources in Zhejiang Province. Archaeological and written evidence both suggest that bayberry existed in Zhejiang Province 7,000 years ago, in Hemudu (Yuyao, Zhejiang Province), and was widely cultivated and grafted in Zhejiang Province with elite cultivars introduced into other provinces of China since the Southern Song Dynasty (1127–1279) [1]. This suggested that Zhejiang would be the centre of diversity of M. rubra. ‘Biqi’ and ‘Fenhong’ would be old landraces, possibly sharing one or a number of common ancestors. However, this remains to be further elucidated.

**Phylogenetic clusters**

Phylogenetic analysis grouped M. rubra accessions into two subgroups consistent with sex type and geographic origin. The results were also in agreement with population structure and PCoA analysis. Collections from Fujian Provinces (Anhaipianzaosheng and Zaoshenganmei) are clustered with those from Zhejiang Provinces (Zaoqimimei and Sanjiaowumei) in SG2-2, indicating gene flow occurred frequently among these two provinces [5,6]. Zhejiang has a long history of bayberry cultivation, and the excellent cultivars have been spread to surrounding provinces [1]. The landraces, ‘Fenhong’, and two best-known main cultivars ‘Biqi’ and ‘Dongkui’, were assigned to the subgroups SG2-1, SG2-2 and SG2-3, respectively, indicative of the rich germplasm in Zhejiang Province. The relatively long genetic distance between ‘Shuijing’ and ‘Shuijing2’ in SG2-1 is probably because ‘Shuijing’ was not a single but a group of cultivars [17], a phenomenon also found in ‘Fenhong’, ‘Yuelian’ (SG2-1) and ‘Biqi’ (SG2-2). The Taiwan and Japanese cultivars, ‘Heiruiin’ and ‘Ruiguangmei’, clustered within SG2-3 and were closely related to cultivars from Zhejiang Province, consistent with the STRUCTURE results and confirming previous findings [5,6,17].

In general, males from geographically closer locations had higher genetic similarities than those from more distant locations, however, the three male plants of Y2012-1, Y2012-2 and Y2012-151 were clustered in SG2-1 and SG2-2, close to female plants collected from the same location, which indicates that these plants may be progeny of the female plants. Though few male accessions are clustered with female accessions, we would speculate that the male and female populations essentially have similar genetic diversity but distinct background associated with sex.

**Sex determination in M. rubra**

Dioecious plants with separate male and female individuals are found in only 6% of the 240,000 angiosperm species, and probably evolved recently from hermaphrodite ancestors [30]. Myricaceae probably evolved from polygamy through monoeisom to dioecism [9]. Clarification of the mechanism for sex determination is important for developmental biology and breeding practices, but studies in dioecious plants have been limited to a few model
plants, such as white campion (Melandrium album) [31-33] and Rumex acetosa [34,35]. Recently, in the study of papaya, bacterial artificial chromosome (BAC) libraries were constructed by sequencing the sex chromosomes [36], and then the X and hermaphrodite-specific region of the Y<sup>b</sup> chromosome (HSY) were sequenced, with the results supporting the model of early sex chromosome evolution [37]. Sex determination and the sex chromosome of M. rubra and other related species have not been reported [9]. The SSR loci ZJU062 and ZJU130 were shown to be outlier (Fig. 5), so appear to be associated with sex segregation, however, excluding the two sex-associated SSR markers, the majority of female and male accessions were still distinct (Figure S3). Linkage status of these two SSR loci is under investigation. More research is needed to localize the precise genetic and genome positions controlling sex traits using segregating and natural populations of both sexes, screening with more SSR markers developed recently [13,16,17].

Conclusions
Analysing male and female genotype data, we were able to estimate the genetic diversity of Chinese bayberry and demonstrate the rich diversity of the population. Phylogenetic cluster and population structure analyses revealed the sex trait effect on genetic structure stratification, and that Zhejiang Province could be the centre of diversity of M. rubra. The genetic diversity of the two cultivars ‘Biqi’ and ‘Dongkui’ and the landrace ‘Fenhong’ makes them an excellent source of variability for Chinese bayberry breeding programs. We also identified two SSR markers putatively associated to sex segregation in the two populations.

Materials and methods
Plant materials and tissue sampling
A total of 213 individuals of Chinese bayberry (M. rubra) and three related species (M. adenophora, M. nana, M. cerifera) were sampled. For M. rubra, 113 females, 98 males, one monoecious plant (C2010-4), as well as
were used as amplified PCR band size control. Among these collections, the male population was collected from five provinces in China, and half of the Yuyao and Cixi accessions from trees planted in avenues located in different areas of Zhejiang Province, with these male trees collected from different regions in Ningbo, Zhejiang Province. Of the female population, 34 were from the China Bayberry Germplasm Repository (CBGR), Yuyao, Zhejiang Province, China, and 62 were local landraces (Additional file 7: Table S4; Additional file 8: Figure S4). We used the Juno SC handheld (Trimble) GPS to obtain the geographic position of each individual. Among the 216 accessions, for 32, genotypic data was obtained from previous reports by Jiao [13], and the genotype of the three accessions ‘Biqi’, ‘Dongku’ and ‘Fenhong’ were used as amplified PCR band size control.

DNA extraction
Young leaves were collected from healthy trees, frozen in liquid nitrogen, and then stored at −40 °C prior to DNA extraction. The genomic DNA was extracted by the optimized CTAB method as previously described [5], and the DNA was quantified in an ultraviolet spectrophotometer (Eppendorf) and diluted to 20 ng/μl for PCR amplification.

Microsatellite marker amplification
A total of 84 primers pairs (Additional file 1: Table S1) were selected according to their high polymorphism in previous studies: six EST-SSR markers with MRU/MYB as prefix were developed from the M. rubra EST database [15], and the remaining 78 were genomic SSR markers [13,14]. Forward primers were labelled with different fluorescence: NED, PET, FAM and HEX (Invitrogen). PCR amplifications were performed in an Eppendorf Mastercycler (Germany), with amplification reactions and temperature cycles according to the protocol described by Terakawa et al., Zhang et al., and Jiao et al. [13-15]. PCR products were checked on 1 % agarose gel at 100 V (m·v⁻¹), and PCR products, with four different colour labelling, were then mixed with the internal size standard LIZ500 (ABI) and separated by capillary electrophoresis in an ABI 3130 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). Allele size was estimated with the GeneMapper v4.0 software (Applied Biosystems, Foster City, CA, USA). The genotypes of 216 accessions were deposited in Dryad (doi: 10.5061/dryad.7qc88).

Data analysis

Analysis of genetic diversity
SSR data were scored as two alleles per locus distinguished by their size. We did not found multi-locus cases. The number of alleles (Na), the effective number of alleles (Ne), observed heterozygosity (Ho), expected heterozygosity (He), genetic diversity (Hs), Shannon’s information index (I) and Wright’s fixation index (F = 1-Ho/He) were calculated using GenAIEx 6.4 [38]. Since the sample size of the two population types was different, the number of distinct alleles and private alleles (not found in other populations) of the male and female population was estimated by ADZE [39], which employs a rarefaction approach to obtain sample-size corrected estimates. Detection of ‘outlier’ loci was performed with FDIST2 software [40,41]. We simulated the neutral distribution of Fst with 50,000 interactions at the 99 % confidence level.

Population structure
Genotype data from 213 M. rubra accessions were used to study population structure using the STRUCTURE v.2.0 software [42], employing an admixture model and correlated alleles frequencies. K values ranging from 1 to 10 were adopted to infer the number of clusters for ten replicate runs, with a 100,000 iterations burn-in period followed by 100,000 iterations MCMC (Markov chain Monte Carlo). The STRUCTURE HARVESTER web-based program was used to estimate the most appropriate K value [43], and CLUMPP (CLUster Matching and Permutation Program) software to calculate the average membership coefficient for each accession [44]. Accessions were assigned to a subpopulation when their membership coefficients were Q ≥ 0.6, and a bar-plot of the results prepared using DISTRUCT software [45]. Nested analysis of each of the subpopulations was using the same software and parameters as in the first STRUCTURE run.

Analysis of molecular variance (AMOVA)
The genetic variation within and among different sub-populations and subgroups of Chinese bayberry accessions and pairwise Fst were calculated by analysis of molecular variance (AMOVA), using Arlequin v3.5 software [46]. Principal coordinate analysis (PCoA) was used to further confirm the cluster analysis results, using GenAIEx 6.4 [38].

Phylogenetic analysis
Cluster analysis was conducted using the Neighbour-joining algorithm as implemented in TREECON ver.1.3 b [47]. Genetic similarity among all the accessions was estimated according to the Nei and Li genetic distance [48]. Bootstrap analysis was performed 1000 times to test the reliability of branches [49], and the tree was rooted using the related species ‘M. cerifera’ as outgroup.
Additional files

**Additional file 1: Table S1.** Diversity parameters of the 213 Chinese bayberry accessions. Na, number of observed alleles; Ne, effective number of alleles; Ho, observed heterozygosity; He, expected heterozygosity; I, Shannon’s information index; Gn, number of genotypes at each locus.

**Additional file 2: Figure S1.** The mean expected number of distinct and private alleles per locus as a function of standardized sample size for the male and female populations.

**Additional file 3: Table S2.** Analysis of molecular variance (AMOVA) based on the 84 SSR loci of 192 M. rubra accessions among six major subgroups inferred from phylogenetic tree analysis (p < 0.05).

**Additional file 4: Table S3.** Pairwise estimates of Fst based on 84 SSRs among the six major subgroups inferred from phylogenetic tree (p < 0.05).

**Additional file 5: Figure S2.** Principal coordinate analysis (PCoA) of 192 M. rubra accessions. The different colours represent the six major subgroups inferred by phylogenetic analysis. The first and second principal coordinates account for 26.78 % and 20.03 % of the total variation, respectively.

**Additional file 6: Figure S3.** Neighbour-joining tree for the 213 M. rubra accessions based on 82 SSRs. The font colour of the accession indicates the sex: red, male plant; black, female plant; blue, monoecious plant. The numbers are bootstrap values based on 1000 iterations. Only bootstrap values greater than 50 are indicated.

**Additional file 7: Table S4.** The 213 bayberry accessions and three related species included in this study. ♂♂, female plant; ♀♀, male plant; ♀♂, monoecious plant. W2011-1 collected from female plant with one branch male mutation. Genotypes of two sex associated markers are included. Accessions belonging to different subpopulations are shown based on STRUCTURE analysis.

**Additional file 8: Figure S4.** Map of China indicating the region of origin of the tree of the Chinese bayberry accessions.

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