

Article

The Musa Marker Database: A Comprehensive Genomic Resource for the Improvement of the Musaceae Family

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Abstract: Molecular markers, including Simple Sequence Repeat (SSR), Single Nucleotide Polymorphism (SNP), and Intron Length Polymorphism (ILP), are widely utilized in crop improvement and population genetics studies. However, these marker resources remain insufficient for *Musa* species. In this study, we developed genome-wide SSR, SNP, and ILP markers from *Musa* and its sister species, creating a comprehensive molecular marker repository for the improvement of *Musa* species. This database contains 2,115,474 SSR, 63,588 SNP, and 91,547 ILP markers developed from thirteen *Musa* species and two of its relative species. We found that 77% of the SSR loci are suitable for marker development; 38% of SNP markers originated from the genic region, and transition mutations (C↔T; A↔G) were more frequent than transversion. The database is freely accessible and follows a ‘three-tier architecture,’ organizing marker information in MySQL tables. It has a user-friendly interface, written in JavaScript, PHP, and HTML code. Users can employ flexible search parameters, including marker location in the chromosome, transferability, polymorphism, and functional annotation, among others. These distinctive features distinguish the Musa Marker Database (MMdb) from existing marker databases by offering a novel approach that is tailored to the precise needs of the *Musa* research community. Despite being an in silico method, searching for markers based on various attributes holds promise for *Musa* research. These markers serve various purposes, including germplasm characterization, gene discovery, population structure analysis, and QTL mapping.

Keywords: molecular markers; microsatellite; SNP; ILP; database; *Musa* spp.



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1. Introduction

The Musaceae family stands as one of the most popular and widely recognized families for fruit crops. This family is composed of three genera (*Musa*, *Musella*, and *Ensete*) with about 91 known species, which are placed in the order Zingiberales. The family is native to the tropics of Africa and Asia [1]. Members of this family, such as bananas, play a significant role in the global economy and contribute to sustainable food security in tropical and sub-tropic regions worldwide. They serve as a primary food staple for millions of people, providing valuable nutrients, calories, fiber, fodder, and raw materials for various industries [2]. There are approximately 135 countries in tropical and sub-tropical regions that cultivate bananas. Among them, India, China, Indonesia, Brazil, Ecuador, the Philippines, Guatemala, Angola, and Tanzania stand out as the top banana producers and exporters. Together, these countries contribute about 80% of the world’s banana production and trade [2–5]. However, the banana industry faces notable challenges, including diseases, pests, and climate change. Traditional methods of plant protection and cultivation are insufficient to meet the demands of the banana industry [6,7]. The triploid nature of cultivated banana cultivars limits the application of conventional breeding techniques for varietal improvements. To address these ongoing challenges

in the banana industry, researchers are exploring various opportunities to enhance crop production [6]. This involves collecting and characterizing wild and cultivated varieties, creating a large germplasm collection using both in vitro and ex vitro methods, developing genomic repositories through genome and transcriptome sequencing, and analyzing and sharing the data in the public domain [6,8]. These efforts aim to improve the resilience and productivity of this important crop.

The collection and characterization of germplasm within the Musaceae family are essential for varietal improvements. The banana research community is consistently engaged in this task, employing mostly phenotypic characteristics and traditional molecular markers such as RAPD and AFLP. Despite the continued use of these legacy markers, they possess several limitations, particularly in terms of reproducibility. Therefore, the comprehensive characterization of banana germplasm faces significant challenges. In addition to legacy molecular markers, SSR and SNP have been employed for banana germplasm characterization. However, there is a notable absence of ILP markers specific to this plant family. While most SSR markers for the Musaceae family have been developed from *M. acuminata* [9–13] with a few in *M. balbisiana* [14,15], it is crucial to expand the collection of molecular markers across all subspecies. Generating markers that cover the entire genome of this family would prove to be a valuable resource for the research community striving to enhance banana cultivars.

Molecular markers are widely used in modern agriculture for crop improvement programs. They have a wide range of applications in plant breeding, including germplasm characterization, parent selection, genetic diversity, population structure, and trait tagging. The easy accessibility of the genome sequence and molecular marker mining tools have revalorized genome-scale marker discovery and their subsequent utility. Among the different types of molecular markers, SSR, SNP, and ILP markers are frequently used in plant breeding due to their genomic abundance, easy assay techniques, and reproducibility.

The developments of genome-scale SSR, SNP, and ILP markers are straightforward, and they have become routine work for molecular breeders and bioinformaticians. Genome-wide molecular markers, especially SSR markers, have been developed for many of the sequenced plant genomes, but the molecular marker databases have been created for only a few of them. For example, the Cotton Microsatellite Database [16] was created to present 5484 microsatellite markers that were developed from nine major cotton microsatellite projects. The Kazusa Marker DataBase [17] was developed to provide linkage maps, physical maps, and information on ~68,000 SSR and ~1,400,000 SNP primers for 14 agronomical important crop species. This database provides forward and reverse primer sequences, as well as a unique primer ID linked with additional information about the primer. The NABIC marker database was developed based on 7250 markers from published articles on different crop plants [18]. Each marker in this database contains information about the marker name, gene definition, general marker information, and expressed sequence tag number. The Pigeon Pea Microsatellite Data Base (PIPEMicroDB) catalogs 123,387 instances of short tandem repeat information from the pigeon pea genome [19]. The Foxtail millet Marker Database (FmMDB) is an online searchable and downloadable database developed from Foxtail millet genome sequences. This database catalogs 21,315 genomic SSRs, 447 genic SSRs, and 96 ILP markers' information [20]. The Chickpea Microsatellite Database (CicArMiSatDB <http://cicarmisatdb.icrisat.org> (accessed on 20 November 2022)) is a relational database created by mining Chickpea genomes, and it provides information on SSRs along with their features including genomic coordination, primer-pairs, annealing temperature, and SSR repeat motifs [21]. The PIP (Potential Intron Polymorphism) marker database, developed from the genome assembly of 59 plant species, does not include any markers from *Musa* and its sister species [22]. The 'SSRome' database represents 45.1 million microsatellite markers across all taxa including plants, metazoa, archaea, bacteria, viruses, fungi, and protozoa. Users can explore microsatellites from 6533 organisms including nuclear, mitochondrial, and chloroplast genomes [23]. All the makers in this database are classified as genic or non-genic. The pineapple genomics database (PGD)

is a core online platform that integrates genomic, transcriptomic, and genetic marker (SSR, SNP, and ILP) data for pineapple [24]. The lily genomic database represents SSR and SNP markers for the lily species (<http://www.genomicsres.org/lilidb/> (accessed on 12 December 2023) [25].

Until today, many online databases have been developed to store genomic data in an accessible platform. Among these, some focus on different types of molecular markers, some store only genomic sequence data, while others integrate both marker and sequence data. Most of the molecular marker databases deal with SSR markers as described in an earlier section of this article. Although some of the existing marker databases integrate molecular markers from *Musa* sp., among them SSRome [23], PMDBase [26], Banana Genome Hub [27], and BanSatDb [28] marker databases are notable. SSRome and PMDBase only integrate SSR marker data from the *Musa acuminata* genome; BGH and TropGENE store only *Musa* SNP markers. The BanSatDb database was created for *Musa* SSR markers, and it allows for the design of SSR markers from the three genomes of *Musa* species: *M. acuminata*, *M. balbisiana*, and *M. itinerans*. BanSatDb does not permit the search of primers from the *M. schizocarpa* genome. All the existing marker databases have many limitations and their major deficiency is: (i) lacking the classification of SSR loci as class I (>20 bp) or class II (≤ 20 bp); (ii) missing polymorphism and transferability information, which is a key attribute for selecting high therapeutic markers in in silico methods; (iii) lack of associated gene function information; and (iv) missing SSR motif information such as motif richness (as AT, GC, or AT/GC balance) or the motif is located in CDS or UTR. Overall, existing marker databases generally present a limited set of information for each marker, such as forward and reverse primer sequences, SSR types, primer annealing temperature, and PCR product size. Moreover, search parameters are often inflexible and narrow.

The objective of this study was to conduct genome-wide mining and characterization of molecular markers (SSR, SNP, and ILP) and to develop a user-friendly database encompassing thirteen Banana and two Ensete species. The database will feature chromosome-wise SSR, SNP, and ILP primers, e-PCR-based polymorphism, and cross-taxa transferability. Additionally, in addressing the limitations of existing marker databases, including *Musa* and other plant species, we aim to provide a more robust platform for marker selection and analysis prior to their use in future research projects such as genetic diversity assessment, genotype characterization, and population structure studies.

2. Materials and Methods

2.1. Data Collection and Processing

The whole genome sequences of 12 banana species (Listed in Table S1) were downloaded from Musa Genome Hub (<https://banana-genome-hub.southgreen.fr/node/50/925> (accessed on 26 July 2020)). A total of 25261 Banana GSS sequences were obtained from the NCBI database. EST sequences of *Musa* spp. were obtained from the EST Tool Kit and NCBI databases. All EST sequences were merged into a single fasta file and then the “est_trimmer.pl” tool (http://pgrc.ipk-gatersleben.de/misa/download/est_trimmer.pl (accessed on 26 July 2020)) was used to remove low-quality sequences, poly A/T, and low-complexity regions at the 5′ and 3′ ends. The resulting sequences were then assembled to remove redundant sequences using CD-Hit [29] with default parameters and the non-redundant sequences were used for subsequent analysis.

2.2. Marker Development

SSR markers: The microsatellite (SSR) markers were developed using a pipeline called 3GTMAT (<https://github.com/mkbcit/3GMAT> (accessed on 26 July 2020)), with a minimum of 12 nt long SSR loci. SSR loci were characterized based on (i) the length of repeat motifs (Class I > 20 bp, Class II ≤ 20 bp) and (ii) the nucleotide composition of repeat motifs (AT-rich, GC-rich, and AT-GC balance). The E-PCR strategy was applied to assess the in silico transferability and polymorphism analysis of the developed SSR markers.

SNP markers: The EST sequences were mapped on the *Musa acuminata* genome using Bowtie2 [30] with default parameters, and then samtools [31] was used to extract SNP and Indels. A Perl script was used to extract flanking sequences of 400 bp in length, including 200 bp upstream and 200 bp downstream of the SNP locus. SNP primer pairs were designed using Primer3 [32] with default parameters. The Functional annotation and gene association of the developed SNP markers were estimated using Blast2Go [33] analysis.

ILP markers: Intron information was extracted from the gff3 file of the annotated *Musa* genomes. Then, a Perl script was used to retrieve intron flanking regions along with the 100 bp on each side of the targeted introns. Subsequently, intron lengths ranging from 100 to 3000 bp were extracted for primer design. Primer3 [32] software with default parameters was used to design the primers. The e-PCR method was used for the in silico transferability assessment of the ILP markers.

2.3. User Interface and Database Construction

The MMdb is a web-based interactive, searchable, downloadable, and relational database server, developed using MySQL 5.0 (www.mysql.com), and it has three tiers: a client-tier, middle-tier, and database tier. All the developed markers with their features have been stored in a MySQL database, which is accessed through PHP and Apache. The user-friendly web interface was designed using HTML5, Bootstrap4, CSS3, JavaScript, and jQuery.

3. Result

3.1. Genome-Wide Marker Developments

We conducted a genome-wide analysis to identify, distribute, and classify simple sequence repeats (SSRs) using 15 whole genome assemblies: thirteen from *Musa* spp. and two from *Ensete* spp., as detailed in Table S1. In total, we identified 2,761,190 SSRs, encompassing various types of desirable repeat motifs ranging from mono- to hexanucleotide repeats in the 15 datasets (see Table S1). The SSR densities exhibited variations across the tested genomes, with a range of 1425 to 5718 base pairs per SSR. An average SSR was found at every 3085 base pairs in the genome. Class II SSRs (motif length less than or equal to 20 base pairs) were more prevalent than Class I SSRs (motif length greater than 20 base pairs) in all *Musa* species. Among the tested genomes, AT-rich SSR motifs were dominant in comparison to other types of SSR motifs, including GC-rich and AT/GC-balanced motifs. All the identified SSRs were utilized for SSR marker development, resulting in 2,115,474 SSR loci successfully developed as SSR markers, representing 77% of the total SSRs. The remaining SSRs failed to develop markers, possibly due to the absence of perfect genome fragments, low-quality base representation, or insufficient flanking regions to generate primer sequences (Table 1).

Table 1. Summary of Genome-wide SSR Mining, Marker Development, and Characterizations in *Musa* and *Ensete* Species.

List of the Genome	Genome Code	Total Number of SSR	Primer Design (Count)	%
<i>Musa acuminata</i> banksii	Mabn2	183,911	135,187	74
<i>Musa acuminata</i> Dwarf_Cavendish	Madc	267,698	235,211	88
<i>Musa acuminata</i> burmannica	Mabu	141,919	125,946	89
<i>Musa acuminata</i> malaccensis V2	Maml2	147,255	118,834	81
<i>Musa acuminata</i> malaccensis V4	Maml4	185,328	132,721	72
<i>Musa acuminata</i> zebrina	Maze1	111,705	100,182	90
<i>Musa balbisiana</i> BGI11	MbBGI1	320,858	130,377	41
<i>Musa balbisiana</i> pkw	Mbpkw	131,403	106,302	81
<i>Musa beccarii</i>	Mbeci	181,767	141,466	78
<i>Musa itinerans</i>	Mitan	151,683	126,107	83
<i>Musa schizocarpa</i>	Mscz1	178,889	135,556	76
<i>Musa textilis</i>	Mtext	215,660	173,840	81
<i>Musa troglodytarum</i>	Mtrog	197,524	155,701	79
<i>Ensete glaucum</i>	Engl	181,817	155,508	86
<i>Ensete ventricosum</i> Bedadeti	Envnb	163,773	142,536	87
Total		2,761,190	2,115,474	77

We systematically characterized all the SSR markers based on their transferability, genomic location, polymorphism, locus type, and the distribution of repeat type, repeat richness, and repeat length among genomic regions. The results are presented in Figure 1a. The findings reveal that 80% of the SSRs are located in intergenic regions. Furthermore, we distributed Class I and Class II SSRs between these two regions, genic and intergenic. However, we did not observe any significant preference for the distribution of Class I and Class II SSR types among these locations in both regions. Class II-type SSRs were more frequent than Class I-type SSRs in both regions (Figure 1a–c). Single-locus markers were found to be prevalent among both *Musa* and *Ensete* species. Based on the e-polymerase chain reaction (e-PCR) results, a significant number of polymorphic markers were identified, indicating their potential utility for further applications (Figure 1d–f).

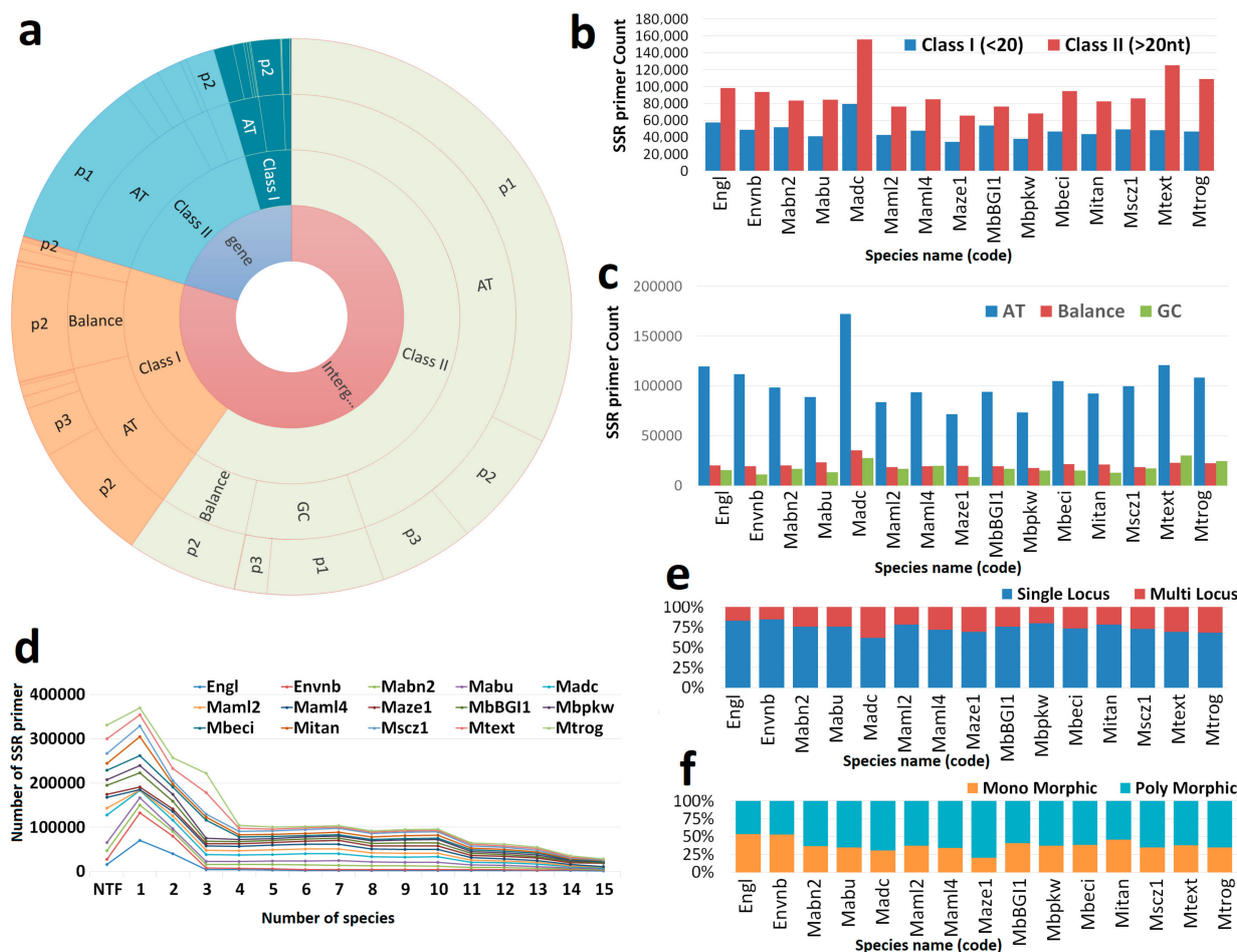


Figure 1. SSR marker development and characterization: (a) shows the distribution of Simple Sequence Repeats (SSRs) in different gene locations, divided into categories like Class I and Class II. We have further categorized these SSRs based on their richness (AT, Balanced, and GC) and displayed the specific motifs (mono to hexa) associated with them; (b) depicts the distribution of Class I and Class II SSR markers among *Musa* and *Ensete* species; (c) demonstrates the distribution of motif richness within SSR markers among *Musa* and *Ensete* species; (d) offers a comparative analysis of SSR marker transferability among different species; (e) illustrates the distribution of single-locus and multi-locus SSR markers among *Musa* and *Ensete* species; and (f) presents the distribution of polymorphic and monomorphic SSR markers among *Musa* and *Ensete* species.

In this study, SNP markers were developed and characterized based on sequences obtained from the public domain. The results are presented in Figure 2 and Supplementary Table S2. A total of 63,588 SNP markers were developed in this study, with 24,375 of

them located in the genic regions. Transition mutations ($C \leftrightarrow T$; $A \leftrightarrow G$) were found to be dominant in both regions, although their frequency was not significantly higher than that of transversion mutations. Indels are more abundant in the non-genic regions than in the genic regions.

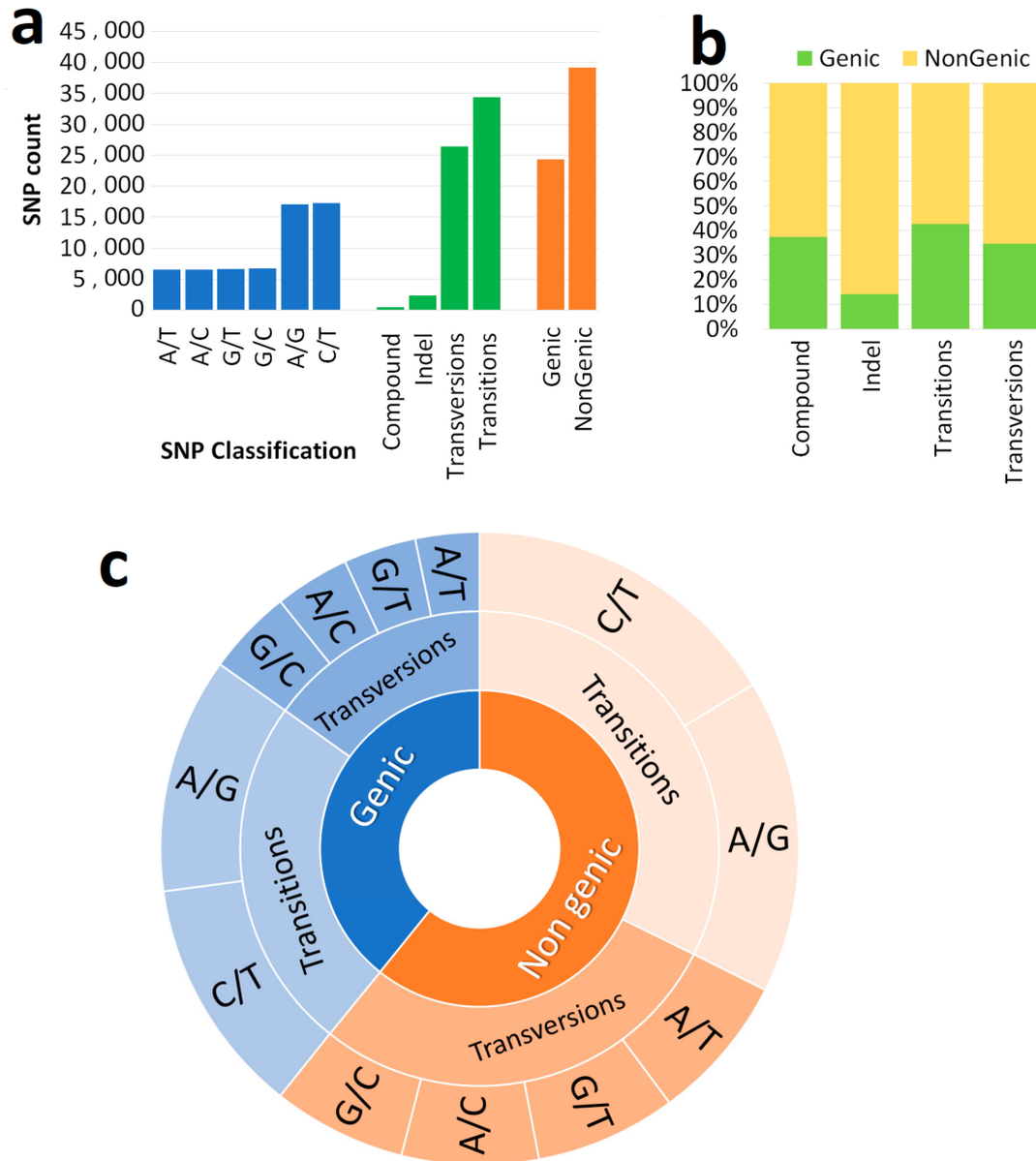


Figure 2. SNP maker mining and characterization. (a) distribution of different class of SNP, (b) distribution of SNP in genetic and non-genic regions, and (c) distribution of SNP types, classed among the genic and non-genic regions.

A total of 91,547 ILP markers were developed from the genomes of Maml2 and Mbpkw. The percentage of single and multi-locus markers significantly varied in both genomes (Maml2 and Mbpkw). The results reveal that single-locus ILP markers constitute more than 98% of both genomes, while multi-locus markers are less common. The majority of the ILP markers show monomorphism, accounting for approximately 82% of the total number (Figure 3a–c). The chromosomal level distributions of ILP markers in both genomes do not exhibit any significant differences.

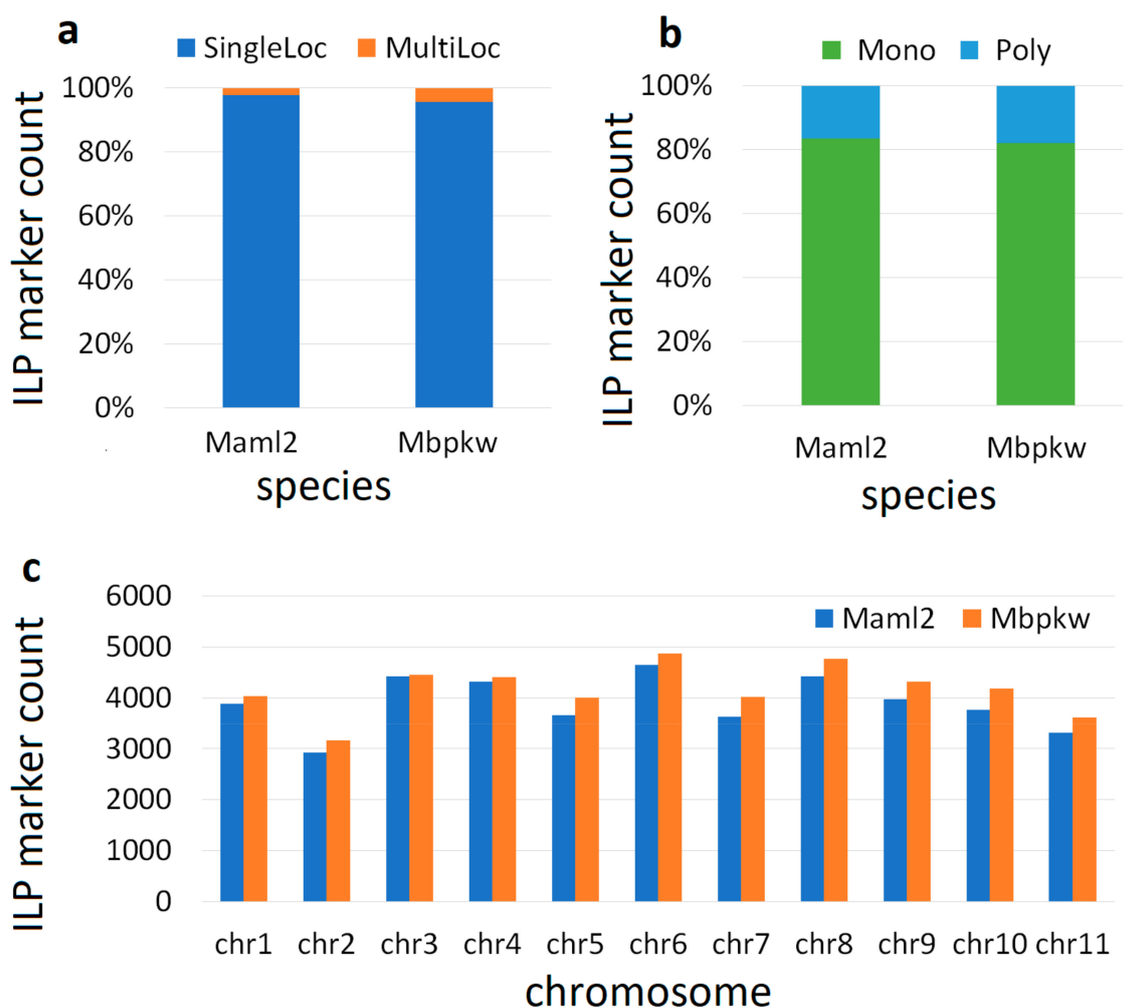


Figure 3. ILP marker development and characterization: (a) comparative distribution of single-locus and multi-locus ILP markers in two banana species; (b) distribution of polymorphic and monomorphic ILP markers in two banana species; and (c) chromosome-level distribution of ILP markers.

3.2. Interface and Search Criteria

The MMdb is a comprehensive and user-friendly resource that enables exploration, searching, downloading, and comparison of molecular markers such as SSR, SNP, and ILP across 12 *Musa* species. It features an interactive web interface accessible via a top navigation bar, enabling users to easily access various sections and tools.

The “Home” page presents an overview of the MMdb and its potential use in banana breeding (Figure 4). The “Marker” menu lists three different marker search interfaces for SSR, SNP, and ILP primers and their information. The “Download” page comprises a list and links to downloadable data, while the “Publication” page lists research articles published by our research group. The “About” page represents information about the group members involved in building this database, while the “Support” page describes the financial support information for this project.

The SSR search interface features three tabbed menus: GW-SSR (Genome-Wide SSR Marker), Novel Functional SSR Marker Search (NFS), and SSR-db v1.0. In the “GW-SSR” tab, there are five different search criteria that users can employ either individually or in various combinations. The “NFS” search page enables users to explore SSR markers from the Functional SSR marker sets developed by Biswas et al. (2020) [34]. This search interface comprises eight individual search criteria, as illustrated in Figure 4. The third tab, SSR-db

V1.0, contains SSR markers published by Biswas et al. (2015) [13]. It offers two distinct search criteria: a basic search with fourteen individual search parameters and an advanced search with seven parameters. Users can select these parameters in various combinations, empowering them to obtain more specific and tailored results.



Figure 4. A comprehensive snapshot of the Musa Marker Database (MMdb) user interface, showcasing key functionalities: (a) main menu with links to various sections, (b) homepage overview, (c) download page, (d) publications, (e) research group members, (f) funding sources, (g) SNP marker search interface with results and details, (h) detailed ILP search page, and (i) SSR marker search interface with results and details (www.genomicsres.org/mmdb/ accessed on 26 December 2023).

The SNP search page features five distinct search criteria, each operating independently based on the specific characteristics of the SNP. It returns specific types of SNP information and presents each search result in a tabulated format. Each entry in the table is linked to detailed information for the associated SNP data.

The ILP search page includes five independent search criteria, aiding users in selecting the best search results from the database. Each search yields results in a tabulated format, with each data entry linked to additional details for each ILP marker.

3.3. Unique Feature of MMdb Compared with Other Existing Marker Databases

There are currently three marker databases for Musa, including the SSRome, TropGENE Database, and BanSatDb Database. A comprehensive overview of their comparative features is provided in Table S3, offering a detailed analysis and insights into the distinct attributes of each database. However, BGH and TropGENE only store Musa SNP markers, while BanSatDb presents SSR markers with very limited data and search parameters. For example, BanSatDb only includes markers from three Musa species (*M. acuminata*, *M. balbisiana*, and *M. itinerans*). In contrast, the MMdb offers several unique features compared to other existing Musa Marker Databases. The MMdb characterizes and classifies each marker based on various parameters, such as SSR motif size, type, location in the genome (genic, non-genic, CDS, and UTR), nucleotide base composition, chromosome position, transferability to other Musa species, and in silico prediction of PCR polymorphism. Moreover, the MMdb contains a much larger number of markers and provides up-to-date marker information for sequenced Musa species. It also offers comprehensive and user-friendly tools for exploring, downloading, and comparing various molecular markers across multiple Musa species.

3.4. Utility and Future Directions

The MMdb is a valuable resource for banana researchers and breeders. It allows users easy access to a large collection of molecular markers, including SSRs, SNPs, and ILPs, along with detailed characterization and classification information. This database can aid in marker-assisted selection, genetic diversity analysis, banana germplasm management,

and genome mapping studies in *Musa* species. Additionally, the in silico transferability and polymorphism feature of the MMdb saves time and resources. This feature allows researchers to select the best marker set without requiring wet lab experiments.

In the future, the database could be extended to include more *Musa* species and additional marker types. The incorporation of functional annotation data and gene expression information could also enhance its usefulness in molecular breeding applications. Additionally, regular updates and improvements to the database interface and search features could improve user experience and make it an even more valuable tool for the *Musa* research community.

4. Discussion

Molecular markers play a crucial role in modern plant breeding by allowing breeders to identify and track desirable traits, evaluate genetic diversity [35,36], and develop more targeted breeding programs. These markers can speed up the breeding process by helping to select plants that are more likely to possess desirable traits, as well as aiding in the understanding of the genetic basis of complex traits and the development of molecular breeding strategies to improve these traits. Molecular markers are particularly useful in crops with long breeding cycles or complex traits that are difficult to measure directly. The availability of whole genome sequences has enabled the development of various types of molecular markers and the creation of molecular marker databases to maximize their utility [37]. In this study, a novel molecular marker database was developed from the whole genome sequences, EST, and GSS sequences of 13 *Musa* species. SSR, SNP, and ILP markers were mined and characterized for optimal marker data quality, resulting in the creation of a searchable *Musa* Marker Database that is freely accessible from the following link (www.genomicsres.org/mmdb/ accessed on 26 December 2023).

In terms of SSR analysis, our study identified a substantial number of SSRs, amounting to 2,761,190, with variable densities in the tested genomes. The prevalence of Class II SSRs (shorter than 20 bp), characterized by shorter motifs, was consistent with the previous findings in plants [37–39]. Furthermore, the dominance of AT-rich SSR motifs aligns with similar observations in *Musa* and other plant species [40–42]. The high percentage of successfully developed SSR markers (77%) suggests the potential for efficient marker development in these genomes. This outcome is in line with previous research demonstrating the utility of SSR markers for genetic mapping and breeding applications.

Our study's exploration of SNP markers in genic and non-genic regions also adds to the growing body of knowledge regarding polymorphism in plant genomes. The higher occurrence of transition mutations (C↔T; A↔G) over transversions observed in both regions corresponds with studies in various plant species [43,44]. Additionally, the prevalence of Indels in non-genic regions resonates with previous reports of the non-coding regions being more prone to structural variations [45]. These findings underscore the importance of considering both genic and non-genic markers for genetic diversity and trait association studies.

Introns, noncoding regions within genes, are transcribed but subsequently spliced out during RNA processing [46]. The variability in the number and length of introns among species highlights their vital role in gene regulation. Throughout evolutionary processes, genes can gain or lose introns, contributing to genomic diversity [46,47]. Intron-based genomic variation can be used for molecular marker development. In the past decade, several studies on plant species have highlighted the utility of Intron Length Polymorphism (ILP) markers as valuable tools in molecular genetics, similar to other markers like SSRs and SNPs [48,49]. However, ILP markers offer distinct advantages, being locus-specific, co-dominant, and facilitating the accurate detection of heterozygosity. By specifically detecting polymorphisms within genic regions, ILP markers capture functionally relevant genetic variation, aiding in the identification and selection of desirable traits for crop improvement and germplasm conservation in plant breeding research [49]. To date, ILP markers have not been available for *Musa* species. Here, we report genome-wide ILP markers from two banana species. Our results show that the dominance of single-locus ILP markers and

their high monomorphism rate (82%) are consistent with studies in other plant species [50]. However, the fact that multi-locus ILP markers are less common raises questions about their potential applications, which may differ from those of single-locus markers. Further research may be needed to explore the specific utility of multi-locus ILP markers in these genomes. It is important to note that the chromosomal distribution of ILP markers showed no significant differences between the genomes of Maml2 and Mbpkw, indicating that the genomic context does not strongly influence the distribution of these markers. This finding contrasts with some studies that have reported uneven distributions of markers across chromosomes [40,50], suggesting the need for further investigation into the underlying factors governing marker distribution.

There are three databases (SSRome, TropGENE, and BanSatDb) containing molecular markers from *Musa* species [23,28]. However, these databases have significant limitations, such as outdated information, inflexibility in data search criteria, and overall they are less user-friendly. They prove time consuming for data retrieval and lack crucial details, such as primer redundancy, transferability, and polymorphism information for other *Musa* or non-*Musa* species. Furthermore, there is no unique identifier for designed primers, potentially impacting reproducibility in downstream marker applications by different researchers. In contrast, the MMdb addresses these limitations, offering a user-friendly interface specifically designed to assist researchers, making it a potential model database for other plant species. The current version of the database contains six times more SSR marker data than the BanSatDb database [28]. The MMdb has a flexible five-search criteria that can use single or indifferent combinations, which leads to the user picking the best primary pairs for their research interest, this facility is absent in the BanSatDb database as well as many other SSR marker databases [25,51].

Although ILP markers have demonstrated their usefulness in plant breeding and genetics, there have been few studies aimed at developing genome-wide ILP markers and databases. One notable example is the PIP marker database, developed by Young et al. in [22], comprising 57,658 ILP markers derived from 59 plant species. However, it is noteworthy that this database does not include markers from *Musa* species, and presently, PIP is not accessible online, rendering data retrieval impossible. In contrast, the Foxtail millet Marker Database (FmMDb) has cataloged 96 ILP markers for the Foxtail millet genome sequences [20]. Currently, the MMdb is the largest ILP marker repository among molecular marker databases. Its ILP marker search page enables users to easily locate and download ILP markers using various flexible search criteria, representing a significant advancement over other existing marker databases.

5. Conclusions

In conclusion, the *Musa* Marker Database (MMdb) is a user-friendly and inclusive genomic resource for the Musaceae research community, that offers its uses to explore, download, and compare 2,115,474 SSR, 63,588 SNP, and 91,547 ILP markers from thirteen *Musa* species and two *Ensete* spp. The unique features of this database such as marker transferability, ePCR validation, and polymorphism information of each marker make it stand out from other existing marker databases. The current version of the MMdb contains a larger number of markers and up-to-date data for *Musa* species. This resource can be used to study genetic diversity and germplasm characterization, develop MAS strategies, perform GWAS, and compare markers across different *Musa* species. The database can be accessed via this link www.genomicsres.org/mmdb/ accessed on 26 December 2023.

Supplementary Materials: The following supporting information can be downloaded at: <https://www.mdpi.com/article/10.3390/agronomy14040838/s1>, Table S1. Summary of data and SSR mining of *Musa* species. Table S2. Summary of the SNP Mining and Characterization. Table S3. Comparison of the MMdb with other related databases.

Author Contributions: The work presented here was carried out in collaboration among all authors. M.K.B. was involved in SSR, SNP, and ILP analysis and database development and drafted the manuscript; D.B. wrote and managed the server to host the database; G.D. participated in SSR data analysis and revised the MS; M.K.B. and G.Y. conceived and designed the experiments. All authors have read and approved the final manuscript.

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Data Availability Statement: The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

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