Genome-wide association study for leaf area, rachis length and total dry weight in oil palm (*Elaeis guineensis*) using genotyping by sequencing

B. Kalyana Babu*, R. K. Mathur, G. Ravichandran, P. Anitha, M. V. B. Venu

ICAR-Indian Institute of Oil Palm Research, Pedavegi, West Godavari District, Andhra Pradesh State, India

* kalyan_biotech@yahoo.co.in, B.Babu@icar.gov.in

Abstract

The marker-trait association for complex traits using genotyping by sequencing (GBS) method is being widely spread in plants. The study aimed to identify significant single nucleotide polymorphism (SNP) associations for rachis length (RL), leaf area (LA) and total dry weight (TrDW) in oil palm among diverse African germplasm. The Illumina NextSeq platform has been used for SNP genotyping and retained 4031 fully informative SNPs after applying the filter criterion. These 4031 SNPs were used for genome wide association study for the above three traits. The LD decay rates of the African germplasm using GBS data of SNP is observed to be 25 Kb at 0.45 of average pair wise correlation coefficient ($r^2$). Association mapping led to the identification of seven significant associations for three traits using MLM approach at a P value of $< 0.001$. Three associations were identified for total dry weight, two each for leaf area index and rachis length. The qtlLA1 was found to be highly significant at a P value of $7.39E^{-05}$ (18.4% phenotypic variance) which is located on chromosome 4. Two QTLs (qtlLA2 and qtlRL1) were located on chromosome 1, which explained 11.9% and 12.4% of phenotypic variance respectively. Three QTLs for total dry weight were located on chromosome 2, 14 and 16, all-together explained 40% phenotypic variance. The results showed that the SNP-trait associations identified in the present study could be used in selection of elite oil palm germplasm for higher yields.

Introduction

Oil palm (*Elaeis guineensis* Jacq.) belongs to the family Arecaceae, which is a major contributor (approximately 40%) of world edible vegetable oil. The genome size of oil palm is 1.8 Gb with haploid chromosome number of 16. The genome sequence of an elite dura palm was published [1] and freely available in the open source. It is a perennial crop having potential oil yield capacity of 10 t/ha, against the current yields which varied between 2 to 6 t/ha [2]. Biotechnological tools like molecular markers facilitates in bridging the gap for genetic improvement in yield, oil quality and other important agro-morphological traits of interest. Leaf area and rachis
length are found to be highly correlated with oil palm yield which is proven in other crops as well. Hence, leaf area and rachis length are considered as important traits influencing the yield in oil palm. Several studies using GBS for genome-wide association study (GWAS) were reported in rice [3], date palm [4] and oil palm [5], but no reports is available for leaf area, rachis length and total dry weight in oil palm using African germplasm. Molecular markers like RFLPs, AFLPs [6], EST-SSRs [7] and SSRs [5] were used for identification of QTLs for important traits in oil palm. However, genome-wide association study (GWAS) is often hindered due to lack of high diverse genetic pool and well structured germplasm [8]. The GWAS analysis using SNPs emerged as a powerful tool initially in human population to identify complex diseases and then widely implemented in several crops including rice [9], foxtail millet [10–11], maize [12] and in some tree species [13]. The genotyping by sequencing (GBS) method based on restriction digestion of DNA samples has been widely used for identification of QTLs. In oil palm, there are few reports on GBS for GWAS studies for different traits like oil to bunch, oil to dry mesocarp etc. which were mostly based on linkage mapping populations. Recently, Ithnin et al. [14] performed GWAS for oil yield and vegetative traits where they identified 19 QTLs for 8 traits. However, all the studies were reported on F1 populations, which lacks clear segregation pattern which can be improved by using advanced segregating generations or highly diverse natural population for association mapping [15]. Till now, no report is available on GWAS for leaf area, rachis length and total dry weight content using SNPs by association mapping.

In perennial crops, natural populations offer advantages like much higher mapping resolution, greater allele number and broader reference population, and less research time in establishing an association [16]. The African sub-continent holds a large diverse germplasm which has highly potential oil yielding and quality traits [17]. In the last 50 years, there was high increase in crude palm oil yield (2.0 to 4.1 t/ha) by conventional breeding, which is still much lower than the theoretical estimates of 18 t/ha [18]. Oil palm yield is highly influenced by leaf area, rachis length and total dry weight content [19]. The shape, leaf area and rachis length of crowns determine light interception and thus influence yield of oil palms (Elaeis guineensis Jacq.) by increasing the photosynthetic efficiency as evidenced by high significant correlation observed by several workers [14]. The long breeding cycle of oil palm makes a complicated process for oil palm improvement by conventional breeding. Hence, molecular markers like SNPs plays important role in identification of markers linked to important traits. Hence, in the present study African germplasm were used for identifying SNP-trait associations using GWAS by GBS method. The objectives of the study are 1) Phenotypic characterization of oil palm germplasm for leaf area, rachis length and total dry matter weight, and 2) GWAS analysis for identification of SNP-trait associations for leaf area, rachis length and total dry weight content in oil palm germplasm.

Methodology

Plant materials and DNA extraction

Oil palm germplasm representing four African countries were used for SNP genotyping (S1 Fig). The germplasm used in the study were obtained from Malaysian Palm Oil Board (MPOB), Malaysia as part of MoU between ICAR-Indian Institute of Oil Palm Research (ICAR-IIOPR) and MPOB. The palms are grown at the ICAR-Indian Institute of Oil Palm Research (ICAR-IIOPR), Pedavegi, Andhra Pradesh, India experimental fields as per the standard procedures (latitude 16° 48’N, longitude 81° 07’E). All the germplasm belongs Dura fruit type. The 96 oil palm germplasm were treated identically in terms of fertilizer application and pest controls. The samples selection was based on the phenotypic variation of selected...
morphological traits. The experiments were conducted with the consent of Director, ICAR-IIOPR, India. The genomic DNA of oil palm genotypes was isolated by standard method as described by Murray and Thomson [20] with few modifications as described in Babu et al. [21]. The genomic DNA quality and quantity was checked on 0.8% agarose gel using lambda DNA as standard.

Phenotypic data analysis
The phenotypic data of leaf area, rachis length and total dry weight content of oil palm were recorded for four consecutive years (2010–2014) according to Corley et al. [22]. The traits used for association mapping are leaf area, rachis length and total dry weight content. Statistical analysis of the traits was estimated for their mean, range, standard deviation and 95% confidence interval. The phenotypic data of rachis length, leaf area and total dry weight is given in S1 Table. The correlation coefficient between the traits and the above statistical parameters calculated using the software JMP [23] and correlation thresholds were considered significant at alpha of 0.001 and 0.01. The distribution of phenotypic data of leaf area and rachis length is given in Fig 1.

Genotyping by sequencing, SNP calling and annotation
SNP genotyping among the oil palm genotypes was done by following the GBS approach as given by Elshire et al. [24] with ApeKI restriction enzyme used for complexity reduction. The library was prepared using paired end method in the Illumina NextSeq 500 platform. The GBS was done by outsourcing to the Bio Serve (India, Pvt. Ltd). Data normalization has been done at minimum allele frequency (MAF) of 0.05, and by removing 80% missing SNP calls. Only SNP calls >90% has been considered for further analysis. The sequences were mapped to oil palm reference genome [1] and SNPs were called using TASSEL ver. 4.3.10 GBS pipe line [25].

Genome-wide association mapping
The population structure analysis was done by model based clustering method [26] as adopted in the STRUCTURE v2.3.4 software [27]. Marker-trait associations were performed by using data of 96 oil palm germplasm, SNP genotypic data by GBS approach and Q matrix obtained from the population structure data by using the software TASSEL [28]. The GWAS was performed using MLM approach embedded in the TASSEL software. The kinship matrix was used in addition to the genotypic, phenotypic and Q matrix data in the MLM approach. The significant threshold for the association was set at P of $1 \times 10^{-4}$. The genome wide patterns of LD decay were estimated by average of $R^2$ in distance intervals across the 16 chromosomes of whole genome of oil palm.

Results and discussion
Genotyping using GBS for SNP discovery
A set of 96 diverse germplasm from four African countries viz., Cameroon, Tanzania, Guinea-Bissau, and Zambia were used for SNP discovery by genotyping by sequencing method. The genomic DNA of 96 African germplasm were subjected to ApeKI digested paired end libraries and sequenced on Illumina NextSeq500 platform. The sequencing resulted in 325 million reads covering 50.78 Gb, at a mean of 3.4 million reads per sample. The sequence data obtained by GBS is lesser than earlier studies [29–31]. Pootakham et al. [13] obtained 524 million reads in F1 oil palm population. Teh et al. [32] obtained 900 million raw reads from 132 oil palm F1 progeny. The GBS raw data was imputed to SNP calling and data normalization.
has been done using MAF of 0.05 and by removing 80% missing SNP calls. Only SNP calls >90% has been considered for further analysis. A total of 325 million raw reads generated, of which 2.4 million SNPs passed bar coding and quality thresholds. Finally, 4031 SNPs were used for mapping to the reference genome with a range between 157 on chromosome 15 to 455 on chromosome 1 (S2 Table). The physical location of SNPs mapped on the oil palm genome was given in Fig 2. The number of SNPs was similar to the earlier reports [14] where they identified 4451 SNPs across the 422 samples. The SNP allele data files were given in supplementary information (S1 Text).

The population structure analysis

The population structure analysis is conducted among African germplasm of oil palm to estimate the population structure and admixture using SNP markers. To estimate the exact population structure among the oil palm germplasm, Ks were ran from 1 to 10 with 10 iterations and grouping was done based on LnP(D) values. The highest peak value of ΔK was observed at K = 4. High amount of admixture also presented among different groups which may be due to involvement of exotic germplasm. The same type of admixture was also found in earlier reports [32]. The structure analysis revealed that there are four structural patterns existed among the African germplasm which divided them into four groups (Fig 3). The grouping pattern observed to be according the geographic origin of the germplasm to some extent. High amount of admixture of alleles also evident in the populations, which showed a high
germplasm exchange between African countries for better breeding programmes. It is also evident that all the present germplasm of oil palm were derived from two Bogor palms, hence high admixture percentage is present.

Fig 2. The location of QTLs on oil palm chromosomes.
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Fig 3. The population structure among the 96 African germplasm as revealed by SNPs by genotyping by sequencing method.
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**Linkage disequilibrium (LD)**

The pattern of LD decay is crucial for identifying significant QTLs of any association mapping studies [33] and for interpretation of association peaks [34]. To estimate the mapping resolution for genome wide association mapping, the average extent of LD decay was quantified. The LD decay rates of the oil palm germplasm using GBS data of SNP was 25 Kb at 0.45 of average pair wise correlation coefficient ($r^2$). The long range LD in the highly diverse populations can be further explained by breeding selections. The process of selective sweep will much enhance the breeding value of the crops like oil palm. The LD of oil palm plantations in general decays considerably faster than in annual crops like rice and foxtail millet. In both the crops LD decays 100 Kb to 200 Kb (2–5), which decays to 50% of its initial value by 1 Kb with in 150 Kb. This difference may be due to low genome coverage of markers and fewer genotypes in the study. Similar LD decay also observed by Teh et al. [32] where they also found 25Kb and 20Kb at 0.12 and 0.15 of average R$^2$ value.

**Genome wide association study (GWAS)**

High yielding oil palm germplasm is an immediate output required for the farming community. There is a big gap in the theoretical CPO yield (18 tons/ha) and global actual average yield (2.0 to 4.0 tons/ha) over the past 50 years [18]. Due to long breeding cycle of oil palm, it became a complicated process in selection of high yielding germplasm. Oil palm yield is highly influenced by leaf area and rachis length. These are complex, quantitative and often controlled by large number of genetic loci [35]. It is important to search for effective QTLs associated with these important traits and utilizing in identification of desirable palms with good yield related traits. Hence, the present investigation aimed to identify significant QTLs using SNPs by genotyping by sequencing method. The study represents the first GWAS study of African germplasm by association mapping approach. Association of leaf area, rachis length and total dry weight using SNPs, resulted identification of seven highly significant QTLs at a P value of ≤ 0.001 by MLM approach. A high significant cut off was given in order to remove the errors generated due to false associations as expected. Three SNP loci were identified to be associated with total dry weight content, followed by two each for leaf area and rachis length. The physical location of QTLs linked to three traits on chromosomes given in Fig 2. All the seven significant QTLs were identified on five chromosomes viz., 1, 2, 4, 14, and 16 which explained a phenotypic variance from 11.5 to 18.5%. Two significant QTLs were identified for leaf area, which together explained 30.5% of phenotypic variance. Among them, qtlLA1 (SGI|741241788|) found significant at P value of 7.39E$^{-05}$ (18.5% phenotypic variance). The qtlLA2 (SGI|741241880|) linked at P of 0.001 and explained 12% R$^2$ value. These two QTLs located on chromosome 4 and 1 respectively. The manhatton plot for leaf area given in Fig 4.

Three QTLs for total dry weight content and two for rachis length were identified. The loci for total dry weight content were located on chromosome 2, 14 and 16. The QTL (qTDM1) located on chromosome 14 found significant at P of 0.0005 which explained 15% phenotypic variance. The three QTLs for total dry weight content together explained 42% of total phenotypic variance. Likewise two QTLs were found significant for rachis length. The manhatton plots for rachis length is given in Fig 5. These two QTLs located on chromosome 1 and 16, together explained 24% of R$^2$ value (Table 1). The present study results are similar to the earlier reports [14–15, 31]. Ithnin et al [14] and Billottee et al [31] found significant SNP loci for rachis length on chromosome 2, 4, 10 and 16 on F$_1$ progeny by linkage mapping studies. In the present study also QTLs for rachis length identified on 16$^{th}$ chromosome along with unique loci on chromosome 1. The results showed that chromosome 16 harboring the major QTL for rachis length which need attention in cloning the whole gene for rachis length. Billottee et al
also identified QTLs for rachis length on 2, 4 and 16 chromosomes. These two reports were based on F1 progeny, however the present study is based on genome-wide association study which gives more polymorphism and high precision in identifying QTLs for specific traits. In addition to the above results, the study also identified unique loci for leaf area and rachis length. The identified QTLs in the present study helps in selection of desired oil palm germplasm for high yield characters in different genetic backgrounds which helps in reducing time and increases accuracy.

**Conclusion**

The genotyping by sequencing method used for GWAS study of African germplasm using GBS method to identify the QTLs for three yield related traits viz., leaf area, rachis length and total dry weight content. Association mapping resulted in identification of seven highly significant QTLs for different traits at a P value of ≤ 0.001 by MLM approach. More genetic loci were identified to be associated with total dry weight, followed by leaf area and rachis length. The QTLs identified in the present study could be used in selection of elite oil palm germplasm for higher yields.
Table 1. The details of significant QTLs identified by GBS method and their P values and phenotypic variances.

| Trait            | QTL name | Linked SNP loci | P value    | R² (%) | Chromosome number |
|------------------|----------|-----------------|------------|--------|-------------------|
| Leaf Area        | qTLA1    | SGI|741241788|         | 7.39E-05 | 18.5 | 4         |
|                  | qTLA2    | SGI|741241880|         | 0.001  | 12   | 1         |
| Total dry weight content | qTTDM1 | SGI|741241610|         | 0.0005 | 15   | 14        |
|                  | qTTDM2   | SGI|741241857|         | 0.0009 | 13.5 | 2         |
| Rachis length    | qTRL1    | SGI|741241880|         | 0.001  | 12.5 | 1         |
|                  | qTRL2    | SGI|741241589|         | 0.001  | 11.5 | 16        |

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Supporting information

S1 Fig. The map showing the distribution of oil palm germplasm in four African countries. (DOCX)

S1 Table. The phenotypic data of the traits under association mapping. (DOCX)

S2 Table. The number of SNPs chromosome wise along with their positions. (DOCX)

S1 Text. Allele calling data files for some markers. (DOCX)

Author Contributions

Conceptualization: B. Kalyana Babu, R. K. Mathur.

Investigation: B. Kalyana Babu, M. V. B. Venu.

Methodology: B. Kalyana Babu.

Resources: B. Kalyana Babu.

Supervision: B. Kalyana Babu.

Writing – original draft: B. Kalyana Babu.

Writing – review & editing: B. Kalyana Babu, R. K. Mathur, G. Ravichandran, P. Anitha.

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