

Identification of main effect and epistatic quantitative trait loci for morphological and yield-related traits in peanut (*Arachis hypogaea* L.)

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Abstract An effort was made in the present study to identify the main effect and epistatic quantitative trait locus (QTL) for the morphological and yield-related traits in peanut. A recombinant inbred line (RIL) population derived from TAG 24 × GPBD 4 was phenotyped in seven environments at two locations. QTL analysis with available genetic map identified 62 main-effect

QTLs (M-QTLs) for ten morphological and yield-related traits with the phenotypic variance explained (PVE) of 3.84–15.06%. Six major QTLs (PVE > 10%) were detected for PLHT, PPP, YPP, and SLNG. Stable M-QTLs appearing in at least two environments were detected for PLHT, LLN, YPP, YKGH, and HSW. Five M-QTLs governed two traits each, and 16 genomic regions showed co-localization of two to four M-QTLs. Intriguingly, a major QTL reported to be linked to rust resistance showed pleiotropic effect for yield-attributing traits like YPP (15.06%, PVE) and SLNG (13.40%, PVE). Of the 24 epistatic interactions identified across the traits, five interactions involved six M-QTLs. Three interactions were additive × additive and remaining two involved QTL × environment (QE) interactions. Only one major M-QTL governing PLHT showed epistatic interaction. Overall, this study identified the major M-QTLs for the important productivity traits and also described the lack of epistatic interactions for majority of them so that they can be conveniently employed in peanut breeding.

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Introduction

Peanut (*Arachis hypogaea* L.) is a legume and oilseed crop with high protein content. Globally, it is grown in

an area of 25.7 million ha with the production of 42.3 million tons (FAOSTAT 2016). Enhancing the overall productivity by improving the traits like resistance to diseases, tolerance to drought, enhanced oil content, and improved quality traits is the main objective in peanut breeding. However, many of these traits are genetically complex, and achieving significant gains therefore requires the applications of genomics-assisted breeding (GAB) (Varshney et al. 2013).

There has been a tremendous advancement over the last decade in understanding the peanut genome. The genomes of the diploid progenitor species of cultivated peanut have been sequenced (Bertioli et al. 2016; Chen et al. 2016b). Recently, the chloroplast genomes of seven species have also been sequenced (Yin et al. 2017) to understand the genetic relationships of the wild species with the cultivated peanut. A molecular phylogeny based on the complete chloroplast genome sequences provided the best resolution of the seven *Arachis* species. A large number of different types of markers have been developed (Zhao et al. 2017), and a high-throughput genotyping assay with 58 K informative single nucleotide polymorphism (SNP) markers was developed (Pandey et al. 2017a). With these markers, the genetic maps of varying saturation levels have been developed and employed for mapping various traits (Vishwakarma et al. 2017). Genome-wide association studies were also conducted to map the traits like disease resistance, quality, drought tolerance and yield components (Pandey et al. 2014a) and agronomic traits (Zhang et al. 2017). Further, QTL-Seq has identified the candidate genes for LLS and rust resistance (Pandey et al. 2017b). Also, the RNA-Seq study has revealed the transcripts that are differentially expressed under LLS infection (Han et al. 2017).

The markers identified for foliar disease resistance were validated (Sujay et al. 2012; Yeri et al. 2014; Sukruth et al. 2015) and employed for molecular breeding (Varshney et al. 2014; Yeri and Bhat 2016). Marker-assisted breeding has also been successful in enhancing the oleic acid content and resistance to nematode. However, the molecular breeding for other traits in peanut demands a thorough detection and analysis of QTL and markers. Identification of QTL with main effects (or individual effects), epistatic effects, and $G \times E$ interactions is very useful in genomics (Carlborg and Haley 2004). In peanut, many researchers have reported additive, non-additive (including epistatic genetic) effects for traits like pod yield, number of pods and seeds per

plant, hundred seed weight, pod length, and shelling out-turn based on the genetic analysis (Layrisse et al. 1980; Dwivedi et al. 1989; Upadhyaya and Nigam 1998). However, identification of M-QTLs, E-QTLs, and $G \times E$ interactions for the agro-morphological and productivity traits will be useful for molecular breeding in peanut.

Availability of an appropriate mapping population with a large number of recombinants, phenotyping over multi-environments, and multi-seasons would allow precise detection of reliable (consistent) QTLs for the target traits. Therefore, the present study employed the RIL mapping population derived from TAG 24 \times GPBD 4 for identifying the M-QTLs, epistatic-QTLs, and QTL \times environment (QE) interactions for the important agro-morphological and yield traits in peanut.

Materials and methods

Plant materials

This study used a mapping population consisting of 268 RILs of TAG 24 \times GPBD 4 developed at the University of Agricultural Sciences (UAS), Dharwad, India. GPBD 4 is a very popular and high-yielding Spanish bunch cultivar with resistance to foliar fungal diseases, early maturity, desirable pod and kernel features, and high oil content (Gowda et al. 2002). TAG 24 is a Spanish bunch cultivar (Patil et al. 1995) which suffers significant yield loss under severe foliar disease infection.

Phenotyping and statistical analysis of mapping populations

The RIL-4 mapping population of F_{9-12} generations was phenotyped for three environments namely post-rainy 2007 (R4EI), rainy 2007 (R4EII), and post-rainy 2008 (R4EIII) at ICRISAT, Patancheru, India (17°30'39.8"N 78°16'30.8"E), and four environments at UAS-Dharwad, India (15°29'30.1"N 74°59'00.9"E), viz., rainy 2007 (R4EIV), rainy 2008 (R4EV), rainy 2009 (R4EVI), and post-rainy 2009 (R4EVII). The RILs were planted in augmented plot design with parents as checks in 15 blocks of 20 rows in R4EI; Alpha design with three replications in R4EII and two replications in R4EIII, and randomized block design (RBD) with two replications in R4EIV, R4EV, R4EVI, and R4EVII. Observations on 14 agro-morphological traits, namely, days to flowering (DF), plant height (PLHT), leaf length

(LLN), leaf width (LWD), pod length (PLN), pod width (PWD), seed length (SLN), number of primary branches (PBR), pods per plant (PPP), yield per plant (YPP), yield in kg per hectare (YKGH), hundred seed weight (HSW), shelling percentage (SLNG), and days to maturity (DM) were recorded at ICRISAT-Patancheru. The RILs were phenotyped for 12 yield-contributing traits at R4EI and all 14 traits measured in R4EII and R4EIII. A total of eight traits viz., PLHT, LLN, LWD, PBR, PPP, YPP, HSW, and SLNG were phenotyped at R4EIV and R4EV; four traits, PLHT, PBR, LLN, and LWD at R4EVI and five yield-contributing traits, PLHT, PPP, YPP, HSW, and SLNG at R4EVII.

Restricted Maximum Likelihood (REML) analysis was performed using GenStat version 12.0 (GenStat Committee 2010) to estimate the variance components for the data collected at ICRISAT, Patancheru. Analysis of variance (ANOVA) was performed using WindowStat, version 8.5 (IndostatServices, Hyderabad, India, <http://members.fortunecity.com/indostat/>) for the data collected from each season at Dharwad location. The parameters like genetic variance (σ^2_g) and broad sense heritability ($h^2_{b.s.}$) were estimated using GenStat version 12.0 and WindowStat version 8.5 for the data collected at Patancheru and Dharwad location, respectively. Broad sense heritability was calculated as $h^2_{b.s.} = (\sigma^2_G / \sigma^2_P)$ for each environment where, σ^2_G represented the genotypic variance, σ^2_P is the phenotypic variance ($\sigma^2_G + (\sigma^2_e \text{ or error residual/number of replications})$). The correlation coefficients (r) among the traits were estimated in each season using an R software (R Core Team 2013).

QTL analysis

The genetic map, carrying 188 mapped markers, constructed previously by Sujay et al. (2012), was used for QTL analysis. Candidate interval selection, putative QTL detection, and QTL effects were estimated from each environment (three from ICRISAT and four from Dharwad) separately. The normalized predicted mean from the REML estimates in Patancheru location and mean phenotypic data across replications in Dharwad location were used to map QTLs by “Composite Interval Mapping (CIM)” approach (Zeng 1994) using WinQTL Cartographer 2.5 (Wang et al. 2007). CIM was carried out using Model 6 with a moving window size of 10 cM and a walking speed of 2 cM. The forward-backward stepwise regression method was

used to select number of marker cofactors for the background selection. To define QTL region automatically, “Locate QTLs” option was used with a minimum distance of 5 cM between QTLs. The highest peak was considered to locate QTLs if the peak distance between the QTL is less than 5 cM. For each trait, a 500-permutation test with a significance level of 0.05 was performed to detect an appropriate LR. Based on significant LR peak for map position under consideration, QTL position, additive effects, favorable allele contribution, and phenotypic variation explained (PVE) were estimated. An estimate of epistatic QTL was studied by mixed linear composite interval mapping (MCIM) using software QTL Network Version 2.0 (Yang et al. 2008). The 2D Genome Scan option was used to map epistatic QTLs. A 1000 permutation was applied to calculate critical F value and the QTL effects were estimated using Monte Carlo Markov chain method with 20,000 Gibbs sample size. The genome scan configuration was performed using testing window size of 10 cM, walk speed of 1 cM, and filtration window size of 10 cM.

Results

Phenotypic variability

REML analysis for the data collected from Patancheru and ANOVA for the data collected from Dharwad showed significant differences among the RILs for majority of the traits except LLN, LWD, YPP, and YKGH in R4EI; DF, SWD, and DM in R4EII; LWD, PBR, SWD, and DM in R4EIII; PBR and LWD in R4EIV; PBR and LWD in R4EV; PBR in R4EVI (Supplementary Table S1). The mean values for all the traits, except PBR in R4EII, were normally distributed (Fig. 1). The phenotypic range of variation was high for all the traits, except for DF in R4EI and R4EIII, PBR in R4EI and R4EII, LWD in R4EII, and PWD and SLN in R4EII and R4EIII. Transgressive segregants were observed for all the traits in all the seven environments at both (Patancheru and Dharwad) the locations. The broad-sense heritability estimates ($h^2_{b.s.}$) ranged from low (0.18, YPP in R4EIV), moderate (0.26, HSW in R4EII; 0.30, LWD in R4EII), high to very high (0.91, PPP in R4EV). Briefly, the heritability estimates were observed for DF (0.36–0.85), PLHT (0.43–0.86), LLN (0.38–0.85), LWD (0.30–0.42), PBR (0.38–0.66), PLN (0.54–0.70), PPP (0.32–0.91), PWD (0.38–0.53), SLN

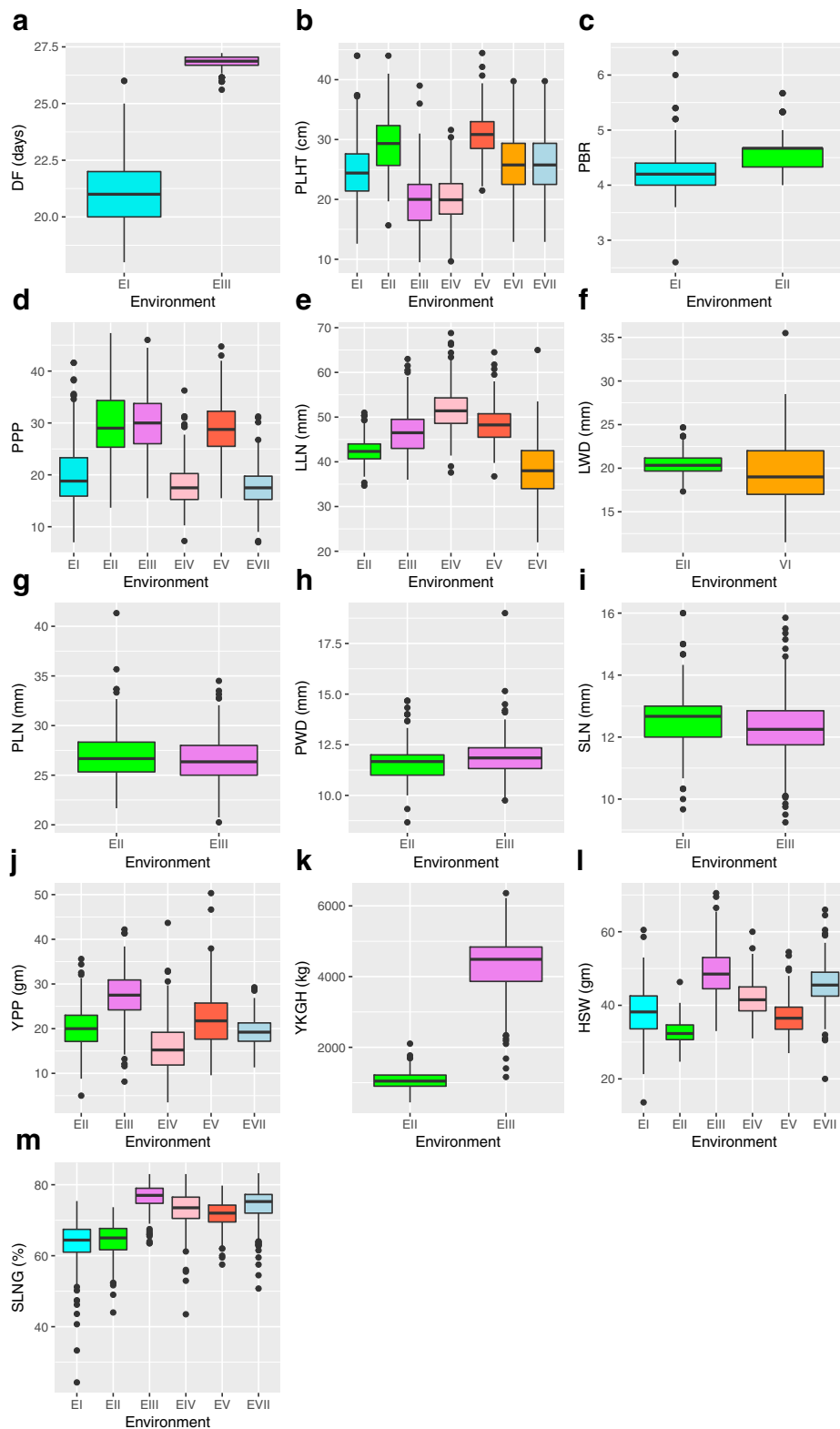


Fig. 1 Boxplot showing the distribution of the means for agro-morphological traits among the RILs over different environments

(0.66–0.77), YPP (0.18–0.51), YKGH (0.63–0.81), HSW (0.26–0.85), SLNG (0.47–0.82), and DM (0.61).

Phenotypic correlations

We studied the degree of relationship between the traits in each season by estimating the Pearson's (r) pairwise correlation coefficients (Supplementary Fig. S1). Across all the seasons, PLHT was found positively correlated with LLN, LWD, PWD, PBR, and YPP ($r=0.13$ to 0.27), however, it was negatively correlated with SLN and SLNG ($r=-0.12$ to -0.15). LLN was positively correlated with pod-attributing traits, namely, PLN, PWD, and SLN ($r=0.12$ to 0.33). Among yield-contributing traits, PPP showed strong positive correlations with PBR, YPP, and YKGH ($r=0.37$ to 0.78) in Patancheru location and low correlations ($r=0.17$ to 0.22) in Dharwad location. However, PPP was negatively correlated with pod-attributing traits, PLN, PWD, and SLN ($r=-0.18$ to -0.24). Among kernel traits, SLNG was positively correlated with HSW, YPP, and YKGH ($r=0.13$ to 0.36), while it was negatively correlated with many agronomic traits, for instance, DF, PLHT ($r=-0.12$ to -0.15), SLN, PWD, PLN ($r=-0.14$ to -0.29), and LLN ($r=-0.14$ to -0.30). HSW showed low to strong positive correlations with SLN, PWD, and PLN ($r=0.26$ to 0.61).

Main effect QTLs (M-QTLs) for agro-morphological traits

The QTL analysis revealed a total of 62 M-QTLs at 39 genomic regions across ten traits, scattered on 15 different LGs. (Table 1; Supplementary Table S2; Fig. 2). M-QTLs could not be detected for LWD, PLN, PWD, and DM. A maximum of 20 M-QTLs were detected for PLHT followed by nine for YPP, eight for HSW, and seven each for PPP and SLNG. DF and YKGH had three M-QTLs each, while LLN and SLN had two M-QTLs each. PBR was detected with a solitary M-QTL. Five M-QTLs controlled two traits each. For instance, a M-QTL mapped at 65.21 cM at GM1955-GM1007 on AhV governed PPP and HSW. M-QTL at 0.01 cM in PM377-TC1A01 region on AhVI governed YPP and YKGH. Similarly, a M-QTL at 7.61 cM within TC1A01-S108 contributed for SLN and HSW. PPP and YPP were governed by a single M-QTL (22.11 cM) in the region GM1097-TC7C06 on AhVI.

PPP and YKGH were governed by an M-QTL at 12.51 cM within GM1536-GM2301 on AhXV.

Contribution of M-QTLs in terms of PVE ranged from 3.84–15.06%. Six M-QTLs were considered to be major since the PVE was more than 10% (Supplementary Fig. S2). The highest PVE of 15.06% was recorded by a M-QTL at 14.51 cM within IPAHM103-GM1954 on AhXV for YPP. Another M-QTL at 20.51 cM with the same marker interval had 13.40% PVE towards SLNG, which was also governed by single M-QTL at 10.01 cM with GM2009-GM1536 on AhXV with 14.04% PVE. YPP was governed by a major M-QTL at 15.11 cM within GM1386-GM1162 on AhXIV with PVE of 13.56%. One M-QTL at 61.71 cM within TC3A12-GM1955 on AhV recorded a PVE of 11.57% for PLHT. Another M-QTL at 60.71 cM within GM1076-Seq5D05 on AhXVIII governed PPP with a PVE of 10.48%. These M-QTLs were detected with high LOD (more than 5.0). Interestingly, the favorable alleles at all the major M-QTLs were contributed by GPBD 4.

The M-QTL at 71.51–73.51 cM within GM633-PM179 on AhV with a maximum PVE and LOD of 9.67 and 7.09%, respectively towards PLHT was detected in five different environments. PLHT was also governed by a stable M-QTL at 0.01–1.91 cM within Seq7H06-IPAHM176 on AhXII with a maximum PVE and LOD of 7.12 and 5.03%, respectively which appeared across four environments. One M-QTL at 63.21–65.21 cM within GM1955-GM1007 on AhV also controlling PLHT appeared in three environments with a maximum PVE and LOD of 6.18 and 3.49%, respectively. The favorable allele at this stable M-QTL was contributed by GPBD 4. Stable M-QTLs appearing in two environments were detected for LLN, (7.61–12.71 cM, TC1A01-S108, AhVI), YPP (56.11–60.11 cM, GM1062-GM2638, AhVII), YKGH (0.01 cM, PM377-TC1A01, AhVI), and HSW (70.51–71.51 cM, GM633-PM179, AhV). GPBD 4 contributed for the favorable allele at the stable M-QTL for LLN, YPP, and YKGH, while TAG 24 contributed the favorable allele at the stable M-QTL for HSW. However, the major M-QTLs appeared in any one environment.

Co-localization of M-QTLs

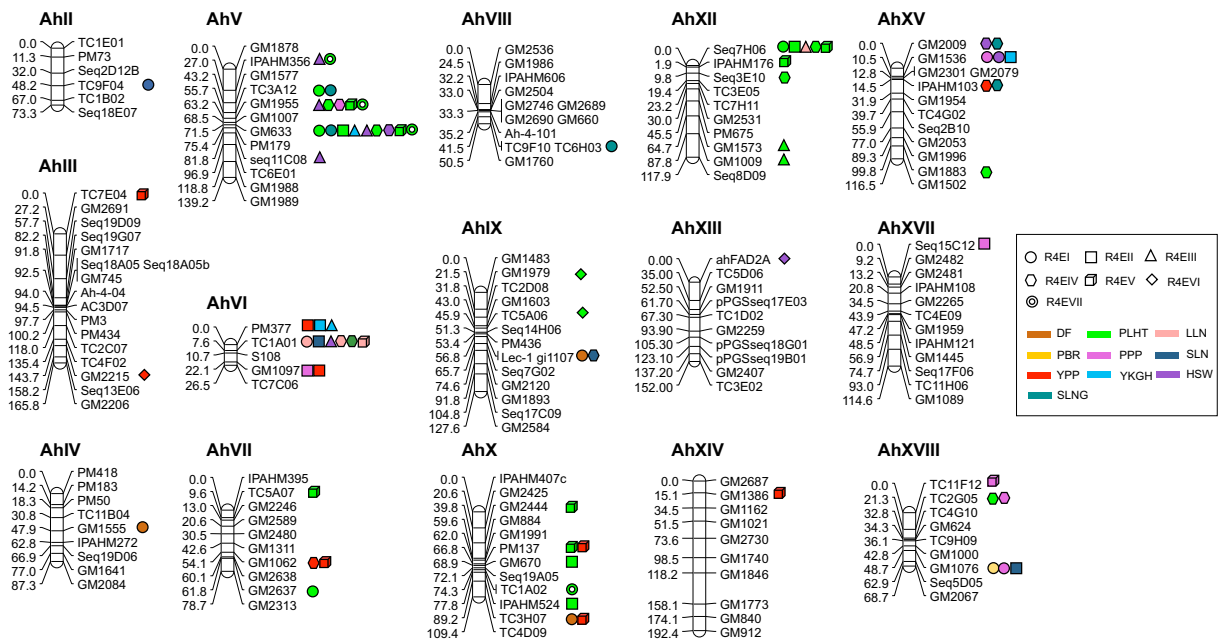
It was interesting to notice that 16 genomic regions consisted of co-localized M-QTLs for various traits (Supplementary Fig. S3). Four genomic regions/

Table 1 Main effect QTLs (M-QTLs) detected in composite interval mapping (CIM) analysis for 14 agro-morphological traits in RIL-4 population

Trait	Number of M-QTLs	PVE %	Environment	Source of favorable allele
Days to flowering (DF) (days)	3	4.92–6.82	R4EI	TAG 24, GPBD 4
Plant height (PLHT) (cm)	20	3.84–11.57	R4EI–R4EVII	GPBD 4, TAG 24,
Leaf length (LLN) (mm)	2	4.14–8.65	R4EIII, R4EIV, R4EV	GPBD 4
Leaf width (LWD) (mm)	–	–	–	–
Primary branching (PBR)	1	5.94	R4EI	GPBD 4
Pods per plant (PPP) (mm)	7	3.91–10.48	R4EI, R4EII, R4EIV, R4EV	GPBD 4, TAG 24
Pod length (PLN) (mm)	–	–	–	–
Pod width (PWD) (mm)	–	–	–	–
Seed length (SLN) (mm)	2	4.30–5.34	R4EII	GPBD 4, TAG 24
Yield per plant (YPP) (gm)	9	5.06–15.06	R4EII, R4EIV, R4EV, R4EVII	GPBD 4, TAG 24
Yield Kg per hectare (YKGH) (kg)	3	4.17–8.20	R4EII, R4EIII	GPBD 4
Hundred seed weight (HSW) (gm)	8	5.68–9.14	R4EI, R4EIII, R4EIV, R4EV, R4EVII	TAG 24, GPBD 4
Shelling percentage (SLNG) (%)	7	5.03–14.04	R4EI, R4EIV	GPBD 4, TAG 24
Days to maturity (DM) (days)	–	–	–	–

clusters were identified where QTLs were detected for more than three traits in the linkage groups AhV, AhVI, AhXV, and AhXVIII. For instance, a 4.9-cM region (GM633-PM179) on AhV mapped at position 70.41–70.51 cM carried four co-localized M-QTLs contributing for PLHT, YKGH, HSW, and SLNG, and was detected in six environments (R4EI, R4EII, R4EIII,

R4EIV, R4EV, and R4EVI) explaining 5.97–9.67% PVE. Similarly, the M-QTL with flanking markers TC1A01-S108 in the LG AhVI (7.61–12.71 cM) was found to be clustered for traits LLN, SLN, and HSW in R4EII, R4EIII, R4EIV, and R4EV and it explained 5.34–8.65% PVE. Another co-mapped M-QTL (GM1536-GM2301) for PPP, YKGH, and HSW detected on

**Fig. 2** Genetic linkage map showing the location of main effect QTLs identified using composite interval mapping (CIM) for agro-morphological traits among the RILs of peanut

AhXV at 10.51–12.51 cM appeared in two (R4EI and R4EII) of the seven environments, which explained 4.17–8.82% PVE. One of the major M-QTL (IPAHM103-Seq19D6/IPAHM103-GM1954) was mapped on AhXV (14.51–20.51 cM), responsible for rust resistance exhibited pleiotropic effects for two yield-contributing traits such as YPP and SLNG explaining 15.06 and 13.40% PVE, respectively. A M-QTL on AhXVIII flanked by markers GM1076-Seq5D05 mapped at position 58.71–62.91 cM was co-localized for the traits like PLHT (R4EIII), PBR (R4EI), PPP (R4EI), and SLN (R4EII) with 4.30–10.48% PVE (Supplementary Table S2). Apart from these clusters, M-QTLs for PLHT, PPP, and HSW were tagged with markers GM1955-GM1007 on LG AhV (63.21–65.21 cM) and were detected in four (R4EIII, R4EIV, R4EV, and R4EVI) of the seven environments that accounted for 5.11–8.06% PVE. A co-localized M-QTL for PLHT and SLNG in the marker interval of TC3A12-GM1955 on LG AhV was detected only in R4EI and accounted for 5.52–11.57% PVE. This M-QTL was detected at high LOD (6.37) and had a major effect on PLHT (11.57% PVE). The M-QTL flanked by GM2009-GM1536 mapped on AhXV (2.01–10.01 cM) co-segregated with HSW and SLNG accounted for 6.17 and 14.04% PVE, respectively.

Of the 16 co-mapped M-QTLs, eight minor effect M-QTLs co-segregated with a combination of two traits with PVE ranged from 3.84–9.73%. These QTL regions included PM137-GM670 (PLHT and YPP), IPAHM356-GM1577 (PLHT and HSW), PM377-TC1A01 (YPP and YKGH), Seq7H06-IPAHM176 (PLHT and LLN), TC2G05-TC4G10 (PLHT and PPP), TC3H07-TC4D09 (DF and YPP), GM1097-TC7C06 (PPP and YPP), and gi1107-Seq7G02 (DF and SLN).

Epistatic QTLs

An effort was made to check the M-QTLs that interacted with each other and to measure the nature and the extent of interaction. In total, 24 E-QTLs were mapped for all traits except LWD, PBR, and PPP (Table 2; Supplementary Table S3; Fig. 3). Of these, 16 E-QTLs showed significant additive \times additive epistasis \times environment (AAE) interactions (0.01–2.81% PVE) for ten traits whereas other eight E-QTLs showed only additive \times additive (AA) effects (0.03–4.49% PVE). Among 24 E-QTLs, one E-QTL pair was mapped for DM and

PWD, two each for DF, PLN, SLN, YPP, YKGH, and SLNG, three for LLN and PLHT, and four for HSW were identified. In general, PVE by E-QTLs was very low due to epistatic interactions. The PVE for E-QTL was observed for SLN (4.28–4.49% PVE) followed by PLN (1.21–3.34% PVE), SLNG (1.75–3.03% PVE), PLHT (1.71–2.91% PVE), YKGH (1.55–2.41% PVE), and HSW (0.36–2.41% PVE).

Of the 16 E-QTLs, only five of them involved the M-QTLs. For instance, an M-QTL at TC3A12-GM1955 on AhV showed an epistatic (additive \times additive) interaction with the M-QTL at GM670-Seq19A05 on AhX governing PLHT with a PVE of 1.21%. Such epistatic interactions were also observed for TC5A06-Seq14H06 (AhIX) \times GM1536-GM2301 (AhXV) influencing PLHT \times PPP, PLHT \times YKGH, and PLHT \times HSW, and TC9F04-TC1B02 (AhII) \times GM1555-IPAHM272 (AhIV) influencing SLNG \times DF. TC3A12-GM1955 \times GM670-Seq19A05 and TC9F04-TC1B02 \times GM1555-IPAHM272 also showed QE (additive \times additive \times environment) interactions. It was interesting to note that the M-QTL at TC3A12-GM1955 which showed epistatic interaction was a major M-QTL.

Discussion

Yield and yield-contributing traits are quantitative in nature and show complex inheritance because their phenotypic expression is dependent on a combination of minor genes, modifiers, and environments (Kover et al. 2009). Genetic dissection of potential genomic regions harboring QTLs associated with these traits is expected to reveal the genetic control of the trait. Identification of QTLs with main effects and epistatic effects is essential for the efficient marker-assisted selection (MAS) aimed at improving breeding efficiency (Bocianowski 2013). An epistatic QTL implies that the effects of single-locus QTLs are mostly dependent on the genotypes of other loci. Thus, breeding programs have to take into account the epistatic effects while employing QTLs.

The classic example for identifying the main-effect QTLs in peanut are for resistance to rust and late leaf spot and high oleic acid which were successfully deployed in developing molecular breeding products in peanut (see Vishwakarma et al. 2017). There have been some reports of using epistatic QTLs in plant breeding to confirm the universality and the importance of epistasis between QTLs and provide useful information for

improving plant height via heterosis and QTL pyramiding in rice (Zhu et al. 2015).

Most of the QTL analysis conducted for agronomic traits in peanut till date reported just M-QTLs and therefore provided no idea on environment interactions (Kolekar et al. 2016; Hake et al. 2017; Luo et al. 2017a; Luo et al. 2017b). However, few studies provided information on both type of QTLs (M-QTLs and E-QTLs) for traits such as drought-related traits (Ravi et al. 2011; Gautami et al. 2012), fatty acids (Pandey et al. 2014b; Wang et al. 2015), and pod- and kernel-related traits (Chen et al. 2016a). The detailed information on main effect and environment interactions for the loci associated with traits provide better understanding on their contribution towards trait development, which helps in devising strategy for improving the studied trait through different available GAB approaches.

In this study, M-QTLs and E-QTLs were identified for the important agro-morphological and productivity traits from a RIL mapping population which was phenotyped for multi-seasons at multiple environments. In general, phenotypic and genotypic variability and heritability estimates were high for majority of the traits indicating to low $G \times E$ interactions. Nevertheless, low heritability (for YPP in R4EIV and HSW in R4EII) indicated low genetic variability due to the influence of environment on these traits. Strong positive correlations between most of the yield-related traits indicated the possibility of common genes/pathways operating at molecular level, thus sharing common genomic regions whereas negative correlation among PLHT with PLN, SLN, and SLNG; PPP with PLN, PWD, and SLN; SLNG with DF, PLHT, SLN, PWD, PLN, and PBR indicates the possibility of independent inheritance of these traits. In addition, the strong positive correlations among traits can help in simultaneous improvement of traits through modern breeding approaches.

The RIL population with high variability and normal distribution of means was considered for QTL analysis using a genetic map reported by our group (Sujay et al. 2012). One or more M-QTLs were detected for ten traits, while no M-QTLs could be detected for four traits (LWD, PLN, PWD, and DM) probably due to non-significant differences among the RILs and skewed distribution of means. Five M-QTLs governed two traits each, and these traits (PPP, SLN, YPP, YKGH, and HSW) were significantly and positively correlated in many environments. Of these 62 M-QTLs, only six were major (PVE > 10%) and they governed YPP

(13.56 and 15.06%), SLNG (13.40 and 14.04%), PLHT (11.57%), and PPP (10.48%). These M-QTLs were detected with at least 5.0 LOD. GPBD 4 contributed the favorable alleles at all the major M-QTLs, indicating a possibility of using this parent as donor in the breeding programs.

Stability of the M-QTL is an important parameter deciding the utility of QTLs. Though none of the major M-QTLs were stable across the environments, a few with relatively high PVE were stable. For PLHT, a M-QTL (71.51–73.51 cM) on AhV with a maximum PVE and LOD of 9.67 and 7.09%, respectively was found stable across five environments. Another M-QTL on AhXII for PLHT with a maximum PVE and LOD of 7.12 and 5.03%, respectively appeared across four environments. Likewise, stable M-QTLs were observed for LLN (7.61–12.71 cM, TC1A01-S108, AhVI), YPP (56.11–60.11 cM, GM1062-GM2638, AhVII), YKGH (0.01 cM, PM377-TC1A01, AhVI), and HSW (70.51–71.51 cM, GM633-PM179, AhV).

Pleiotropic M-QTLs (QTLs governing multiple traits) and co-localized M-QTLs (QTLs located in adjacent genomic regions) are important for the simultaneous improvement of multiple traits. In this study, five pleiotropic M-QTLs governing PPP + HSW, YPP + YKGH, SLN + HSW, PPP + YPP, and PPP + YKGH were identified. At sixteen genomic regions, the M-QTLs governing two to four productivity traits were co-localized. Marker-assisted breeding for these traits could be favored due to pleiotropy and co-localization of the M-QTLs. Such co-localized QTLs were also observed in the previous studies (Ravi et al. 2011; Gautami et al. 2012; Sujay et al. 2012; Pandey et al. 2014b; Wang et al. 2015) for traits like drought, foliar diseases, and fatty acid contents.

A previously identified marker linked to rust resistance (Khedikar et al. 2010; Sujay et al. 2012) was found to be associated with YPP (15.60% PVE) and SLNG (13.40% PVE) in R4EIV in this study. The introgression of this genomic region in the elite peanut varieties has not only enhanced rust resistance but increased the yield by 30–70% (Varshney et al. 2014; Yeri and Bhat 2016; Kolekar et al. 2017). This genomic region was dissected to identify genes (Pandey et al. 2017b), and the transcriptomic analysis showed differential expression of these genes among the genotypes differing for LLS resistance (Han et al. 2017).

Inter-allelic interactions among two QTLs/genes lead to E-QTLs, which play a key role in controlling trait

expression, and are considered major components controlling qualitative and quantitative traits (Yu et al. 1997). In the present study, 24 E-QTLs were identified by two-locus analysis accounting for small effects (< 5%) compared to single locus main effect QTLs. This showed that E-QTLs associated with agro-morphological traits encompass QTL interactions QTL \times environment (QE) interactions. QTL with more than one digenic interactions was observed in the present study; of which, 16 also showed QE interactions. Five epistatic QTLs involved the M-QTLs identified in this study, and they influenced PLHT \times PLHT, PLHT \times PPP, PLHT \times YKGH, PLHT \times HSW, and SLNG \times DF. However, in all these cases, the PVE was relatively low (0.64–15.5% PVE), and the major M-QTLs were not involved in any epistatic interactions, indicating that they can be considered for marker-assisted breeding without considering the need for transferring their interacting QTLs, which could simplify the breeding program.

Conclusions

The present study identified several major and minor effect M-QTLs for various economically important agro-morphological and productivity traits. Six major QTLs were detected for yield-attributing traits that can be validated using either mapping populations with different genetic background or germplasm lines and later can be utilized in peanut improvement programs using marker-assisted selection (MAS). In the present study, the favorable alleles at most of the M-QTLs and E-QTLs were contributed by GPBD 4, thereby easing the process of their transfer to the elite genotypes. The epistatic effects were not very high for the small effect M-QTLs and major effect M-QTLs, which again would suggest a relatively simple backcross breeding scheme. With the current next generation sequencing (NGS) and next generation mapping (NGM), the genomic regions contributing significantly for the productivity traits can be dissected for the candidate gene(s).

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Compliance with ethical standards

Competing interests The authors declare that they have no competing interests.

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