Discovery of genomic regions and candidate genes controlling shelling percentage using QTL-seq approach in cultivated peanut (*Arachis hypogaea* L.)

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Summary

Cultivated peanut (*Arachis hypogaea* L.) is an important grain legume providing high-quality cooking oil, rich proteins and other nutrients. Shelling percentage (SP) is the 2nd most important agronomic trait after pod yield and this trait significantly affects the economic value of peanut in the market. Deployment of diagnostic markers through genomics-assisted breeding (GAB) can accelerate the process of developing improved varieties with enhanced SP. In this context, we deployed the QTL-seq approach to identify genomic regions and candidate genes controlling SP in a recombinant inbred line population (Yuanza 9102 × Xuzhou 68-4). Four libraries (two parents and two extreme bulks) were constructed and sequenced, generating 456.89–790.32 million reads and achieving 91.85%–93.18% genome coverage and 14.04–21.37 mean read depth. Comprehensive analysis of two sets of data (Yuanza 9102/two bulks and Xuzhou 68-4/two bulks) using the QTL-seq pipeline resulted in discovery of two overlapped genomic regions (2.75 Mb on A09 and 1.1 Mb on B02). Nine candidate genes affected by 10 SNPs with non-synonymous effects or in UTRs were identified in these regions for SP. Cost-effective KASP (Kompetitive Allele-Specific PCR) markers were developed for one SNP from A09 and three SNPs from B02 chromosome. Genotyping of the mapping population with these newly developed KASP markers confirmed the major control and stable expressions of these genomic regions across five environments. The identified candidate genomic regions and genes for SP further provide opportunity for gene cloning and deployment of diagnostic markers in molecular breeding for achieving high SP in improved varieties.

Keywords: peanut, shelling percentage, QTL-seq, genomic regions, candidate genes.

Introduction

The cultivated peanut, *Arachis hypogaea* L., is an important oilseeds crop grown in >100 countries and is consumed worldwide in form of high-quality cooking oil, edible nut, peanut butter and candy. During 2016, the annual production of peanut pods (with shell) was more than 43.98 million tonnes throughout the world (FAOSTAT). Peanut is an allotetraploid grain legume (AABB, 2n = 4x = 40) with genome size of ~2.7 Gb (Kochert et al., 1996). It is also known as the groundnut due to underground pod development. Peanut pod has two parts: kernel and shell (Figure 1). During post-harvested processing, pods are passed through peanut shelling machines to collect kernels (seeds) which provide rich nutrients such as high-quality oil, proteins and a variety of healthy bioactive compounds for human (Davis and Dean, 2016). As by-product, shells cannot be effectively and economically utilized and may create waste disposal problem around growing or processing areas (Zhao et al., 2012). Therefore, shelling percentage (SP), a ratio in percentage of the weight of kernels to the weight of pods, is an important economic trait in peanut production. High SP varieties provide more kernels for human nutrients and produce fewer by-products than low SP varieties. In recent studies, significant variations for SP were found among peanut varieties, for example ranging from 59.9% to 81.0% in the Chinese core collection (Jiang et al., 2013) and from 45.3% to 73.1% in the 256 accessions evaluated in Mediterranean Basin (Yol et al., 2018), suggesting that there is a great potential in the enhancement of SP through breeding.

Genomics-assisted breeding (GAB) can accelerate the process of developing new varieties with improved traits of interest through precise selection of favourable alleles across generations (Varshney et al., 2005). However, diagnostic markers of the targeted traits are the prerequisite for deploying GAB and currently no such diagnostic marker is available for SP. The identification of linked markers to major and stable quantitative trait loci (QTLs) for SP is required to develop improved peanut varieties. Limited efforts were made to identify QTLs associated with SP in cultivated peanut, however, none of them could develop markers for use in breeding (Faye et al., 2015; Huang et al., 2015). Also these QTLs were detected in single environment, hence, do not provide information on their stability and consistency. Association analysis identified one SSR allele linked with SP in three field trials but had only 1.49%–2.98% phenotypic variation explained (PVE) (Jiang et al., 2014) which...
cannot be exploited in GAB. Recent study from our research group identified two major and consistent QTLs for SP, that is cqSPA09 (10.47%–17.01% PVE) and cqSPB02 (8.01%–11.20% PVE), in a recombinant inbred line (RIL) population (Yuanza 9102 × Xuzhou 68-4) across four consecutive years in Wuhan, China (Luo et al., 2017). Despite providing idea on genomic regions, none of these studies provided precise localization in the peanut genome, which hampered further candidate gene discovery and diagnostic marker development for SP.

Sequencing-based trait mapping approaches facilitate faster discovery of genomic regions and candidate genes for target traits (Pandey et al., 2016). The QTL-seq approach (Takagi et al., 2013) is one of the successful sequencing-based mapping approaches which deals with sequencing of pooled samples for RILs with extreme phenotypes and parental genotypes. As a result, the approach rapidly locates the candidate genomic regions and underlying genes for QTLs in plants with reference genomes. This approach is cost-effective as sample number is less for sequencing and facilitates development of cost-effective markers in less time for deployment in GAB. This approach has been deployed in several crops including rice (Wambumgu et al., 2018), cucumber (Lu et al., 2014), tomato (Illya-Berenguer et al., 2015), rapeseed (Hua et al., 2016), chickpea (Singh et al., 2016a), pigeonpea (Singh et al., 2016b), peanut (Pandey et al., 2017) and soybean (Zhong et al., 2018), and proved to be very successful in RIL populations with phenotyping data generated in multiple environments. Since the reference genome sequence of cultivated peanut is still unpublished, the available reference genome sequences of its two diploid ancestors, A. duranensis (AA, 2n = 2x = 20) and A. ipaensis (BB, 2n = 2x = 20) (Bertioli et al., 2016) provided a foundation to find candidate genes and SNPs present in QTL regions. In summary, this study reports the deployment of QTL-seq approach that successfully located the genomic regions on the peanut genome and facilitated discovery of candidate genes and marker development for SP to use in GAB for faster development of new varieties.

**Results**

**Construction of extreme bulks for shelling percentage**

The RIL population (Yuanza 9102 × Xuzhou 68-4) had high phenotypic variability for SP (Figures 1 and S1). Based on the
phenotyping data generated in Wuhan in four consecutive years (Luo et al., 2017), the SP of Yuzanza 9102 was 81.65 ± 0.44%, whereas that of Xuzhou 68-4 was 75.81 ± 0.29% (Table S1). The SP of the RIL population showed continuous distributions skewed towards higher values in all of the four environments (Figure S2), ranging from 67.83% to 93.36% on average. Therefore, 15 extreme RILs with high SP (81.71%–93.36%) and 15 extreme RILs with low SP (67.83%–72.87%) were selected to construct the two bulks (Figure 1, Table S1), that is high SP bulk (HB) and low SP bulk (LB). The whole-genome resequencing (WGRS) data were generated for these extreme bulks in addition to parental genotypes followed by their analysis using the QTL-seq pipeline as shown in Figure 2.

Whole-genome resequencing, mapping of reads and identification of SNPs

WGRS data were generated for Xuzhou 68-4, Yuzanza 9102, HB and LB samples using the Illumina NovaSeq. A total of 544.70 million reads for Xuzhou 68-4, 456.89 million reads for Yuzanza 9102, 790.32 million reads for HB and 717.72 million reads for LB were generated (Tables 1 and S2). The maximum sequencing data were obtained for HB (119.34 GB) followed by LB (108.38 GB), Xuzhou 68-4 (82.25 GB) and Yuzanza 9102 (68.99 GB).

The reads of Xuzhou 68-4 were aligned to the genome sequences of A. duranensis and A. ipaensis, and resulted in 92.61% genome coverage and 16.18 mean read depth. A reference-guided assembly was then generated for the male parent Xuzhou 68-4 and referred as Xuzhou assembly hereafter (Figure 2). The reads of the two extreme bulks were mapped to the Xuzhou assembly, and achieved 93.18% mapping coverage and 21.37 mean read depth for HB, whereas 93.12% coverage and 20.96 mean read depth for LB (Tables 1 and S2). The comprehensive sequence analysis between the high and low bulks identified 172 715 genome-wide SNPs (Table S3).

Similarly, the reads of Yuzanza 9102 were aligned to the genome sequences of A. duranensis and A. ipaensis, and resulted in 91.85% genome coverage and 14.04 mean average read depth. A reference-guided assembly was then generated for the female parent Yuzanza 9102 and referred as Yuzanza assembly hereafter (Figure 2). The reads of the two extreme bulks were mapped to the Yuzanza assembly and achieved 93.12% coverage and 21.37 mean read depth for HB, whereas 93.06% coverage and 20.96 mean read depth for LB (Tables 1 and S2). The comprehensive sequence analysis between the high and low bulks identified 241 078 genome-wide SNPs (Table S3). This resulted in discovery of a common set of 75 087 SNPs when either of the parents’ assembly was used as reference (Figure 3).

Candidate genomic regions for shelling percentage

With the Xuzhou assembly as reference, the SNP-index of the genome-wide SNPs was calculated for each bulk (Figure S3). The ΔSNP-index was then calculated by subtracting SNP-index of HB from SNP-index of LB. If more alleles of one parent than the other were presented in the two extreme bulks, the SNP-index will significantly deviate from 0.5 and the ΔSNP-index will significantly deviate from 0 (zero). Based on the sliding window analysis for SNP-index and ΔSNP-index plots, a 3.20 Mb (66.75–69.90 Mb) interval on chromosome A09 and a 1.30 Mb (5.65–6.95 Mb) interval on chromosome B02 were identified for SP at a statistical confidence of P < 0.01 (Figures 4 and S4–S6). The ΔSNP-index of the two genomic regions were negative, indicating that more alleles were from the reference parent Xuzhou 68-4 in the low SP bulk but from non-reference parent Yuzanza 9102 in the high SP bulk.

Similarly, a 2.75 Mb (66.75–69.50 Mb) interval on A09 and a 1.50 Mb (5.85–7.35 Mb) interval on B02 were identified at P < 0.01 level, using the Yuzanza assembly as reference (Figures S7–S9). The ΔSNP-index of the two genomic regions were positive, indicating that more alleles were from the non-reference parent Xuzhou 68-4 in the low SP bulk but from reference parent Yuzanza 9102 in the high SP bulk. These intervals were significantly overlapped with the genomic regions identified with the Xuzhou assembly (Table 2). Therefore, the overlapped regions were selected as the candidate regions for SP, that is, 66.75–69.50 Mb on A09 and 5.65–6.95 Mb on B02 (Figure 5, Table 2).

Candidate SNPs and putative candidate genes in the genomic regions for shelling percentage

The genomic region spanning 2.75 Mb on chromosome A09 had 455 effective SNPs with read depth of ≥10, SNP-index significantly deviated from 0.5 and ΔSNP-index higher than the statistical confidence at P < 0.01 (Figure 3, Tables S4 and S5). Of the 455 SNPs, 454 SNPs were intergenic and one was non-synonymous. The non-synonymous SNP could be identified using both parents as reference and affected the candidate gene encoding histone-lysine N-methyltransferase SUVR2-like isoform X2 (Aradu.A5UR9) (Table 3).

The genomic region spanning 1.1 Mb on chromosome B02 had 148 effective SNPs with read depth of ≥10, SNP-index significantly deviated from 0.5 in addition to ΔSNP-index higher than the statistical confidence at P < 0.01 (Figure 3, Tables S4 and S5). Of the 148 SNPs, 134 SNPs were intergenic and 14 SNPs were genic, including four intronic, seven non-synonymous (one resulted in stop gained), one synonymous and two in UTRs. The seven non-synonymous SNPs and the two SNPs in UTRs affected eight candidate genes encoding four unknown proteins (Arap.G7S23, Arapi.HEW90, Arapi.Y03J3 and Arapi.X1ALQ), two serine/threonine-protein phosphatase homologs (Arapi.R3ORS and Arapi.N4BSE), one vacuolar protein sorting-associated protein (Arapi.J157B) and one Ulp1 protease family carboxy-terminal domain protein (Arapi.V4DX) (Table 3).

Validation of identified genomic regions

To validate the genomic regions identified for SP, 10 SNPs (One SNP from A09 and nine SNPs from B02) were targeted for development of KASP markers. Primers were successfully developed for the SNP at position 66949737 on chromosome A09. Of the nine SNPs on chromosome B02, primers were developed for three SNPs, whereas no primers could be designed for remaining six SNPs (Table 4, Figure S10). The RIL population (Yuzanza 9102 and Xuzhou 68-4) in F9 generation were genotyped with the four KASP markers (Table S6).

To test the stability of the identified QTLs, the phenotyping data of the RIL population in F9 generation was generated in another geographical location in 2017 (Yangluo2017) (Table S7), in addition to the phenotyping data used to construct extreme bulks. The phenotypic distribution in the Yangluo2017 environment was continuous and had two peaks (Figure S2). Single-marker QTL analysis with the phenotyping data in the five
Mapping of shelling percentage in peanut by QTL-seq

Figure 2  QTL-seq approach used for trait mapping in peanut for shelling percentage.

- Development of mapping population
- Phenotyping of mapping population
- Selection of RILs with extreme phenotype
- Construction of high and low pools
- Sequencing of pools and parents
- Analyzing sequence data with QTL-seq pipeline
- Identification of genomic regions, SNPs and candidate genes
- Validation of newly developed markers

**A. duranensis (A09):**
- 66.75-69.50 Mb (2.75 Mb)
- Confidence: ~99%
- 455 effective SNPs with the read depth of 10
- 1 nonsynonymous SNPs affecting 1 candidate genes

**A. ipaensis (B02):**
- 5.85-6.95 Mb (1.1 Mb)
- Confidence: ~99%
- 148 effective SNPs with the read depth of 10
- 7 nonsynonymous SNPs and 2 in UTR affecting 8 candidate genes

Validation using KASP markers:
- Two parents, two bulks and the RIL population
- The SNP explained 11.44%-15.48% PVE with LOD score of 5.63%-7.19%

- Two parents, two bulks and the RIL population
- Out of 9 SNPs, 3 validated for 3 genes
- Up to 10.91%-20.26% PVE with LOD scores of 3.73%-8.52%
environments showed high significances ($P < 0.001$) for all of the four KASP markers (Table 5). The marker, Aradu_A09_6694 9737, on chromosome A09 explained high phenotypic variation of 11.44%–15.48% with LOD (Likelihood of odds) scores of 5.63–7.19 across five environments. The genotyping data of the three KASP markers (Araip_B02_6155951, Araip_B02_6770282 and Araip_B02_6776001) on chromosome B02 were used to estimate genetic distances, and a map length of 2.76 cM was obtained. Among the three candidate markers on chromosome B02, Araip_B02_6776001 explained highest phenotypic variation 10.91%–20.26% with LOD scores of 3.73–8.52, whereas Araip_B02_6155951 explained 8.18%–19.20% with LOD scores of 3.54–8.42. These marker-trait associations confirmed the identified genomic regions harbouring candidate genes for two major and stable QTLs for SP.

To evaluate the combined effect of the identified QTLs, RILs were classified into four groups using the four KASP markers as diagnostic markers. The alleles of the identified QTLs on A09 and B02 from Yuanza 9102 were designated as ‘AA’ and ‘BB’, respectively, whereas those from Xuzhou 68-4 were designated.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean shelling percentage</th>
<th>Number of reads generated</th>
<th>Total bases</th>
<th>Genome coverage (%)</th>
<th>Mean depth (X)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xuzhou 68-4*</td>
<td>75.81</td>
<td>544 700 200</td>
<td>82 249 730 200</td>
<td>92.61</td>
<td>16.18</td>
</tr>
<tr>
<td>LB†</td>
<td>70.96</td>
<td>717 723 902</td>
<td>108 376 309 202</td>
<td>93.12</td>
<td>20.96</td>
</tr>
<tr>
<td>HB‡</td>
<td>82.42</td>
<td>790 320 814</td>
<td>119 338 442 914</td>
<td>93.18</td>
<td>21.37</td>
</tr>
<tr>
<td>Yuanza 9102*</td>
<td>81.65</td>
<td>456 893 014</td>
<td>68 990 845 114</td>
<td>91.85</td>
<td>14.04</td>
</tr>
</tbody>
</table>

*The short reads of parental lines were aligned to the publicly available genome of diploid progenitors A. duranensis and A. ipaensis (PeanutBase: http://peanutbase.org).
†The short reads of the extreme bulks were aligned to the Xuzhou ‘reference sequence’ developed by replacement of SNPs between Xuzhou 68-4 and diploid progenitors.
‡The short reads of bulks were aligned to the Yuanza ‘reference sequence’ developed by replacement of SNPs between Yuanza 9102 and diploid progenitors.
as ‘aa’ and ‘bb’. RILs with the AABB genotype showed significant higher SP than the other three genotypes (aaBB, AAbb or aabb) in all five environments (Figure S11, Table S8). The average SP of RILs with the aaBB genotype was slightly higher than that with the AAbb genotype but not significant. Furthermore, RILs with the aaBB or AAbb had significant higher shelling percentages than those with the aabb genotype.

In addition, the four KASP markers were also genotyped in diverse panel consisting of 337 cultivars whose shelling percentages successfully evaluated in Wuhan in 2016 and Nanchong in 2017. Eight cultivars showed the AA genotype and their shelling percentages were significantly higher than the others’ (Figure S12a). A total of 61 cultivars showed the BB genotype, however, their shelling percentages were significantly lower than the others’ on average (Figure S12b). Six cultivars showed the AABB genotype, and their shelling percentages were significantly higher than the 274 cultivars with aabb genotype (Figure S12c).

### Discussion

The genetic mapping in cultivated peanut started in 2009 with only 135 SSR loci (Varshney et al., 2009a). The last decade witnessed development and availability of genomic and genetic resources including reference genomes of diploid progenitors and several thousands of genetic markers in peanut (Bertioli et al., 2016; Chen et al., 2016; Luo et al., 2017; Shirasawa et al., 2012). The genetic markers were widely used in the QTL mapping of important traits such as yield, kernel, oil quality, disease resistance and drought tolerance (Vishwakarma et al., 2017).

However, genotyping of these markers in mapping population is laborious and time-consuming, and also do not lead to discovery of candidate genes and tightly linked markers. Additionally, low level of polymorphism poses a great challenge in developing high-density genetic maps and conducting high-resolution genetic mapping (Luo et al., 2017). In the recent past, NGS-based technologies were deployed to overcome these issues because of

### Table 2 Genomic regions identified for shelling percentage

<table>
<thead>
<tr>
<th>Reference assembly</th>
<th>Chr</th>
<th>Genomic region (Mb)</th>
<th>Length (Mb)</th>
<th>ΔSNP-index</th>
<th>U99</th>
<th>L99</th>
<th>Allele source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xuzhou</td>
<td>A09</td>
<td>66.70–69.90</td>
<td>3.20</td>
<td>−0.71</td>
<td>0.62</td>
<td>−0.63</td>
<td>Yuanza 9102</td>
</tr>
<tr>
<td></td>
<td>B02</td>
<td>5.65–6.95</td>
<td>1.30</td>
<td>−0.66</td>
<td>0.62</td>
<td>−0.63</td>
<td>Yuanza 9102</td>
</tr>
<tr>
<td>Yuanza</td>
<td>A09</td>
<td>66.75–69.50</td>
<td>2.75</td>
<td>0.69</td>
<td>0.63</td>
<td>−0.61</td>
<td>Yuanza 9102</td>
</tr>
<tr>
<td></td>
<td>B02</td>
<td>5.85–7.35</td>
<td>1.50</td>
<td>0.69</td>
<td>0.63</td>
<td>−0.61</td>
<td>Yuanza 9102</td>
</tr>
<tr>
<td>Overlapped</td>
<td>A09</td>
<td>66.75–69.50</td>
<td>2.75</td>
<td>0.69</td>
<td>0.63</td>
<td>−0.61</td>
<td>Yuanza 9102</td>
</tr>
<tr>
<td></td>
<td>B02</td>
<td>5.85–6.95</td>
<td>1.10</td>
<td>0.69</td>
<td>0.63</td>
<td>−0.61</td>
<td>Yuanza 9102</td>
</tr>
</tbody>
</table>
the affordable cost of sequencing and large number of genome-wide SNPs identified by NGS-based technologies for trait mapping and molecular breeding (Varshney et al., 2009b; Zhou et al., 2014). The recently completed genome sequences of the diploid ancestors of cultivated peanut were reported to be highly similar to cultivated peanut’s A and B subgenomes (Bertioli et al., 2016), which may due to its short evolutionary history after two diploid genomes merged ~3500–4500 years ago (Kochert et al., 1996). The genome sequences of the diploid ancestors were used as reference and proved to be successful in rapid trait mapping by NGS-based technologies in peanut (Agarwal et al., 2018; Bertioli et al., 2016; Pandey et al., 2017; Wang et al., 2018). One of such approach namely QTL-seq only requires whole-genome resequencing of extreme bulks and parent of mapping
population, making it cost effective (Takagi et al., 2013). This approach has been proved to be very successful with RIL populations in several crops including rice (Takagi et al., 2013), peanut (Pandey et al., 2016), chickpea (Singh et al., 2016a) and pigeonpea (Singh et al., 2016b). Therefore, we have applied the QTL-seq approach, in the present study, to identify genomic regions and candidate genes for the important economic trait namely SP in a RIL population (Yuanza 9102 × Xuzhou 68-4).

According to the QTL-seq approach, only one parent was sequenced and used as reference to compare the two extreme bulks (Takagi et al., 2013), and recent reports in cucumber (Lu et al., 2014), tomato (Illa-Berenguer et al., 2015), rapeseed (Hua et al., 2016), chickpea (Singh et al., 2016a), pigeonpea (Singh et al., 2016b), peanut (Pandey et al., 2017), soybean (Zhong et al., 2018) and rice (Wambu et al., 2018) used one parent as reference. Whether the choice of different parent would affect the results were not reported. In this study, both parents were sequenced and used together with the same bulks to generate two sets of results in parallel using the default parameters of the QTL-seq pipeline developed by the Iwate Biotechnology Research Center, Japan (Takagi et al., 2013), and this is the main methodological difference between this study and other QTL-seq reports in different crops. The numbers of identified genome-wide SNPs showed significantly difference between the two sets of results (Figure 3a). The large differences might due to multiple factors, for example: (i) the existing similarity between A and B subgenomes of the cultivated peanut; (ii) the high percentage of repetitive content of peanut (~64%); (iii) the reference-guided assembly of parents were constructed based on the genome sequences of both the diploid progenitor species A. duranensis and A. ipaensis, which might be still distant from the cultivated peanut; and (iv) complexity posed due to complex polyploid genome. These factors might cause different alignment errors and detection of false-positive SNPs upon using different parental reference genomes for sequence analysis of pooled samples. For example, 22 reads (20C + 2G) of LB were mapped at the 6470236 position of chromosome B02 when Xuzhou 68-4 was used as reference; however, only eight reads (6C + 2G) of LB were mapped at this position when Yuanza 9102 was used as reference.
was used as reference. This SNP was identified when Xuzhou 68-4 was used as reference while it was treated as spurious and filtered out when Yuanza 9102 was used as reference (Table 3). The validated SNPs Araip.B02.6155951 and Araip.B02.6770282 (Table 4) were filtered out when Xuzhou 68-4 was used as reference because their consensus qualities (18 and 12 respectively) were lower than the default cutoff (20) of the QTL-seq pipeline. However, in order to address the reason for the large differences, a large-scale testing needs to be conducted on species with different ploidy levels and on different combinations of internal parameters of the QTL-pipeline. Nevertheless, such large differences did not affect the identification of genomic regions for SP (Table 2). With different parent as reference, two overlapped genomic regions on A09 (2.75 Mb) and B02 (1.1 Mb) were identified as candidates controlling SP. Notably, some candidate SNPs in these regions were unique to either of the parent-based reference, that is, they were less significant than the selected region (Figure 4), which might be the result of recombination suppression; (ii) they were significant (P < 0.01) when Xuzhou 68-4 used as reference (Figures 5a and S6) but not significant (P > 0.01) when Yuan 9102 used as reference (Figures 5d and S9). As shown in Figure 5, we selected the overlapped genomic regions of two sets of results as candidates controlling SP, which were validated by KASP markers. This is a benefit of sequencing both parents.

The genomic regions identified with the QTL-seq approach were robust and more precise than the classical SSR marker-based QTL mapping. The present study successfully identified two genomic regions responsible for SP with stable expressions across five environments. The genomic region (66.75–69.50 Mb) on chromosome A09 was mapped on the same location with the previously identified major QTL cqSPA09 through classical SSR marker-based genetic mapping (Luo et al., 2017). This study has also successfully narrowed down the QTL region mapped on A09 from 44 Mb (26–70 Mb, identified through genetic mapping) to 2.75 Mb. Similarly, the genomic region (5.85–6.95 Mb) on chromosome B02 identified in this study also harboured the previously identified major QTL cqSPB02 through classical SSR marker-based genetic mapping (Luo et al., 2017). This study successfully narrowed down the second genomic region mapped on B02 from 3.18 Mb (5.95–9.13 Mb) to 1.1 Mb. Furthermore, the genomic regions for the other 23 QTLs, reported by Luo et al. (2017) with lower PVE and less stable across environments, could not be identified based on the QTL-seq approach. These results illustrated that the QTL-seq approach could be successful in identifying candidate regions for major and stable QTLs but not for minor QTLs.

The QTL-seq approach facilitated discovery of 12 putative genes in the 2.75 Mb genomic region on A09, encoding histone-lysine N-methyltransferase SUVR2-like isoform X2, oxysterol-binding protein-related protein, ankyrin repeat family, zeta-carotene desaturase, serine carboxypeptidase-like, RNA-binding protein, pentatricopeptide repeat (PPR) superfamily protein and function unknown proteins (Table S9). Notably, there was only one non-synonymous SNP in this region which affected the candidate gene (Araip. ASUVR9) encoding histone-lysine N-methyltransferase SUVR2-like isoform X2, which was reported to be involved in transcriptional gene silencing by RNA-directed DNA methylation in Arabidopsis thaliana (Han et al., 2014; Liu et al., 2018). The possible role of RNA-directed DNA methylation in controlling of SP in peanut needs to be further investigated. Similarly, the genomic region identified in 1.1 Mb region on B02 harboured 78 putative genes mainly related to catalytic activity, binding, metabolic process and cellular process (Table S9). Notably, there were seven non-synonymous SNPs and two SNPs in UTR in this region and they affected eight candidate genes (Table 3). Araip.R30RS and Araip.N485E were predicted to code for serine-threonine protein phosphatases, which might be

<table>
<thead>
<tr>
<th>Marker</th>
<th>Environment</th>
<th>PVE (%)</th>
<th>LOD</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aradu.A09.66949737</td>
<td>Yangluo2017</td>
<td>11.44</td>
<td>4.92</td>
<td>2.23E-06</td>
</tr>
<tr>
<td></td>
<td>Wuhan2016</td>
<td>13.50</td>
<td>6.16</td>
<td>1.21E-07</td>
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<tr>
<td></td>
<td>Wuhan2015</td>
<td>15.48</td>
<td>7.19</td>
<td>1.10E-08</td>
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<tr>
<td></td>
<td>Wuhan2014</td>
<td>12.14</td>
<td>5.63</td>
<td>4.21E-07</td>
</tr>
<tr>
<td></td>
<td>Wuhan2013</td>
<td>13.78</td>
<td>6.41</td>
<td>6.60E-08</td>
</tr>
<tr>
<td>Araip.B02.6155951</td>
<td>Yangluo2017</td>
<td>19.20</td>
<td>8.42</td>
<td>&lt;1.0E-09</td>
</tr>
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<td></td>
<td>Wuhan2016</td>
<td>14.09</td>
<td>6.31</td>
<td>8.40E-08</td>
</tr>
<tr>
<td></td>
<td>Wuhan2015</td>
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<td>6.90</td>
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PVE, phenotypic variation explained.
the regulators of signal transduction cascades and has been reported to be crucial factor for correct cell division and differentiation in *A. thaliana* (Uthiken et al., 2014). *Araip.J1578* would code for vacuolar protein sorting-associated proteins, which form endosomal the sorting complex required for plant development (Cai et al., 2014). *Araip.V24DX* might code for Ulp1 protease family protein, which regulate protein desumoylation and flower development (Murta et al., 2003). The SNP *Araip.B02_6776001* showed significant association with SP and was found located in the 5′-UTR of *Araip.X1ALQ* encoding unknown protein. Therefore, the unknown proteins coded by *Araip.G7S23*, *Araip.HEW90*, *Araip.Y0313* and *Araip.X1ALQ* may possess function in the determination of SP as well. Based on these findings, these genes should be targeted as candidates for fine mapping and gene cloning to reveal the genetic basis of SP in peanut.

The main advantage of the sequencing-based trait mapping approaches is development of markers in addition to gene discovery for target traits (Pandey et al., 2016). The QTL-seq approach has earlier helped in identification of genomic regions and discovery of candidate genes in addition to development of diagnostic markers for foliar disease resistance in peanut (Pandey et al., 2017). The QTL-seq approach deployed in present study also successfully facilitated development and validation of four KASP markers (for one SNP on A09 and for three SNPs on B02) for used in molecular breeding to improve SP. When these KASP markers were used to screen 337 cultivars, five cultivars possessing both alleles of the identified QTLs from Yuanza 9102 showed high shelling percentages, ranging from 76.78% to 81.42% in two environments. When used in selection programs, genotyping with KASP makers is a cost-effective approach than SSR markers (Steele et al., 2018) in addition to being amenable to high-throughput genotyping. These results indicated the potential of these KASP markers in tracking the favourable alleles for SP in the breeding programs.

In summary, the QTL-seq approach is powerful to refine genomic regions and identify candidate genes for major and stable QTLs for SP. Utilization of both parents as reference genome generated more comprehensive information for the exploration of candidate SNPs and genes for traits of interest. The present study successfully not only significantly narrowed down the genomic regions for both QTLs but also identified candidate genes and markers for SP for using in breeding.

**Materials and methods**

**Plant materials and phenotyping**

A RIL population consisting of 195 lines (*F₉* generation) was developed from the cross Yuanza 9102 × Xuzhou 68-4 by single seed decent (SSD) method (Luo et al., 2017). The shelling percentage (SP = \( \frac{\text{Weight of kernels}}{\text{Weight of pods}} \times 100\% \)) of female parent, Yuanza 9102, is significantly higher than that of the male parent, Xuzhou 68-4 (Figure 1). Phenotyping data for SP of the RIL population were generated in four consecutive years in the experimental field in Wuhan city, China and designated as Wuhan2013, Wuhan2014, Wuhan2015 and Wuhan2016 (Luo et al., 2017). To test the stability of the identified QTLs, the RIL population in *F₉* generation and the two parents were planted in another experimental field in the rural area in Yangluo, Wuhan, China in 2017, designated as Yangluo2017. They were planted and phenotyped as explained in the earlier study (Luo et al., 2017). Statistical analysis was performed using IBM SPSS Statistics Version 22 software (Armonk, NY).

**Construction of extreme bulks**

Genomic DNA of the RILs in *F₉* generation and the two parents were extracted from the young leaves collected in the Yangluo2017 environment using CTAB method (Doyle, 1990). For developing the extreme bulks, 15 RILs with high SP and 15 RILs with low SP were selected based on the mean phenotypic values of precise phenotyping data obtained in four environments, that is, Wuhan2013, Wuhan2014, Wuhan2015 and Wuhan2016 (Luo et al., 2017). High SP bulk (HB) were pooled with same amount of DNA from the 15 selected RILs with high phenotypic values, and similarly low SP bulk (LB) were pooled with same amount of DNA from the 15 selected RILs with low mean phenotypic values.

**Construction of libraries and Illumina sequencing**

A total of four Illumina libraries were prepared for the HB and LB bulks mentioned above as well as the two parents, using the NEBNext Ultra II DNA Library Prep Kit for Illumina. For each library, two mg DNA was sheared, end repaired, adapter ligated and separated using 2% agarose gel. The fragments around 350 bp were purified and enriched using the adaptor compatible PCR primers. After size checking with Agilent Technologies 2100 Bioanalyzer (Santa Clara, CA), pair-ended 150 bp short reads of the four amplified libraries were generated on NovaSeq platform with NovaSeq 6000 54 Reagent Kit.

**Identification of genomic regions for shelling percentage**

The QTL-seq pipeline (QTL-seq_framework1.4.4), developed by the Iwate Biotechnology Research Center, Japan (Takagi et al., 2013), was downloaded and used for the mapping of QTLs for SP with few modification (Figure 2). The raw sequencing reads with more than 10% of nucleotides showing Phred quality scores less than 30 were filtered out to generate cleaned reads.

To construct reference-based assembly for the parent Xuzhou 68-4, its cleaned reads were first aligned to the genomic sequences of *A. duranensis* and *A. ipaensis* (Bertioli et al., 2016) using the BWA software (Li and Durbin, 2009). The Coval software (Kosugi et al., 2013) was used to refine the short-read sequence alignments. Then, SNPs were identified with the Samtools software and refined with the Coval software. Finally, the Xuzhou assembly was developed for male parent Xuzhou 68-4 by substituting the reference bases with alternative bases at the positions of confidence SNPs in the genomic sequences of *A. duranensis* and *A. ipaensis*. Similarly, a reference-based assembly, the Yuanza assembly, was constructed for the other parent Yuanza 9102.

To discover the genomic regions for QTLs controlling SP, the reads from high and low bulbs were first aligned to the Xuzhou assembly, and SNPs were called for both the bulks. SNP-index for each SNP position was calculated for both the bulks using the formula: SNP-index (at a position) = \( \frac{\text{Count of alternate base}}{\text{Count of reads aligned}} \) (Figure S3). ΔSNP-index was then calculated by subtracting SNP-index of high bulk from SNP-index of low bulk for each SNP position except those with read depth <10 in both the bulks and SNP-index <0.3 in either of the bulks. Sliding window analysis was conducted with 2 Mb interval and 50 kb increment for SNP-index and ΔSNP-index plots (Figure S3–S9). Only SNPs with ΔSNP-index significantly higher than 0.5 or lower than −0.5 at the 0.01
confidence level were considered as the effective SNPs for SP. The effects of these SNPs were analysed with the SnpEff v3.0 software (Cingolani et al., 2012). On the other hand, the reads from high and low bulks were aligned to the Yuanza assembly to identify genomic regions for SP in the same way. The overlapped genomic regions identified with both parental assemblies were finally identified as the genomic regions controlling SP in the RIL population.

**KASP marker development and validation of identified candidate genomic regions**

In order to validate the identified genomic regions for SP in the RIL population, 100 bp upstream and downstream sequences of genic SNPs were used for the development of KASP markers (He et al., 2014). For each SNP, two allele-specific forward primers and one common reverse primer were designed and synthesized in LGC Genomic Ltd. Hoddesdon, UK. All KASP primers were listed in Table 4. These markers were used to genotype all individuals of the RIL population. The association between genotyping data and phenotyping data were calculated by single-marker analysis (SMA) with the Windows QTL Cartographer software (Wang et al., 2012). The phenotyping data generated in the Yangluo2017 environment and the four environments used for making the pool as mentioned above were used for this analysis.

**Acknowledgements**

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**Conflict of interest**

The authors declare that there is no conflict of interest.

**Author contribution**


**References**


Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1 Phenotypic differences in the representative RILs.

Figure S2 Phenotypic distribution of shelling percentage in the RIL population across five environments.

Figure S3 Alignment, SNP identification and calculation of SNP index for shelling percentage.

Figure S4 SNP-index plots for 20 pseudomolecules of low bulk with parent Xuzhou 68-4 as reference.

Figure S5 SNP-index plots for 20 pseudomolecules of high bulk with parent Xuzhou 68-4 as reference.

Figure S6 The Δ(SNP-index) plot obtained by subtraction of high pool SNP-index from low pool SNP-index using parent Xuzhou 68-4 as reference.

Figure S7 SNP-index plots for 20 pseudomolecules of low bulk with parent Yuanza 9102 as reference.

Figure S8 SNP-index plots for 20 pseudomolecules of high bulk with parent Yuanza 9102 as reference.

Figure S9 The Δ(SNP-index) plot obtained by subtraction of high pool SNP-index from low bulk SNP-index using parent Yuanza 9102 as reference.

Figure S10 Genotypes of the four KASP markers in the RIL population displayed in the SNPviewer software.

Figure S11 Boxplot of shelling percentage for RILs with different combinations of the two identified QTLs in the present study.

Figure S12 Boxplots of shelling percentages for diverse cultivars screened with KASP markers.

Table S1 Details on the recombinant inbred lines (RILs) selected for construction of extreme bulks.

Table S2 Details on whole genome re-sequencing data generated on parental genotypes and bulks.

Table S3 Pseudomolecule-wise SNPs distribution between extreme bulks for shelling percentage.

Table S4 Identification of SNPs for shelling percentage between extreme bulks using the Xuzhou assembly.

Table S5 Identification of SNPs for shelling percentage between extreme bulks using the Yuanza assembly.
Table S6 Genotypes of the developed KASP markers in the RIL population.
Table S7 Phenotyping data generated in the Yangluo2017 environment for the RIL population.
Table S8 Phenotypic effect of the two major and stable QTLs for shelling percentage in the RIL population.
Table S9 Putative genes and their functional annotations within the identified regions on chromosomes A09 and B02.