QTL for Yield Traits and Their Association with Functional Genes in Response to Phosphorus Deficiency in *Brassica napus*

Taoxiong Shi1,2, Ruiyuan Li1, Zunkang Zhao1,2, Guangda Ding1,2, Yan Long1, Jinling Meng1, Fangsen Xu1,2, Lei Shi1,2*

1 National Key Laboratory of Crop Genetic Improvement and National Centre of Plant Gene Research, Huazhong Agricultural University, Wuhan, China, 2 Key Laboratory of Arable Land Conservation (Middle and Lower Reaches of Yangtze River), Ministry of Agriculture, Huazhong Agricultural University, Wuhan, China

Abstract

**Background:** Oilseed rape (*Brassica napus* L.) is one of the most important oil crops. A primary limitation to the cultivation of this crop is the lack of available phosphorus (P) in soils. To elucidate the genetic control of P deficiency tolerance in *Brassica napus*, quantitative trait locus (QTL) for seed yield and yield related-traits in response to P deficiency were identified using a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ningyou 7 and a P-inefficient cultivar, Tapidor.

**Results:** Three field trials were conducted to determine seed yield (SY), plant height (PH), number of primary branches (BN), height to the first primary branch (FBH), relative first primary branch height (RBH), pod number per plant (PN), seed number per pod (SN) and seed weight of 1,000 seeds (SW) in 188 lines of TN DH population exposed to low P (LP) and optimal P (OP) conditions. P deficiency decreased PH, BN, SN, PN and SY, and increased FBH and RBH with no effect on SW. Three reproducible LP-specific QTL regions were identified on chromosomes A2, A3 and A5 that controlled SN, PN and SW respectively. In addition, six reproducible constitutive regions were also mapped with two each for SY-LP on A2, and FBH-LP on C6 and one each for PH-LP and SW-LP on A3. About 30 markers derived from 19 orthologous genes involved in *Arabidopsis* P homeostasis were mapped on 24 QTL regions by comparative mapping between *Arabidopsis* and *Brassica napus*. Among these genes, GPT1, MGD2 and SIZ1 were associated with two major loci regulating SY-LP and other yield-related traits on A2 between 77.1 and 95.0 cM.

**Conclusion:** The stable QTLs detected under LP conditions and their candidate genes may provide useful information for marker-assisted selection in breeding high-yield *B. napus* varieties with improved P efficiency.

Introduction

Phosphorus (P) is one of the essential macronutrients for plants and is a major limitation for plant development worldwide due to its low availability and inaccessibility in soil [1–3]. To mitigate this, P fertilizers are applied to improve P availability in soils. However, cost and low recovery rates of P fertilizers not only increase the cost of crop production but also lead to serious environmental pollution and exhaust non-renewable phosphate resources. Therefore, a sustainable strategy for crop production would be to breed crops with high efficiency in acquiring P from native soil reserves or fertilizer sources.

Nutrient-efficient plants produce higher yields per unit of nutrient applied or absorbed compared with other plants grown under similar agroecological conditions [4]. One way to determine the genetic basis of tolerance to nutrient deficiency is to map quantitative trait loci (QTL) based on molecular markers. A number of QTLs associated with P efficiency traits have been detected in rice [5–7], maize [8], wheat [9,10], common bean [11,12] and soybean [13,14]. Using a soybean *F₂* recombinant inbred lines (RILs), 13 QTLs associated with root traits and 18 associated with P-efficiency were identified under low P and high P conditions, and three QTL clusters were found to be located with traits for root and P-efficiency at low P levels [14]. In wheat, each QTL detected for P use efficiency (PUE) was found to be linked to QTLs regulating agronomic traits [10]. *Pup1* is a major locus that confers P deficiency tolerance in rice [5,7,15,16]. Overexpression of a *Pup1*-specific protein kinase gene (*PSTOL1*) in phosphorus-starvation-intolerant varieties significantly enhanced grain yield in phosphorus-deficient soil [17]. These results provide strong evidence for the hypothesis that enhancing P efficiency would improve agronomic performance of crops.

*E-mail: leishi@mail.hzau.edu.cn*
Brassica napus is one of the most important oilseed crops worldwide and is extremely sensitive to P availability in soil. Since high seed yield is one of the desired targets in B. napus breeding, various studies have reported the importance of identifying P-efficient germplasm in B. napus [18]. QTLs for seed yield related traits in response to P deficiency were first identified in B. napus using an F10 RIL population in two crop seasons. In these experiments, a number of low-P-specific QTLs were detected across seasons. Four markers developed from genes involved in Arabidopsis P homeostasis were mapped to the confidence intervals of QTLs associated with a P efficiency coefficient (the ratio of seed yield at low P level to that at adequate P level) or low-P-specific QTLs [19]. Identification of QTLs associated with root traits and P-efficiency parameters partially revealed the genetic basis of tolerance to P deficiency and also confirmed the importance of root morphology traits in adaption to low P levels [20,21]. High colinearity and synteny between the Brassica species and Arabidopsis genomes have allowed in silico mapping of multiple Arabidopsis thaliana genes involved in P metabolic pathway as well as genes controlling agronomical traits in B. napus QTLs. These include QTLs for shoot mineral concentrations [22], seed mineral concentrations [23], root morphological traits [20,21], flowering time [24] and seed yield traits [19,25]. The sequence of B. rapa genome [26] and B. oleracea genomes [http://www.ocri-genomics].

Figure 1. Trait differences between Tapidor and Ningyou 7 under low (LP) and optimal phosphorus (OP) conditions in three field trials. Field trial conducted from Sept. 2008 to May 2009 (Tri.1), field trial conducted from Sept. 2009 to May 2010 (Tri.2), field trial conducted from Sept. 2010 to May 2011 (Tri.3).
doi:10.1371/journal.pone.0054559.g001
org/) has further facilitated identification of conserved genome segments and prediction of homologous genes for target traits in *B. napus*, in addition to confirming QTLs detected previously for shoot P content and P uptake efficiency related traits in *Brassica* species [21–23,27–30].

In the present study, a double haploid (DH) population derived from a cross between P-efficient cv. Ningyou 7 and P-inefficient cv. Tapidor (TN DH population) was used to determine seed yield and yield-related traits under low P (LP) and optimal P (OP) conditions in three trials conducted during four years. In addition, QTLs controlling seed yield and yield-related traits conferring P-deficiency tolerance were also determined. Comparative mapping QTLs controlling seed yield and yield-related traits conferring P-deficiency tolerance were also determined. Comparative mapping QTLs for phosphorus efficiency in *Brassica* species was also performed to associate QTLs with orthologous genes involved in tolerance to P deficiency. QTLs that confer P efficiency and their candidate genes would provide useful information to better understand P efficiency and to develop molecular markers to improve seed yield in *B. napus* cultivated in environments with low P.

**Results**

**Phenotypic analysis of the tested traits in TN DH population**

Phenotypic differences in the tested traits between the two parental lines under low P (LP) and optimal P (OP) conditions are presented in Figure 1. Compared to the P-inefficient parent ‘Tapidor’, the P-efficient parent ‘Ningyou 7’ had higher number of primary branches (BN), seed weight of 1,000 seeds (SW), seed number per pod (SN) and seed yield (SY), and lower height to the first primary branch (FBH) and relative first primary branch height (the ratio of FBH to plant height (PH), RBH) under both P conditions. However, average pod number per plant (PN) in Ningyou 7 was lower than Tapidor under LP conditions. There was no significant difference in PH between the TN DH parents under two P conditions in Tri.1 (field trial conducted from Sept. 2008 to May 2009) and Tri.2 (field trial conducted from Sept. 2009 to May 2010), while PH in Ningyou 7 was higher than Tapidor in Tri.3 (field trial conducted from Sept. 2010 to May 2011). Each parent experienced lower PH, BN, SN, PN and SY in P deficient than optimal P conditions. In contrast, each parent had greater FBH and RBH in low than optimal conditions and experienced no different in SW across P levels. 

TN DH population exposed to P deficiency also showed lower PH, BN, SN, PN and SY, and higher FBH and RBH with no significant influence of P deficiency was on PN (decreasing by 20.7% in Tri.1, 39.3% in Tri.2 and 20.1% in Tri.3) and RBH (increasing by 21.4% in Tri.1, 39.3% in Tri.2 and 20.1% in Tri.3) and RBH (increasing by 21.4% in Tri.1, 39.3% in Tri.2 and 20.1% in Tri.3). Each parent experienced lower PH, BN, SN, PN and SY in P deficient than optimal P conditions. In contrast, each parent had greater FBH and RBH in low than optimal conditions and experienced no different in SW across P levels.

TN DH population exposed to P deficiency also showed lower PH, BN, SN, PN and SY, and higher FBH and RBH with no effect on SW (Table 1 and Table S1; Figure 2). The most significant influence of P deficiency was on PH (decreasing by 67.2% in Tri.1, 61.6% in Tri.2 and 48.9% in Tri.3) followed by BN (decreasing by 32.2% in Tri.1, 28.1% in Tri.2 and 34.3% in Tri.3) and RBH (increasing by 21.4% in Tri.1, 39.3% in Tri.2 and 46.4% in Tri.3). Genotypic variation in each tested trait also existed in TN DH population under both P conditions. Moreover, variation coefficients for BN, SN, PN and SY in LP conditions were larger than in OP conditions in all three trials (Table 1). In addition to the strong effects of environment, genotype, environment x P level and genotype x P level interactions for most traits, there were also strong effects of P level on all traits (Table 2). Transgressive segregation and continuous distribution was observed for each trait in TN DH population exposed to both P conditions in all three trials (Figure 2).

Pearson’s correlation coefficient between traits was also calculated (Table 3). The phenotypic correlation between SY and yield-related traits under LP was similar to OP conditions in the TN DH population. SY was positively correlated to PH, BN, SN and PN, and negatively correlated to RBH. However, higher correlation coefficients between SY and PH, RBH, SN, and lower correlation coefficients between SY and PN were observed under LP conditions than under OP conditions (Table 3).

**QTL for seed yield and yield-related traits**

A total of 155 significant QTLs were identified for the eight tested traits from all three trials, including 79 QTLs detected under LP conditions and 76 under OP conditions (Figure 3 and Table S2). Intervals of several QTLs overlapped with each other resulting in clustering of the 155 QTLs into 81 regions on 15 chromosomes. A majority (80.0%) of these regions was distributed on A2, A3, A5, A9, C6 and C9 (Figure 3). Of the 81 QTL regions, 29 were specifically detected under LP condition, and 29 specifically under OP condition. The remaining 23 QTL regions were detected under both P conditions (Figure 4 and Table S2).

The 29 LP-specific regions were derived from 42 QTLs detected under LP conditions, with individual phenotypic contributions of QTL ($R^2$) ranging from 6.1 to 12.9% (Figure 4 and Table S2). Three of the 29 LP-specific regions were consistently associated with the same trait in two of the three trials and mapped between 56.4–66.4 cM on A2, 80.0–92.1 cM on A3 and 0.0–28.2 cM on A5 and controlled SN, PN and SW respectively. The consistency of PN locus on A3 also coincided with QTLs for SY-LP and SW-LP. Furthermore, there were four LP-specific regions where QTLs for different traits were also observed. For example, qPN-LP2-A3a and qSW-LP1-A3c were co-located between 27.8–52.3 cM on A3; qSN-LP3-A3 and qPH-LP2-A3c were observed between 58.5–61.5 cM on A3; qRBH-LP1-A4, qPN-LP3-A1 and qBN-LP1-A1 were clustered between 57.5–84.9 cM on A4; and qSY-LP1-C1a was associated with qSN-LP2-C1 located between 40.8–48.6 cM on C1.

The 29 OP-specific QTL regions were integrated from 40 QTLs detected under OP conditions, with individual $R^2$ ranging from 5.9–17.6% (Figure 4 and Table S2). Among the 29 OP-specific regions, two regions were consistently mapped between 59.8–63.4 cM on A3 and 36.9–43.7 cM on A9 that regulate SW and PH respectively in two of the three field trials. In region 56.9–64.5 cM on C1, qSY-OP1-C1 coincided with qSN-OP2-C1 and the QTLs for FBH and RBH were co-located in five regions: 0.0–5.7 cM on A3, 18.2–28.1 cM and 128.2–141.2 cM on A9, 28.9–46.4 cM on C6 and 54.1–70.5 cM on C9. In addition, QTL for FBH also overlapped with QTL for PH in two regions: 28.4–34.8 cM on A3 and 28.7–31.0 cM on A9.

The 23 constitutive QTL regions detected under both P conditions were integrated from 73 QTLs, with individual $R^2$ varying from 5.7 to 20.2% (Figure 4 and Table S2). Six regions were considered as major loci that respond to LP deficiency because these QTLs were consistently associated with the same trait under LP conditions in at least two trials. QTLs for SY-LP were detected in two of the three trials and mapped in the 77.1–91.3 cM and 91.3–95.0 cM regions on A2 accounting for 10.8–13.0% and 8.4–14.2% of SY-LP variation, respectively, and both also coincided with QTLs for SN-OP. The reproducible locus for PH-LP was located between 50.2–59.9 cM on A3, accounting for 7.9–9.2% of PH-LP variation. The reproducible locus for SW-LP was located in 70.2–77.0 cM region on A3, explaining 7.8–11.5% of SW-LP variation. The two major regions affecting FBH-LP were detected in all three field trials and mapped in the intervals of 46.0–50.8 cM and 51.6–61.5 cM on C6, explaining 9.2–17.8% and 12.4–18.5% of FBH-LP variation respectively.
Association of QTLs with functional genes involved in P homeostasis

In total, 30 gene-based markers (GBMs) associated with 19 P homeostasis genes in Arabidopsis were found to be located in the QTL confidence intervals. These 30 GBMs were grouped into six functional categories according to the recent functional classification of the corresponding Arabidopsis genes (Table 1). P transporter category was the largest group with 10 GBMs. Eleven GBM markers including BnAP1/1-A1, BnLPK2-A4, BnPHT1-A4, BnPT1-A5, BnLO1-1A/d/e, BnSOD2-A10a/c, BnGPT2-C1 and BnRNS1-C7 derived from eight genes were associated with LP-specific QTL regions. Eight GBM markers (BnPHR1-A1b, BnPH1-A3-A4/A6, BnBHLH32-A6, BnLPR1-A9a, BnDG-2-A9 and BnPHO1-C1) derived from six genes were related to OP-specific QTL regions. Eleven GBM markers (BnPHR1-A1a, BnPHT1-A2, BnSIZ1-A2, BnMGD2-A2a/b, BnWRKY75-A3, BnPH13-A3, BnPYK10-A3a/b, BnSQD1-A3 and BnPH1-C6) derived from nine genes were located in the constitutive QTL regions. Twelve of the 19 genes distributed in the QTL confidence intervals were mapped to corresponding genetic positions on B. napus or B.oleracea genomes included genes related to P transport (PHR1, PHO1, GPT1 and GPT2), transcriptional control (SIZ1, PRR1, WRK73 and BHLH32), phospholipid metabolism (MGD2, SQD2), metal binding (LPR1), and C-compound and carbohydrate metabolism (PIT10) (Figure 4 and Table 4). Among the twelve genes, SQD2 and GPT2 were associated with LP-specific QTL region between 72.8–87.3 cM on A10 regulating SW-LP, respectively. PHR1, BHLH32 and PHO1 were associated with the OP-specific QTL region between 43.6–49.8 cM on A1, 83.1–102.8 cM on A6 and 56.9–64.5 cM on C1 that regulate FBH-OP, BN-OP and SY-OP and SN-OP, respectively. PHT1 and PEP10 were linked to the constitutive QTL region between 0.0–8.4 cM on C6 regulating SN-LP/OP and 70.2–77.0 cM on A5 controlling SW-LP/OP, respectively. PTT1 and MGD2, and SQZI were repeatedly associated with the constitutive QTL regions regulating SY-LP mapped at 77.1–91.3 cM and 91.3–95.0 cM on A2, respectively.

Discussion

The purpose of this study was to detect P efficiency related QTLs regulating seed yield and yield-related traits using the TN DH population of B. napus derived from a cross between P-efficient cv. Tapidor and P-efficient cv. Ningyou 7, and to associate QTLs with functional genes involved in P homeostasis.

Effect of P deficiency on seed yield and yield-related traits

Both parents, Tapidor and Ningyou 7, showed significant differences in the tested traits under both low P (LP) and optimal P (OP) conditions (Figure 1). Ningyou 7 had higher number of primary branches (BN), seed weight of 1,000 seeds (SW), seed number per pod (SN) and seed yield (SY) and lower height to the first primary branch (FBH), relative first primary branch height (RBH) and pod number per plant (PN) than Tapidor under both P conditions. QTL mapping revealed that majority of the Ningyou 7 alleles among the identified QTLs were associated with increased BN, SW, SN and SY but decreased PN under both LP and OP conditions (Table 4), indicating that the contribution of Ningyou 7 seed yield was higher than Tapidor under both P conditions. P deficiency significantly decreased PN, BN, SN and plant height (PH), increased FBH and RBH, and had no effect on SW in the double haploid mapping population (TN DH) derived from a double-haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ningyou 7 and a P-efficient cultivar, Tapidor under low (LP) and optimal phosphorus (OP) conditions in three field trials.

Table 1. Trait differences in a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ningyou 7 and a P-inefficient cultivar, Tapidor under low (LP) and optimal phosphorus (OP) conditions in three field trials.

| Trait | P level | Tri.1 | Tri.2 | Tri.3 |
|-------|---------|-------|-------|-------|
|       | mean    | range | cv%   | mean  | range | cv%   | mean  | range | cv%   |
| FBH   | LP      | 38.3  | 13.7–61.2 | 23.5 | 41.2  | 19.2–59.9 | 19.6 | 49.2  | 28.9–69.4 | 15.6 |
|       | OP      | 38.5  | 8.0–66.0   | 25.5 | 39.0  | 10.0–69.0 | 25.0 | 35.0  | 14.0–62.3 | 24.3 |
| PH    | LP      | 116.0 | 84.5–143.3 | 9.6  | 108.2 | 74.5–137.2 | 10.9 | 124.1 | 84.0–153.6 | 9.5  |
|       | OP      | 140.7 | 106.8–172.5 | 9.0  | 124.6 | 89.0–161.6 | 11.0 | 138.2 | 84.0–174.5 | 9.1  |
| RBH   | LP      | 0.34  | 0.15–0.61  | 24.7 | 0.39  | 0.17–0.59 | 21.1 | 0.41  | 0.23–0.66 | 16.9 |
|       | OP      | 0.28  | 0.09–0.47  | 21.2 | 0.28  | 0.14–0.48 | 20.8 | 0.28  | 0.07–0.55 | 27.1 |
| BN    | LP      | 4.0   | 2.3–6.0    | 17.0 | 4.1   | 2.6–7.0   | 19.4 | 4.5   | 2.5–7.5   | 19.2 |
|       | OP      | 5.9   | 4.4–8.8    | 22.5 | 5.7   | 2.8–8.8   | 18.3 | 6.9   | 2.5–10.0  | 16.2 |
| SW    | LP      | 2.88  | 2.04–4.18  | 14.1 | 3.25  | 2.10–4.91 | 15.1 | 2.83  | 1.88–4.77 | 16.2 |
|       | OP      | 2.93  | 2.10–4.30  | 13.9 | 3.16  | 2.09–4.95 | 14.6 | 2.78  | 1.84–4.85 | 15.0 |
| SN    | LP      | 13.3  | 3.0–25.3   | 28.4 | 13.5  | 6.3–20.3  | 21.6 | 14.8  | 5.3–22.9  | 26.6 |
|       | OP      | 17.8  | 7.9–26.2   | 19.9 | 14.9  | 7.3–25.5  | 20.5 | 15.5  | 5.7–23.5  | 23.0 |
| PN    | LP      | 61.4  | 19.0–148.5 | 40.2 | 57.9  | 12.0–115.5| 37.8 | 71.3  | 20.7–142.3| 37.3 |
|       | OP      | 187.0 | 43.0–340.0 | 29.9 | 150.6 | 33.0–321.0| 34.3 | 139.6 | 35.5–259.1| 34.8 |
| SY    | LP      | 484.1 | 19.7–1155.6| 44.2 | 434.9 | 48.0–1007.3| 42.4 | 539.0 | 61.9–1450.0| 49.7 |
|       | OP      | 1649.7| 455.7–2832.8| 28.2 | 1271.0| 326.7–2572.9| 33.0 | 1163.9| 239.5–2135.7| 39.2 |

Note: Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN), seed weight of 1,000 seeds (g per 1,000 seeds; SW), seed number per pod (N; SN), pod number per plant (N; PN), seed yield per hectare (kg ha$^{-1}$; SY). Field trial conducted from Sept. 2008 to May 2009 (Tri.1), field trial conducted from Sept. 2009 to May 2010 (Tri.2), field trial conducted from Sept. 2010 to May 2011 (Tri.3).

doi:10.1371/journal.pone.0054559.t001
The cross between Ningyou 7 and Tapidor (Tables 1 and 2) is consistent with previous studies [19,25] indicating that SW was genetically more tightly controlled and less affected by environmental factors than other yield-related traits. Of the seven yield-related traits, PN, BN and RBH were most severely affected by P-deficiency (Table 1) in addition to having high correlation with SY (Table 3). Therefore, higher levels of PN and BN, and lower levels of RBH should be highly considered in the breeding of B. napus cultivars tolerant of environments with low P.

Dissection of stable QTLs associated with seed yield and yield-related traits conferring P-deficiency tolerance

In this study, a total of 155 putative QTLs associated with seed yield and seven yield-related traits were identified under LP and OP conditions in all three field trials (Table 4). These QTLs were clustered and integrated into 29 LP-specific, 29 OP-specific and 23 constitutive QTL regions (Figure 4 and Table S2). The genetic architecture of seed yield in B. napus was previously analyzed by QTL mapping of seed yield-related traits [25,31–35]. About 85 QTLs associated with seed yield were identified along with 785 QTLs for eight yield-associated traits using TN DH population from 10 natural environments with optimal P [25]. In the present study, a majority of these OP-specific QTLs were identified in only one of the three trials and 10 OP-specific QTLs, i.e. all QTLs for PH-OP mapped on A9, were located in the QTL associated with traits detected in semi-winter environments by Shi et al. [25]. These results indicated that quantitative traits were influenced both by heredity and growth conditions.

In recent studies on B. napus LP-specific QTLs were reported for yield-related traits [19], root morphological traits [20] and shoot traits at the vegetative stage [21]. Further analysis of these QTLs suggested that the presence of specific genes or differential expression of some genes regulated the response to P deficiency. In this study, nine QTL regions were detected for the same traits under LP conditions in at least two field trials, including three LP-specific regions and six constitutive QTL regions (Figure 4; Table S2). These stable QTL regions associated with the response to P deficiency have been compared with QTLs for P efficiency identified in a BE RIL population by comparative mapping [19–21,23]. Of the three reproducible LP-specific QTL regions regulating SN-LP (56.4–66.4 cM on A2), PN-LP (80.0–92.1 cM on A3) and SW-LP (0.0–28.2 cM on A5) (Figure 4 and Table S2), the regions for SN-LP and PN-LP were co-located with the QTLs

### Table 2. Significance of three-way ANOVA analysis for the eight traits among a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ningyou 7 and a P-inefficient cultivar, Tapidor under low and optimal phosphorus conditions in three field trials.

| Source                | FBH     | PH      | RBH    | BN     | SW     | SN     | PN     | SY     |
|-----------------------|---------|---------|--------|--------|--------|--------|--------|--------|
| Genotype              | 152331  | 262362.9| 8.8    | 1029.7 | 396.1  | 1671.3 | 126059 | 3915.2 |
| d.f.s                 | 187     | 187     | 187    | 187    | 187    | 187    | 187    | 187    |
| significant           | ***     | ***     | ***    | ***    | ***    | ***    | ***    | ***    |
| P level               | 19382.3 | 200903.8| 5.60   | 2838.0 | 0.4    | 2403.1 | 386117.7| 9484.5 |
| d.f.s                 | 1       | 1       | 1      | 1      | 1      | 1      | 1      | 1      |
| significant           | ***     | ***     | ***    | ***    | **     | **     | **     | **     |
| Environment           | 12902.1 | 109287.7| 0.4    | 415.1  | 67.2   | 918.1  | 141888.4| 591.8  |
| d.f.s                 | 2       | 2       | 2      | 2      | 2      | 2      | 2      | 2      |
| significant           | ***     | ***     | ***    | ***    | ***    | ***    | ***    | ***    |
| Genotype ×P level     | 9690.8  | 35805.5 | 1.3    | 230.0  | 25.6   | 2357.8 | 590475.3| 1075.8 |
| d.f.s                 | 186     | 186     | 186    | 186    | 187    | 187    | 187    | 187    |
| significant           | ***     | ***     | ***    | ***    | ***    | ***    | ***    | ***    |
| Environment ×P level  | 15061.4 | 16992.1 | 0.4    | 77.3   | 2.4    | 1375.3 | 233904.2| 789.2  |
| d.f.s                 | 2       | 2       | 2      | 2      | 2      | 2      | 2      | 2      |
| significant           | ***     | ***     | ***    | ***    | ***    | ***    | ***    | ***    |
| Environment ×Genotype | 38349.0 | 67629.7 | 2.7    | 584.0  | 103.6  | 4389.6 | 712967.7| 1094.2 |
| d.f.s                 | 342     | 341     | 343    | 343    | 350    | 344    | 345    | 345    |
| significant           | ***     | ***     | ***    | ***    | ***    | ***    | ***    | ***    |

Note: Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN), seed weight of 1,000 seeds (g per 1000 seeds; SW), seed number per pod (N; SN), pod number per plant (N; PN), seed yield per hectare (kg·ha⁻¹; SY).

Sums of squares (S.S), degrees of freedom (d.f.s).

**p<0.001**,

**p<0.01.**

doi:10.1371/journal.pone.0054559.g002
for SN and PN identified in the BE RIL population exposed to LP, respectively [19].

Among the six reproducible constitutive QTLs under LP condition, SY-LP was mapped between 77.1–91.3 cM and 91.3–95.0 cM on A2, FBH-LP between 46.0–50.8 cM and 51.6–61.5 cM on C6, PH-LP between 50.2–59.9 cM on A3 and SW-LP between 70.2–77.0 cM on A3 (Figure 4 and Table S2). The constitutive region from 77.1–91.3 cM on A2 played a major role in controlling SY-LP in Tri.1 and Tri.2 explaining 10.8–13.0% of SY-LP variation and also was detected for SN-OP in Tri.1 and Tri.3 accounting for 7.2–12.0% of SN-OP variation. The region from 91.3–95.0 cM on A2 was another reproducible locus for SY-LP in Tri.2 and Tri.3 explaining 10.8–13.0% of SY-LP variation and also played a major role in regulating SN-OP in Tri.1 and Tri.3 accounting for 7.2–12.0% of SN-OP variation. This region also coincided with QTLs for SN and PN in BE population exposed to LP [19]. The two major loci associated with FBH-LP that were repeatedly detected in all three trials was located on C6 between 46.0–50.8 and 51.6–61.5 cM, accounting for 9.2–17.8% and 12.4–18.5% of FBH-LP variation, respectively. The stable region for FBH-LP (51.6–61.5 cM on C6) associated with PH-LP (51.6–61.5 cM on C6) was co-localized with the reproducible QTL for PH in BE RIL population exposed to LP. The reproducible locus for SW-LP located between 70.2 and 77.0 cM on A3 also coincided with the stable loci for SW in BE RIL population exposed to LP [19]. These results indicated that common mechanisms may be involved in the response of different *B. napus* cultivars to P deficiency. In addition, stable QTL regions, 50.2–59.9 cM on A3 regulating PH-LP and PB-LP, co-localized with QTL regions for root morphology traits conferring P-efficiency in BE RIL population [20,21]. This indicated that selection of early vigor plants with more developed root system at the seedling stage could enhance P acquisition, improve seed yield-related traits and therefore increase the seed yield at low P.

![Figure 3. Distribution of quantitative trait loci (QTLs) associated with seed yield and yield-related traits on fifteen linkage groups under low (LP; above x-axis) and optimal phosphorus (OP; below x-axis) conditions in three field trials. Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN); seed weight of 1,000 seeds (g per 1000 seeds; SW); seed number per pod (N; SN); pod number per plant (N; PN); seed yield per hectare (kg·ha<sup>−1</sup>; SY).](doi:10.1371/journal.pone.0054559.g003)

**Table 3.** Pearson’s correlation coefficients among traits in a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ningyou 7 and a P-inefficient cultivar, Tapidor under low (below diagonal) and optimal phosphorus (above diagonal) conditions.

|       | FBH  | PH   | RBH  | BN   | SW   | SN   | PN   | SY  |
|-------|------|------|------|------|------|------|------|-----|
| FBH   | 0.52** | 0.89** | −0.22** | −0.25** | ns   | ns   | ns   | ns  |
| PH    | 0.45** | 0.15*  | 0.25** | 0.19*  | 0.16* | 0.22** | 0.22** | 0.22** |
| RBH   | 0.74** | ns    | −0.38** | −0.23** | ns   | −0.21** | −0.22** | 0.22** |
| BN    | −0.22** | 0.26** | −0.36** | 0.01   | ns   | 0.36** | 0.39** | 0.39** |
| SW    | −0.16*  | −0.08  | ns    | ns    | ns   | −0.19*  | ns    | ns  |
| SN    | ns    | 0.28** | ns    | 0.20*  | ns   | −0.05  | ns    | 0.52** |
| PN    | ns    | 0.22** | −0.21** | 0.30** | −0.19* | 0.24** | 0.70** | 0.70** |
| SY    | ns    | 0.31** | −0.33** | 0.39** | ns   | 0.58** | 0.55** | 0.55** |

Note: Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN); seed weight of 1,000 seeds (g per 1000 seeds; SW); seed number per pod (N; SN); pod number per plant (N; PN); seed yield per hectare (kg·ha<sup>−1</sup>; SY).

**P,** *P* < 0.01, *P* < 0.05.

doi:10.1371/journal.pone.0054559.t003
Figure 4. Location of quantitative trait loci (QTLs) on the linkage groups of *B. napus* and the genes aligned with *B. rapa* and *B. oleracea*. Hollow bars on the left represent the genome of *B. rapa* and *B. oleracea*, and horizontal-crossing lines indicate gene positions. Hollow bars on the right represent the linkage groups of *B. napus* and horizontal-crossing lines indicate the position of molecular markers. To the right of the linkage groups, the vertical lines with arrows indicate the QTL, with the length of each line indicating the confidence interval and circles indicating the peak position. Upward arrows represent alleles from Tapidor, and downward arrows represent alleles from Ningyou 7. Gray and black bars represent QTL detected under low (LP) and optimal phosphorus (OP) condition, respectively. Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN), seed weight of 1,000 seeds (g per 1000 seeds; SW), seed number per pod (N; SN), pod number per plant (N; PN), seed yield per hectare (kg ha⁻¹; SY). Trait name and the related trial number are labeled near each QTL line. Markers shown in red and oriented on the right of the *B. napus* hollow bars represent gene-based markers (GBM) mapped in QTL intervals in *B.napus* and the corresponding homologous genes aligned to *B.rapa* and *B.oleracea* are showed in the left.

doi:10.1371/journal.pone.0054559.g004

Table 4. Gene-based markers (GBM) associated with Quantitative trait loci (QTLs) regulating seed yield-related traits and their functions in *Arabidopsis*.

| GBM | Chr. | QTL region | Trait | Gene name  | Function in *Arabidopsis*                                      |
|-----|------|------------|-------|------------|---------------------------------------------------------------|
| *P* assimilation                      |      |            |        |            |                                                               |
| BnPAP17-A1                            | A1   | 74.0–82.5  | BN-LP | PAP17      | Encodes acid phosphatase                                      |
| BnRNS1-C7                             | C7   | 19.1–38.5  | FBH-LP| RNS1       | Encodes ribonuclease                                          |
| *Metal binding*                       |      |            |        |            |                                                               |
| BnLPR2-A4                             | A4   | 57.5–84.9  | RBH-LP| LPR2       | Controlling low Pi–triggered root growth inhibition           |
| BnLPR1-A9c/d/e                        | A9   | 66.0–71.5  | PH-LP | LPR1       | Controlling low Pi–triggered root growth inhibition           |
| BnLPR1-A9a                            | A9   | 55.1–63.1  | PH-OP | LPR1       | Controlling low Pi–triggered root growth inhibition           |
| *P* transporter                       |      |            |        |            |                                                               |
| BnPHF1-A4                             | A4   | 57.5–84.9  | RBH-LP| PHF1       | Enables the endoplasmic reticulum exit of a high-affinity phosphate transporter |
| BnPHT1-A5                             | A5   | 30.9–37.7  | SW-LP | PHT1;4     | Encodes phosphate (Pi) transporters                          |
| BnPHT1-A3                             | A3   | 28.4–34.8  | FBH-OP| PHT1;4     | Encodes phosphate (Pi) transporters                          |
| BnPHT1-A4                             | A4   | 21.4–29.7  | SW-OP | PHT1;8     | Encodes phosphate (Pi) transporters                          |
| BnPHT1-C6                             | C6   | 0.0–8.4    | SN-LP | PHT1       | Encodes phosphate (Pi) transporters                          |
| BnPHT3-A3                             | A3   | 12.2–18.6  | FBH-LP| PHT3;1     | Transmembrane phosphate transportor                          |
| BnGPT1-A2                             | A2   | 77.1–91.3  | SW-OP | PHT1       | Glucose-6-phosphate transporter                               |
| BnGPT2-C1                             | C1   | 50.2–57.2  | SY-LP | PHT2       | Glucose-6-phosphate transporter                               |
| BnPHO1-C1                             | C1   | 56.9–64.5  | SY-OP | PHO1       | Yxle loading of inorganic phosphate                          |
| *Phospholipid metabolism*             |      |            |        |            |                                                               |
| BnSQD1-A3                             | A3   | 114.0–127.3| PN-OP | SQD1       | Sulfortransferase activity                                    |
| BnSQD2-A10a/c                         | A10  | 72.8–87.3  | SW-LP | SQD2       | Sulfortransferase activity                                    |
| BnMGD2-A2a/b                          | A2   | 91.3–95.0  | RBH-OP| MGD2       | Encodes Type-B monogalactosyldiacylglycerol synthases        |
| BnDGD2-A9                             | A9   | 128.2–141.2| FBH-OP| DGD2       | Encodes a UDP-galactose-dependent digalactosyldiacylglycerol synthase |
| *Carbohydrate metabolism*             |      |            |        |            |                                                               |
| BnPYL10-A3a/b                         | A3   | 70.2–77.0  | SW-LP | PYK10      | Encodes beta-glucosidase, hydrolyzing O-glycosyl compounds    |
| *Transcriptional factor*              |      |            |        |            |                                                               |
| BnSIZ1-A2                             | A2   | 77.1–91.3  | RBH-OP| SIZ1       | SUMO ligase activity, a focal controller of Pi starvation-dependent response |
| BnPHR1-A1a                            | A1   | 30.1–38.2  | FBH-OP| PHR1       | MYB transcriptional activator of Pi starvation-responsive genes |
| BnPHR1-A1b                            | A1   | 43.6–48.8  | FBH-OP| PHR1       | MYB transcriptional activator of Pi starvation-responsive genes |
| BnBHL32-A6                            | A6   | 93.1–102.8 | BN-OP | BHL32      | Cellular response to phosphate starvation                     |
| BnPHT1-A6                             | A6   | 93.1–102.8 | BN-OP | PHT1;8     | Encodes phosphate (Pi) transporters                          |
| BnWRKY75-A3                           | A3   | 12.2–18.6  | FBH-OP| WRY75      | Transcription factor modulates PHT1 expression                |

Note: Height to the first primary branch (cm; FBH), plant height (cm; PH), relative first primary branch height (the ratio of FBH to PH; RBH), number of primary branches per plant (N; BN), seed weight of 1,000 seeds (g per 1000 seeds; SW), seed number per pod (N; SN), pod number per plant (N; PN), seed yield per hectare (kg ha⁻¹; SY), low phosphorus (LP), optimal phosphorus (OP).
doi:10.1371/journal.pone.0054559.g004
Detection of these reproducible loci in different populations that respond to P deficiency would provide useful information to conduct marker-assisted selection to breed *B. napus* with improved P-efficiency.

**Gene prediction in Brassica by in silico comparative mapping**

Unlike anonymous molecular markers which were widely used to construct *B. napus* genetic maps such as SSR and AFLP [24,36], GBMs developed from specific genes that are closely related to phenotypic traits could be useful in associating target traits with function [37,38]. For example, Yang et al. [20] found that the two mapped GBMs, BnIPS2 and BnPHT3, were linked to the QTL controlling root morphology in response to P starvation. The RI lines with favorable alleles related to BnIPS2 and BnPHT3 could develop larger root system, produce higher dry weight, acquire more P under LP conditions and produce more seeds [19,20]. In this study, a total of 30 GBMs derived from 19 genes involved in P homeostasis in *Arabidopsis* were found to be linked to 24 QTL regions associated with yield traits (Figure 4 and Table 4). BnWRKY75-A3 and BnPHT3-A3 designed from the phosphate-starvation-response genes *AtWRKY75* and *AtPHT3*; respectively, were associated with the constitutive QTL region regulating FBH in LP and OP conditions, and RH and SW in OP condition and located between 12.2–18.6 cM on A3 (Table 4). They are also associated with primary root length (PRL) in LP and OP conditions, and total root length (TRL) in OP condition (data not shown). The association between functional genes and QTLs may accelerate identification of candidate genes regulating target traits.

Comparative genome analysis is commonly performed between *Brassica* species and *Arabidopsis* genome to align conserved genomic blocks and associate genes with QTLs identified in *Brassica* species [24,39–41]. For example, 40 synteny blocks and 84 islands were aligned between *A. thaliana* [24,39–41] and *B. rapa* and *B. oleracea* genomes (http://www.ocri-genomics.org/) along with the single nucleotide polymorphism linkage maps of *B. napus* [42,43] would greatly facilitate the genomic alignment of *Brassica* species and allow candidate gene prediction in *B. napus*. In this study, a comparative genetic mapping was performed for the first time between the TN linkage map and the *B. rapa* and *B. oleracea* genomes. Most chromosomal regions in TN linkage groups can be aligned linearly with the *B. rapa* and *B. oleracea* genomes (Figure 4), which provided useful sequence information to develop molecular markers for QTL fine mapping in the future. Of the 19 P homeostasis genes in *Arabidopsis* located in the QTL intervals determined in this study, twelve were refracted to the corresponding segments of *B. rapa* or *B. oleracea* genomes (Figure 4 and Table 4). Among these genes, *GPT1*, *MGD2* and *SIZ1* were associated with two major loci for SY-LP, which co-located with yield-related traits between 77.1–95.0 cM on A2. *GPT1* functioning as an importer of glucose-6-phosphate in *Arabidopsis* [44] was associated with PN detected under LP conditions by Ding et al. [19]. *MGD2* encoded a type-B monogalactosyldiacylglycerol synthase involved in phospholipid metabolism and was induced under phosphate (P) deprivation [45]. *SIZ1* encoded a plant small ubiquitin-like modifier (SUMO) E3 ligase and was a focal controller of P starvation dependent responses [46]. These three genes would be the most likely candidates for target QTLs that confer P deficiency tolerance and near-isogenic lines were developed to confirm the function of these genes. With deep sequencing already performed on the parents of TN DH population, sequence of orthologous genes in *B. rapa* and *B. oleracea* will provide useful information in the assembly and dissection of alleles between the parents, Tapidor and Ningyou 7.

**Materials and Methods**

**Plant materials and Field trials**

*A. napus* DH population with 188 lines was used in this study. The DH population was derived from a cross between Tapidor and Ningyou 7 by microspore culture [47]. Ningyou 7 was characterized as a P-efficient cultivar with better growth and higher P acquisition than Tapidor under low P (LP) and optimal P (OP) conditions in pot culture [48].

Three field trials were conducted at an experimental agricultural research station in Qichun, Hubei Province in China (115°45′N latitude, 30°19′E longitude) which has sandy paddy soil. The first trial coded Tri.1 was from Sept. 2008 to May 2009, the second trial coded Tri.2 was from Sept. 2009 to May 2010 and the third trial coded Tri.3 was from Sept. 2010 to May 2011. Soil properties were as follows: pH (1:1 H₂O) 4.8, organic matter 34.9 g kg⁻¹, total nitrogen 0.22 g kg⁻¹, available nitrogen 74 mg kg⁻¹, Olsen-P 3.32 mg kg⁻¹, available potassium 42 mg kg⁻¹ and HWSB (hot water soluble boron) 0.09 mg kg⁻¹.

The basal fertilizers included 60% N (urea, N 46%), 100% P₂O₅ (sodium metaphosphate, P₂O₅ 12%), 100% K₂O (potassium chloride, K₂O 60%), 100% ZnSO₄·7H₂O and 100% Borax (Na₂B₄O₇·10H₂O) according to the following nutrient rates: N 120 kg ha⁻¹, P₂O₅ 9 (LP) or 90 (OP) kg ha⁻¹, K₂O 150 kg ha⁻¹, ZnSO₄·7H₂O 45 kg ha⁻¹ and Borax (Na₂B₄O₇·10H₂O) 15 kg ha⁻¹. Rest of the urea was applied before the green bud stage. 90 and 9 kg ha⁻¹ of P₂O₅ (sodium metaphosphate, P₂O₅ 12%) were applied to create OP and LP conditions before transplantation. Three replications for 188 TN DH lines and their parents were planted in a randomized plot design with each plot comprising 18 plants, separated by a distance of 0.20 m between plants and 0.28 m between rows.

Seeds were sown in a nursery bed in the field in middle September and seedlings were transplanted 30 d after sowing. Plants were harvested in the following middle May. Standard agricultural practices were followed for field management.

**Measurement of phenotypic traits**

In each plot, six individuals from the middle row were used to determine height to the first primary branch (FBH) measured from ground level to the base of the lowest primary branch, plant height (PH) measured from ground level to the tip of the main inflorescence, number of primary branches (BN) measured as the number of primary branches arising from main shoot and seed number per pod (SN) measured as the average number of well-filled seeds from 100 well-developed pods sampled from the primary branch in the middle of each plant studied. All representative individuals from each plot were harvested by hand at maturity stage to investigate seed yield per plant (SY) and seed
weight of 1,000 seeds (SW). Pod number per plant (PN) was calculated using the following formula: \( PN = SY \times 10000/ (SW \times SN) \). Relative first primary branch height (RBH) was defined as the ratio of FBH to PH in each sampled plant. Tested traits were denoted with 'trait name and treatment name' such as PH-LP which indicates plant height under low P conditions.

**Statistical analysis and QTL detection**

Data analysis was conducted using SAS 8.1 (SAS Institute, Cary, NC, USA). Fisher's least significant difference (LSD) was used to test the significance of means at 0.05 levels. Correlation analysis was conducted to determine the relationship between the tested traits.

QTLs were detected by composite interval mapping (CIM) using WinQTL cartographer 2.5 software (http://statgen.ncsu.edu/qtlcart/WQTLCart.htm; [52]). For each trait, QTL threshold (\( P \leq 0.05 \)) was estimated from 1,000 permutations [53]. Each QTL was denominated as ‘q’ (abbreviation of QTL) + trait name + serial number + chromosome name + serial letter. For example, gBN-LP1-A1a and gBN-LP1-A1b denote two QTLs for number of primary branches (BN) detected on chromosome A1 under LP condition in Tri. 1.

**Supporting Information**

---

**Table S1** Seed yield and yield-related traits in 188 lines of a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ninyou 7 and a P-inefficient cultivar, Tapidor under low (LP) and optimal phosphorus (OP) conditions in three field trials. (XLSX)

**Table S2** Quantitative trait loci (QTLs) associated with seed yield and yield-related traits under low (LP) and optimal phosphorus (OP) conditions and distribution of gene-based markers (GBM) in the QTL intervals. (DOC)

**Author Contributions**

Obtained permission for use of TN DH population: JM. Conceived and designed the experiments: TS LS FX JM. Performed the experiments: TS ZZ. Analyzed the data: TS RL. Contributed reagents/materials/analysis tools: TS RL YI GD. Wrote the paper: TS LS GD.

---

**References**

1. Holford ICR (1997) Soil phosphorus: its measurement, and its uptake by plants. Ann J Soil Res 35: 237–239
2. von Eckardt HR, Mutert E (1995) Global extent, development and economic. Plant Soil 171: 1–15
3. Vance CP, Ude-Stone C, Allan DL (2003) Phosphorus acquisition and use: critical adaptations by plants for securing a renewable resource. New Phyto 157: 423–447
4. Fageria NK, Baligar VC, Li YC (2008) The role of nutrient efficient plants in improving crop yields in the twenty first century. J Plant Nutr 31: 1121–1157
5. Ni J, Wu P, Sendahra D, Huang N (1996) Mapping QTLs for phosphorus deficiency tolerance in rice (Oryza Sativa L.). Theor Appl Genet 97: 1361–1369
6. Wissuwa M, Yano M, Ae N (1998) Mapping QTLs for phosphorus deficiency tolerance in rice (Oryza Sativa L.). Theor Appl Genet 97: 777–783
7. Ding G, Zhao Z, Liao Y, Hu Y, Shi L, et al. (2010) Quantitative trait loci for phosphorus efficiency in Brassica napus. J Plant Nutr 32:1148–1163
8. Duan HY, Shi L, Ye XS, Wang YH, Xu FS (2009) Identification of phosphorous efficient germplasm in oilseed rape. J Plant Nutr 32:1148–1163
9. Ding G, Zhao Z, Liao Y, Hu Y, Shi L, et al. (2012) Quantitative trait loci for seed yield and yield-related traits, and their responses to reduced phosphorus supply in brassica napus. Ann Bot 109: 747–759
10. Yang M, Ding GD, Shi L, Feng J, Xu FS, et al. (2010) Quantitative trait loci for root morphology in response to low phosphorus stress in brassica napus. Theor Appl Genet 121: 181–193
11. Luo Y, Wang Y, Hu Y, Liao Y, Shi L, et al. (2011) Flowering time quantitative trait loci analysis of oilseed brassica in multiple environments and genome-wide alignment with Arabidopsis. Genetics 177: 2433–2444

---

**Table S1** Seed yield and yield-related traits in 188 lines of a double haploid mapping population (TN DH) derived from a cross between a P-efficient cultivar, Ninyou 7 and a P-inefficient cultivar, Tapidor under low (LP) and optimal phosphorus (OP) conditions in three field trials. (XLSX)

**Table S2** Quantitative trait loci (QTLs) associated with seed yield and yield-related traits under low (LP) and optimal phosphorus (OP) conditions and distribution of gene-based markers (GBM) in the QTL intervals. (DOC)

**Author Contributions**

Obtained permission for use of TN DH population: JM. Conceived and designed the experiments: TS LS FX JM. Performed the experiments: TS ZZ. Analyzed the data: TS RL. Contributed reagents/materials/analysis tools: TS RL YI GD. Wrote the paper: TS LS GD.

---

**References**

1. Holford ICR (1997) Soil phosphorus: its measurement, and its uptake by plants. Ann J Soil Res 35: 237–239 2. von Eckardt HR, Mutert E (1995) Global extent, development and economic. Plant Soil 171: 1–15 3. Vance CP, Ude-Stone C, Allan DL (2003) Phosphorus acquisition and use: critical adaptations by plants for securing a renewable resource. New Phyto 157: 423–447 4. Fageria NK, Baligar VC, Li YC (2008) The role of nutrient efficient plants in improving crop yields in the twenty first century. J Plant Nutr 31: 1121–1157 5. Ni J, Wu P, Sendahra D, Huang N (1996) Mapping QTLs for phosphorus deficiency tolerance in rice (Oryza Sativa L.). Theor Appl Genet 97: 1361–1369 6. Wissuwa M, Yano M, Ae N (1998) Mapping QTLs for phosphorus deficiency tolerance in rice (Oryza Sativa L.). Theor Appl Genet 97: 777–783 7. Ding G, Zhao Z, Liao Y, Hu Y, Shi L, et al. (2010) Quantitative trait loci for phosphorus efficiency in Brassica napus. J Plant Nutr 32:1148–1163 8. Duan HY, Shi L, Ye XS, Wang YH, Xu FS (2009) Identification of phosphorous efficient germplasm in oilseed rape. J Plant Nutr 32:1148–1163 9. Ding G, Zhao Z, Liao Y, Hu Y, Shi L, et al. (2012) Quantitative trait loci for seed yield and yield-related traits, and their responses to reduced phosphorus supply in brassica napus. Ann Bot 109: 747–759 10. Yang M, Ding GD, Shi L, Feng J, Xu FS, et al. (2010) Quantitative trait loci for root morphology in response to low phosphorus stress in brassica napus. Theor Appl Genet 121: 181–193 11. Luo Y, Wang Y, Hu Y, Liao Y, Shi L, et al. (2011) Flowering time quantitative trait loci analysis of oilseed brassica in multiple environments and genome-wide alignment with Arabidopsis. Genetics 177: 2433–2444
34. Chen W, Zhang Y, Liu X, Chen B, Tu J, et al. (2007) Detection of QTL for six
32. Quijada PA, Udall JA, Lambert B, Osborn TC (2006) Quantitative trait analysis
30. Hammond JP, Mayes S, Bowen HC, Graham NS, Hayden RM, et al. (2009) Shoot
31. Basunanda P, Radoev M, Ecke W, Friedt W, Becker HC, et al. (2010)
28. Zhao JJ, Jamar DCL, Lou P, Wang YH, Wu J, et al. (2008) Quantitative trait
35. Li YY, Shen JX, Wang TH, Chen QF, Zhang XG, et al. (2007) QTL analysis of
36. Lowe A, Moule C, Trick M, Edwards K (2004) Efficient largescale development
38. Zhang WK, Wang YJ, Luo GZ, Zhang JS, He CY, et al. (2004) QTL mapping of
ten agronomic traits on the soybean (Glycine max L. Mor.) genetic map and
their association with EST markers. Theor Appl Genet 108: 1131–1139.
39. Parkin IA, Golden SM, Sharpe AG, Lukens L, Trick M, et al. (2005) Segmental
structure of the Brassica napus genome based on comparative analysis with
Arabidopsis thaliana. Genetics 171: 765–81.
40. Schranz ME, Lysak MA, Mitchell-Olds T (2006) The ABC's of comparative
genomics in the Brassicaceae: building blocks of crucifer genomes. Trends Plant
Sci 11: 535–542.
41. Panjabi P, Jagannath A, Bish N, Padmaja KL, Sharma S, et al. (2008) Comparative
mapping of Brassica juncea and Arabidopsis thaliana using Intron Polymorphism
IJP: markers: homoeologous relationships, diversification and evolution of the A, B
and C Brassica genomes. BMC Genomics 9: 113.
42. Trick M, Long Y, Meng J, Bancroft I (2009) SNP discovery in the polyploid
Brassica napus using Solexa transcriptome sequencing. Plant Biotechnol J 7: 334–
346.
43. Bancroft I, Morgan C, Fraser F, Higgins J, Wells R, et al. (2011) Dissecting the
genome of the polyploid oilseed rape by transcriptome sequencing. Nat
Biotechnology 29: 762–766.
44. Niewiadomski P, Knapp S, Geimer S, Fischer K, Schulz B, et al. (2005) The
Arabidopsis plastidic glucose 6-phosphate/phosphate translocator GPT1 is
essential for pollen maturation and embryo sac development. Plant Cell 17:
760–775.
45. Awai K, Maréchal E, Block MA, Bruin D, Masuda T, et al. (2001) Two types of
MGDG synthase genes, found widely in both 16:3 and 18:3 plants, differentially
mediate galactolipid syntheses in photosynthetic and nonphotosynthetic tissues
in Arabidopsis thaliana. Proc Natl Acad Sci USA 98: 10960–10965.
46. Miura K, Rus A, Shakhhabi A, Yokes S, Karthikeyan AS, et al. (2005) The
Arabidopsis SUMO E3 ligase SIZ1 controls phosphate deficiency responses. Proc
Natl Acad Sci USA 102: 7760–7765.
47. Qi D, Morgan C, Shi J, Long Y, Liu J, et al. (2006) A comparative linkage map
of oilseed rape and its use for QTL analysis of seed oil and erucic acid content.
Theor Appl Genet 114: 67–80.
48. Shi TX, Wang SS, Shi L, Meng JL, Xu FS (2010) Effects of different nitrogen
and phosphorus levels on seed yield and quality parameters of double high and
double low Brassica napus. Plant Nutrition and Fertilizer Science 16: 959–964. (in
Chinese with English abstract)
49. Ding GD, Liao Y, Yang M, Zhao ZK, Shi L, et al. (2011) Development of gene-
based markers from Arabidopsis thaliana functional genes involved in phosphorus
homeostasis and mapping in Brassica napus. Euphytica 181: 305–322.
50. Long Y, Xia W, Li R, Wang J, Shao M, et al. (2011) Epigenetic QTL mapping
in Brassica napus. Genetics 189: 1093–102.
51. Van Ooijen JW (2006) JoinMap® Software for the calculation of genetic
linkage maps in experimental populations. Kyazma B. V., Wageningen,
Netherlands.
52. Wang SC, Eastern J, Zeng ZB (2006) Windows QTL Cartographer 2.5.
Department of Statistics.
53. Churchill GA, Doerge RW (1994) Empirical threshold values for quantitative
trait mapping. Genetics 138: 903–971.