QTL Alignment for Seed Yield and Yield Related Traits in *Brassica napus*

Nadia Raboanatahiry1,2, Hongbo Chao1,2, Hou Dalin1,2, Shi Pu1, Wei Yan1, Longjiang Yu1, Baoshan Wang3 and Maoteng Li1,2*

1 Department of Biotechnology, College of Life Science and Technology, Huazhong University of Science and Technology, Wuhan, China, 2 Hubei Key Laboratory of Economic Forest Germplasm Improvement and Resources Comprehensive Utilization, Hubei Collaborative Innovation Center for the Characteristic Resources Exploitation of Dabie Mountains, Huanggang Normal University, Huanggang, China, 3 College of Life Sciences, Shandong Normal University, Jinan, China

Worldwide consumption of oil is increasing with the growing population in need for edible oil and the expansion of industry using biofuels. Then, demand for high yielding varieties of oil crops is always increasing. *Brassica napus* (rapeseed) is one of the most important oil crop in the world, therefore, increasing rapeseed yield through breeding is inevitable in order to cater for the high demand of vegetable oil and high-quality protein for live stocks. Quantitative trait loci (QTL) analysis is a powerful tool to identify important loci and which is also valuable for molecular marker assisted breeding. Seed-yield (SY) is a complex trait that is controlled by multiple loci and is affected directly by seed weight, seeds per silique and silique number. Some yield-related traits, such as plant height, biomass yield, flowering time, and so on, also affect the SY indirectly. This study reports the assembly of QTLs identified for seed-yield and yield-related traits in rapeseed, in one unique map. A total of 972 QTLs for seed-yield and yield-related were aligned into the physical map of *B. napus Darmor-bzh* and 92 regions where 198 QTLs overlapped, could be discovered on 16 chromosomes. Also, 147 potential candidate genes were discovered in 65 regions where 131 QTLs overlapped, and might affect nine different traits. At the end, interaction network of candidate genes was studied, and showed nine genes that could highly interact with the other genes, and might have more influence on them. The present results would be helpful to develop molecular markers for yield associated traits and could be used for breeding improvement in *B. napus*.

**Keywords:** *Brassica napus*, quantitative trait loci, alignment map, seed-yield, yield-related traits, candidate genes

**INTRODUCTION**

Rapeseed (*Brassica napus*) is an important crop utilized globally for vegetable oil production, animal feed and biofuel. *B. napus* (AACC, 2n = 38) is a polyploid species, which was derived from the hybridization between turnip rape (*B. rapa*, AA, 2n = 20) and cabbage (*B. oleracea*, CC, 2n = 18) (Nagaharu, 1935; Al-Shehbaz et al., 2006; Bailey et al., 2006; Chalhoub et al., 2014).

**Abbreviations:** BH, branch height; BN, branch number; BY, biomass yield; DTS, development time of seeds; FBH, first branch height; FBN, first effective branch number; FT, flowering time/days to flowering; LML, length of main inflorescence; MT, maturity time; NPB, number of primary branches; NRV, Napus root vigor; PH, plant height; PN, pod number/silique per plant; PNM, pod number of main inflorescence; PYP, pod yield/100pods; SB, silique breadth; SD, seed density; SID, silique density; SL, silique length; SN, seed number per pod/Seeds per silique; SPUA, Silique per unit area; ST, silique thickness; SV, silique volume; SW, seed weight/thousand seed weight; SY, seed yield; TW, test weight.
In terms of evolutionary age of plants in *Brassica* genera, *B. napus* registers the shortest age and history of domestication (Jesske et al., 2013; Chalhoub et al., 2014). Spring type *B. napus* are typically cultivated in Canada, Asia, Eastern Europe, and Australia, whereas winter type *B. napus* are usually grown in Western Europe; those winter type *B. napus* generally gives higher SY than spring type (Nesi et al., 2008). Currently, the most important task for plant breeders is to develop a high yielding variety in order to attain sustainable production in future. Rapeseed yield (SY per plant) is determined by the number of siliques per plant, the number of seeds per silique and the SW. The total number of seeds are determined by the number of seeds per silique and the number of siliques per plant (Chen et al., 2007).

Genetic approach has become the most powerful tool for examining the genetic traits in many crops including *B. napus*. Application of molecular marker techniques for quantitative trait loci (QTL) is a trending strategy (Paran and Zamir, 2003). The major objective of QTL studies is to find genes that could be used in breeding via marker assisted selection (MAS). QTL mapping is also a potent method which is widely used to decipher regulatory loci and genetic mechanism of traits. Identification of QTLs is necessitated by molecular markers coupled with mapping populations. The magnitude and presence of QTLs varies between experiments, so the sensitivity of the QTL can easily be underscored. Also, the number of QTLs identified in different populations varies with the genetic background of parental lines, the type and size of the population, and the number of environments that were used in different studies (Zhu and Zhao, 2007).

Seed yield is a very complex trait. In an open-pollinated and hybrid type crop, such as rapeseed, high SY is partly attributed to epistatic interaction between the loci and possible multiple alleles at the loci which control SY trait (Engqvist and Becker, 1991; Radoev et al., 2008). Moreover, the effects of the QTL alleles increasing SY on a particular genetic background could produce a different phenotypic effect on another genetic background, and the alleles may be responsible for very poor yield in some cases (Kramer et al., 2009). Therefore, the fact that QTL alleles may exhibit an effect in a certain combination of alleles that involves one or several loci, and another different effect in another set of allele combinations might suffice the need for researchers to discover and determine the effects of different genes in varying environment. This will enable breeding programs to be effective, since the different individual alleles from each locus, as well as allelic combinations from different loci, will be utilized in identification of the most favorable gene combinations that will produce maximum benefit. Nevertheless, this task is not simple considering that more than one QTL regions and multiple variants of alleles at every locus have the ability to control quantitative traits, such as SY and yield associated traits. Also, as mentioned earlier, it is clear that QTL vary with genetic × environment background (Zhu and Zhao, 2007), therefore, combination of multiple QTLs from different genetic × environmental background into one map would allow an easy comparison between these QTLs. Establishing a consensus map allows comparison between these multiple QTLs and valuable findings might be obtained. For example, Wang et al. (2013) constructed a consensus map of 2395.2 cm genome length with 1335 markers to compare QTL for oil content in seven different populations of *B. napus*. They affirmed that combining multiple QTLs from different population into one consensus map could be a potential tool not only to study the genetic diversity of loci with complex traits, but also to develop molecular breeding and map-based cloning of genes (Wang et al., 2013). Additionally, Zhou et al. (2014) performed in silico integration of QTLs for SY and yield related trait from 15 *B. napus* mapping populations by using *B. rapa* and *B. oleracea* genome sequences, and 736 SY and yield related QTLs on 283 loci of *B. napus* A and C genome were mapped, they were distributed across 19 chromosomes and most of them were located on A03. Also, Zhao et al. (2016) built a consensus map to compare QTLs for SY and yield related traits from five different populations of *B. napus*, 34 SY-QTLs could be projected onto the “KenC-8 × N53-2” map, half of them were placed on chromosome A02, A05, A06, C02, and C06. Nevertheless, difference in markers used with these QTLs makes difficult this combination into one consensus map and their comparison (Raman et al., 2013). Therefore, a QTL alignment map would help to overcome this difficulty. In fact, a QTL alignment map combines diverse QTLs from different genetic and environmental background. Therefore, QTLs from different populations which were developed in the same environments and which could overlap in one region might be defined as fixed QTLs for that environment, and QTLs that could not overlap with other QTLs might be specified QTLs. In this situation, fixed QTLs might be more stable and more important in further fine-mapping for gene cloning; firstly, the variation underlying the fixed QTL was more common in different varieties. If this variation is rare, it is impossible to detect it in many populations. Secondly, it can be detected in many populations and detected in different planting environments where each population can adapt to, which indicated that it is an ubiquitous variation (derived from early evolution) and has strong environment stability. Thus, conservative and variant regions could be uncovered with this QTL alignment map. Such technique was earlier used to compare multiple QTLs for seed oil content and fatty acid profile using *B. napus Darmor-bzh* as reference genome (Liu et al., 2016; Chao et al., 2017; Raboanatahiry et al., 2017). To date, establishment of QTL alignment map for SY and yield related traits has not been performed yet.

Identification of candidate genes implies the detection of important genes for agricultural and economic quantitative traits. Candidate genes are present within QTL region and they are responsible for phenotype variation. The effect of these genes on variation of phenotype could be elucidated through investigation on the gene arrangement and interaction of loci affecting this variation (Zhu and Zhao, 2007). Moreover, study on QTLs and related candidate genes could help in knowledge of their control on the phenotype (Remington and Purugganan, 2003; Zhu and Zhao, 2007). In *B. napus*, several studies on candidate genes for yield related traits have been performed before: for instance, Zhao et al. (2016) found four, two and one candidate genes for SY, SW and PH, respectively in a study performed with a
double-haploid (DH) population of 348 lines derived from a cross between KenC-8 × N53-2. Also, in a study executed with 333 diverse rapeseed accessions (20 winter type, 308 semi-winter type and 5 unknown accessions), candidate genes for PH (31 genes), branch initiation height (15 genes), and BN (17 genes) were found (Zheng et al., 2017). More, candidate genes for SN (six genes), PN per plant (seven genes), branch PN (seven genes), branch yield (three genes) and thousand seed weight (one gene) were discovered from a study made with 520 rapeseed accessions (Lu et al., 2017). In the model plant Arabidopsis thaliana, 425 genes were found to be related to BN, FT, MT, PH, PN, seed number, seed weight, and SY (Shi et al., 2009). In B. napus, homologous of these 425 genes of A. thaliana have not been searched before.

Therefore, the present study aimed to compare yield and yield-related QTLs in 22 populations which were developed in four different countries (Table 1) via arrangement into the physical map of B. napus “Darmor-bzh”, the candidate genes for SY and yield-related traits in B. napus were also searched inside the overlapping QTLs regions and a genetic network was constructed to understand the interaction of candidate genes.

MATERIALS AND METHODS

QTL Alignment Map for Seed-Yield and Yield Related Traits in B. napus
Quantitative trait loci for SY and yield related traits obtained from 22 populations were aligned to the physical map of B. napus winter type Darmor-bzh. To perform this alignment, QTL flanking markers were positioned on the map by using of E-PCR (Schuler, 1997; Rotmistrovsky et al., 2004). One marker could be detected on the genome owing to corresponding primer. So, primer sequences were searched on Darmor-bzh to spot the markers. Because some flanking marker sequences could not be found in some literatures or they are found but could not be placed on the map (misplaced to another chromosome or location not found on Darmor-bzh), some QTLs could not be aligned on the map. Also, it happened that only one flanking marker could be placed, and some literatures gave only one flanking marker; therefore, we took a uniform area of 1 cM to delimit all the QTLs, so that QTLs region could be located on the map even with one marker. Note that apart from related literatures, marker sequences could also be found on some database, such as Brassica.info 1, NCBI 2, Brassica IGF Project 3, Gramene Markers Database 4. The map was build using Circos software (Krzywinski et al., 2009).

Candidate Genes Identification
As mentioned above, Shi et al. (2009) found 425 genes in A. thaliana which were related to BN, FT, MT, PH, PN, seed number, seed weight, and SY in their study; so in the current study, candidate genes were identified based on this study performed by Shi et al. (2009) First, genes related to yield and yield related traits in B. napus were identified based on homology to these 425 yield related genes in A. thaliana, by using Browse Data tool in http://www.genoscope.cns.fr/brassicanapus/. Then, we aligned these B. napus genes with the previous QTL alignment map of the current study. Consequently, candidate genes were genes located inside a QTL region.

Rapeseed genome showing beneficial stable loci with related candidate genes was then proposed. For that, overlapping QTLs identified previously, with candidate genes found inside the region of these overlapping QTLs were highlighted. A simple construction was made using an excel based tool called E-maps (Ulitsky et al., 2008).

Construction of Candidate Genes Interaction Network
A network was constructed to understand the interaction between candidate genes. Interaction of proteins was predicted using STRING 5 (Szklarczyk et al., 2015), clustered using Mapman software (Thimm et al., 2004) and then visualized with Cytoscape 3.6.0 (Shannon et al., 2003). Orthologous genes in A. thaliana were used to perform the analysis, because B. napus is still not available on STRING database.

RESULTS

Overlapping QTLs for Seed Yield and Yield-Related Traits Were Found Aligned in Physical Map of Darmor-bzh
From the 22 populations, 972 QTLs for 26 yield traits could be aligned on the physical map of B. napus Darmor-bzh (Supplementary Table 1): 182 (FT), 168 (SW), 124 (PH), 93 (SY), 73 (MT), 65 (SN), 53 (NPB), 46 (PN), 28 (BY), 23 (PY), 21 (SL), 15 (TW), 13 (SPUA), 12 (FBBN), 10 (BH and LMI, each), 7 (SB, SID, and ST, each), 4 (PNMI and SV, each), 3 (SD) and 1 (BN, DTS, FBH and NRV, each). QTLs from different populations on the map displayed different positions. Several QTLs for the same traits were found across the map, for example, in KenC-8 × N53-2 population, QTLs for SW were both present on A02 (21.52 and 22.11 Mb), on C07 (2.53 Mb) and on C08 (36.35 Mb). They could be located in separate regions or they could overlap with other populations’ QTLs. Thus, 92 regions that contained 198 overlapping QTLs were observed, they were located on 16 chromosomes and involved 11 traits and 17 populations which were developed in China, America and Germany (Figure 1 and Supplementary Table 2).

The present results revealed that QTLs from the Canadian population Regent × Lagoda did not overlap with any of the QTLs from other populations. In fact, while observing the genetic background of parental lines in these populations (known relatives are presented on Table 1), it is obvious that Express617 × R53 and Express617 × V8 shared a common

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1http://www.brassicace.info/resource/maps/lg-assignments.php
2http://www.ncbi.nlm.nih.gov/
3http://brassicace.info/IGF/
4http://archive.gramene.org/markers/
5http://string-db.org/
TABLE 1 | List of populations used for the alignment map.

| Populations | Known relatives | Lines | Environments | Traits | Reference |
|-------------|-----------------|-------|--------------|--------|-----------|
| 04-1139 × 05-1054 | 05-1054 | 221 F2 | China | PN, SN | Wang and Guan, 2010 |
| 448 lines | | 448 IL | China | FT | Wang N. et al., 2016 |
| 8008 × 4942C-5 | 8008 (Eyou Long Pod) | 181 DH | China | SN | Qi et al., 2014 |
| Express × SWU07 | | 261 DH/ | China | SL, SW | Fu et al., 2015 |
| | | 233 RC-F2 | | | |
| Express617 × R53 | Express617 (Express) | 250 DH | Germany | PH, SW, SY, SPUA | Basunanda et al., 2010 |
| Express617 × V8 | Express617 (Express) | 250 DH | Germany | SY, SN, SPUA, SW | Raschev et al., 2008 |
| | | | | | Basunanda et al., 2010 |
| HZ96 × Y106 | | 140 DH | China | SL, SW | Zhang et al., 2011 |
| J7046 × J7005 | | 190 F2 | China | SW | Fan et al., 2010 |
| KenC-8 × N53-2 | N53-2 (Midas, SE8) | 348 DH | China | SD, SL, SN, ST, SV | Wang X. et al., 2016 |
| | N53-2 (Midas, SE8) | 348 DH | China | BH, BY, FE, BN, LMI, PH, PNMI, SW, SY | Zhao et al., 2016 |
| MF216 × P1804 | MF216 (Stellar, Major) | 150 DH | America | FT, PH, SY, TW | Quijada et al., 2006 |
| MF216 × P1804 | | 160 DH | America | FT, PH, SY, TW | Quijada et al., 2006 |
| Quantum × NO.2127-17 | NO. 2127-17, (resynthesized B. napus line NO. 7076) | 258 DH | China | NPB, PH, SId, SL | Chen et al., 2007 |
| Regent × Lagoda | | 93 F2 | Canada | FT, NRV | ArifUzZaman et al., 2016 |
| RV128 × P1804 | RV128 (BC252, Westar, Samourai, Bienvenu, Major) | 150 DH | America | FT, PH, SW, SY, TW | Quijada et al., 2006 |
| RV289 × P1804 | RV289 (Hua-obl2) | 148 DH | America | PH, SW, SY, TW, FT | Udall et al., 2006 |
| RV289 × P1804 | | 160 DH | America | PH, FT, PH, SW, SY, TW | Quijada et al., 2006 |
| S1 × S2 | S1 (breeding line 92-B10) | 186 RIL | China | SL, SW | Yang et al., 2012 |
| Sollux × Gaoyou | | 282 DH | China, Germany | FT, PH, MT, PH | Zhao et al., 2005 |
| Tapidor × Ningyou7 | | 182 DH | China | BN, BY, DTS, FBH, FT, MT, PH, PN, SN, SW, SY | Luo et al., 2017 |
| | | | | | Shi et al., 2009 |
| | | 202 DH | China | NPB, BY, FT, MT, PH, PN, SN, SW, SY | Shi et al., 2011 |
| TO1141 × P1804 | TO1141 (B. rapa cv. Reward and B. oleracea TO1000) | 160 DH | America | FT, SL, PH, SW, SY, TW | Udall et al., 2006 |
| Westar(Mamo) × Ceres | | 128 IBL | America | PH, SW, FT | Butruille et al., 1999 |
| Zhongshuang11 × No.73290 | | 182 F2 | China | SN | Shi et al., 2015 |
| | | 184 RIL | China | SN | Yang et al., 2016 |
| | | 184 F2/ | China | PN | Ye et al., 2017 |

...parental line in Express617, as well as MF216 × P1804, RV128 × P1804, RV289 × P1804, and TO1141 × P1804 which had P1804 in common. Thus, overlapping QTLs from these populations were not surprising (e.g., overlapping SPUA QTLs DH-slpa from Express617 × V8 and Sil/dm2N16-2 from Express617 × R53 on C6, overlapping SW QTLs sw7.3 from RV289 × P1804 and sw7.1 from RV128 × P1804 on A7). Further analysis revealed that regions of overlapping QTLs were mainly located on chromosome A03 (24/92), then on C06 (17/92), A10 and C01 (8/92, each), A7 and C06 (7/92, each), A01 (6/92), A02 (5/92), A05 (3/92), A06, A09, C03, C05, C07 and C09 (2/92, each), and C08 (1/92). The most frequently detected overlapping QTLs for each trait are presented on Figure 2A, among 92 regions of overlapping QTLs, 30 regions were for FT, 20 regions were for SW, 13 regions were for PH, 11 regions were for SY, 4 regions were for SPUA, 3 regions each were for NPB, PN, SN and TW, and 1 region each for BY and SL.

In addition, 74 of 92 regions of overlapping QTLs involved two populations (e.g., RV289 × P1804 and Tapidor × Ningyou7 for FT trait on A02), eight regions of overlapping QTLs involved...
five populations (e.g., Tapidor × Ningyou7, MFDH × P124, RV128 × P1804, RVDH × P124 and 448 lines for FT trait on A03), six regions of overlapping QTLs involved three populations (e.g., Express617 × V8, Express617 × R53 and S1 × S2 for SW trait on A01), and four regions of overlapping QTLs involved four regions (e.g., Tapidor × Ningyou7, RV128 × P1804, RVDH × P124 and 448 lines on A03) (Figure 2B). Moreover, 53 regions of overlapping QTLs were of environmental fixed QTLs: 28 among them were from the mapping population of Chinese environments, 13 and 12 among them were from German and American environments, respectively. The remaining 39 regions of overlapping QTLs were of mixed populations: QTLs from China and America overlapped the most (27/39 overlapping QTLs), and eight and two implied QTLs from China and Germany, and America and Germany, respectively, also since “Sollux × Gaoyou” was developed both in China and Germany, two overlapping QTLs implied the populations developed from America, China and Germany (Figure 2C). Finally, 13 regions of overlapping QTLs might affect multiple traits simultaneously: three regions were found on each of A03 (FT/PH, PH/PN, and FT/PH/PN) and C02 (FT/PH/PN, PN/SW, and SPUA/SY), two regions were found on each of A10 (FT/SW and SY/TW) and C01 (both SN/SW), and one region was found on each of A05 (NPB/PH/SN), A07 (PH/SW/SY) and C06 (BY/PH/SW) (Figure 2D and Supplementary Table 3). In overall, 57 regions of overlapping QTLs were located on A genome and 35 regions were on C genome. These results indicated that despite the difference in genetic × environment backgrounds, stable loci could be conserved in multiple populations of rapeseed. In addition, some loci might affect more than one beneficial trait, simultaneously.

Regions of Overlapping QTLs Contained Potential Candidate Genes

A total of 1562 genes in B. napus were homologous of 425 yield related genes in A. thaliana (Supplementary Table 4). Then, 1398 among these genes were used to uncover potential candidate genes in overlapping QTLs, the remaining 164 genes could not be placed since their chromosome location were unknown. As result, a total of 147 candidate genes were found inside regions of overlapping QTLs for nine traits: FT, NPB, PH, PN, SN, SPUA, SW, SY, and TW (Figure 2E and Supplementary Table 5). Candidate genes for single trait were respectively of 34 (FT), 30 (SW), 18 (PH), 16 (SY), 8 (SPUA), 4 (PN), and 2 (NPB), and candidates for multiple traits were respectively of 8 (FT/SY), 6 (FT/PH/PN), 5 (SY/TW), 3 (NPB/PH, NPB/PH/SN, and SN/SW, each), 2 (PH/SY, SPUA/SY, and FT/PH, each), and 1 (FT/SW). Thus, 53 (FT), 34 (PH and SW, each), 33 (SY), 10 (PN and SPUA, each), 9 (SN), 8 (NPB), and 5 (TW) were discovered in this analysis. Further observation revealed that most of these candidate genes were on chromosome A03 (33/147), the other chromosomes contained 17 (A10), 15 (C02), 12 (A01, A02 and C03, each), 10 (A07), 9 (C08), 6 (A05 and C07, each), 5 (C01), 4 (A09), 3 (C06), 2 (A06) and 1 (C09) candidate genes, respectively. In this study, nine candidate genes were found inside region of overlapping QTLs which involved three traits: three genes on
A05 (BnaA05g07510D, BnaA05g07580D, and BnaA05g07740D) that might affect NPB, PH and SN traits, six other genes on C2 (BnaC02g01710D, BnaC02g45080D, BnaC02g45090D, BnaC02g02210D, BnaC02g45250D and BnaC02g02440D) that possibly affect FT, PH and PN traits. Besides, candidate genes detected in *B. napus* often fell into QTL intervals of traits which were different from related function in *A. thaliana* orthologous genes. For example, *B. napus* BnaC09g36060D is homologous of *A. thaliana* RAX2 gene AT2G36890 which was related to BN, however, *B. napus* BnaC09g36060D fell into overlapping QTLs of FT. These findings showed genes that might be responsible for one or multiple traits. Besides, homologous genes might affect dissimilar traits.

Assumption from above-cited findings, genome of *B. napus* that might offer maximum benefit might correspond to a genome enriched with regions where QTLs of multiple traits overlapped, as seen on chromosomes A03, A05, A07, A10, C01, C02, and C06 where QTLs for two or three traits could overlap in the same regions, as mentioned previously (Figure 3 and Supplementary Table 3).

**Candidate Genes Interaction Network Investigation Revealed Nine More Influential Genes of Diverse Metabolisms**

Interaction analysis between 147 candidate genes of *B. napus* were conducted using their 117 orthologous genes in *A. thaliana* (Supplementary Table 6). Gene ontology analysis indicated that these candidate genes could be divided into ten clusters, corresponding to 12 distinct functions which were clearly classified in the interaction network analysis (Figure 4 and Supplementary Table 7): cell metabolism, transport, signaling,
hormone metabolism, RNA regulation, development, protein metabolism, others (stress, nucleotide metabolism, polyamine metabolism, secondary metabolism), miscellaneous group, and an unknown group. The network contained 105 nodes and 449 edges. Obviously, nine genes that have dissimilar function might have higher influence over the other genes (degree layout, DL ≥ 20): AUX1 (transport, DL = 34), CO and FT (development, DL = 32 and 30, respectively), AGL20 and FLC (RNA regulation, DL = 26 and 24, respectively), CRY2 and PHYA (signaling, DL = 25 and 24, respectively), and BRI1 and GAI (hormone metabolism, DL = 24). In fact, functions of these nine influential candidate genes are related to flowering process, light response
and plant hormones. In B. napus, these candidate genes fell into ten regions of overlapping QTLs, and might affect six traits, including SY, SW, PN, FT, PH, and NPB. These findings indicated that yield traits might be affected at multiple set of metabolisms.

DISCUSSION

Unified Seed Yield and Yield-Related QTLs in One Map Revealed Important Stable Loci for Yield Improvement

In B. napus, SY depends on the number of PN or number of siliques per plant, the number of seeds per silique and the seed weight, these traits affect SY QTLs in a direct manner (Ozer et al., 1999; Quarrie et al., 2006; Chen et al., 2007). PN is a component in rapeseed that is targeted by breeders because it shows the highest correlation with yield, it is a sensitive quantitative trait, which shows extensive variation among cultivars and environment factors, thus it is important in genetic variation (Shi et al., 2015). SL and seed weight are both quantitative trait with high inheritance (Zhang and Zhou, 2006), and represent crucial keys for yield potentiality; thus, they are also targeted for breeding selection in rapeseed (Ali et al., 2003; Samizadeh et al., 2007). Furthermore, SY is indirectly influenced by traits, such as PH, first effective BH, first effective BN, LMI, PNMI, FT, MT, BY (Qiu et al., 2006; Shi et al., 2009; Zhou et al., 2014; Zhao et al., 2016). These traits are important to breeders because they show interaction with the SY (Zhang and Zhou, 2006); but also, they are environment sensitive traits, few stable QTLs were found in different environments in previous study (Shi et al., 2009).

In the present study, position of QTL in different populations were different, probably due to the genetic difference between parental lines, the type and size of the populations used, and the environments where they were developed (Zhu and Zhao, 2007). Overlapping QTLs emphasized regions on the genome which were responsible for trait variation and well conserved across the populations. Overlapping QTLs emphasized regions on the genome which were responsible for trait variation and well conserved across the populations. Overlapping QTLs within Express617 × R53 and Express617 × V8 populations, MF216 × P1804, RV128 × P1804, RV289 × P1804 and TO1141 × P1804 populations were not surprising, as they shared common ancestry which were Express617 and P1804, respectively. Besides, Regent was among the 448 lines studied by Wang N. et al. (2016), it is surprising
that no overlapping QTLs were found between QTLs detected in this study with the Canadian line Regent × Lagoda. Similarly, RV128 and Westar were related, and no overlapping QTLs were found between RV128 × P1804 and Westar (Marnoo) × Ceres. The remaining populations mentioned in this study had no common ancestry, in our knowledge. Otherwise, 53 regions of environment fixed QTLs could be deduced from the alignment map (e.g., 26/53 for China), despite the genotype difference of populations, similar QTL regions could be found due to the environment impact. Nevertheless, overlapping QTLs could be observed from populations of different environments (e.g., China/America), these regions might be enriched with related gene variations, but it is also possible that they share common ancestry. Besides, we could find QTLs which could not overlap with other QTLs (e.g., QTLs from Canadian populations Regent × Lagoda), they were specified QTLs which could not be on the same region as the others, probably due to the above-mentioned factors (difference in parents, environments, size of populations), but also the density of genetic map of which they were deduced. A region on A03 (from 4.25 to 6.05 Mb) had overlapping QTLs for FT, implying five populations, which were stable QTLs from populations of two different environments. Also, several regions of overlapping QTLs with multiple traits were found, these regions might be advantageous to develop for rapeseed breeding in targeting multiple traits simultaneously.

Besides, our analysis supported the importance of quantitative traits analysis in identification of SY and yield components. However, identification of the right gene combination in various loci is a challenging task, particularly when several loci with multiple alleles are associated. In this case, molecular plant breeding and genomics rather than conventional breeding will play a very significant role in increasing breeding efficiency. For example, sequencing of the Brassica genomes, candidate gene identification for controlling SY, and knowledge of the location of these specific genes in the genome and their roles will extend their importance in molecular-marker technology in plant breeding. This is because development of allele molecular markers from the gene itself can be synthesized. Findings of the current analysis are valuable to enhance breeding programs. Genes found within these QTLs intervals could be potential genes that might affect these traits and consequently could be manipulated to get the desired traits, notably producing high SY.

**Candidate Genes Revealed in the Region of Overlapping QTLs Could Be Used as Tool to Get Desired Traits**

In the current study, a total of 147 candidate genes were found in 65 regions of overlapping QTLs, and several candidate genes might affect multiple traits at the same time. As mentioned above, three independent works were published on candidate genes for yield traits in B. napus: first, Zhao et al. (2016) found candidate genes for SY (n = 4), seed weight (n = 2), and PH (n = 1), by comparative mapping between A. thaliana, B. rapa, B. oleracea and B. napus, in 348 DH lines from cross between KenC-8 and NS3-2, they were grown in Dali, Shanxi for 5 years (Winter type) and Sunan, Gansu Province for 3 years (Spring type). Second, Zheng et al. (2017) discovered 31, 15, and 17 candidate genes for PH, branch initiation height and BN traits, respectively, by using of 333 diverse rapeseed accessions (20 are winter type, 308 are semi-winter type and 5 are unknown as referred to Sun et al., 2016), which were grown during 2012–2015 in Wuhan, China. They identified candidate genes by using gene sequences corresponding to A. thaliana and located on flanking regions of up to 450 kb on either side of significant SNPs markers, and then used them to GO annotations, with auxin, GA, IAA, SL, CK, and FT taken as the most likely candidate. Finally, Lu et al. (2017) identified candidate genes for SN (n = 6), PN per plant (n = 7), branch PN (n = 7), branch yield (n = 3), and thousand seed weight (n = 1), from a study led on 520 rapeseed accessions grown in Chongqing and Yunnan, by using GWAS and RNA-seq tools. Comparing our results with these cited papers, BnaA02g02560D was a candidate for SY in both our analysis and Zhao et al. (2016) analysis. Besides, three candidate genes for different traits: BnaA02g02560D was a candidate for SY in our analysis, however, it was a candidate for PH in Zheng et al. (2017) analysis, also BnaA02g12130D and BnaA02g12260D were candidates for FT in our analysis, and they were candidates for PH and branch initiation height in Zheng et al. (2017) analysis. However, none of candidate genes in Lu et al. (2017) analysis were similar to our findings. Since one gene might influence many traits and one trait might be influenced by many genes, as established in our analysis, hypotheses from our study and from these other studies might be both exact if proven by experimental approaches.

Identification of candidate genes in earlier time needed comparative mapping using A. thaliana as a model plant to map and discover genes in related species. For instance, Long et al. (2007) found BnFLC010, candidate gene underlying qFT10-4 QTL, which was the key gene that controls differentiation of spring or winter rapeseed types based on these comparative mapping analyses. Furthermore, Ding et al. (2010) located the specific genes which controlled phosphorus concentration in the seeds, while the candidate genes controlling FT was discovered by Shi et al. (2011) using comparative mapping with the genome of A. thaliana. Advancement of these studies in order to ascertain the effects of the QTLs on different agronomic traits by use of comparative mapping technique have been undertaken by many research groups (Cai et al., 2014). Adoption of this technique is very significant in breeding for high yielding traits in B. napus cultivars. This is because it has made available abundant information that are useful in further understanding the genetic mechanisms of SY and yield related traits.

Nowadays, since the release of B. napus genome sequence, it is possible to identify candidate genes through B. napus physical map. Location of QTL flanking markers were searched on B. napus physical map via Blastn and E-PCR (Schuler, 1997; Rotmistrovsky et al., 2004), then the genes found inside the QTL regions are potential candidate genes. This technique has already been used to identify potential candidate genes in B. napus, for instance, Chao et al. (2017) used this technique to identify potential candidate genes for seed oil content trait, and found 448 genes underlying 41 oil content QTLs. Such kind of technique is also beneficial to identify candidate genes of QTLs from different linkage maps that could be could be combined.
In our analysis, QTLs from different genotype × environment background then we identified 82 regions of overlapping QTLs and 218 candidate genes in 162 QTLs regions, 76/218 candidate genes were found inside 57 regions of overlapping QTLs, which were majorly for oil content traits (Raboanatahiry et al., 2017). Candidate genes of overlapping QTLs were supposed to be more stable rather than from independent QTL, different results might be obtained if candidate genes of independent QTLs were studied. Candidate genes are responsible for QTL and cause trait variation (Tabor et al., 2002). Thus, in our analysis, these candidate genes that might affect one or multiple traits could be used as tool to acquire any chosen traits in order to get high SY. They also can be used to get the most beneficial gene combination to get maximum profit, especially those genes which were found in region of overlapping QTLs involving many traits.

**Candidate Genes Interaction Network Indicated That Yield Might Be Affected at Different Set of Metabolisms**

In our analysis, QTLs from different genotype × environment background had different location on the physical map of *Darmor-bzh*, and this affect also the candidate genes discovered during this study. Thus, the study of genetic network allows to comprehend the interaction between genes and the mechanism that works for the biological system. Besides, quantitative traits are influenced by dynamism and structure of genetic network (Frank, 2003). In this study, candidate genes interaction network was made from homologous genes in *A. thaliana*, since *B. napus* is not accessible on STRING Database. Actually, *B. rapa* (A genome) is available on STRING database but is not enough to provide more information for this study compared to *A. thaliana*. First, homologous genes in *B. rapa* would represent only a part of detected candidate genes in *B. napus* and this would distort the results. Second, annotated and cloned genes were rare in *B. rapa* than in *A. thaliana*, thus taking *A. thaliana* as reference would help greatly in comprehension of the interaction of candidate genes based on their functions. This strategy was used previously to investigate on proteins interaction in other studies (Gu et al., 2016; Chao et al., 2017; Raboanatahiry et al., 2017).

Thus, nine genes (AUX1, CO, FT, FLC, PHYA, CRY2, AGL20, BRI1, GA1) which are involved in important functions (flowering process, light response, and plant hormones) might have higher influence on the other genes which possibly dependent on these nine genes. CO, FT, FLC and AGL20 were four key genes in the network interaction of our analysis and they are both implicated in plant flowering development. CONSTANS (CO) encodes a transcription factor that induces flowering in long exposure to daylight and stimulates the expression of FLOWERING LOCUS (FT) in phloem of leaves (Koornneef et al., 1991; Putterill et al., 1995; Suárez-López et al., 2001; An et al., 2004; Smit and Weijers, 2015). It has been shown recently that CO interacted with two VASCULAR PLANT ONE-ZINC FINGER (VOZ1 and VOZ2) and indorsed transition of photoperiodic flowering (Kumar et al., 2018). Note that FT is induced in long daylength and promotes flowering (Li C. et al., 2011). In fact, FT is the major component of florigen and promotes the transmission of flowering signal from leaf to shoot, where FT forms a complex with FD (bZIP transcription factor) and activate meristem genes to promote flowering (Abe et al., 2005; Wigge et al., 2005; Corbesier et al., 2007; Zeevaart, 2008; Nakamura et al., 2014). However, FLOWERING LOCUS C (FLC), which is a dominant transcription factor gene implied in vernalization, is antagonistic to FT (Michaels and Amasino, 1999; Sheldon et al., 1999; Hepworth et al., 2002; Hellwell et al., 2006). In fact, FLC blocks expression of both FT, FD and SUPPRESSOR OF CONSTANS 1 (SOC1) in meristem, which consequently abort flower genesis (Lee et al., 2000; Onouchi et al., 2000; Samach et al., 2000; Searle et al., 2006). Nevertheless, a gene that encodes a MADS-domain protein, named AGAMOUS-LIKE 20 (AGL20) is positively regulated by unsolicited vernalization, and independent flowering and photoperiodic pathways, and could relieve late flowering in plants that hold FLC functional alleles (Lee et al., 2000). PHYTOCHROME A (PHYA) and CRYPTOCHROME2 (CRY2) were also key genes in interaction network of our analysis, they could also encourage flowering by CO stabilization via CONSTITUTIVE PHOTOMORPHOGENIC 1 (COP1) and suppressor of PHYA1 (SPA1) ubiquitin ligase complex in nightfall (Sarid-Krebs et al., 2015). Light-labile PHYA is the most abundant phytochrome in etiolated seedlings and could absorb red, far-red light and blue/ultraviolet-A wavelengths (Sharrock and Quail, 1989; Li J. et al., 2011). PHYA could interact with SPA and TANDEM ZINC-FINGER/PLUS3 (TZP), disrupting COP1/SPA complex to induce photomorphogenesis, and regulating PHYA abundance and phosphorylation in far red light, respectively (Sheerin et al., 2015; Zhang S. et al., 2018). However, CRY2 are blue light receptors and intercede in response to photoperiodic flowering, leaf senescence and cell development, hypocotyl elongation and flowering regulations (Ahmad and Cashmore, 1993; Guo et al., 1998; Lin et al., 1998; El-Assal et al., 2001; Cashmore, 2003; Meng et al., 2013). AUX1, BRI1 and GA1 are genes related to plant hormones which were detected as highly influential in our interaction network analysis. AUXIN INFLUX TRANSPORTER1/AUXIN RESISTANT1 (AUX1) brings auxin hormone to the root apex and intercedes in repression of root elongation in presence of heavy metals and alkalinity, and in control of cytokinin on root gravitropism (Swarup et al., 2001; Sun et al., 2010; Li et al., 2015; Pernisova et al., 2016; Yuan and Huang, 2016). Besides, BRASSINOSTEROID INSENSITIVE1 (BRI1) recognizes brassinosteroids hormone (BRs) in growth and development regulation, and endocytosis of BRI1 is important to reduce signaling; if incorporation of BRI1 is inhibited, BRs response is enhanced, and in case of ubiquitination loss, degradation of BRI1 is hindered (Li and Chory, 1997; Irani et al., 2012; Zhou et al., 2018). Also, GIBBERELLIC ACID INSENSITIVE (GAI), which is a DELLA motif containing protein, is involved in regulation of plant growth and in inhibition of GA-mediated plant responses (Yamaguchi, 2008; Davière and Achard, 2013). With REPRESSOR OF GA1-3 (RGA),
GAI might have overlapping roles in senescence regulation by weakening transcription factor WRKY6 activity on senescence related genes (Zhang Y. et al., 2018).

Actually, appropriate FT greatly influences the potential of rapeseed yield (Luo et al., 2014), it could have impact on hybrid vigor, and consequently on reproductive success (Srikanth and Schmid, 2011). FT is significantly related to maturity (Miguel et al., 1998; Searle et al., 2006), and the optimal time depends on the cultivar (i.e., winter or spring type) and the environment of cultivation. The identification of genes which control FT might offer sustainable agriculture according to environmental variation. Multiple genetic pathways that respond to different development and environment conditions control the floral transition (Baurle and Dean, 2006; Srikanth and Schmid, 2011; Andres and Coupland, 2012). In A. thaliana, five main pathways were found to control flowering: the gibberellin pathway in which flowering needs the activity of gibberellic acid; the endogenous pathway which modulates flowering according to the age of plant; the photoperiod and the vernalization pathways which control flowering according to day length and temperature fluctuation, respectively; and the autonomous pathway which works independently of gibberellin and photoperiod pathways (Amasino and Michaels, 2010; Srikanth and Schmid, 2011). The keys genes of our interaction network were involved in these pathways. Note that in a study which aimed to evaluate rapeseed for yield performance, it has been demonstrated that early flowering promoted better SY because it gave enough time for the seeds to fill (Ali et al., 2013).

Otherwise, these nine above-mentioned genes might affect variation of traits, and quantitative traits are affected by structure and dynamism of genetic regulatory networks (Tabor et al., 2002; Frank, 2003). In the current interaction network analysis, candidate genes from overlapping QTLs of multiple population were taken to the study to find the nine above-mentioned pivotal genes. Other findings might be found if investigation was made on single population. At the end, these findings indicated that yield traits might be affected at multiple set of metabolisms. Unfortunately, in our analysis, we used A. thaliana but not B. napus, which indicated that findings might be imprecise. Nevertheless, construction of this network gave new insight on interaction of candidate genes.

CONCLUSION

This study reported QTLs for yield and yield-related traits in B. napus, which could be aligned and combined in one physical map. Unluckily, some QTLs could not be aligned in the map. However, stable QTLs which corresponded to overlapping QTLs could be predicted, but need confirmation. Nevertheless, region enriched with QTL were found which could help to understand the process of close linkage in traits and to identify varied or conserved area. Then, this map could serve as reference to understand rapeseed genome characteristic, especially while studying yield traits. Additionally, new candidate genes could be detected in these hypothetical stable QTLs, but still need to be confirmed as well, even if they were carefully chosen due to their presupposed yield related function and their position within QTL regions. Based on our current study, markers could be developed and potential candidate genes could be tested for breeding improvement purpose.

AUTHOR CONTRIBUTIONS

NR and HC wrote the manuscript. NR, HC, HD, SP, and WY performed the analysis of QTL alignment map. NR analyzed the candidate genes and interaction network. ML, BW, and LY supervised the work, and revised the manuscript.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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