

ORIGINAL ARTICLE

Multienvironment QTL analysis delineates a major locus associated with homoeologous exchanges for water-use efficiency and seed yield in canola

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Abstract

Canola varieties exhibit variation in drought avoidance and drought escape traits, reflecting adaptation to water-deficit environments. Our understanding of underlying genes and their interaction across environments in improving crop productivity is limited. A doubled haploid population was analysed to identify quantitative trait loci (QTL) associated with water-use efficiency (WUE) related traits. High WUE in the vegetative phase was associated with low seed yield. Based on the resequenced parental genome data, we developed sequence-capture-based markers and validated their linkage with carbon isotope discrimination ($\Delta^{13}\text{C}$) in an F_2 population. RNA sequencing was performed to determine the expression of candidate genes underlying $\Delta^{13}\text{C}$ QTL. QTL contributing to main and QTL $\times$ environment interaction effects for $\Delta^{13}\text{C}$ and yield were identified. One multiple-trait QTL for $\Delta^{13}\text{C}$, days to flower, plant height, and seed yield was identified on chromosome A09. Interestingly, this QTL region overlapped with a homoeologous exchange (HE) event, suggesting its association with the multiple traits. Transcriptome analysis revealed 121 significantly differentially expressed genes underlying $\Delta^{13}\text{C}$ QTL on

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A09 and C09, including in HE regions. Sorting out the negative relationship between vegetative WUE and seed yield is a priority. Genetic and genomic resources and knowledge so developed could improve canola WUE and yield.

KEYWORDS

carbon isotope discrimination, drought avoidance, gene expression, genetic analysis, physiology, water use

1 | INTRODUCTION

Drought is the major abiotic stress that reduces the yield potential of various crops including canola (*Brassica napus* L.), especially in arid and semiarid regions, of which 89% of regions are prevalent in Oceania (Koochafkan & Stewart, 2008). No doubt the impact of drought stress on crop productivity can be alleviated through irrigation at the “critical” stages of plant development. However, in recent years, freshwater, suitable for irrigation, is becoming scarce for crop production, required to meet the demand of a burgeoning human population (Gleick, 2000). Predicted climatic patterns such as debilitating drought and heat-wave episodes and their possible increased frequency further pose a significant threat to crop production (Mills et al., 2018; S. Smith et al., 2012). The proportion of arable land per capita is also decreasing at a significant rate due to population growth and land degradation (<http://www.fao.org/sustainability/>). Therefore, improving crop varieties that have high yield potential and utilize water more effectively or require less water could provide a part of the solution to reduce the negative impacts of drought stress and increase productivity and food security (Bertolino et al., 2019; Blum, 2009; Kijne et al., 2003; Leakey et al., 2019; Passioura, 1977).

Canola is an allotetraploid (AACC, $2n = 38$) member of the family Brassicaceae derived from interspecific hybridization between the diploid progenitors, *Brassica rapa* (AA, $2n = 20$) and *Brassica oleracea* (CC, $2n = 18$). Genomes of these diploid and allotetraploid species have undergone extensive evolutionary events, including homoeologous exchange (HE; Bayer et al., 2021; Chalhoub et al., 2014; Cheung et al., 2009; Hurgobin et al., 2018; Liu et al., 2014; Lysak et al., 2005; Parkin et al., 2005; Schranz et al., 2006; Town et al., 2006; Wang et al., 2011). Currently, canola is the second most important oilseed crop, grown worldwide with a global production of 75 million tons and used to produce 15% of vegetable oil (FAO STAT, <http://www.fao.org/>). Water stress reduces canola seed yield by up to 60% (Champolivier & Merrien, 1996; Gan et al., 2004; Qian et al., 2018). However, its effect varies depending upon the stage of plant development, and the severity and frequency of water stress (Hall et al., 1994). Despite its economic significance to the oilseed industry as well as being an essential rotational crop in agricultural production systems, little research has been conducted on traits contributing to drought avoidance and drought tolerance in canola (Fletcher et al., 2016, 2015; Hossain et al., 2020; Knight et al., 1994; Matus et al., 1995; McVetty et al., 1989; Pater et al., 2017; Raman, Raman, Mathews, et al., 2020; Raman, Raman, McVittie, et al., 2020).

In nature, to cope with water-deficit conditions, plants have evolved three strategies known as drought escape, drought avoidance, and drought tolerance (Levitt, 1980; Ludlow, 1989; Rodrigues et al., 2019; Zhu et al., 2016). The drought escape strategy enables plants to complete their life cycle before the onset of drought. Drought avoidance involves maintaining internal water status under drought conditions by minimizing water loss/or maximizing water uptake (water-use efficiency, WUE) while the drought tolerance strategy enables the plant to survive under internal water deficits (Ludlow, 1989; McKay et al., 2003). Through tiny microscopic pores on the surface of leaves called stomata, plants assimilate CO_2 for photosynthesis by trading-off water, required for transpiration and other biological processes. This close intimacy between productivity and water use contributes to the adaptation of plants to their growing environments. Therefore, genetic variation in WUE and transpiration efficiency (TE, biomass production/transpirational water loss) that occurs as a result of intentional (via breeding/selection) and unintentional selection in nature provides an opportunity to identify and assemble useful alleles for improving the productivity of various crops.

Genetic variation in WUE has been described variously at the single leaf level as intrinsic WUE (iWUE), defined as the ratio of the photosynthetic CO_2 assimilation rate (A) over stomatal conductance, g_{sw} , as TE, defined as the ratio of A over transpirational water loss, as whole-plant vegetative WUE, as the ratio of total dry matter production to total water transpired (dmWUE) or as an integrated whole-plant WUE, as the ratio of above-ground shoot biomass (sbWUE) or seed yield (syWUE) to evapotranspiration (Farquhar et al., 1982; Farquhar & Richards, 1984; Leakey et al., 2019; Raman et al., 2019; Zhengbin et al., 2011). However, iWUE assessments using the gas-exchange method are very challenging to be accurately performed, particularly in large breeding populations, as WUE is regulated by a myriad of plant development, physiological, biochemical, and molecular networks (Moore et al., 2009; Takahashi et al., 2018). Farquhar and Richards (1984) proposed $\Delta^{13}\text{C}$ (the discrimination against ^{13}C which is negatively related to $\delta^{13}\text{C}$) as a time-integrated surrogate trait for assessing WUE both at the single leaf level (mmol C/mol H_2O) and the whole plant level (dmWUE, dry matter/kg H_2O), as C_3 plants discriminate less against ^{13}C during photosynthesis with increased water-deficit stress. The negative relationship between TE and $\Delta^{13}\text{C}$ (mostly in the vegetative growth periods) has been verified in *Arabidopsis thaliana* (Campitelli et al., 2016; Masle et al., 2005) and some agricultural crop plants,

including canola (Des Marais et al., 2014; Ehleringer, 1993; Farquhar & Richards, 1984; Fischer et al., 1998; Hall et al., 1994; Hossain et al., 2020; Rebetzke et al., 2008), with some exceptions where nil or weak positive relationships were observed between syWUE and $\Delta^{13}\text{C}$ (Devi et al., 2011; Hammer et al., 1997; Mekonnen et al., 2020; Monneveux et al., 2007; Raman, Raman, McVittie, et al., 2020).

Earlier genetic analysis studies suggest that drought tolerance is a complex trait, controlled by multiple quantitative trait loci (QTL) in canola. So far, QTL linked to drought avoidance traits; $\Delta^{13}\text{C}$, lateral root number, root mass, root diameter, taproot length, and root pulling force have been identified in canola by traditional simple-, interval-, and composite QTL mapping approaches (Fletcher et al., 2016, 2015; Mekonnen et al., 2020; Raman, Raman, Mathews, et al., 2020; Raman, Raman, McVittie, et al., 2020). Some of these QTL associated with drought avoidance traits also map in the vicinity or colocalize with QTL for flowering time, shoot biomass, plant height, and seed yield in canola, and with QTL for iWUE, photosynthetic capacity, leaf thickness, nitrogen content, and stomatal density in *B. oleracea* (Fletcher et al., 2015; Hall et al., 2005; Mekonnen et al., 2020). It is yet to be established which canola genotypes and loci show association with phenotypic plasticity and genotype (G) $\times$ environment (E) interaction in $\Delta^{13}\text{C}$ and other integrated WUE traits in response to different environmental conditions. Thus, a comprehensive understanding of the genetical and physiological bases, and identification of stable loci, and also subjected to G $\times$ E interaction across environments underlying WUE is central to developing strategies for resilience to water-deficit conditions. Recently, it was shown that two Australian canola inbred lines, BC1329, and BC9102, differ by ~2% in their $\Delta^{13}\text{C}$ signatures with corresponding opposite differences in TE (Hossain et al., 2020). However, the genetic and environmental determinants underlying variation in $\Delta^{13}\text{C}$ and other integrated WUE traits, such as plant biomass, flowering time, and seed yield were not deciphered. To uncover environmentally stable and environment-specific loci associated with WUE traits, we carried out multienvironment QTL analyses based on extensive phenotypic and physiological measurements and genomic and transcriptomic studies. This multienvironment QTL approach was not used to uncover the genetic basis of drought avoidance and drought escape traits in earlier studies (Fletcher et al., 2016, 2015; Mekonnen et al., 2020; Raman, Raman, Mathews, et al., 2020; Raman, Raman, McVittie, et al., 2020). We demonstrate that multiple genetic and environmental determinants underlie phenotypic variation in drought avoidance traits such as $\Delta^{13}\text{C}$, early vigour, and plant height and drought escape traits such as flowering time in canola. We show that one QTL which is associated with multiple traits; $\Delta^{13}\text{C}$, days to flower, plant height, and seed yield on chromosome A09 was subjected to homoeologous recombination. Our research also describes candidate genes and differentially expressed genes (DEGs) underlying two stable QTL for $\Delta^{13}\text{C}$ on chromosomes A09 and C09 that account for the majority of genotypic variance in the canola doubled haploid (DH) mapping population.

2 | MATERIALS AND METHODS

2.1 | Plant materials

A DH population of 223 lines (06-5101DH) derived from the F_1 cross between advanced breeding lines “BC1329” (maternal parent) and “BC9102” (paternal parent) were evaluated in different genetic analysis experiments. In addition, an F_2 population comprising 733 lines derived from a single F_1 plant from BC1329/BC9102 was employed for validation of QTL associated with $\Delta^{13}\text{C}$.

2.2 | Experimental trials

Six experiments were conducted to (i) determine the genetic basis of variation in WUE-related traits (Experiments 1–4), (ii) determine the relationship between physiological ($\Delta^{13}\text{C}$, iWUE) and integrated WUE-related traits under wet and dry conditions (Experiment 5), and (iii) identify DEGs/candidate genes underlying QTL for $\Delta^{13}\text{C}$ variation (Experiment 6) under field, glasshouse, and rain-out shelter conditions. Details of the experiments and their designs are presented (Supporting Information: Method S1). Monthly weather statistics for average atmospheric temperatures and rainfall are also presented (Supporting Information: Figure S1).

2.3 | Phenotypic trait measurements

Several plant development, agronomic, gas exchange, and other physiological traits were measured for genetic analysis. A summary of the experiments in terms of their aim, genetic material evaluated, and traits measured are presented (Table 1, Supporting Information: Figure S2). Details of trait measurement protocols are given in our recent study (Raman, Raman, McVittie, et al., 2020) and summarized in Methods S1. A brief description of the traits measured is given below.

2.4 | Plant development and agronomic traits

$\Delta^{13}\text{C}$, days to flower, plant height, and seed yield were measured for Experiments 1–3 and 5, and normalized difference in the vegetative index (NDVI) was measured only for Experiment 2. The $\delta^{13}\text{C}$ composition ($^{13}\text{C}/^{12}\text{C}$) was determined at laboratory phase experiments for dried leaf samples collected from field/pot phase experiments with appropriate multiphase experimental designs (Supporting Information: Method S1) to account for the variations attributed to field/pot and laboratory conditions (Smith et al., 2006), as described previously (Raman, Raman, McVittie, et al., 2020). $\Delta^{13}\text{C}$ was determined from the leaf $\delta^{13}\text{C}$ composition and that of the source CO_2 in the air (taken as -7.8‰) as proposed by Farquhar and Richards (1984). For Experiment 4, $\Delta^{13}\text{C}$, days to flower, specific leaf weight, and leaf-water content (LWC) were measured.

TABLE 1 Description of phenotyping experiments conducted to uncover the genetic basis of carbon isotope discrimination ($\Delta^{13}\text{C}$) and water-use efficiency-related traits in canola (*B. napus* L). The plant development stage at which traits were measured is detailed (Method S1)

Experiment	Research question	Material	Phenotyping environment	Traits measured
1	Genetic basis of $\Delta^{13}\text{C}$ variation	223 DH lines from BC1329/BC9102, plus parental lines	Field (2017)	$\Delta^{13}\text{C}$ Days to flower (DTF) Plant height (PH) Seed yield (SY)
2		223 DH lines from BC1329/BC9102, plus parental lines and commercial cultivars	Field (2018)	$\Delta^{13}\text{C}$ Normalised difference in the vegetative index (NDVI) DTF PH SY
3		217 Doubled haploid lines from BC1329/BC9102, plus parental lines	Pots under rain-out shelter (2017)	$\Delta^{13}\text{C}$ DTF PH
4		733 F ₂ lines derived from BC1329/BC9102, plus parental lines	Pots in glasshouse (2020) under wet (field capacity) conditions	$\Delta^{13}\text{C}$ DTF Specific leaf weight (SLW) Leaf-water content (LWC)
5	Relationship between physiological (intrinsic water-use efficiency, $\Delta^{13}\text{C}$) and agronomic water-use efficiency (canola productivity) related traits	Selected 70 DH lines representing extreme (low and high $\Delta^{13}\text{C}$ values) plus parental lines under wet and dry conditions	Rain-out shelter (2019)	$\Delta^{13}\text{C}$ DTF PH SY Photosynthesis (A) Stomatal conductance (g_{sw}) Intrinsic water-use efficiency (iWUE) SLW LWC
6	Identify differentially expressed genes underlying QTL for $\Delta^{13}\text{C}$ variation	Parental lines of DH population (BC1329 and BC9102)	Pots in glasshouse (2020) under wet and dry conditions	None

Note: The plant development stage at which traits were measured is detailed in Supporting Information: Method S1.

Abbreviations: DH, doubled haploid; QTL, quantitative trait loci.

2.5 | Physiological traits

The gas exchange measurements were taken at the single leaf level for the plots under wet conditions in Experiment 5, as the relationship between $\Delta^{13}\text{C}$ and WUE varies under different water-deficit levels. We determined iWUE by measuring light-saturated assimilation rate (A) and stomatal conductance to the diffusion of water vapour (g_{sw}). The fifth fully expanded leaf of each of the 70 lines of the DH population (representing extreme phenotypes), including parental lines, was tagged and utilized for gas exchange measurements. Details of parameters used for gas exchange measurements are given in our earlier study (Raman, Raman, McVittie, et al., 2020).

2.6 | Light microscopy

A leaf disc (9.08 cm² size) was taken from each of the two replicate plots in the wet block from Experiment 5, fixed, and stored in 70% ethanol as detailed (Supporting Information: Method S1). Leaf

sections were stained using a method modified from Rae et al. (2020) and were imaged using 488 nm excitation and 500–560 nm emission on a Leica SP8 confocal microscope.

2.7 | Genotyping and linkage map construction

Genotyping of DH lines was carried out using the *PstI*-*MseI* restriction enzymes based genotyping-by-sequencing DArTseq approach (Raman et al., 2014). Sequence polymorphisms were used for linkage map construction, following the DArT P/L's mapping software OCDmap[®] detailed in Raman et al. (2014). Recombination frequencies were converted to centiMorgan (cM) map distances using the Kosambi function. The markers that showed cosegregation were "binned" into a unique locus and the resulting "bin" map was used to identify trait-marker associations. To obtain the physical position of markers, DArTseq sequences were aligned with the Darmor-bzh reference assembly version 4.1 using the default parameter settings with the Bowtie programme.

2.8 | Statistical methods

Commensurate with the aims of the experiments and the structure of the datasets, for Experiments 1–3 whole genome, single-step QTL analyses were performed on each trait using an extension of the approach developed by Verbyla and Cullis (2012) within a multi-environment trial (MET) analysis framework using factor analytic linear mixed models (FA-LMM) (Smith et al., 2015). A genome scan within a linear mixed model (LMM) framework was performed for $\Delta^{13}\text{C}$ measured at Experiment 4 to validate the QTL identified from Experiments 1 to 3. Each trait measured for Experiment 5 was analysed individually using appropriate LMM. A detailed description of the methods is presented (Supporting Information: Method S1).

All analyses were performed in ASReml-R (Butler et al., 2018), which provides residual maximum likelihood (REML) estimates of variance parameters, empirical best linear unbiased predictions (EBLUPs) of random effects, and empirical best linear unbiased estimates of fixed effects. The extent of genetic control of traits in each environment was investigated by calculating line mean H^2 (broad-sense heritability) as the mean of the squared accuracy of the predicted line effects as described previously (Cullis et al., 2006) and found to be dependent on the environment. The across environment summary measure of overall performance (OP) proposed by Smith and Cullis (2018) was used to identify lines of interest. OP is one of the so-called factor analytic selection tools to summarize the predictions from the fit of an FA-LMM for a MET dataset in a concise yet informative manner. OP is defined for a line using the EBLUP of the first-factor score in the FA-LMM representing the generalized genotype main effect. We examined the genetic correlations between pairs of traits, especially $\Delta^{13}\text{C}$ with agronomic traits (seed yield, days to flowering, plant height, and NDVI) using bivariate analyses where data for each pair of traits were combined and analysed together within the LMM framework. Three sets of bivariate analyses were performed: (1) Experiments 1–3, (2) Experiment 4, and (3) Experiment 5. Correlation between traits was modelled using an unstructured variance model and the inference on the genetic correlation between traits was obtained (see Supporting Information: Method S1).

2.9 | Resequencing and structural variation analysis of parental lines

Libraries from high-quality genomic DNA from both parental lines, BC1329, and BC9102, were constructed using the Illumina TruSeq DNA Preparation Kit, following the manufacturer's instructions (Illumina). Whole-genome resequencing (2×150 bp) was performed at the Novogene facility (Novogene Co., Ltd.) using the Illumina HiSeq 2000 sequencing platform. The coverage of the parental lines ranged from 77.6 (BC1329, 102.6 Gb) to 83.8 $\times$ (BC 9102, 112.4 Gb). Read mapping to the "Darmor-bzh" reference assembly (version 4.1, <http://www.genoscope.cns.fr/brassicapapus/data/>), single nucleotide polymorphisms (SNP) and InDel (<50 bp) calling, structural

variation (SV, ≥ 50 bp) detection, and identification of HE event (≥ 10 -kb windows) were performed as described in Raman et al. (2021). Details of HE analysis are described in Supporting Information: Method S1.

2.10 | Development of sequence-capture-based DArTag markers

We processed resequence data for target $\Delta^{13}\text{C}$ QTL regions on A09 and C09 chromosomes and selected 148 SNPs for DArTag oligo synthesis. Oligos were synthesized by IDT (Ultramer DNA Oligos, <http://idtdna.com>) at a 200 pmol scale, pooled in the equimolar amount into a single assay, and used for processing eight plates (96 well/plate) of DNA with the F_2 population and a control canola sample using a proprietary DArTag assay (Targeted Genotyping–Diversity Arrays Technology) using 384 plate format. For each plate, a sample of the pooled product was also run on agarose gel and compared against positive control before proceeding with the sequencing process. The libraries were sequenced on Illumina HiSeq 2500 with an average volume of sequencing per sample at 43 225 sequencing reads (median at 46 389) and average read depth per assay at 280. Marker data were extracted using DArT PL's proprietary algorithm deployed as a plugin in the KDCompute application framework (<https://www.kddart.org/kdcompute.html>).

2.11 | RNA sequencing and differential gene expression analysis

Parental lines, BC1329 and BC9102, of the DH population were grown with three replicates under both wet (100% field capacity) and dry (50% field capacity) conditions in a glasshouse (Experiment 6). Details of the layout of the experiment, sampling, RNA isolation, and sequence analysis are presented (Supporting Information: Method S1). A generalized linear model approach was then used to quantify the differential expression between the groups. The DEGs were obtained using a false discovery rate (<0.05). The p -value was adjusted for multiple hypothesis testing using the Benjamini–Hochberg adjustment ($p < 0.05$). Heatmaps showing the expression pattern of genes in A09 and C09 QTL regions were produced using the ComplexHeatmap R package (Gu et al., 2016).

2.12 | Identification of candidate genes for WUE

Arabidopsis thaliana genes which had been annotated with various WUE-related terms were retrieved from the TAIR 10 database (<https://www.arabidopsis.org/>). These genes were then used to identify putative homologs in canola. We implemented an improved syntenic approach to exclude genes as causal candidates in syntenic QTL regions combining genomic and transcriptomics data (Zhang et al., 2019).

3 | RESULTS

3.1 | Substantial genetic variation in $\Delta^{13}\text{C}$ and other productivity traits

We observed high levels of genetic variation in $\Delta^{13}\text{C}$ and other WUE-related traits in the DH population. The significant source of genetic variation was from the additive component (genetic markers), where the mean values ranged from 21.5% for NDVI to 79.1% for days to flower (Supporting Information: Table S1, Additive M1, %). Broad sense heritability estimates for $\Delta^{13}\text{C}$ and other integrated WUE-related traits (plant height, NDVI, flowering time, and seed yield) were variable, ranging from low (56%) to high (98%), depending on the nature of the trait and growing environment (Supporting Information: Table S2). Across traits, pot experiments had lower H^2 values compared to field experiments. The frequency distributions of the traits from DH lines evaluated across environments are presented (Supporting Information: Figure S3). Estimated additive and total (additive plus non-additive) genetic correlations between environments revealed that there are strong correlations between environments for both additive and total genetic variance with values greater than 0.89 and 0.83, respectively, for all traits (Supporting Information: Table S3). Total (additive plus non-additive) OP estimates for $\Delta^{13}\text{C}$ ranged from 18.73‰ to 21.25‰ and displayed transgressive segregation among DH lines across environments (Figure 1a, Supporting Information: Table S4). Up to 2.52‰ variation in $\Delta^{13}\text{C}$ was observed among DH lines which equates to a fivefold increase compared with the parental lines.

3.2 | Genetic correlations between WUE and productivity traits at plot level

To determine the relationships between $\Delta^{13}\text{C}$ and other WUE-related traits, between trait total (additive plus non-additive) genetic correlations were obtained using bivariate analyses (Figure 1b). The $\Delta^{13}\text{C}$ showed a negative correlation with days to flower ($r = -0.54$), while positive correlations were observed with NDVI, a proxy for plant vigour ($r = 0.30$), plant height ($r = 0.45$), and seed yield ($r = 0.63$). Flowering time showed a negative correlation with seed yield ($r = -0.53$). The promising DH lines that had high seed yield (high $\Delta^{13}\text{C}$) for use in canola breeding programmes based on the OP estimates are presented in Figure 1c. DH line 06-5101-137 had the maximum $\Delta^{13}\text{C}$ (21.25‰) among the DH progenies.

3.3 | Genetic correlations between physiological WUE (single leaf level) and integrated WUE (whole plant level)

Significant genetic variation for both A and g_{sw} was observed, although the H^2 estimate of $i\text{WUE}$ was low (Supporting Information: Table S5). Genotype EBLUPs for A and g_{sw} ranged from 4.97 to

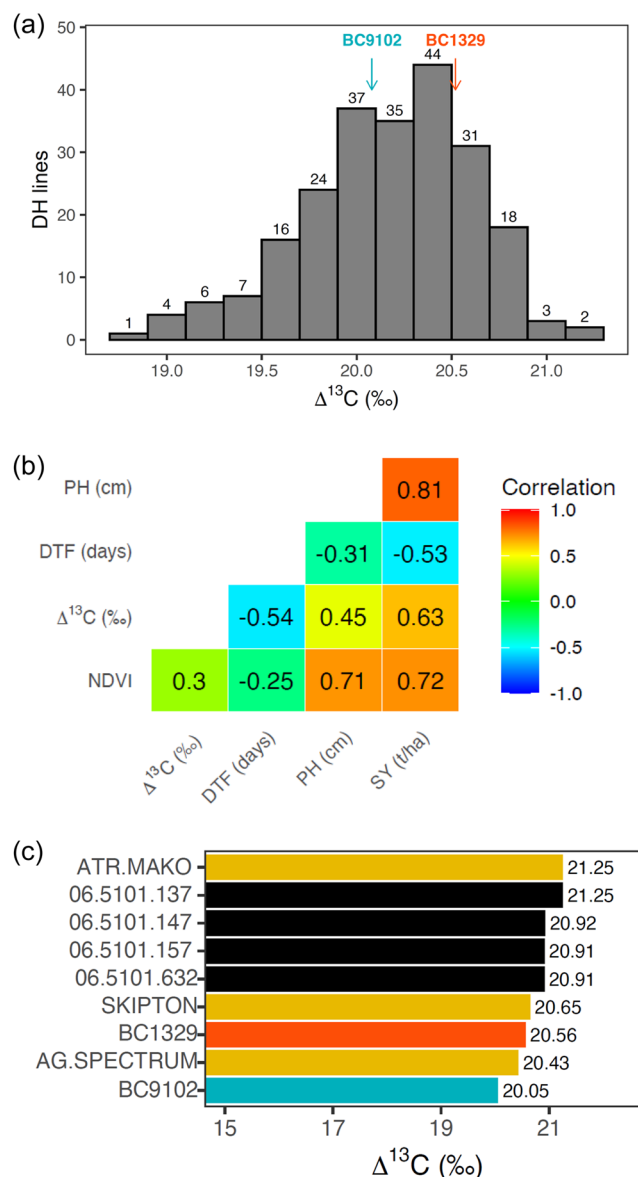


FIGURE 1 Genetic variation in WUE traits and their relationships among doubled haploid lines derived from the cross, BC1329/BC9102 (Experiments 1–3). (a) Frequency distribution of the OP for $\Delta^{13}\text{C}$. Estimates for the parental lines are shown with arrows; (b) genetic correlations between $\Delta^{13}\text{C}$ (%) and other WUE-related traits; and (c) the top four DH lines that showed the highest $\Delta^{13}\text{C}$ based on OP across environments in relation to control commercial varieties of canola and the parental lines are shown. DH, doubled haploid; DTF, days to flower; NDVI, normalised difference in the vegetative index; OP, overall performance; PH, plant height; SY, seed yield; WUE, water use efficiency.

17.15, and 0.11 to 0.38, respectively (Supporting Information: Table S5). The estimated between trait genetic (non-additive) correlations revealed that both A and g_{sw} are dependent on each other with a correlation of 0.61 (Figure 2a). We observed a negative correlation between $\Delta^{13}\text{C}$ and $i\text{WUE}$ ($r = -0.38$), indicating that DH lines with low $\Delta^{13}\text{C}$ have higher $i\text{WUE}$, consistent with the findings made earlier (Adiredjo et al., 2014; Farquhar & Richards, 1984). There

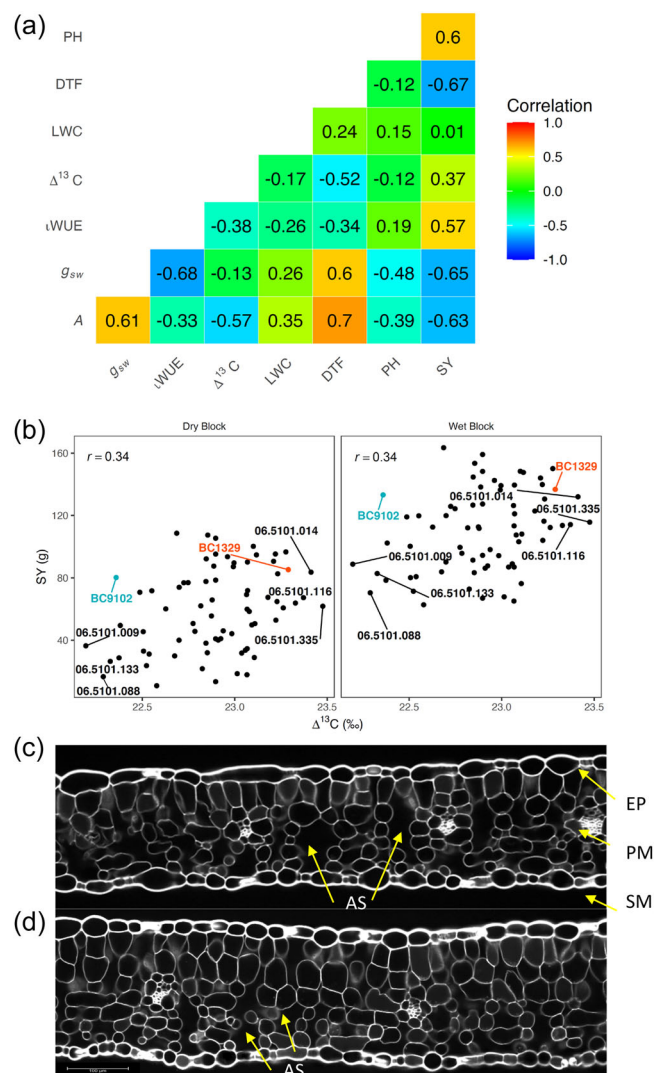


FIGURE 2 Relationships between $\Delta^{13}\text{C}$, gas exchange measurements (CO_2 assimilation (A), stomatal conductance (g_{sw}), intrinsic water-use efficiency (iWUE), and plant developmental and agronomic traits of selected 70 DH lines of the BC1329/BC9102 population, representing extremes (high and low values) in $\Delta^{13}\text{C}$ and their parents (Experiment 5). (a) Genetic correlations from the bivariate analyses are shown. (b) Relationships between $\Delta^{13}\text{C}$ and seed yield (SY) for wet and dry blocks. Genotype EBLUPs for $\Delta^{13}\text{C}$ and SY are plotted. Parental lines and the DH lines with high and low $\Delta^{13}\text{C}$ are labelled. Leaf sections showing differences in air spaces (AS, marked with arrow) between parental lines BC1329 (c) and BC9102 (d). DH, doubled haploid; DTF, days to flower; EBLUP, empirical best linear unbiased predictions; EP, epidermis; LWC, leaf-water content; PH, plant height; PM, palisade mesophyll (comparatively regular elongated cells); SM, spongy mesophyll (irregular cells).

was a more negative correlation between iWUE and g_{sw} ($r = -0.68$) in comparison to A ($r = -0.33$), suggesting that g_{sw} is the predominant driver for variation in iWUE parameters.

This study showed that $\Delta^{13}\text{C}$ correlates negatively with iWUE but it ($\Delta^{13}\text{C}$) correlates positively with seed yield (Figure 2a). Under well-watered conditions, there were negative correlations between

$\Delta^{13}\text{C}$ and days to flower, A, and iWUE. We further investigated relationships between LWC at a single leaf level and WUE traits at the whole plant level and found that LWC shows a negative relationship with $\Delta^{13}\text{C}$, but it did not show any relationship with seed yield (Figure 2a). Furthermore, the estimated genetic correlations between wet and dry blocks for seed yield (Figure 2b) and plant height, the only two traits measured after imposing water stress at the first flowering stage, were very high (0.93 for both traits). This suggests that genotype by irrigation block interaction is small. High $\Delta^{13}\text{C}$ lines revealed higher yield across irrigation blocks compared to low $\Delta^{13}\text{C}$ lines (Figure 2b). Our anatomical analysis of leaf discs revealed both parental lines BC1329 and BC9102 differ in thickness and arrangement of palisade and spongy mesophyll cells: BC1329 (192 μm) had high porosity with large air spaces compared to BC9102 (184 μm , Figure 2c,d), which may facilitate gas exchange, thus leading to efficient water use.

3.4 | Genetic basis underlying $\Delta^{13}\text{C}$ and WUE-related traits

We constructed a linkage map that includes 8985 DArTseq markers onto 24 linkage groups, representing all the 19 chromosomes of *B. napus* (Supporting Information: Table S6a). To reduce computation time for genetic analysis, we produce a “bin” map of 1793 markers that spanned a total of 1965.29 cM, with an average interval of 1.10 cM between adjacent loci (Supporting Information: Table S6c). Multi-environment QTL analyses identified a total of 29 QTL (15 QTL for main effects); corresponding to genotypic (G) effects, and 14 for QTL (Q) $\times$ environment (E) interactions; corresponding to G $\times$ E effects for variation in leaf $\Delta^{13}\text{C}$ and other WUE-related traits (Supporting Information: Table S7). For $\Delta^{13}\text{C}$, three QTL main effects that showed statistically significant (logarithm of the odds [LOD] ≥ 3) associations were identified on chromosomes A08, A09, and C09, while one “suggestive” QTL (LOD > 2.5 but less than 3) was located on chromosome A07 (Supporting Information: Table S7, Figure 3a, Supporting Information: Figure S5). We identified QTL for phenotypic variation in different traits between three growing environments (Q $\times$ E interaction) on A02, A05, A08, A09, A10, C02, C03, C06, C07, and C09 chromosomes (Supporting Information: Table S7). For $\Delta^{13}\text{C}$, two QTL for Q $\times$ E interactions were identified on chromosomes A02 and C06 (Figure 3a). Collectively, QTL explained 38% of genotypic variation in $\Delta^{13}\text{C}$ (Supporting Information: Table S1, VAF_m) of which the QTL delimited with DArTseq marker 3153720 accounted for the most R^2 (Supporting Information: Table S7). Environment-specific QTL showed a range of allelic effects, varying from consistent (in the same direction with same/similar magnitude) to opposite (in opposite direction with similar/variable magnitude, Supporting Information: Table S7). The strongest Q $\times$ E interactions with opposite allelic effects (different parental alleles) were observed for $\Delta^{13}\text{C}$ (A02, FT18), days to flower (A05, FT18; C02-FT17), plant height (C07 and C09; rainout shelter17, FT18), with variable allelic effects; depending on the environment

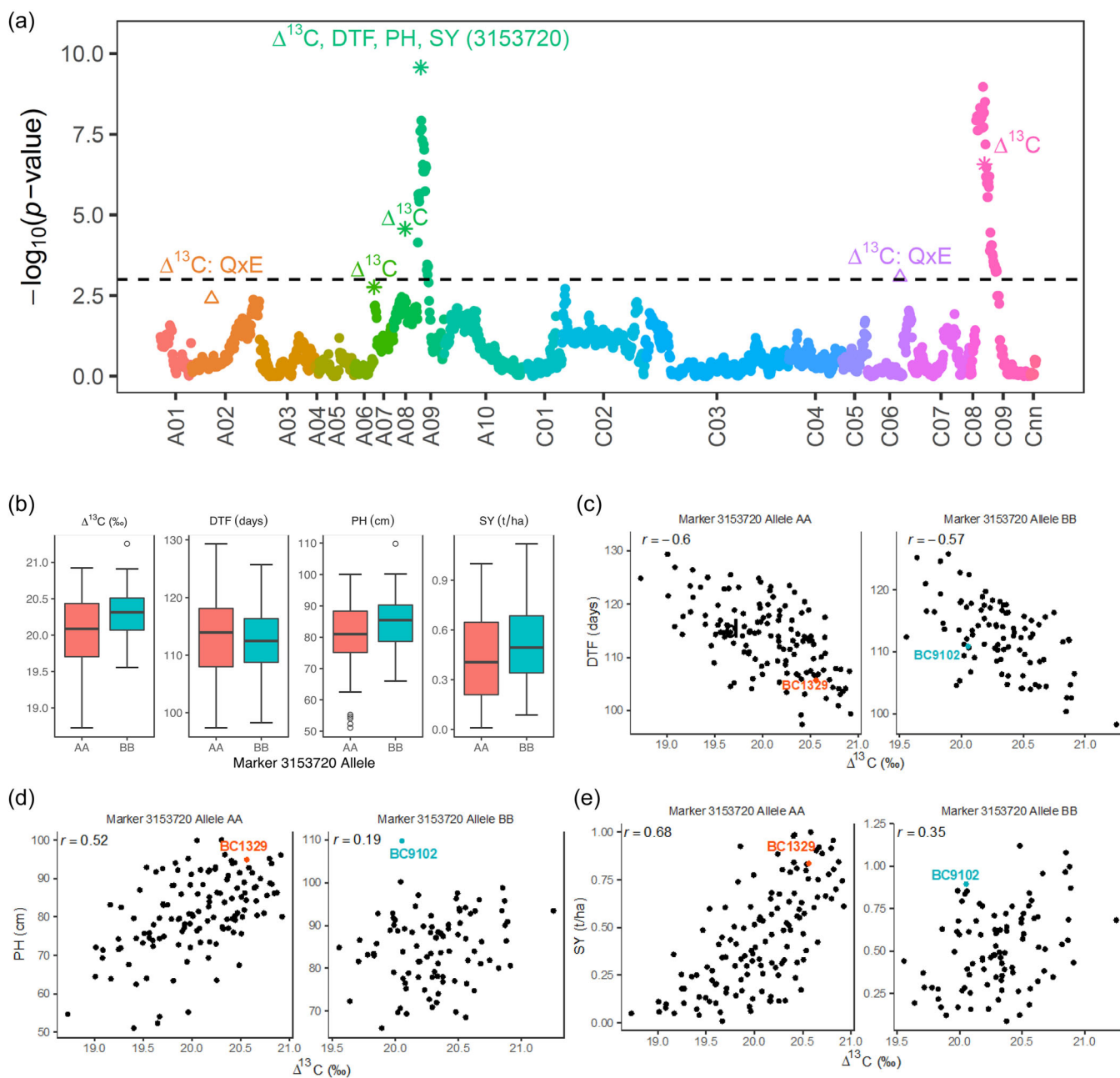


FIGURE 3 Distribution and relationships between $\Delta^{13}\text{C}$, days to flower (DTF), plant height (PH) and seed yield (SY), and DArTseq marker alleles for the QTL (3153720) that colocalized in the same genomic region on chromosome A09. (a) Manhattan plot showing LOD scores for associations between DArTseq markers and $\Delta^{13}\text{C}$. QTL main effects (depicted by *) are labelled with the respective trait (for DTF, PH, and SY only the 3153720 QTL is shown) and QTL $\times$ environment (E) interactions (depicted by Δ) are labelled with the trait followed by "Q $\times$ E" (only shown for $\Delta^{13}\text{C}$). LOD scores presented in the Manhattan plot are from the genome scan for the QTL main effects where the LOD scores of the significant QTL are replaced with the ones from the final model. The black dash line indicates the threshold value for significant SNPs at LOD ≥ 3 . Manhattan plot for each trait, are presented in Supporting Information: Figure S4. (b) Box plots showing the distribution of the OP estimates for $\Delta^{13}\text{C}$, DTF, PH, and SY partitioned into allele combinations, "AA (BC1329)" and "BB (BC9102)", for the SNP marker 3153720. Pair-wise correlations of OP estimates between $\Delta^{13}\text{C}$ versus DTF (c), $\Delta^{13}\text{C}$ versus PH (d), and $\Delta^{13}\text{C}$ versus SY (e) are partitioned into different allelic combinations. LOD, logarithm of the odds; OP, overall performance; QTL, quantitative trait loci; SNP, single-nucleotide polymorphisms.

(Supporting Information: Table S7). This implies that $G \times E$ interactions contribute to observed trait variation. Reaction norm of $\Delta^{13}\text{C}$ estimates from DH lines evaluated under two rainfed (field-2017, 2018, Experiments 1 and 2) and two irrigated (pot-2017, wet block

of rainout shelter-2019, Experiments 3 and 5) environments showed a change in scale (non-crossover interaction) and as well as ranks (crossover interaction), suggesting that variation in $\Delta^{13}\text{C}$ is also due to $G \times E$ interactions (Figure 4).

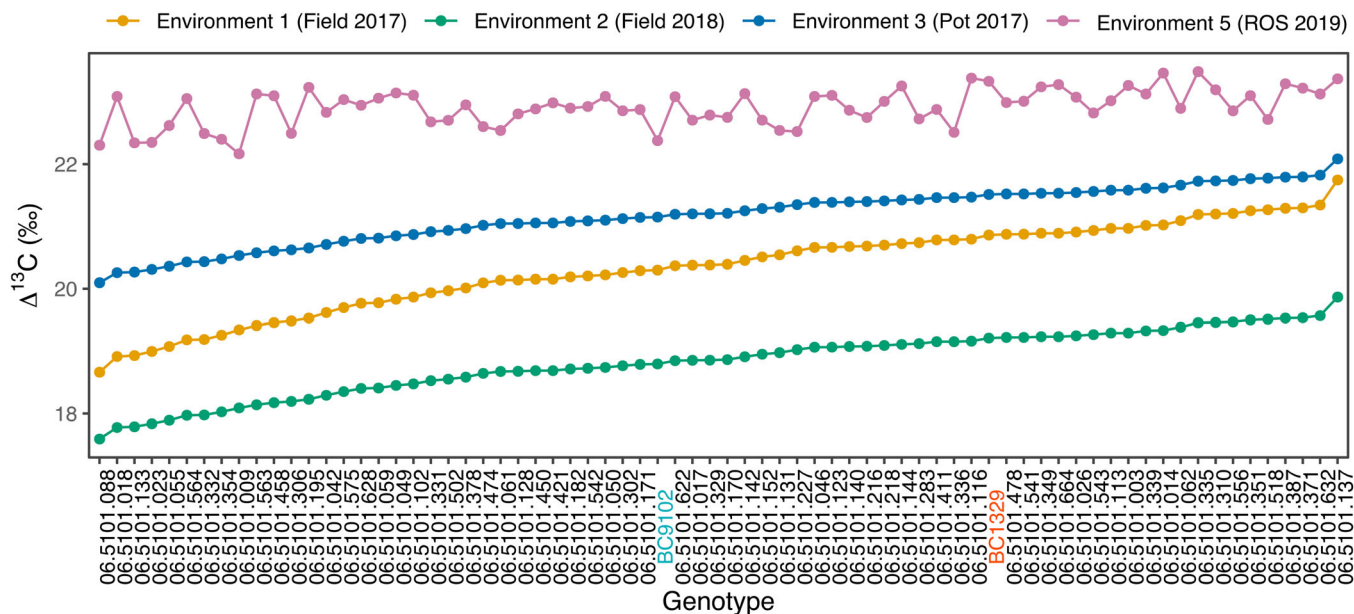


FIGURE 4 Reaction norm and phenotypic variation in $\Delta^{13}\text{C}$ (‰) among a subset of 70 DH lines and the parental lines evaluated under different environments (Experiments 1–3, 5). CGE EBLUPs for Experiments 1–3 and genotype EBLUPs for Experiment 5 are plotted to show the change in scale and ranks. Genotypes are ordered based on the ranks for Environment 1. DH, doubled haploid; EBLUP, empirical best linear unbiased predictions; CGE, common genotype by environment.

3.5 | Comparative localization of QTL

Three QTL for multiple traits on chromosomes A01, A08, and A09 were colocalized to the same genomic regions (Supporting Information: Table S7, Supporting Information: Figure S5). One QTL delimited with marker 3153720 for variation in $\Delta^{13}\text{C}$ was colocalized with QTL for days to flower, plant height, and seed yield on chromosome A09 (Supporting Information: Table S7, Figure 3a). We further sought a correlation between allelic effects of markers and variation in $\Delta^{13}\text{C}$, days to flower, plant height, and seed yield (Figure 3b–e). Up to 68% of allelic effects were explained by the same marker allele (Figure 3e), suggesting pleiotropic relationships between these traits and/or tight genetic linkage between them.

3.6 | Verification of $\Delta^{13}\text{C}$ QTL in an F_2 population

We validated the genetic control, the linkage between 148 DArTseq markers and $\Delta^{13}\text{C}$ (in DH population), and focused on the identification of candidate gene(s) underlying most of the genetic variation in $\Delta^{13}\text{C}$ at QTL regions on chromosomes A09 and C09 (Supporting Information: Table S7). The $\Delta^{13}\text{C}$ values showed normal segregation among F_2 lines (Figure 5a). Consistent with DH lines, $\Delta^{13}\text{C}$ exhibited a negative correlation with flowering time and SLW, and a positive correlation with LWC (Figure 5b). Genetic analysis revealed that the majority of polymorphic DArT markers (83.3%, 65/78) show significant segregation distortion (deviating from the normal segregation consistent with a 1:2:1 ratio for codominance, or 3:1 ratio for dominance) on chromosomes A09 and C09 (Supporting Information: Table S8) suggesting that the corresponding multiple trait QTL

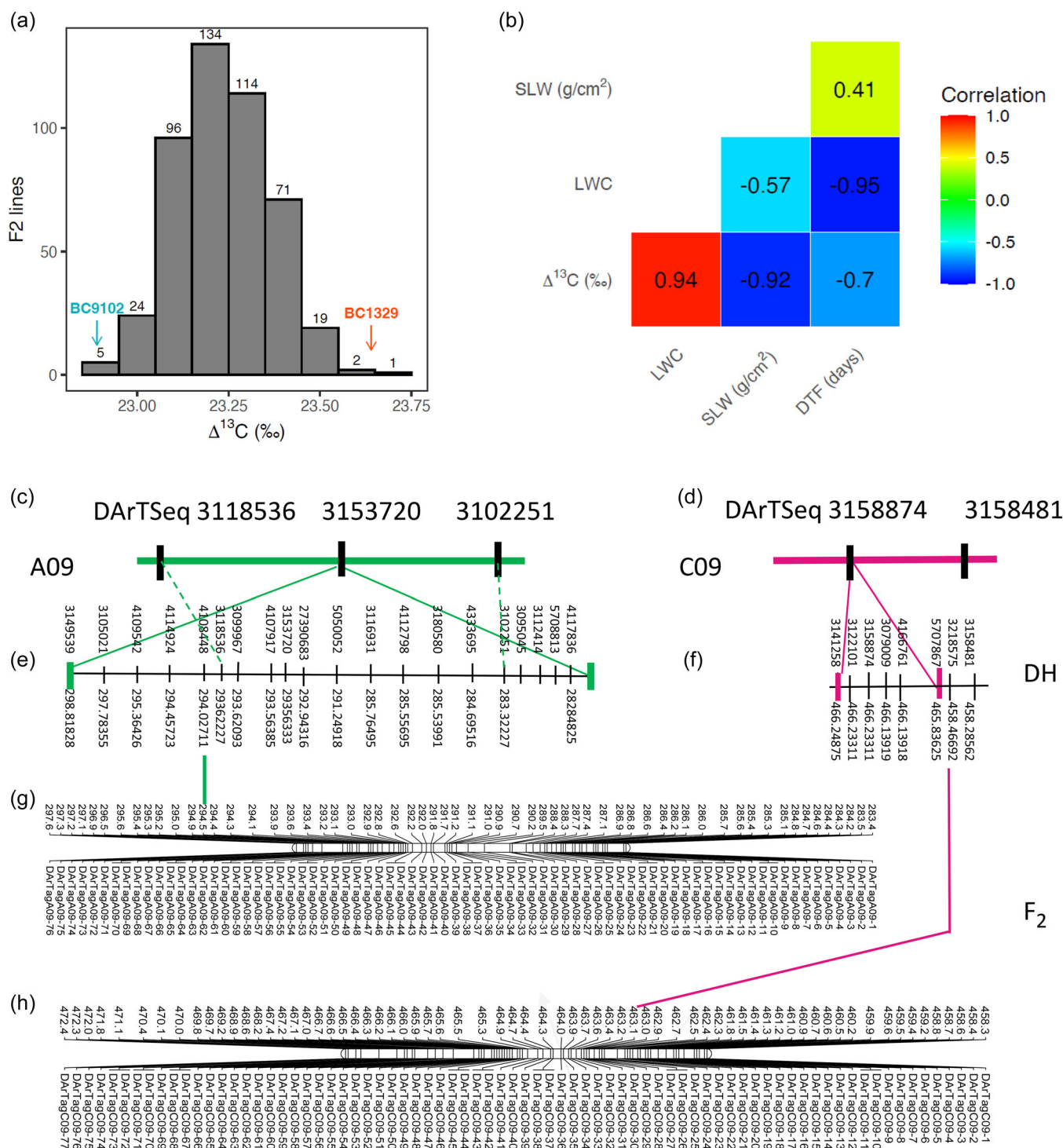
regions could be subjected to SV. Genome scan within LMM framework revealed that DArT markers positioned at 29 499 141 bp on chromosome A09, and 46 318 271 bp on C09 of the *Darmor-bzh* genome exhibit statistically significant association with $\Delta^{13}\text{C}$ in an F_2 population (Supporting Information: Figure S4). Physical mapping verified that the genomic regions for $\Delta^{13}\text{C}$ variation are localized to the same genetic intervals in both DH and F_2 populations (Figure 5c–f).

3.7 | $\Delta^{13}\text{C}$ QTL region on chromosome A09 is subjected to HE

To investigate whether significant segregation distortion among marker alleles on chromosomes A09 and C09 (Supporting Information: Table S8) is due to SV, we resequenced the genomes of parental lines of mapping/validation populations (NCBI, Accession No. PRJNA 743730). Sequence mapping revealed 26 genomic regions undergone HE events, varying from 90 to 870 kb, including the A09 multiple trait QTL regions (29.1 to 29.3 Mb) from the C08 chromosome in BC9102, as a result of homoeologous recombination (Figure 6a, Supporting Information: Table S9). However, in the maternal line BC1329, no such events were identified (Figure 6b).

3.8 | Candidate genes associated with WUE near $\Delta^{13}\text{C}$ QTL

We first interrogated genomic regions underlying the significant marker associations in both the mapping (DH) and validation (F_2)



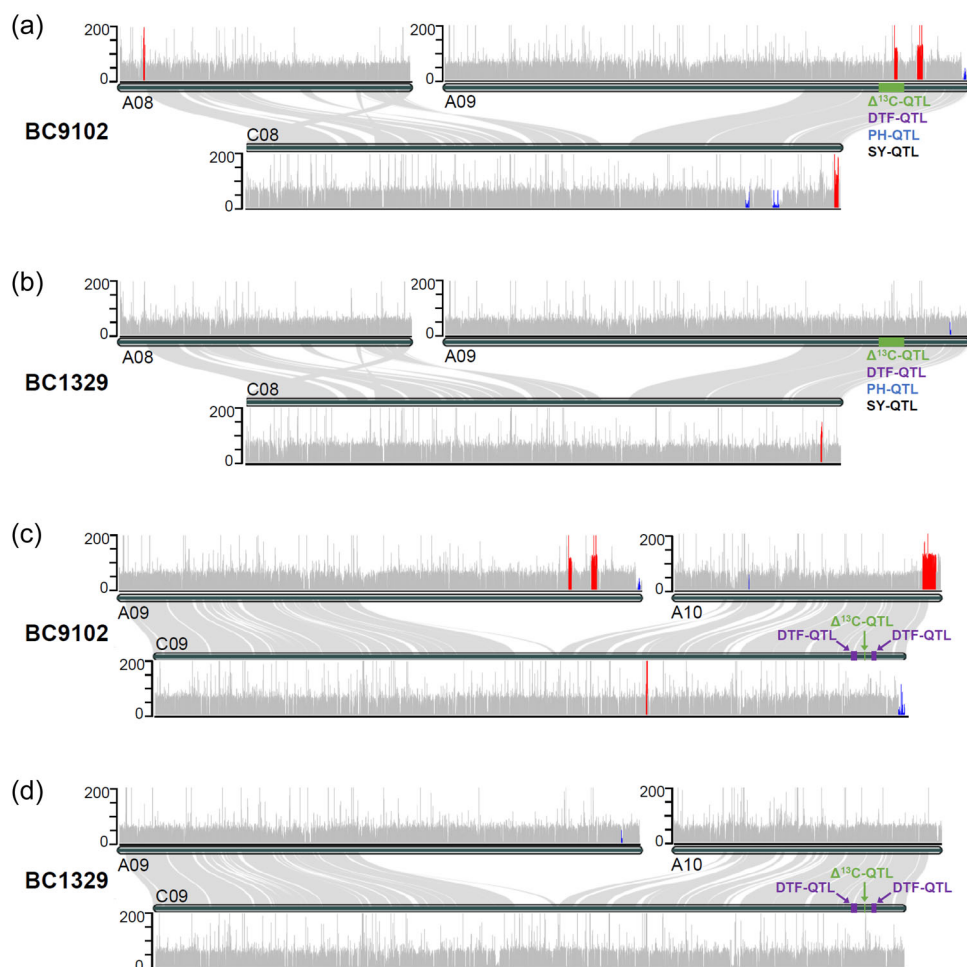


FIGURE 6 Homoeologous exchange (HE) events were detected between parental lines of the doubled haploid population derived from the BC1329/BC9102 Coverage depth (vertical axis) obtained for each chromosomes was calculated by mapping the Illumina resequencing reads to the reference genome of *B. napus* Darmor-bzh. The height of each line (a 10-kb window) from the horizontal axis represents the coverage depth. Genomic sequences (regions) that have undergone HE is shown in Supporting Information: Table S9. Substituted and “translocated” reads are highlighted in blue and red colour, respectively. DTF, days to flower; PH, plant height; QTL, quantitative trait loci; SY, seed yield.

populations (Figure 3c–h). DArTseq “bin” marker 3153720 on A09 revealed the complete linkage with another 18 markers in the DH population, which were localized within the 1.54 Mb region, spanning 28.28 to 29.82 Mb (Supporting Information: Tables S6 and S9). To identify potential candidate genes involved in the $\Delta^{13}\text{C}$ variation, we first excluded the syntenic genes in QTL regions and their corresponding homologs without any SNP and gene expression ($\log\text{FC} \geq 1.5$) variations between parental lines (Supporting Information: Table S10). In addition, we also excluded the genes in QTL regions with synonymous SNPs and InDels. We then focused on the remaining genes especially non-syntenic genes in QTL regions with different expression levels between parental lines. Finally, we identified a total of eight genes as candidates for $\Delta^{13}\text{C}$ variation at three QTL regions of chromosomes A08, A09, and C09 (Supporting Information: Table S10). Annotation of genomic intervals revealed that haloacid dehalogenase-like hydrolase superfamily protein (BnaA08g17740D) on chromosome A08/C03, myelin transcription factor-like protein (BnaA09g40350D), C3HC4-type RING finger protein (BnaA09g40860D), serine

carboxypeptidase-like 23 (BnaA09g41720D), poly(A) binding protein 4 (BnaA09g42030D), and cation-transporting P-type ATPase (BnaA09g42040D) and cytochrome P450 (BnaA09g42330D) on A09 and its homoeologous chromosome C08, and serine carboxypeptidase-like 19 (BnaC09g47000D) on C09/A10 chromosomes are likely candidates to be involved in $\Delta^{13}\text{C}$ variation (Supporting Information: Table S10, Supporting Information: Figure S4). The leaf tissue-specific transcriptome analyses revealed a total of 60 genes on A09 and 51 genes on C09 underlying $\Delta^{13}\text{C}$ QTL regions that were significantly differentially expressed between the two parental lines: BC1329 and BC9102 (Supporting Information: Table S11, Figure 7). Several of the DEGs, such as myelin transcription factor-like protein, casein kinase 2 $\alpha 4$, cation-transporting P-type ATPase, C3HC4-type RING finger protein, and serine carboxypeptidase, are related to water use, WUE, and response to water stress (<https://www.arabidopsis.org/>). We also found that the expression levels of genes in BC9102 (with HE event) such as BnaA09g41850D, BnaA09g41970D (wall-associated receptor kinase-like 14), BnaA09g41990D (cyclin-dependent kinase inhibitor),

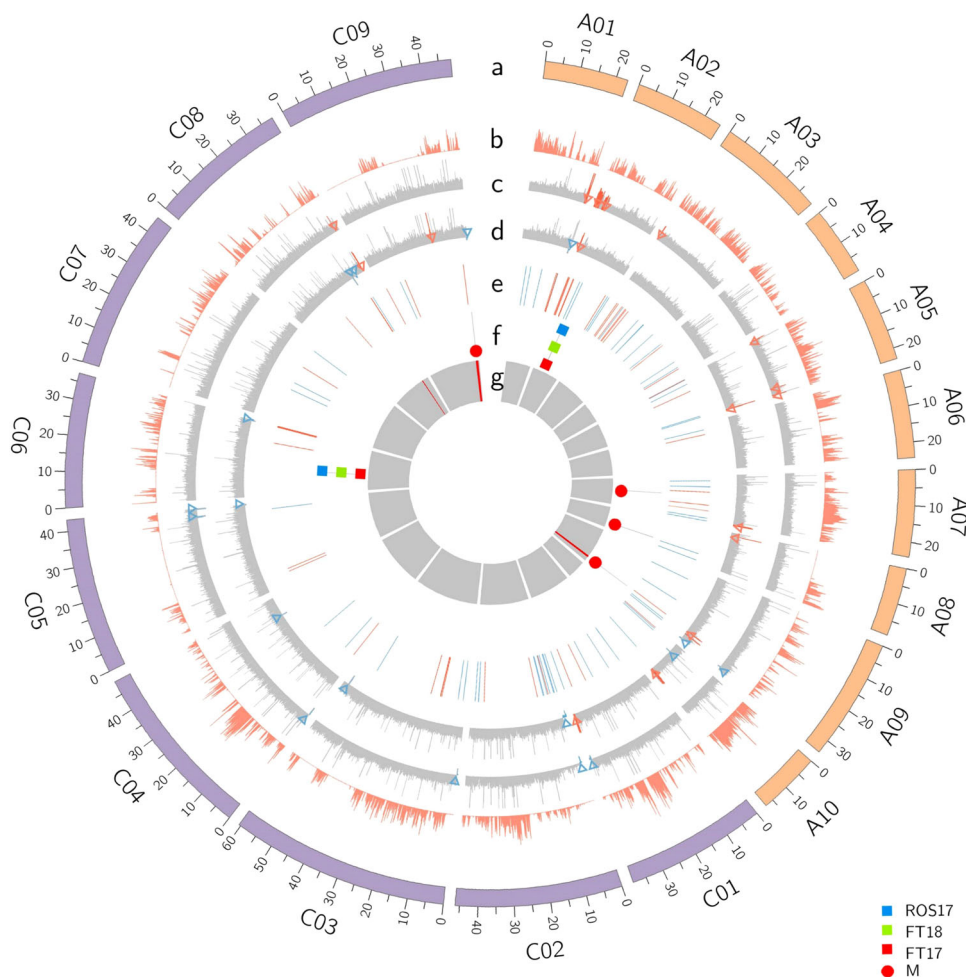


FIGURE 7 Physical map positions of QTL controlling $\Delta^{13}\text{C}$ variation in a doubled haploid population derived from the BC1329/BC9102. a: the outermost circle represents the physical length of the chromosomes (A genome in orange, C genome in purple), b: polymorphic SNPs between BC1329 and BC9102, c: homoeologous exchanges identified in BC1329, d: homoeologous exchanges identified in BC9102, e: DEGs identified between BC1329 and BC9102 (under dry and wet conditions), f: QTL associated with $\Delta^{13}\text{C}$ variation; blue, green and red, g: DEGs located within QTL regions. DEG, differentially expressed gene; QTL, quantitative trait loci; SNP, single nucleotide polymorphism.

BnaA09g42000D (nicotinate phosphoribosyltransferase 2), BnaA09g42030D (RNA recognition motif domain), and BnaA09g42040D were significantly higher (at least twofold) than those of BC1329 (without HE event) (Figure 8, Supporting Information: Table S11), suggesting that HE may be responsible for expression variation at the $\Delta^{13}\text{C}$ -QTL region on A09.

4 | DISCUSSION

Delineating the genetic and environmental determinants underlying variation in WUE traits is essential for the optimized breeding of high-yielding varieties in water-limited environments. In this study, we explored the genetic basis of variation in $\Delta^{13}\text{C}$ and related WUE traits in the biparental population derived from a cross between BC1902 and BC9102 lines that differ in $\Delta^{13}\text{C}$ and TE signatures (Hossain et al., 2020). We employed multi-environment-based QTL analysis which is considered a more powerful approach to dissecting

complex traits, controlled by G, E, and G $\times$ E interactions than the traditional QTL approaches (Zhang et al., 2010). We show that the growing environment plays an important role in the phenotypic expression of trait variation; thus, in the identification of environmental stable and environment-specific QTL effects. To reveal candidate genes underlying stable QTL for $\Delta^{13}\text{C}$, our work also identifies SV (HE), and DEGs between the parental lines of the mapping population, suggesting that homoeologous recombination may contribute to phenotypic variation for multiple traits at the chromosome A09 QTL.

4.1 | Canola reveals considerable variation for $\Delta^{13}\text{C}$

We found substantial genotypic variation in $\Delta^{13}\text{C}$, from 18.78% to 21.23% among DH, and 22.89% to 23.67% among F₂ lines. An earlier study has shown that an increase of 0.5‰ in $\delta^{13}\text{C}$ can lead to

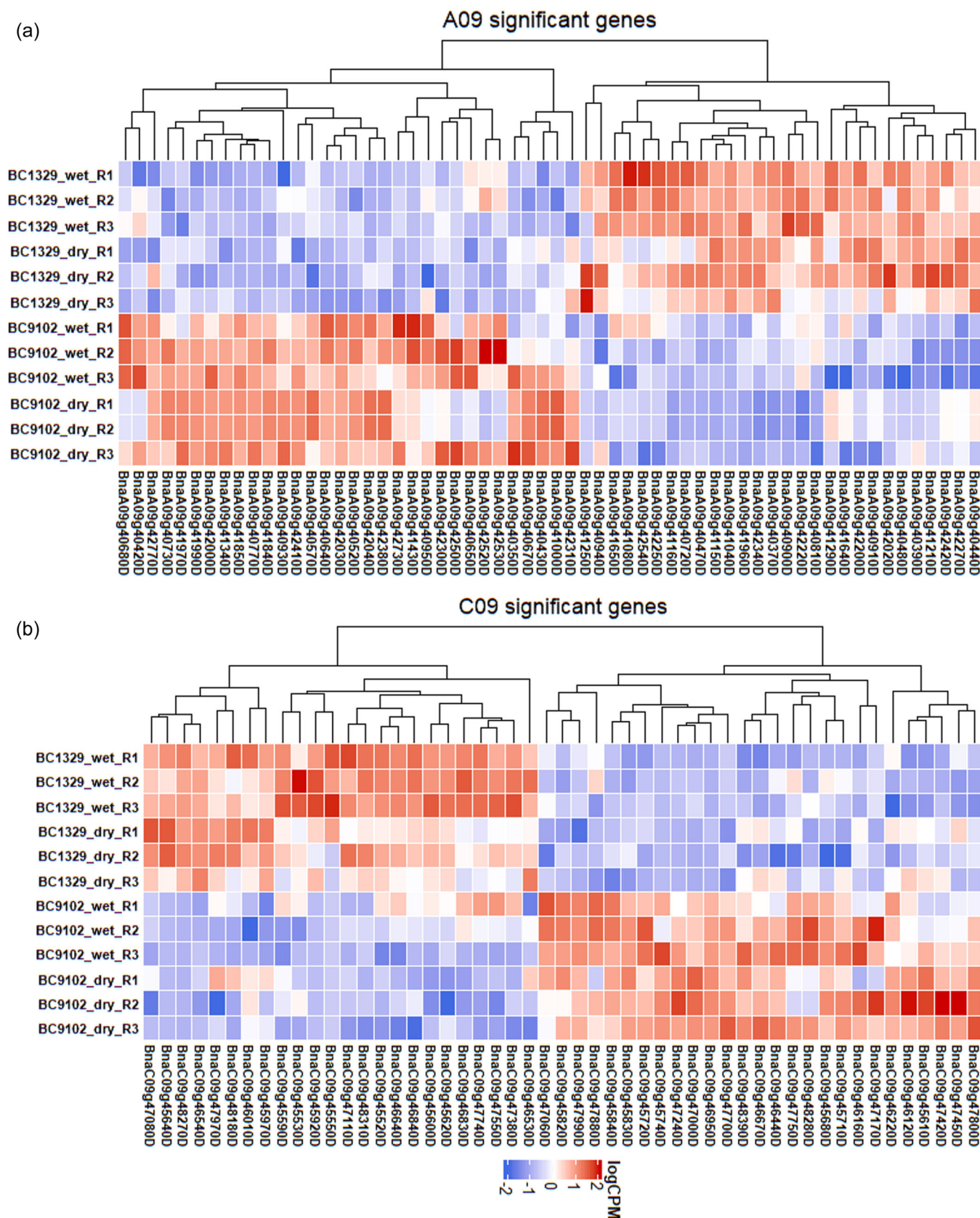


FIGURE 8 Expression profiles of differentially expressed genes (DEGs) in A09 (a) and C09 (b) QTL regions under water-deficit and water non-deficit conditions (Experiment 6) of the parental lines of the doubled haploid population derived from the BC1329/BC9102. The normalized read counts were plotted as a heatmap and genes were clustered according to the basis of their expression pattern. Detailed information on DEGs of parental lines (in triplicate: R1, R2, and R3) is presented in Supporting Information: Table S11. QTL, quantitative trait loci.

25% more TE in *Arabidopsis* (Juenger et al., 2005). Extrapolating this relationship, which is positive between $\Delta^{13}\text{C}$ and syWUE, and negative between $\Delta^{13}\text{C}$ and iWUE (Figure 2A), canola lines with 2.52‰ higher $\Delta^{13}\text{C}$ values than parental lines should reduce WUE theoretically by 126%. It reflects the dependence of the sensitivity on the general level of $\Delta^{13}\text{C}$. For example, Masle et al. (2005) found that at the level they saw in *Arabidopsis*, an increase in $\Delta^{13}\text{C}$ of 1‰ was associated with a 15% decrease in TE.

4.2 | Integrated WUE is partly driven by fitness traits

This study showed that DH lines that discriminate less between ^{12}C and ^{13}C as a carbon source for photosynthesis (low $\Delta^{13}\text{C}$) show higher iWUE at the single leaf level (Figure 2a). However, low $\Delta^{13}\text{C}$ lines did not produce high seed yield) reiterating that $\Delta^{13}\text{C}$ estimates TE at the leaf level which is not syWUE. Our results also suggest that selection for high iWUE at a single leaf level is useful for improving seed yield ($r=0.57$, Figure 2a), rather than using low $\Delta^{13}\text{C}$ as a surrogate trait for predicting high seed yield in canola, consistent with our earlier findings made in the Skipton/Ag-spectrum DH population as well as in diverse canola accessions (Raman, Raman, McVittie, et al., 2020). The positive relationship observed between $\Delta^{13}\text{C}$ and seed yield could be due to genotypic variation in WUE being driven by variation in soil water capture and water use rather than by variation in assimilation per unit of water applied (Blum, 2005; Kobata et al., 1996; Sinclair, 2018). $\Delta^{13}\text{C}$ estimates TE at the leaf level which is not syWUE. In this study, we did not establish the relationship between $\Delta^{13}\text{C}$ and total dry matter production. Canola plants often senesce their leaves before the crop reaches maturity, and, therefore, it was not possible to estimate WUE accurately without measuring leaf mass. WUE, being a multidimensional trait can also be driven by other “fitness” traits that reduce evapotranspiration rate and crop water use. For example, high $\Delta^{13}\text{C}$ lines with faster growth (NDVI, a proxy for plant vigour and plant height) could provide quicker canopy cover, which enables plants to reduce water loss from soil evaporation, thus increasing seed yield ($r=0.63$ to 0.81 , Figure 1b). This is partly supported by this study showing a high correlation between plant fitness and seed yield and tight linkage of corresponding QTL (Figures 1–3). In addition, $\Delta^{13}\text{C}$ exhibited negative correlations with flowering time ($r=-0.53$; DH population), and a positive correlation with NDVI, plant height, and seed yield (Figure 1b) suggesting that high $\Delta^{13}\text{C}$ lines tend to “escape” via accelerating growth and flowering—an evolutionary trait for adaptation to terminal drought stress. Our results showed that genotypes with low $\Delta^{13}\text{C}$ had less canopy cover, late flowering, and lower seed yield; these characteristics are typical for plants with a drought avoidance strategy. However, under terminal water-deficit situations, low $\Delta^{13}\text{C}$ lines could yield poorly due to the shorter seed filling period, accompanied by high temperatures. It remains to establish how low $\Delta^{13}\text{C}$ lines which require a longer season for seed filling, perform in climates that are not prone to environmental constraints

(non-water deficit/heat stress). Research findings suggest that canola breeding programmes should focus on adaptative traits such as early flowering, and drought avoidance traits such as early vigour for effective water use for achieving high yield under water-limited conditions.

4.3 | Genetic and environmental determinants affect phenotypic trait expression

Our comprehensive multienvironment QTL analysis showed that by using well-designed multiphase experiments across three environments (Experiments 1–3, Table 1) and accounting for sources of variation in field/pot and laboratory conditions (Supporting Information: Method S1), both genetic and environmental determinants underpinning phenotypic variation can be deciphered for traits of interest (Supporting Information: Table S7). For example, we substantiated phenotypic plasticity for $\Delta^{13}\text{C}$ /WUE-related traits (Figure 4) and identified QTL for the main effects and $Q \times E$ interaction effects underlying variation in $\Delta^{13}\text{C}$, days to flower, seed yield, and plant height under field conditions (Supporting Information: Table S7). In addition, the magnitude and inconsistent direction of allelic effects also hint that observed variation in $\Delta^{13}\text{C}$, days to flower, seed yield, and plant height across environments is due to $Q \times E$ interaction. A substantial proportion of QTL (51.7%, 15/29) that exhibited $Q \times E$ interaction, suggests their contribution to phenotypic trait variation across different environments (Supporting Information: Table S7). Similar estimates for $Q \times E$ interaction effects are reported in other studies (Borevitz et al., 2002; El-Soda et al., 2014; Ungerer et al., 2003).

Consistent detection of $\Delta^{13}\text{C}$ -QTL (QTL main effects on A07, A08, A09, and C09) across three environments suggests that these loci contribute to the adaptive capacity of DH lines from BC9102/BC1902) to water-deficit stress conditions and thus translate to economic seed yield (~1 t/ha). Across field environments, DH lines were subjected to water-deficit conditions, right from stem elongation to seed maturity (rainfall ranged from 225 to 235 mm over 7 months of the growing season, Supporting Information: Figure S1). Colocation of QTL for seed yield, $\Delta^{13}\text{C}$, days to flower and plant height at the same genomic regions, and stable allele (BC9102), contributing to trait variation suggest multiple trait QTL on chromosome A09 is associated with effective water use. Early flowering showed a negative relationship with seed yield (Figure 1b), reiterating the crosstalk between drought stress signalling and flowering time pathways (Des Marais et al., 2012).

It was interesting that none of the $\Delta^{13}\text{C}$ QTL that we identified for the main effect and $Q \times E$ interactions (Supporting Information: Table S7) was detected in the Skipton/Ag-spectrum population (Raman, Raman, McVittie, et al., 2020). In an independent study, Mekonnen et al. (2020) identified three QTL for $\delta^{13}\text{C}$ on chromosomes A02, A09, and C08 in the North American *B. napus* mapping population. However, none of the QTL was consistently detected across environments. It is yet to establish whether the

genomic region on chromosome A09 or its homoeologous counterpart C08 (QTL for root pulling force, plant height, and $\delta^{13}\text{C}$) is the same as found in our study, as the authors did not report the physical positions of QTL marker intervals. In addition, there was a poor marker coverage on chromosome C08 in our genetic mapping population due to low recombination frequency (13 markers, Supporting Information: Table S6c), which may have led to QTL (if any) being undetected in the unmapped regions, especially in the HE region. Nevertheless, these studies suggest that several genomic regions on A02, A03, A07, A09, C03, C06, C08, and C09 control variation in $\Delta^{13}\text{C}$; thus, the genetic architecture of $\Delta^{13}\text{C}$ is rather complex.

4.4 | A priori genes regulating WUE and efficient water use underlie QTL for $\Delta^{13}\text{C}$

Genetic and genomic approaches utilized herein facilitated the identification (in a DH population from BC1902/BC9102) and validation (in an F_2 population from BC1902/BC9102) of genomic regions for $\Delta^{13}\text{C}$ variation and delimited candidate genes in canola, which are implicated in leaf-level WUE (Cutler et al., 2010; Hersen et al., 2008; Menéndez et al., 2019; Tao et al., 2018; Youn et al., 2016). For example, this study identified a QTL that influences multiple traits; $\Delta^{13}\text{C}$, days to flower, plant height, and seed yield on chromosome A09 that map within 92 kb of the *ERECTA* gene (see Masle et al., 2005) (Supporting Information: Table S12). We also localized several stress-responsive genes, including DEGs that may contribute to drought avoidance strategies via signal transduction pathways, encoding functional proteins (LEA18, RD20, glycine metabolism, and CAT) and regulatory proteins, including transcription factors (bHLH, MYB, TINY2, ATHB6), protein kinases (tyrosine-protein kinase, wall-associated receptor kinase-like 14, MAPK, SNF1-related protein kinase), and receptors (ABA receptor PYL12), phosphatases (PP2C), and calmodulins (CPK17) (Des Marais et al., 2014; Jagodzick et al., 2018; Jonak et al., 2002; Yong et al., 2019) within QTL intervals associated with $\Delta^{13}\text{C}$ variation (Table 1, Supporting Information: Tables S11 and S12). Plant expressing *PYL12*, *SRK2C*, and *AtTPPF* genes are shown to improve the water use and drought tolerance (Lin et al., 2019; Yang et al., 2016) whereas ABC transporter (*ABCG22*) and ABA-responsive kinase gene, *MPK12* reduced the WUE (Des Marais et al., 2014). Our data hint that genes affecting stomatal characteristics (*RD20*, *ERECTA*), leaf thickness, and water-deficit responsive genes described above likely underlie WUE and drought avoidance traits, while $Q \times E$ interactions are likely driven by environmental cues such as water stress and temperature (phytochrome C was mapped with 6.2 kb from $\Delta^{13}\text{C}$ -QTL on C06, Supporting Information: Table S12). Our results suggest that a QTL region underlying $\Delta^{13}\text{C}$, flowering time, plant height, and seed yield on chromosome A09 may be subjected to HE. Homoeologous recombination is associated with presence-absence variation (Hurgobin et al., 2018; Nicolas et al., 2007). Recently, a major QTL for homoeologous recombination, *BnaPh1* was mapped on A09

(Higgins et al., 2021) and this was located within 5 Mbp of the QTL region that is associated with multiple traits. The same genomic region may be involved in regulating WUE in diverse canola accessions and require further research.

The most pressing need for research is to clarify the bases of the observation of a positive relation between seed yield and leaf $\Delta^{13}\text{C}$, despite the negative relation between shoot growth and leaf $\Delta^{13}\text{C}$. Condon et al. (1987) reported the same phenomenon in wheat and discussed some possible biophysical explanations, including the idea that the major source of variation might be stomatal conductance rather than photosynthetic capacity. The negative association between time to flowering and $\Delta^{13}\text{C}$ may also have contributed, but the most revealing observation (Condon & Richards, 1993) was that wheat crop yields were positively correlated with $\Delta^{13}\text{C}$ in wet environments, but negatively correlated in dry environments, with no association under intermediate levels of water limitation.

In summary, this current study demonstrates that measures of iWUE, $\Delta^{13}\text{C}$, and integrated WUE (dmWUE and syWUE) are complex and modulated by genetic and environmental determinants, including that subject to HE. Our findings identify useful variation in $\Delta^{13}\text{C}$ (up to 2.52‰) and its underlying genetic basis of variation in WUE traits, including their plasticity across the natural field and rainout shelter environments. Identification of favourable alleles for increasing WUE would provide potential resources for developing new drought-tolerant varieties for drier environments to continue making genetic gains in the breeding programmes via genomic-assisted selection strategies. Resources developed in this study could also enable us to understand molecular mechanisms underlying phenotypic plasticity and $G \times E$ interactions.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

The raw sequence data reported in this paper have been deposited in the National Center for Biotechnology Information Sequence Read Archive (Accession No. PRJNA743730 for RNA-sequencing data, PRJNA743989 for whole-genome resequencing data).

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