

# Genetic dissection of photosynthetic performances in maize under drought-stressed and well-watered environments

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## Research article

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# Abstract

## Background

Maintaining photosynthetic capacity is a critical function that allows maize (*Zea mays* L.) to adapt to drought stress. The elucidation of genetic controls of photosynthetic performances, and tightly linked molecular markers under water stress are thus of great importance in marker-assisted selection (MAS) breeding. Meanwhile, little is known regarding their genetic controls under drought stress. Two F<sub>4</sub> populations were developed to identify quantitative trait loci (QTLs) and dissect the genetic variation underlying six photosynthetic-related traits, namely, net photosynthetic rate (Pn), stomatal conductance (Gs), intercellular CO<sub>2</sub> concentration (Ci), transpiration rate (Tr), ribulose 1,5-biphosphate carboxylase activity (RuBP), and water use efficiency (WUE) under drought-stressed and well-watered environments.

## Results

For two populations, we detected 54 QTLs under drought-stressed and well-watered environments by single-environment mapping with composite interval mapping (CIM), approximately 81.8~100 % QTLs displayed non-additive effects, and 43 of the 54 QTLs were identified under drought-stressed environment. We also dissected 54 QTLs via joint analysis of all environments with mixed-linear-model-based composite interval mapping (MCIM), 24 QTLs involved in QTL × environment interactions (QELs), approximately 87.5 % QELs were identified under drought-stressed environments, as well as 14 pair epistasis exhibited dominance-by-additive/dominance (DA/DD) effects under constraining environments. We further identified 8 constitutive QTLs (cQTLs) across two populations by CIM/MCIM under multiple environments. Remarkably, bin 1.07\_1.10 (cQTL2), bin 6.05 (cQTL5), bin 7.02\_7.04 (cQTL6), bin 8.03 (cQTL7), and bin 10.03 (cQTL8) exhibited 5 pleiotropic cQTLs that were consistent with phenotypic correlations among all photosynthetic-related traits. Additionally, 17 candidate genes were validated in above cQTLs.

## Conclusions

Photosynthetic performances in maize were predominantly controlled by non-additive and QELs effects, where more QELs effects occurred in drought stress. 8 cQTLs affecting six photosynthetic-related traits could be useful for genetic improvement of these traits via QTL pyramiding, corresponding 5 QTLs clusters indicated tight linkage or pleiotropy in the inheritance of these traits, and 17 candidate genes involved in leaf morphology and development, photosynthesis, and stress reponse coincided with above corresponding cQTLs.

## Background

Water stress is one of the most important environmental limiting factors for maize (*Zea mays* L.) productivity in tropical and subtropical regimes [1], and the global climate change scenario and growing population exerts great pressure tend to increase the problems of food insecurity. Therefore, the

improvement of resistance to water scarcity and breeding drought tolerant varieties are crucial for maize survival, growth, and biomass production living in water scarce environments [2-3]. Maize as an important  $C_4$  crop, and its highly complex mechanism of photosynthetic performance is one of the main targets for improving maize grain yield (GY) and drought resistance. He et al. [4], Liu et al. [5] and Zhao et al. [6] reported that net photosynthetic rate (Pn), chlorophyll relative content (SPAD), chlorophyll a content (FCa), chlorophyll b content (FCb), total chlorophyll content (FCT), ribulose 1,5-bisphosphate carboxylase activity (RuBP), stomatal conductance (Gs), and transpiration rate (Tr) in maize were significantly reduced, meanwhile chlorophyll a/b (FCa/b), intercellular  $CO_2$  concentration (Ci) and water use efficiency (WUE) were significantly increased under drought stress, and compared with drought sensitive materials, strong drought resistant maize could maintain higher photosynthetic capacity in drought land. Zhang et al. [7] suggested moderate and severe drought stress caused an obvious decrease in Pn, Gs, and RuBP, caused a significant increase in Ci, damaged photosystem II (PSII), reduced electron transport in diverse maize varieties. Tezara et al. [8] and Liu et al. [9] also analyzed the activities of photosystem I (PSI), PSII, and photosynthetic electron transport chain (PETC) between PSII and PSI in response to drought stress in maize, and of which showed that a significant increase in absorption of antenna chlorophyll per PSII reaction center (ABS/RC) was found, a significant decrease in quantum yield for electron transport ( $\phi E_0$ ), efficiency of an electron beyond primary quinone acceptor of PSII ( $Q_A$ ), efficiency of an electron beyond  $Q_A$  ( $\Psi_0$ ), and performance index for energy conservation from photons absorbed by PSII to the reduction of intersystem electron acceptors ( $PI_{ABS}$ ) were observed, and no significant change in the fast 820 nm modulated reflection (MR) phase and the amplitude of delayed fluorescence under drought stress. In this regard, once photosynthesis performance is inhibited under drought environments, it is likely to result in maize plant absorbing more light energy than can be consumed through photosynthetic carbon fixation, and even damage photo-oxidation, finally cause more GY losses.

Using identified molecular markers and quantitative trait loci (QTLs) for photosynthetic-related traits under different watering treatments in marker-assisted selection (MAS) breeding is a promising way to maximize the productivity of maize grown in drought land. Until now, genetic studies on wheat (*Triticum aestivum* L.) [10], rice (*Oryza sativa* L.) [11-13], sunflower (*Helianthus annuus* L.) [14], rapeseed (*Brassica napus* L.) [15], cotton (*Gossypium* spp) [16-17] etc. for photosynthetic-related traits have been extensively conducted using QTL mapping during drought, high temperature, salt and alkali stresses. In maize, however, only several experiments have been conducted for mapping QTLs for photosynthetic-related traits, e.g., Wang and Zhang [18] mapped 32 QTLs for FCa, FCb, FCT, and other chlorophyll content (FCc) in one  $F_2$  population derived from A150-3-2  $\times$  Mo17. Trachsel et al. [19] found 7 QTLs related to quantum efficiency of photosystem II ( $\Phi_{PSII}$ ), and SPAD in the intermated CML444  $\times$  SC-Malawi recombinant inbred lines (RILs) populations. Yu et al. [20] detected 32 QTLs associated with FCa, FCb, FCT, Pn, Gs, Ci, and Tr in two  $F_2$  populations derived from Y114  $\times$  Y115 and Y105  $\times$  Y106. In addition, only two mapping studies for photosynthetic performance in maize have been reported under drought environments, e.g., Pelleschi et al. [21] identified 19 major QTLs controlling net  $CO_2$ -uptake (CO), ADPglucose pyrophosphorylase (AGP), Gs, and Tr in a RILs population derived from the cross F-2  $\times$  MBS847 under drought-stressed and well-watered regimes. Prado et al. [22] also assessed 16 robust QTLs

affecting Gs with a diversity panel of maize hybrids under water deficit. Even the photosynthetic-related *ZmPPDK*, i.e., *pyruvate, orthophosphate dikinase 1* gene, the transgenic wheat [23], potato (*Solanum tuberosum*) [24], and *Arabidopsis thaliana* [25] plants with *ZmPPDK* could significantly improve their photosynthetic efficiency and GY. Despite these researches, the molecular mechanisms for photosynthetic-related trait in response to drought remains poorly understood. Thus, unraveling in-depth investigations on the genetic mechanisms controlling for photosynthetic performances in contrasting watering treatments, which may be used to speculate the possible genetic locations of corresponding candidate genes. Moreover, a better understanding of the genotype × environment interaction (GEI) will provide a foundation for the genetic improvement and optimization of genotypes across different environments [26].

The overall objective of this study were to identify QTLs responsible for Pn, Gs, Ci, Tr, RuBP, and WUE in two maize F<sub>4</sub> populations subjected to both watering regimes as well as estimate their effects by single environment mapping with composite interval mapping (CIM), then to further dissect joint QTLs, QTL by QTL interaction (epistasis), as well as QTL × environment interactions (QEIs) among all watering environments via mixed-linear-model based composite interval mapping (MCIM). By combining these experiments, we hope to identify hot spots, areas of co-localization experiments, as well as corresponding candidate genes, which may further contribute to mapping QTLs, revealing genetic mechanisms for photosynthetic performances under diverse watering environments, and developing maize MAS breeding for photosynthetic performances to improve drought tolerance and GY in the future.

## Methods

### Plant materials

The two F<sub>4</sub> mapping populations included 218 and 202 families (POP-CT and POP-LT) derived from Chang7-2 × TS141 (CH × TS) and Langhuang × TS141 (LH × TS) hybrids. The corresponding two F<sub>2</sub> families derived from the corresponding crosses self-pollinated to generate F<sub>3</sub> families at Pingliang maize breeding station of Gansu Agricultural University, China (106.93°N, 35.43°E; 1,204 m altitude; loessial soil) in 2013, and then each F<sub>3</sub> plant was again selfed to derive the corresponding F<sub>4</sub> families at Zhangye breeding base of Yuyuan Co., LTD., China (38.83°N, 106.93°E; 1,785 m, altitude; sandloam) in 2015 and Jingtai farm of Tiaoshan Nongken corporation, China (37.18°N, 104.03°E; 1,640 m, altitude; sandloam) in 2015, respectively [27]. In addition, Chang7-2 (drought-tolerant line), Langhuang (drought-tolerant line), and TS141 (drought-sensitive line) are foundation inbred lines in China maize breeding and are representatives of Tangsipingtou (TSPT), TSPT, and Reid yellow dent (Reid) heterotic groups, respectively [28-29].

### Field experiment and trait evaluation

The phenotype data of the two F<sub>4</sub> families (POP-CT and POP-LT) and their parents (CH, LT, and TS) were evaluated in a completely randomized block design with two replications and 10 plants at a density of

55,580 plant ha<sup>-1</sup> for each plot at Huangyang, Gansu Academy of Agricultural Sciences Proving Ground, China ([http://hyc.gsagr.ac.cn/channels/channel\\_503\\_1.html](http://hyc.gsagr.ac.cn/channels/channel_503_1.html)) (37.67°N, 102.85°E; 1,740 m, altitude; sandloam) in 2019. The mean temperature, total sunshine duration, total rainfall, total evaporation capacity, mean relative humidity, and mean wind velocity at the experimental site was 17.9 °C, 1,427.6 h, 118.7 mm, 1,039.7 mm, 52.3 %, and 2.2 m·s<sup>-1</sup> during the growing seasons (from April to September) in 2019, respectively (Additional file 1: Fig. S1). Then the experimental field was divided into well-watered (WW) and drought-stressed (DS) treatments. The DS treatment was equivalent to rainfed conditions with rainfall of 118.7 mm during the growing season (from April to September). The WW treatment involved irrigation with 4,500 m<sup>3</sup>·ha<sup>-1</sup> water supply at each of V18, R1, and R3 stage. In addition, because of the abundant annual evaporation capacity (2,164.7 mm), low annual relative humidity (45.8 %), scarce annual rainfall (172 mm), and low annual mean temperature (9.3 °C) in this experimental site (Additional file 1: Fig. S1), plastic film (0.02 mm thick, 140 cm wide) was laid out by land over the fields and covered the soil surface before sowing.

Until spreading pollen stage, the six photosynthetic-related traits, i.e., Pn (μmol CO<sub>2</sub>·m<sup>-2</sup>·s<sup>-1</sup>), Gs (mol H<sub>2</sub>O·m<sup>-2</sup>·s<sup>-1</sup>), Ci (μmol CO<sub>2</sub>·mol<sup>-1</sup>), Tr (mol H<sub>2</sub>O·m<sup>-2</sup>·s<sup>-1</sup>), RuBP (mol·m<sup>-2</sup>·s<sup>-1</sup>), and WUE (μmol CO<sub>2</sub>/mol H<sub>2</sub>O) were measured in the corresponding populations and parents, using a portable photosynthesis system, namely LI-6400 XT (LI-COR Inc. Lincoln, Nebraska, USA). Specifically, Pn, Gs, Ci, and Tr were measured at a light intensity of 1,500 μmol m<sup>-2</sup> s<sup>-1</sup> photosynthetic active radiation (PAR), a leaf temperature of 30 °C and a constant CO<sub>2</sub> concentration of 380±5 μmol CO<sub>2</sub> mol<sup>-1</sup> in the sample chamber provided with buffer volume [13]. All measurements for each treatment were made on the fully expanded leaves between 9:00 and 11:00 a.m. on sunny days to avoid effects of photo-inhibition and were repeated at least 5 times using different plants. RuBP was estimated in response to apparent mesophyll conductance (AMC) as follows [3]: (1). And WUE was also calculated as follows [10, 17]: (2). Then the corresponding F<sub>4</sub> plants were harvested and subsequently air-dried to evaluate ear weight (EW), grain weight per ear (GW), and 100-kernel weight (KW). Relative to the WW treatment, the average rate of change (RC) of each trait under DS treatment was also estimated according to Zhao et al. [28] as follows: (3). Here, was the average value of each trait under DS treatment, and was the average value of corresponding trait under WW treatment.

### Statistical analysis of phenotypic data

The all phenotypic data obtained from two F<sub>4</sub> populations and three parents under each watering environment were statistically analyzed using IBM-SPSS Statistics v. 19.0 (SPSS Inc., Chicago, IL, USA) (<https://www.ibm.com/products/spss-statistics>). The significance of total and residual variances of each trait in two F<sub>4</sub> populations were estimated by the general linear model for univariate (GLM-Univariate) with one-way analysis of variance (ANOVA), respectively. The broad-sense heritability ( ) and GEI heritability ( ) for combined environments (i.e., WW and DS) were estimated as follows [30]: /nr (4) and /nr (5). Here, was genotypic variance, was environmental variance, was error variance, was GEI variance, n (n = 2) was the No. of environments, and r was the No. of replications for the experiment (r = 10). Possible

associations between corresponding traits were tested using several methods, based either on phenotypic Pearson correlation or on principal component Analysis (PCA) via IBM-SPSS Statistics v. 19.0 [21].

### **Genetic map construction and QTLs identification**

A total of 205 and 199 genome-wide polymorphic simple sequence repeats (SSRs) obtained from the Maize Genetics and Genomics Database (MaizeGDB, <http://www.maizegdb.org/>) were applied to develop two F<sub>2</sub> populations genetic maps, using JionMap v. 4.0 (<https://www.kyazma.nl/index.php/JoinMap/>), and the total maps length were 1,648.8 and 1,542.5 cM, with an average interval of 8.0 and 7.8 cM, respectively [27-29].

In a single watering environment, QTL mapping for each photosynthetic-related trait was performed using CIM implemented with Windows QTL Cartographer software v. 2.5 (<http://statgen.ncsu.edu/qtlcart/winqtlcart.htm>). For CIM, model 6 of the Zmapqtl module was used to analyze QTLs. Window size was 10 cM, and cofactors were selected through forward and backward regressions with the in and out thresholds at a  $P < 0.05$ . A genome-wide critical threshold value was estimated for experiment-wise type I error rate of 0.05 with 1,000 random permutations [31]. The identified QTLs effects were estimated according to criteria suggested by Stuber et al. [32] as follows: |dominance/additive| (d/a); additive (A), |d/a| = 0.00~0.20; partial dominance (PD), |d/a| = 0.21~0.80; dominance (D), |d/a| = 0.81~1.20; over-dominance (OD), |d/a| > 1.20.

Additionally, among all watering environments, the MCIM via QTL Network v. 2.0 (<http://ibi.zju.edu.cn/software/qtlnetwork>) was used to dissect joint QTLs, epistatic QTLs, and QEI for each photosynthetic-related trait based on the all watering environments. The testing window, walk speed, and filtration window of the genome scan were set at 10, 2, and 10 cM, respectively. A total of 1,000 permutations were performed to determine the threshold logarithm (base 10) of odds ratio (LOD), for the traits for declaring a significant QTL at a  $P < 0.05$  probability level. The name of QTL was assigned according to the modifying nomenclature of Zhao et al. [29]. Furthermore, the letter “J” was added into the middle of the QTL name (i.e., inserted between trait abbreviation and No. of chromosome), whereas one QTL was only detected in joint analysis with MCIM but not in a single environment through the CIM program. The LOD confidence interval (CI) of the QTL were estimated according to Zhao et al. [26] as follows: (6). Here, N was the population size, was the value of the phenotypic variation contributed by the QTL. Additionally, the photosynthetic-related traits QTLs were identified within same marker interval or within overlapping Cis, the corresponding loci were assumed to be common QTLs with pleiotropic effects.

### **Constitutive QTLs (cQTLs) detection and candidate genes dissection**

cQTLs refer to the QTL stably and repeatedly detected with CIM/MCIM across different mapping populations under two or more watering environments [33]. Then the corresponding cQTLs intervals were projected on the physical reference map B73 RefGen\_v4 ([https://maizegdb.org/gbrowse/maize\\_v4](https://maizegdb.org/gbrowse/maize_v4)), and the corresponding candidate genes were further validated in corresponding cQTL regimes [34], and corresponding genes functions of which even were analyzed via the AgBase v. 2.00

(<http://agbase.arizona.edu/>) online software and public databases, namely, National Center for Biotechnology Information (NCBI, <http://www.ncbi.nlm.nih.gov/pubmed>), MaizeGDB (<http://www.maizegdb.org/>), and China National Knowledge Infrastructure (CNKI, <http://www.cnki.net>).

## Results

### Photosynthetic performances variations under drought-stressed environments

We analyzed the six tested photosynthetic performance values across three parents and two  $F_4$  populations under both experimental watering treatments. There were significant difference at  $P < 0.05$  or  $P < 0.01$  level on six photosynthetic-related traits in three parents under well-watered and drought-stressed environments (Additional file 2: Table S1). Compared with well-watered environment, the Pn, Gs, Tr, and RuBP showed significant decreases in Chang7-2, Langhuang, TS141, POP-CT, and POP-LT, Ci and WUE, however, displayed significant increases under drought-stressed environments (Fig. 1). Further analysis indicated that drought-sensitive line TS141 had larger RC in Pn, Gs, Ci, and RuBP, but had less RC in Tr compared to drought-tolerant lines Chang7-2 and Langhuang, and even the average RC of the three parents and two  $F_4$  populations in Pn, Gs, Ci, Tr, RuBP, and WUE were 16.02, 17.05, -9.34, 10.80, 18.96, and -8.62 %, respectively (Fig. 1). These phenotypic analyses demonstrated that the drought-induced limitation of photosynthesis in maize was primarily due to  $CO_2$  diffusion efficiency from sub-stomatal interval internal cavities to carboxylation site in chloroplasts and degree of stomatal closure (SC), therefore, the RuBP and Gs decreased remarkably in different maize materials under drought stress. Moreover, except for RuBP, other measured photosynthetic performance values in two  $F_4$  populations were continuously distributed, with absolute values of skewness and kurtosis being less than 1.0 (Additional file 2: Table S1, Additional file 3: Fig. S2), indicative of continuous variation and a quantitative genetic basis in these maize photosynthetic performances. For all photosynthetic performances in two  $F_4$  populations, results of ANOVA showed that there were significant variation among genotypes (Table 1), which further allowed their genetic dissection under drought stress. The environmental and GEI variances were also significant ( $P < 0.01$  or  $P < 0.05$ ) (Table 1), which implied that the populations performed quite inconsistently between the both watering environments. Additionally, the estimated and values of the all photosynthetic performance in two populations were 46.429 (Gs in POP-CT)~91.030 % (Ci in POP-LT)/4.531 (Ci in POP-LT)~19.770 % (Tr in POP-CT) because of the large GEI contribution to phenotypic variances (Table 1).

### Phenotypic framework of photosynthetic performances

Examination of phenotypic correlations between photosynthetic-related traits may be useful before interpreting the co-locations between QTLs which were more likely to reveal genetic relationships. PCA and Pearson correlations were performed in two populations and three parents under both watering environments. PCA, under well-watered and drought-stressed environment, displayed two significant PCs (PC1 and PC2) with eigenvalues greater than 1 were extracted that together explained 65.032 % and 75.602 % of the variance, respectively (Fig. 2). These PCs were linear combinations of the original photosynthetic-related traits that were independent of each other, and represented different combinations

of the traits based on their variable loadings under different watering environments. Among them, PC1 primarily represented variances in Pn, Gs, and Tr under both contrasting environments. PC2, however, mainly captured variance in RuBP or Ci under WW/DS environment (Fig. 2). Further analysis showed that pairwise Pearson correlations for six photosynthetic performances and three yield components yielded complementary information under both watering environments, and each corresponding trait positively/negatively ( $P < 0.01$  or  $P < 0.05$ ) correlating with four to eight other traits under single watering environment (Fig. 3), thus indicated that yield formation in maize was the result of the synergistic or inhibited effects of multiple photosynthetic performances under well-watered and drought-stressed environments, and the influence degree of drought stress to drought-sensitive maize plants photosynthesis and yield were larger. Moreover, Pn, Gs, Ci, RuBP and WUE of the two  $F_4$  progeny populations depicted significantly positive correlation to female parent ( $P < 0.05$  or  $P < 0.01$ ), respectively, GW and KW were significantly positive correlated to male parent ( $P < 0.05$  or  $P < 0.01$ ), as well as Tr and CW showed significantly positive correlation to both parents ( $P < 0.05$  or  $P < 0.01$ ) (Additional file 4: Table S2), indicative that the effects of both parents on different photosynthetic performances and yield components in  $F_4$  progeny populations were inconsistent.

### **QTL analysis for photosynthetic performances under single watering environment**

To dissect the genetic control underlying six corresponding photosynthetic performances using single environment mapping with CIM, we totally mapped 54 QTLs (13 for Pn, 6 for Gs, 10 for Ci, 9 for Tr, 10 for RuBP, and 6 for WUE) across two  $F_4$  populations (POP-CT and POP-LT) based on both watering environments (WW and DS), and the phenotypic variance explained by individual QTL ranged from 2.37 (for RuBP in POP-LT under DS) to 18.21 % (for Ci in POP-CT under DS) within each watering environment (Additional file 5: Fig. S3, Additional file 6: Table S3, Fig. 4). A total of 43 (79.63 %) of the identified QTLs affecting six photosynthetic-related traits were detected under DS environments (Additional file 7: Fig. S4). For these identified QTLs, Pn, Gs, and RuBP displayed both additive (A) and non-additive (including PD, D, and OD) effects, however, QTLs for Ci, Tr, and WUE only showed no-additive effects (Additional file 5: Fig. S3). Moreover, approximately 33.33, 18.18, 37.50, 43.75, 35.29, and 44.44 % alleles for increase in Pn, Gs, Ci, Tr, RuBP, and WUE were contributed by male parent TS141, respectively (Additional file 6: Table S3).

### **Joint analysis and QEIs for photosynthetic performances in multiple watering environments**

The joint QTL analysis of all watering environments with MCIM revealed 54 QTLs controlling six photosynthetic-related traits in POP-CT and POP-LT, being absolutely equal to detected No. of QTLs based on single environment, and 25 of the identified QTLs were consistent with those of Pn, Gs, Ci, Tr, RuBP, and WUE through single environment mapping with CIM. These identified QTLs explained 2.07 (for Ci in POP-CT) to 13.21 % (for Pn in POP-LT) phenotypic variance contributed by  $h^2(A)$  (Fig. 4, Additional file 8: Table S4). Moreover, 24 QTLs (6 for Pn, 3 for Gs, 6 for Ci, 3 for Tr, 4 for RuBP, and 2 for WUE) were involved in QEI in two  $F_4$  populations, and accounted for 1.34 (for Ci in POP-LT) ~6.86 % (for Pn in POP-LT) of the phenotypic variance explained by  $h^2(AE)$  (Additional file 8: Table S4). These QEI may thus impart stronger effects on photosynthetic performances in maize under contrasting environments. In addition, for the two

F<sub>4</sub> populations, totally 7 stable bin intervals may be mediated by environmental factors, i.e., bin 1.08\_1.10 (mmc0041-phi308707) exhibited a constitutive QEI involved in Pn, Ci, and RuBP, bin 3.07\_3.08 (umc1286/umc2275-umc2081) mapped a constitutive QEI for Gs, bin 6.05 (umc2040-bnl1174a) found a constitutive QEI associated with Ci and RuBP, bin 7.00 (umc2177\_umc1378) detected a constitutive QEI controlling Ci and RuBP, bin 7.02\_7.04 (umc2057-bnl1666/umv1708) identified a constitutive QEI for Pn, bin 8.03 (bnl1863-umc2075) dissected a constitutive QEI affecting WUE, bin 10.03 (bnl1655-umc2016) validated a constitutive QEI involved in Pn and Tr (Additional file 8: Table S4).

### **cQTLs and candidate genes dissection for photosynthetic performances**

Further analysis showed that 8 cQTLs were simultaneously identified by single environment mapping with CIM and joint analysis through MCIM in two F<sub>4</sub> populations, which were located on chromosome 1 (Chr. 1), Chr. 6, Chr. 7, Chr. 8, and Chr. 10, and each cQTL accounting for 3.21~15.78 % of the average observed phenotypic variance (Table 2). Except for the cQTL1 for WUE, other 7 cQTLs were detected under DS environments, indicative that these cQTLs regimes may have several stable alleles that were involved in photosynthesis under stress environments (Table 2). Moreover, 5 cQTLs, i.e., cQTL2 for Pn, Ci, Tr, and WUE, cQTL5 associated with Pn, Tr, Ci, and RuBP, cQTL6 controlling Pn and RuBP, cQTL7 affecting Ci, Tr, and WUE, as well as cQTL8 involved in Pn, Tr, and RuBP (Table 2), imparted pleiotropic effect on two or four photosynthetic-related traits, suggesting that these cQTLs intervals control two or more tightly linked photosynthetic-related traits. In addition, the corresponding identified 8 cQTLs for all six photosynthetic-related traits were projected on the physical map B73 RefGen\_v4 ([http://www.maizegdb/gbrowse/mazie\\_V4](http://www.maizegdb/gbrowse/mazie_V4)), resulting in the identification of 17 candidate genes involved in leaf morphology and development, photosynthesis, and stress response (Table 2).

### **Epistasis for photosynthetic performances**

Out of all QTLs identified, 6 and 8 pairs of epistatic interactions for six photosynthetic-related traits exhibited dominance-by-additive (DA) and dominance-by-dominance (DD) effects based on different watering environments in two F<sub>4</sub> populations, and one for each pair epistatic interaction explained 2.11 (for Tr under WW)~4.96 % (for Gs under WW) of the observed phenotypic variance contributed by  $h^2(DA)$ , and 2.09 (for Ci under DS)~5.87 % (for Ci under DS) of the observed phenotypic variance contributed by  $h^2(DD)$ , respectively (Table 3, Fig. 5), indicating that the main effects of significant QTLs may be stronger on these six photosynthetic performances. Additionally, two pairs of the epistatic interactions were consistently detected in two F<sub>4</sub> populations under different watering environments, i.e., simultaneously located between bin 1.07\_1.10 (bnl1025-mmc0041-phi308707) and bin 10.03 (bnl1655-umc2016/umc1345) for Pn, and between bin 1.08\_1.10 (mmc0041-phi308707) and bin 6.05 (umc2040-bnl1174a) for Ci (Table 3, Fig. 5). The two epistasis may thus be critical for MAS.

## **Discussion**

### **Maize photosynthetic performances variations in response to drought stress**

Approximately 95 % of the organic matter accumulated by crops comes from its own photosynthesis, and photosynthesis efficiency could directly determine the GY. Drought stress can significantly affect crop photosynthesis, and the responses of leaf photosynthesis to drought were mediated by two different physiological processes [35-36]. Firstly, SC and AMC decrease were recognized as the main driver of the photosynthetic response to drought stress, in order to reduce transpiration under water deprivations, the plant stomatal can close, and SC could limite CO<sub>2</sub> diffusion efficiency from the atmosphere to the substomatal cavities to slow photosynthesis [37-38], and AMC rapidly decrease could further limite CO<sub>2</sub> diffusion efficiency from the substomatal cavities to the chloroplast stroma during water stress [39-40]. Secondly, photosynthesis could be limite by biochemical processes resulting in photosynthetic enzyme activity inhibition, Pn, Rubisco and RuBP activity decrease, etc. [7-9, 36]. Fortunately, above these findings were also supported by our results in this study, namely, drought-stressed Chang7-2, Langhuang, TS141, POP-CT, and POP-LT exhibited obvious decrease in Pn, Gs, Tr, and RuBP, their Ci and WUE, however, significantly displayed the increase relative of corresponding plants with sufficient water (Fig. 1, Additional file 2: Table S1). Furthermore, RuBP (average RC 18.96 %) and Gs (average RC 17.05 %) in response to drought stress were more sensitive than the response to Pn (average RC 16.02 %), Ci (average RC -9.34%), Tr (average RC 10.80 %), and WUE (average RC -8.62 %) (Fig. 1). It could be concluded that the drought-induced limitation of photosynthesis in maize was primarily due to CO<sub>2</sub> diffusion efficiency from sub-stomatal interval internal cavities to carboxylation site in chloroplasts and degree of SC, and increasing evidences in maize also supported our results by Liu et al. [5], Veroneze-Júnior et al. [41], Perdomo et al. [42], and He et al. [4]. Therefore, genetic improvement of photosynthetic performances in maize can be applied to MAS breeding to improve drought tolerance and high-yielding in the future.

### **Genetic architectures for photosynthetic performances**

Although a wealth of information from previous researches considerably improved our understanding of leaf photosynthetic performances [43-45], as well as applications in maize MAS breeding [5, 9, 18, 35], few studies considered the genetic basis of maize photosynthetic-related traits under water deficit at the molecular level [20-22]. Based on the above considerations, in this study we detected 54 QTLs for six photosynthetic-related traits across two F<sub>4</sub> populations via single watering environment mapping with CIM (Fig. 4, Additional file 6: Table S3), and for the identified QTLs, Pn, Gs, and RuBP showed both additive and non-additive effects under drought and non-drought stressed environments, but non-additive effects were largely responsible for the genetc basis of these three traits, of which accounted for 95.24, 81.82, and 94.12 %, respectively (Additional file 5: Fig. S3). However, all identified QTLs for Ci, Tr, and WUE displayed non-additive effects under both contrasting watering environments (Additional file 5: Fig. S3). These results were consistent with Wang and Zhang [18] and Li et al. [46] involved in photosynthetic-related traits in maize. Further Pearson correlation analysis among F<sub>4</sub> progencies and their parents showed that Pn, Gs, Ci, RuBP, and WUE of the two F<sub>4</sub> populations depicted significantly positive correlation to female parent, as well as Tr showed significantly positive correlation to both parents (Additional file 4: Table S2). Thereby, breeders should pay more attentation to the evaluation of F<sub>1</sub> cross combinatons to make good use of their prominent non-additive effects and specific combining ability for

above six photosynthetic-related traits, as well as should carefully select parents with elite photosynthetic-related traits, finally achieving the aim of improving these traits under drought and un-stressed environments. Additionally, 43 of the 54 identified QTLs were found in drought-stressed environments (Additional file 8: Fig. S4). Suggesting that these identified QTLs controlling Pn, Gs, Ci, Tr, RuBP, and WUE could be changes under both contrasting environments, and the QTLs were identified under drought stress that may directly lead to differences in Pn, Gs, Ci, Tr, RuBP, and WUE.

GEI is critical in determining the adaption and fitness of genotypes in adverse environments, resulting in phenotypic variations [47-48], QEI information thus obtained was of great value for breeders and genetic researchers [47]. Furthermore, the variations of photosynthetic-related traits in two F<sub>4</sub> populations showed that the wide variations, i.e. , , and were observed for six photosynthetic-related traits ( $P < 0.01$  or  $P < 0.05$ ), as well as their ranged from 1.466 % to 19.770 % (Table 1). Further analysis of QEI in two F<sub>4</sub> populations with MCIM among all watering environments also suggested that totally 24 QEIs (44.44 %) of the identified joint QTLs controlling Pn, Gs, Ci, Tr, RuBP, and WUE were mapped, and each QEI explained 1.34~6.86 % of the phenotypic variance by  $h^2(AE)$  in present study (Additional file 8: Table S4). In this regard, photosynthetic-related traits are similar to other traits, such as yield-related, and leaf and inflorescence architecture traits in maize [27-29], of which showed extensive GEI. As a result, GEI may be a major challenge to MAS breeding for photosynthetic-related traits in maize.

Epistasis, i.e. the interactions between genetic loci, is also thought to contribute to photosynthetic performances variations [49-51]. In maize, Li et al. [46] reported that SPAD was controlled by 2 major genes with AD-epistatic effects and polygene with AD effects by mixed major gene plus polygene genetic model. Liu et al. [52] also suggested that SPAD was in agreement with the AD-epistatic model using diallel cross II (Griffing). As expected, in accord with previous studies [46, 52-53], totally 14 pairs of epistatic interactions with DA and DA controlling Pn, Gs, Ci, Tr, RuBP, and WUE based on different watering environments in two F<sub>4</sub> populations, and which accounted for 2.09~5.87 % of the observed phenotypic variance contributed by  $h^2(DA/DD)$  that were clearly lower than those from A effects for all photosynthetic-related traits (Table 3, Fig. 5). It could be concluded that low contributions to phenotypic variance explained by DA/DD effects were due to large No. of DA/DD-QTLs with minor genetic effects, which would significantly influence the efficiency of MAS breeding for photosynthetic-related traits. Notably, two stable DD-epistatic interactions were also validated in our study, i.e., controlling Pn between bin 1.07\_1.08\_1.10 (bnlg1025-mm0041-phi308707) and bin 10.03 (bnlg1655-umc2016/umc1345) was repeatedly detected in POP-CT under both contrasting watering environments/POP-LT under stressed environment, and affecting Ci between bin 1.08\_1.10 (mm0041-phi308707) and bin 6.05 (umc2040-bnlg1174a) in two F<sub>4</sub> populations under both all four environments (Table 3, Fig. 5). These results were supported by Zhao et al. [26] and Sa et al. [54], who indicated that an AD/additive-additive (AA) epistatic interaction for KW, plant height, and ear length, etc., located on Chr. 1 and Chr. 6/10 under multiple drought and non-drought environments. Hence, these regions of chromosomes may be considered as epistatic regulators that influence maize plant development, photosynthesis, and ear formation under multiple watering regimes.

## cQTLs comparison and candidate genes validation for photosynthetic performances

The identified cQTLs for photosynthetic-related traits in a broad genetic background under adverse watering environments could provide guidance for fine mapping and maize MAS in the future. We totally identified 8 cQTLs for six photosynthetic-related traits using CIM/MCIM across POP-CT and POP-LT under multiple watering environments, of which accounting for 3.21~15.78 % of the average observed phenotypic variance in this study (Table 2).

Af these, cQTL2 in bin 1.07\_1.10 (bnlg1025/mmc0041-phi308707/umc1847) was simultaneously associated with Pn, Ci, Tr, and WUE, sQTL5 in bin 6.05 (umc2141-umc2040-bnlg1174a) was simultaneously involved in Pn, Tr, Ci, and RuBP, sQTL6 in bin 7.02\_7.04 (umc2057-bnlg1666-umc1708) was simultaneously affected Pn and RuBP, sQTL7 in bin 8.03 (bnlg1863-umc2075) was simultaneously responsible for Ci, Tr, and WUE, and sQTL8 in bin 10.03 (bnlg1655-umc2016/umc1345) was simultaneously associated with Pn, Tr, and RuBP in two  $F_4$  populations under both contrasting watering environments, suggestive of pleiotropic cQTLs, which were highly agreed with the Pearson correlations analysis among six photosynthetic-related traits under both watering environments (Fig. 3). In bin 1.08\_1.10 (umc83a-umc39c), in bin 6.05 (near gsy298e\_pmg), bin 7.03\_7.04 (gsy113\_cs-gsy107\_pr), and bin 10.03 (gsy321\_aba-gsy329\_pp) intervals, Pelleschi et al. [21] also detected multiple QTLs associated with photosynthesis (i.e., sucrose-P synthase (SPS), hexoses (HEX), sucrose (SUC), AGP, Tr, and CO) and leaf morphologies (i.e., number (LN), width (LW), length (LL), and relative water content (RWC)) across 120 F-2  $\times$  MBS847 RILs under drought and non-drought conditions. Li et al. [50] also mapped multiple QTLs involved in SPAD in bin 1.08 (umc1013-umc2047), 7.02 (umc1585-bnlg1305), bin 8.03 (bnlg1863-bnlg2046/umc2075-phi100175), and bin 10.03 (bnlg1655) regions in 172 Xu172  $\times$  Zong3 single segment substitution lines (SSSLs) under both high and low N treatments. Wang and Zhang [18] also identified two pleiotropic QTLs were simultaneously responsible for FCa, FCb, and FCt in bin 1.08 (mmc0041-bnlg1556) and bin 8.03 (umc1457-umc2199) using 189 A150-3-2  $\times$  Mo17  $F_2$  plants under a single environment. Peng et al. [33] also found a QTL for grain yield per plant (GYPP) in near phi308707 (bin 1.10), a stable QTL controlled GYPP and kernel number per plant (KNPP) located in bnlg1094-bnlg1579 (bin 7.02\_7.03), and a QTL affecting KW in bin 8.03 (bnlg1352-umc1778) cross 230 Qi319  $\times$  Huangzaosi and 235 Ye478  $\times$  Huangzaosi  $F_{2:3}$  families during six environments. These results indicated that pleiotropic cQTLs in the bin 1.07\_1.10, 6.05, 7.02\_7.04, 8.03, and 10.03 regions may play critical roles in leaf development, photosynthesis, and yield formation in maize under contrasting watering environments, and some important genes may be located in these bin regions. As expected, totally 11 candidate genes were also validated in above 5 pleiotropic cQTLs (cQTL 2, cQTL5, cQTL6, cQTL7, and cQTL8) intervals. *GRMZM2G018627 (LHCB9)* is a light harvesting chlorophyll binding (LHCB) protein, and Zhao et al. [55] proved that LHCB protein was required for the maintenance of photosystem I and specific protein-chlorophyll complexes especially under certain stress conditions. *GRMZM2G162672 (chlg1)*, i.e., *chlorophyll synthase G1* gene [56], acted as a chlorophyll biosynthetic process in biological progress via the AgBase v. 2.00 (<http://agbase.arizona.edu/>) online software with gene ontology (GO) analysis. *GRMZM2G039113 (tan1)*, i.e., *tangled 1* gene was required for spatial control of cytoskeletal arrays

associated with cell division during maize leaf development [57], as well as the *tan1* mutation could alter cell division orientations through leaf development without altering leaf shape, and even affected maize photosynthesis capacity [58]. *GRMZM2G013657 (dwil1)*, i.e., *dwarf & irregular leaf 1* gene may regulate leaf base, tip, and sheath development in maize ([https://maizegdb.org/gene\\_center/gene](https://maizegdb.org/gene_center/gene)). *GRMZM5G809292 (PYG7)*, i.e., tetratricopeptide repeat domain-containing protein PYG7 [59], was the components of the chloroplast and thylakoid membrane in mesophyll component and involved in photosystem I assembly in biological processes by GO analysis. *GRMZM2G042592*, encoded the thioredoxin-like 6, chloroplast contained a variety of thioredoxin systems [60], while the thioredoxin could interact with CHL1 subunits of Mg<sup>2+</sup> chelatase to regulate the chelation of Mg<sup>2+</sup> chelatase and chlorophyll synthesis precursor protoporphyrin IX [61]. Wang et al. [62] also predicted *GRMZM2G042592* located in the same region of bnlg1863-umc2075 (bin 8.03), and its mutant with abnormal chloroplast, lacking pigment and reducing of PSII. *GRMZM2G163437 (agps11)*, i.e., *ADP glucose pyrophosphorylase small subunit leaf 1* gene, could be expressed in maize grain and source leaf during grain filling [63]. *GRMZM2G033885 (psb29)*, encoded photosystem II subunit29, which was reversibly phosphorylated in maize upon exposure to high light in the cold condition, as well as its phosphorylation was controlled by the redox state of the plastoquinone pool, and may be part of a novel mechanism of photoprotection [64]. Another 2 candidate genes, i.e., *GRMZM2G045431 (bHLH150)* and *GRMZM2G058451 (bHLH164)*, belonging to *bHLH* transcription factors, which played important roles in maize growth and development, such as, root differentiation [65], photomorphogenesis and light signal transduction [66], and stress response [67].

In addition, We also noted that cQTL3 affecting Gs in bin 3.07\_3.08 (umc1286/umc2275-umc2081) with CIM/MCIM cross two F<sub>4</sub> populations under drought and non-drought environments. Li et al. [50] also found a QEI for SPAD in bin 3.08 (umc1844-bnlg1182) under high N condition, and even Guo et al. [68] reported a meta-QTL (mQTL) for LW in bin 3.08 region from 28 original populations under multiple environments via mQTL analysis. Then further analysis validated two candidate genes, i.e., *GRMZM2G159937 (bHIH57)* and *GRMZM2G117851 (bZIP1)* in the bin 3.07\_3.08 intervals in our study. *GRMZM2G159937* may have all the functions of *bHLH* transcription factor family [65-67]. Walsh et al. [69] reported *liguleless2 (lg2)* gene encoded a bZIP protein, involved in maize ligule and auricle development of leaf during vegetative and reproductive growth periods. *GRMZM2G117851* was also a bZIP transcription factor, may play an important role in maize leaf development.

Additionally, 2 new cQTLs (cQTL1 and cQTL4) information were also found in our study. cQTL1 controlling WUE in bin 1.00\_1.01 (bnlg149-bmc1014/umc1177) with CIM/MCIM cross both populations, of which *GRMZM2G042250 (rld2, rolled leaf 2)* was predicted, and its orthologous gene *ATHB23* (i.e. a phytochrome B-interacting protein), was important for phytochrome B-mediated red light signaling in *Arabidopsis thaliana* [70]. cQTL4 responsible for Ci in bin 4.08\_4.09 (umc2041-umc2188/umc2287) with CIM cross both populations under stressed and un-stressed environments. *GRMZM2G446426 (MADS52)*, *GRMZM2G038479 (bHLH8)*, and *GRMZM2G074122 (pep3)* were identified in cQTL4. *GRMZM2G446426* as the *MADS-transcription factor 52*, could have the similar as *ZmMADS4* in influencing chlorophyll content, and involve in response to osmotic stress [71]. *GRMZM2G074122* was *phosphoenolpyruvate*

*carboxylase isoform 1* gene, which involved in carbon fixation and tricarboxylic acid cycle in biological process and influenced phosphoenolpyruvate carboxylase activity in molecular function via GO analysis. As a result, the 2 cQTLs intervals may provide new information for genetic basis dissection in maize photosynthetic performances under both watering environments in the future.

## Conclusions

Photosynthetic performances in maize were predominantly controlled by non-additive and QEIs effects, where more QEIs (87.5 %) effects occurred in drought stress. 8 cQTLs (bin 1.00\_1.01, bin 1.07\_1.10, bin 3.07\_3.08, bin 4.07\_4.08, bin 6.05, bin 7.02\_7.04, bin 8.03, and bin 10.03) affecting six photosynthetic-related traits could be useful for genetic improvement of these traits via QTL pyramiding, corresponding 5 cQTLs (cQTL2, cQTL5, cQTL6, cQTL7, and cQTL8) clusters indicated tight linkage or pleiotropy in the inheritance of these traits, and 17 candidate genes (*GRMZM2G042250*, *GRMZM2G018627*, *GRMZM2G159937*, *GRMZM2G117851*, *GRMZM2G446426*, *GRMZM2G038479*, *GRMZM2G074122*, *GRMZM2G162672*, *GRMZM2G039113*, *GRMZM2G013657*, *GRMZM2G045431*, *GRMZM2G033885*, *GRMZM2G058451*, *GRMZM5G809292*, *GRMZM2G042592*, *GRMZM2G042895*, and *GRMZM2G163437*) involved in leaf morphology and development, photosynthesis, and stress response coincided with above corresponding cQTLs.

## Abbreviations

A, additive effect; AMC, apparent mesophyll conductance;  $C_i$ , intercellular CO<sub>2</sub> concentration; CI, confidence interval; CIM, composite interval mapping; cQTLs, constitutive QTLs, D, dominance effect; DA, dominance-by-additive effect; DD, dominance-by-dominance effect; Gs, stomatal conductance; GY, grain yield; MAS, marker-assisted selection; OD, over-dominance effect; PCA, principal component analysis; PD, partial dominance effect; Pn, net photosynthetic rate; QEIs, QTL × environment interactions; QTLs, quantitative trait loci; RuBP, ribulose 1,5-biphosphate carboxylase activity; SC, stomatal closure; Tr, transpiration rate; WUE, water use efficiency.

## Declarations

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## Availability of data and materials

All relevant data are available within the manuscript and its additional files.

## Authors' contributions

XZ and YZ designed the experiments; XZ wrote the manuscript; XZ, WL, DZ, and YZ performed the experiments and analyzed the data. All authors have read and approved the final manuscript.

## Competing interests:

The authors declare that they have no competing interests.

## Consent for publication

Not applicable.

## Ethics approval and consent to participate

Not applicable.

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## Tables

**Table 1** Variance analysis, broad-sense heritability (  $h^2$  ), and  $G \times E$  genotype  $\times$  environment interaction (GEI) heritability (  $h^2_{GEI}$  ) of the photosynthetic-related traits in the POP-LT and POP-CT, respectively

Item	Pn	Gs	Ci	Tr	RuBP	WUE
-----POP-CT-----						
Corrected model	279.722**	0.025*	9400.878**	4.289**	0.033**	3.883**
Intercept	5687.46**	0.262**	212617.821**	232.181**	0.713**	329.913**
	102.911**	0.013*	3214.928**	5.129**	0.032**	3.728**
	71.213**	0.007*	837.813**	2.259**	0.016**	1.039*
	25.598**	0.008*	320.051**	2.293**	0.009*	0.126*
	2.514**	0.002NS	31.353NS	0.217NS	0.002NS	0.101NS
Total	6573.387	0.293	222050.052	236.688	0.749	333.897
	88.939	48.148	91.030	69.683	68.817	86.778
	9.977	14.815	4.531	15.576	9.677	1.466
-----POP-LT-----						
Corrected model	395.355**	0.124*	6224.776**	8.165**	0.024**	3.480**
Intercept	6174.345**	0.268**	182666.913**	288.320**	1.138**	407.828**
	121.67**	0.016*	3014.544**	6.087**	0.020**	3.401**
	53.986**	0.009*	731.405**	6.573**	0.012**	1.020*
	29.726*	0.010*	363.211**	3.266**	0.004*	1.059*
	3.687NS	0.009 NS	24.168NS	0.108NS	0.002NS	0.199NS
Total	6573.387	0.301	188915.857	296.593	1.163	411.507
	78.513	46.429	90.882	73.692	62.500	69.049
	9.591	8.929	5.475	19.770	6.250	10.750

Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphospate carboxylase activity, WUE water use efficiency, the genotypic variance, the environmental variance, the 'genotypic  $\times$  environment' interaction variance, the error variance, heritability, the genotype by environment interaction heritability. \*\*/\* indicated the significant difference at the  $P < 0.01/P < 0.05$  probability level, respectively

**Table 2** Summary of constitutive QTLs (cQTLs) and candidate genes for photosynthetic-related traits in the POP-LT and POP-CT, respectively

cQTL	Marker interval	Trait	Population (Environment)	QTL	Bin	R <sup>2</sup> (Average) (%)	Candidate gene (Annotation)
cQTL1	bnlg149- bmc1014/umc1177	WUE	POP-CT (WW), POP- LT (Joint)	qWUE- Ch.1-1, qWUE-J1- 1	1.00-1.01	2.53-3.88 (3.21)	<i>GRMZM2G042250 (rid2)</i>
cQTL2	bnlg1025/mmc0041- phi308707/umc1847	Pn Ci Tr WUE	POP-CT (WW, DS, Joint), POP-LT (DS) POP-CT (WW, DS, Joint), POP-LT (DS, Joint) POP-CT (DS, WW), POP-LT (DS) POP-CT (DS, Joint), POP-LT (WW, DS)	qPn-Ch.1- 2 qCi-Ch.1-1 qTr-Ch.1-1 qWUE- Ch.1-2, qWUE-J1- 2	1.07/1.08_1.10	5.68-9.33 (7.06) 5.98- 10.13 (7.16) 3.77-9.35 (5.88) 2.92-7.99 (4.95)	<i>GRMZM2G018627 (LHCB9)</i>
cQTL3	umc1286/umc2275- umc2081	Gs	POP-CT (WW, DS, Joint), POP-LT (WW, DS, Joint)	qGs-Ch.3- 1	3.07-3.08	4.14- 10.10 (7.93)	<i>GRMZM2G159937 (bHLH57), GRMZM2G117851 (bZIP1)</i>
cQTL4	umc2041- umc2188/umc2287	Ci	POP-CT (WW, DS), POP-LT (WW, DS)	qCi-Ch.4-1	4.08_4.09	5.98- 13.40 (10.68)	<i>GRMZM2G446426 (MADS52), GRMZM2G038479 (bHLH8), GRMZM2G074122 (pep3)</i>
cQTL5	umc2141-umc2040- bnlg1174a	Pn Tr Ci RuBP	POP-CT (WW, DS) POP-CT (WW, DS, Joint) POP-CT (Joint), POP- LT (Joint) POP-LT (WW, DS, Joint)	qPn-Ch.6- 1 qTr-Ch.6-1 qCi-J6-1 qRuBP- Ch.6-1	6.05	6.76-9.53 (8.15) 3.58-8.69 (5.43) 3.83-3.95 (3.89) 2.64-3.38 (3.00)	<i>GRMZM2G162672 (chlg1), GRMZM2G039113 (tan1), GRMZM2G013657 (dwil1)</i>
cQTL6	umc2057- bnlg1666/umc1708	Pn RuBP	POP-CT (WW, DS, Joint), POP-LT (WW, DS, Joint) POP-CT (WW, DS, Joint), POP-LT (Joint)	qPn-Ch.7- 2 qRuBP- Ch.7-1, qRuBP-J7- 1	7.02_7.04	5.03- 17.26 (9.95) 3.73-7.84 (5.56)	<i>GRMZM2G045431 (bHLH150), GRMZM2G033885 (psb29), GRMZM2G058451 (bHLH164), GRMZM5G809292 (PYG7)</i>
cQTL7	bnlg1863-umc2075	Ci Tr WUE	POP-CT (WW, DS), POP-LT (WW, DS, Joint) POP-CT (WW, DS), POP-LT (WW, DS, Joint) POP-CT (Joint), POP- LT (Joint)	qCi-Ch.8-1 qTr-Ch.8-1 qWUE-J8- 1	8.03	11.74- 18.21 (15.78) 3.48-8.69 (5.42) 4.05-7.53 (5.79)	<i>GRMZM2G042592 (Similar to Thioredoxin-like 6)</i>
cQTL8	bnlg1655- umc2016/umc1345	Pn Tr RuBP	POP-CT (Joint), POP- LT (WW, Joint) POP-CT (WW, DS, Joint), POP-LT (WW, DS, Joint)	qPn-J10-1, qPn- Ch.10-1 qTr-Ch.10- 1	10.03	6.10- 11.17 (8.78) 2.92-5.25 (3.65)	<i>GRMZM2G042895 (bHLH116), GRMZM2G163437 (agps11)</i>

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Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphospatе carboxylase activity, WUE water use efficiency

**Table 3** Epistatic interactions for photosynthetic-related traits were detected in POP-LT and POP-CT with a mixed linear model based on composite interval mapping (MCIM)

Trait	Environment	QTL(i)	Marker interval(i)	Bin(i)	QTL(j)	Marker interval(j)	Bin(j)	DA	DD	h <sup>2</sup> (DA) (%)	h <sup>2</sup> (DD) (%)
-----POP-CT-----											
Pn	WW	qPn-Ch.1-2	mmc0041-phi308707	1.08_1.10	qPn-J10-1	bnlg1655-umc2016	10.03		-0.61		4.62
	DS	qPn-Ch.1-2	mmc0041-phi308707	1.08_1.10	qPn-J10-1	bnlg1655-umc2016	10.03		-0.74		5.35
Gs	WW	qGs-Ch.3-1	umc1286-umc2081	3.07	qGs-J4-1	umc2041-umc2188	4.08	0.05		4.96	
Ci	WW	qCi-Ch.1-1	mmc0041-phi308707	1.08_1.10	qCi-J6-1	umc2040-bnlg1174a	6.05		-1.18		3.56
	DS	qCi-Ch.1-1	mmc0041-phi308707	1.08_1.10	qCi-J6-1	umc2040-bnlg1174a	6.05		-1.40		4.01
Tr	WW	qCi-J6-1	umc2040-bnlg1174a	6.05	qCi-J7-1	umc2177-umc1378	7.00	-0.98		2.79	
	DS	qCi-J6-1	umc2040-bnlg1174a	6.05	qCi-J7-1	umc2177-umc1378	7.00	-1.07		3.01	
Tr	WW	qTr-Ch.5-1	umc2216-umc1072	5.06_5.07	qTr-Ch.10-1	bnlg1655-umc2016	10.03	0.06		2.11	
	DS	qTr-Ch.5-1	umc2216-umc1072	5.06_5.07	qTr-Ch.10-1	bnlg1655-umc2016	10.03	0.08		2.64	
RuBP	WW	qRuBP-J1-1	mmc0041-phi308707	1.08_1.10	qRuBP-J7-1	umc2177-umc1378	7.00		-0.05		3.73
	DS	qRuBP-J1-1	mmc0041-phi308707	1.08_1.10	qRuBP-J7-1	umc2177-umc1378	7.00		-0.09		4.80
WUE	WW	qWUE-Ch.1-2	mmc0041-phi308707	1.08_1.10	qWUE-J8-1	bnlg1863-umc2075	8.03	-0.38		3.13	
	DS	qWUE-Ch.1-2	mmc0041-phi308707	1.08_1.10	qWUE-J8-1	bnlg1863-umc2075	8.03	-0.32		2.94	
-----POP-LT-----											
Pn	DS	qPn-J1-1	bnlg1025-mmc0041	1.07_1.08	qPn-Ch.10-1	bnlg1655-umc1345	10.03		-0.43		2.46
Ci	DS	qCi-Ch.1-1	mmc0041-phi308707	1.08_1.10	qCi-J6-1	umc2040-bnlg1174a	6.05		-1.96		5.87
	WW	qCi-J4-1	umc1963-umc1031	4.04_4.05	qCi-J6-1	umc2040-bnlg1174a	6.05		1.25		3.39
	DS	qCi-J4-1	umc1963-umc1031	4.04_4.05	qCi-J6-1	umc2040-bnlg1174a	6.05		-1.03		2.09
Tr	WW	qTr-J6-1	bnlg238-umc2310	6.00	qTr-J9-1	dupssr29-umc2359	9.07		-0.10		5.04
	DS	qTr-J6-1	bnlg238-umc2310	6.00	qTr-J9-1	dupssr29-umc2359	9.07		-0.11		5.18
	WW	qTr-Ch.10-1	bnlg1655-umc1345	10.03	qTr-J10-1	bnlg1839-bnlg2162	10.07_10.08		0.09		4.75
	DS	qTr-Ch.10-1	bnlg1655-umc1345	10.03	qTr-J10-1	bnlg1839-bnlg2162	10.07_10.08		0.06		3.10
RuBP	WW	qRuBP-J1-1	mmc0041-phi308707	1.08_1.10	qRuBP-Ch.6-1	umc2040-bnlg1174a	6.05	0.05		2.47	
	DS	qRuBP-J1-1	mmc0041-phi308707	1.08_1.10	qRuBP-Ch.6-1	umc2040-bnlg1174a	6.05	0.06		2.53	
WUE	DS	qWUE-J1-2	phi308707-umc1847	1.10	qWUE-J8-1	bnlg1863-umc2075	8.03	-0.34		2.39	

Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphosphate carboxylase activity, WUE water use efficiency, WW well-watered environment at Wuwei in 2019, DS drought-stressed environment at Wuwei in 2019, DA the dominance by additive epistatic interaction effects, DD the dominance by dominance epistatic interaction effects,  $h^2(DA)$  percentage of phenotypic variance explained by the dominance-by-additive epistatic interaction effects,  $h^2(DD)$  percentage of phenotypic variance explained by the dominance-by-dominance epistatic interaction effects

## Figures

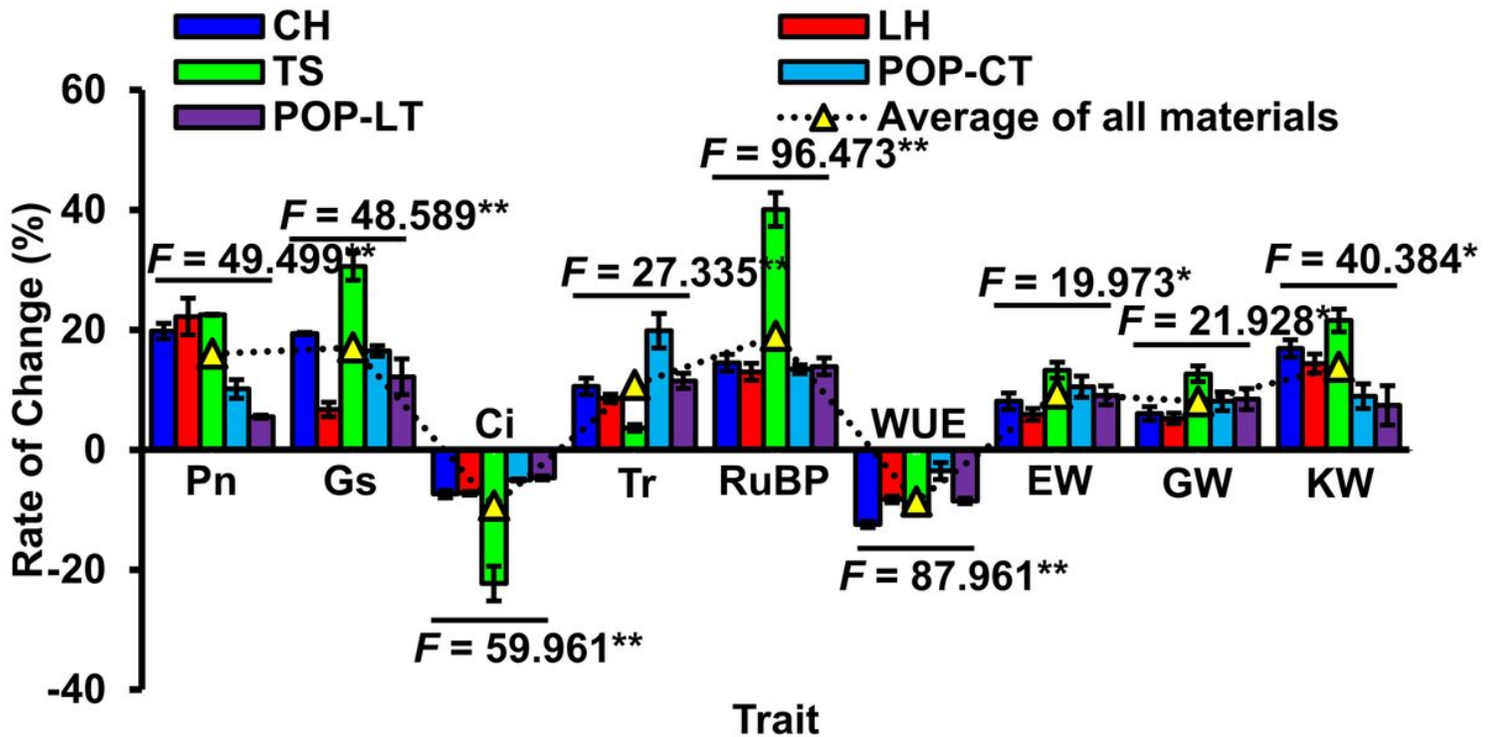
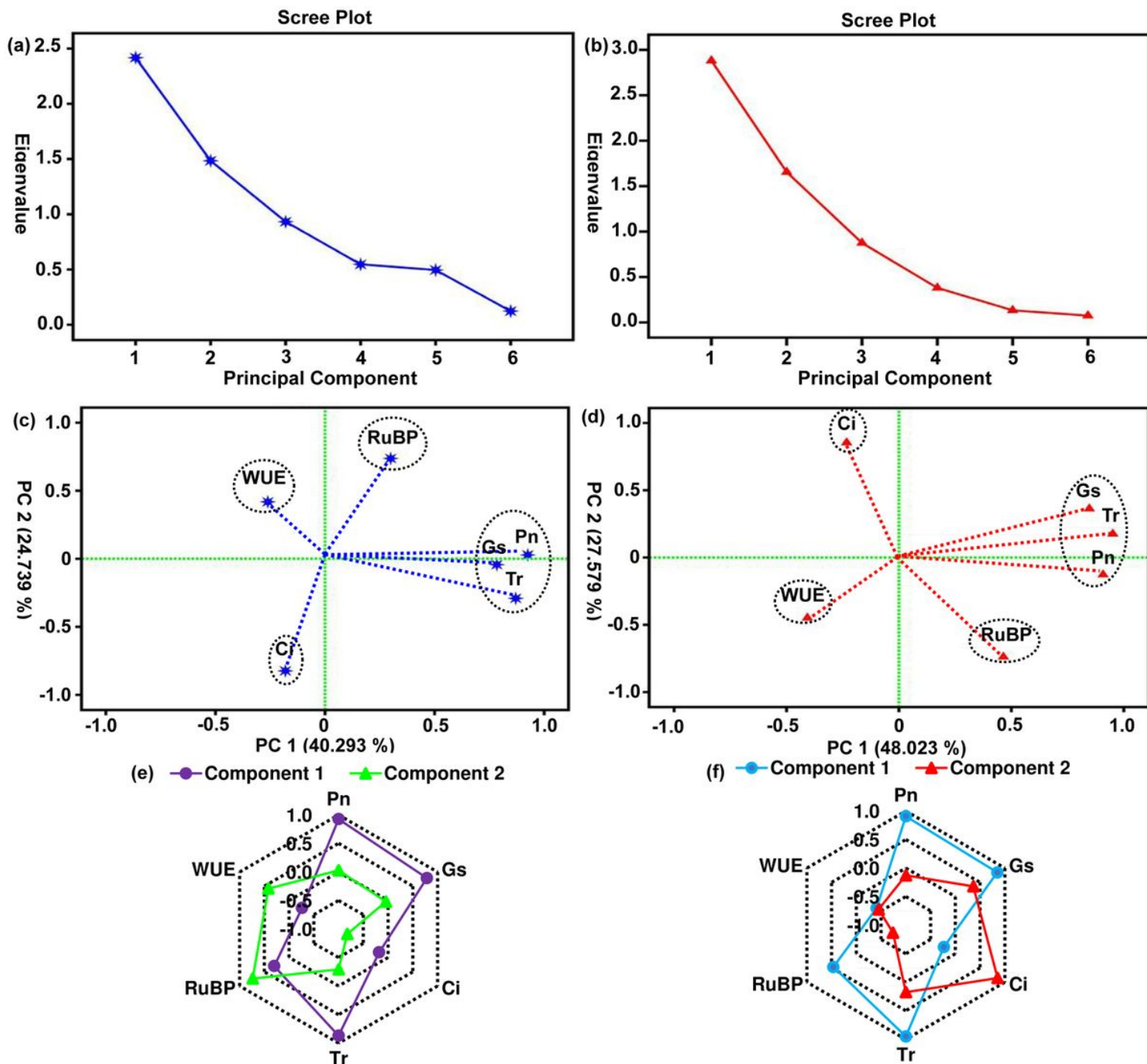


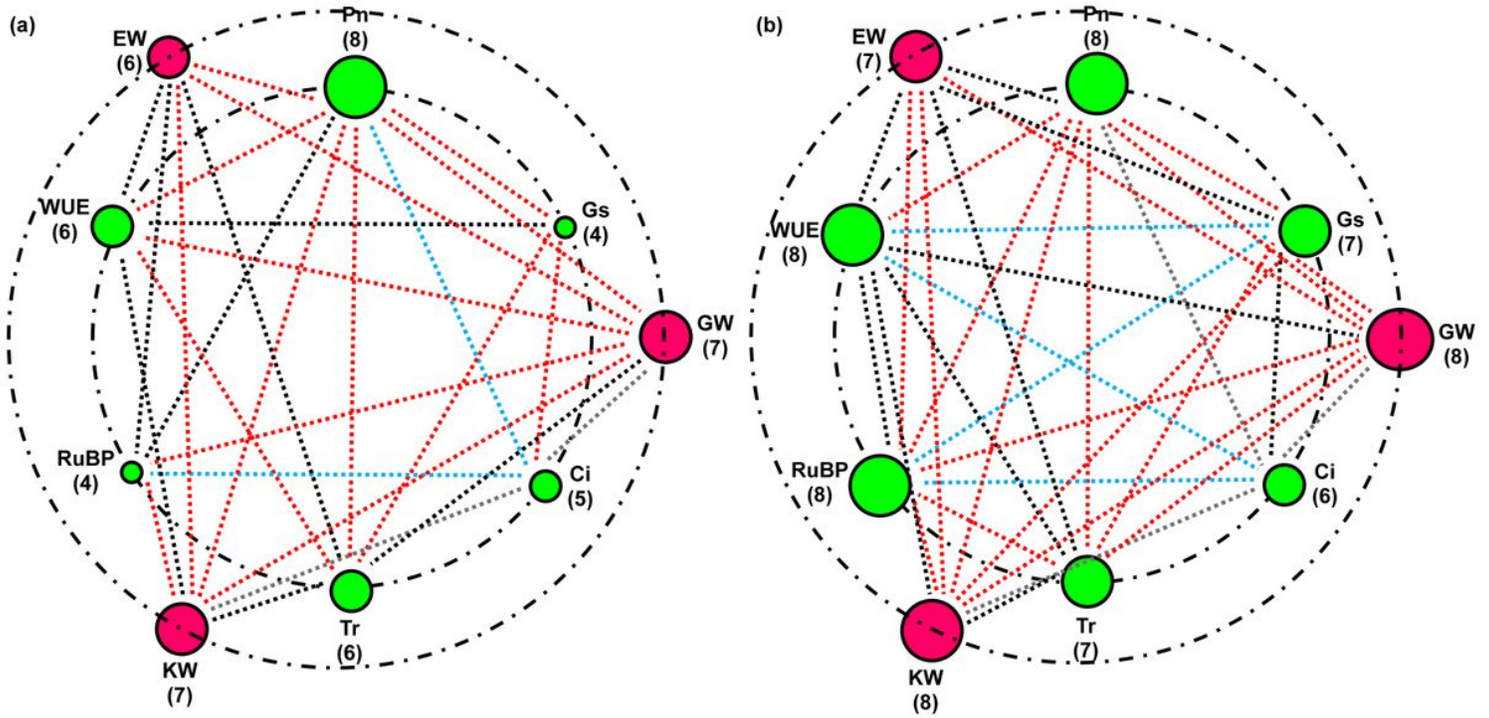
Figure 1

The rate of changes of photosynthetic-related traits (Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphosphate carboxylase activity, WUE water use efficiency) and yield component traits (EW ear weight, GW grain weight per ear, KW 100-kernel weight) in three parents (LH Langhuang, CH Chang7-2, TS TS141) and two F4 families (POP-LT, POP-CT) under different watering conditions (WW well-watered environment at Wuwei in 2019, DS, drought-stressed environment at Wuwei in 2019), F-value\*\*/\* indicated the significant difference at  $P < 0.01$  or  $P < 0.05$  level via one-way analysis of variance (ANOVA)



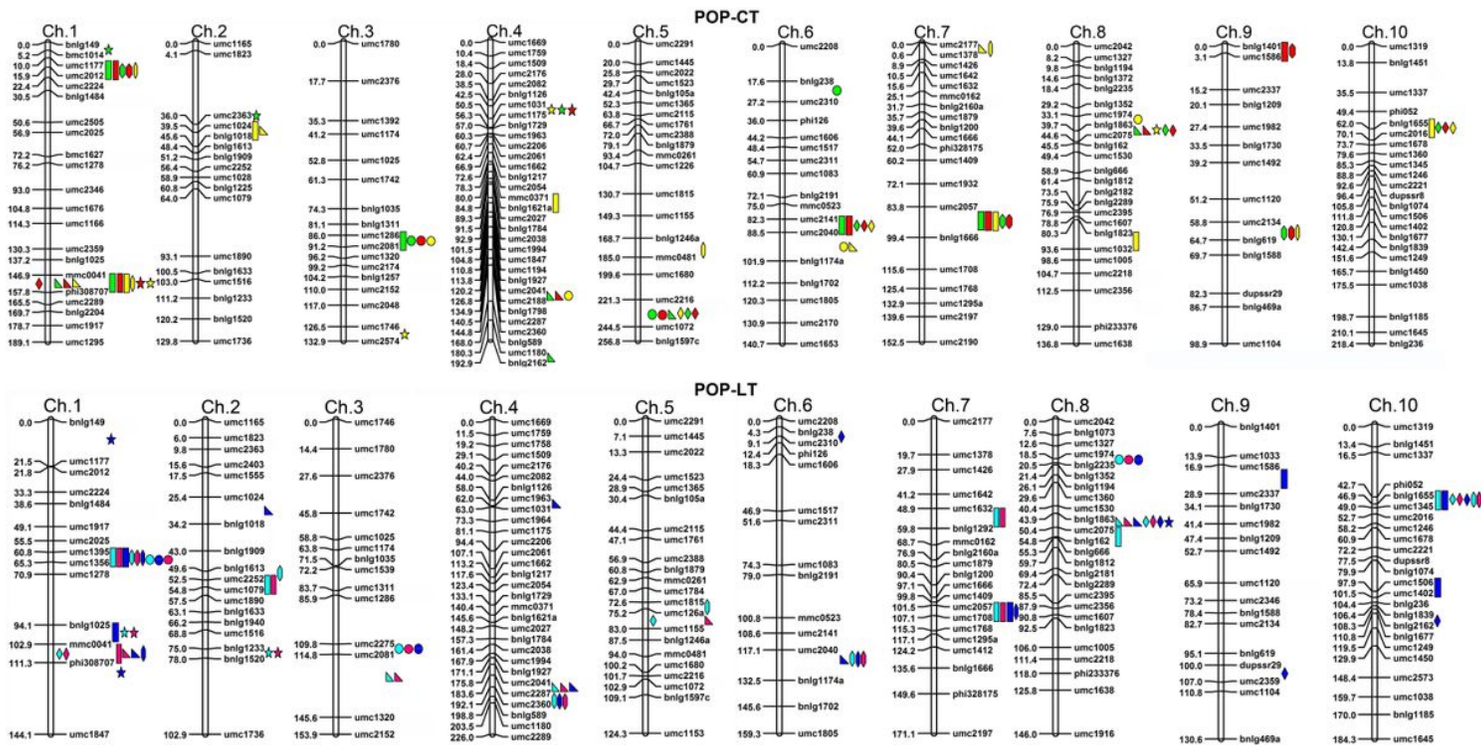
**Figure 2**

Principal component analysis (PCA) of photosynthetic-related traits (Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphospate carboxylase activity, WUE water use efficiency) in the both F4 populations (POP-LT, POP-CT) under different watering environments, (a, b) Eigenvalues of principal components (PCs) under contrasting watering environments, respectively, PCs with eigenvalue greater than 1 were retained, (c, d) The effects of corresponding photosynthetic-related traits in PC1 and PC2 under contrasting watering environments, (e, f) Eigenvectors of photosynthetic-related traits in PC1 and PC2 under contrasting watering environments



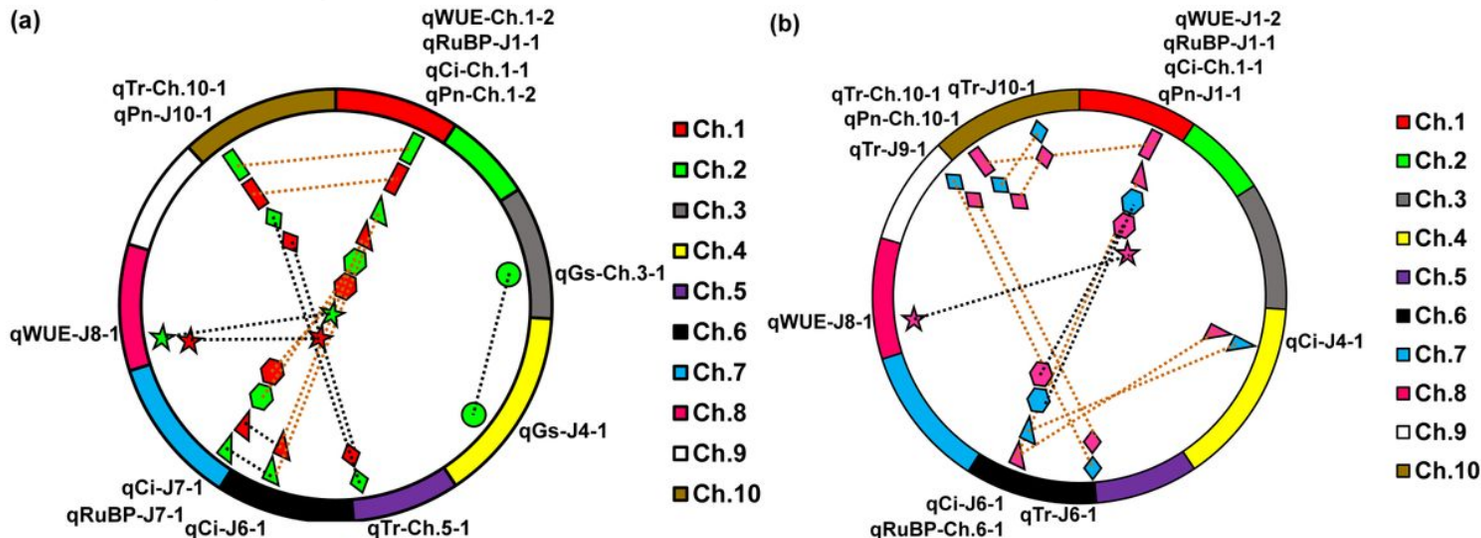
**Figure 3**

Pearson correlation among corresponding tested traits (Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphosphate carboxylase activity, WUE water use efficiency; EW ear weight, GW grain weight per ear, KW 100-kernel weight) under well-watered (a) and drought-stressed environments. Red/sapphire dotted lines designated positive /negative correlations between both traits ( $P < 0.01$ ), and black/gray dotted lines designated positive/negative correlations between both traits ( $P < 0.05$ ), respectively. Circles of different sizes reflected No. of corresponding tested traits



**Figure 4**

Genetic map and identified QTLs for photosynthetic-related traits (Pn net photosynthetic rate, Gs stomatal conductance, Ci intercellular CO<sub>2</sub> concentration, Tr transpiration rate, RuBP ribulose 1,5-biphosphate carboxylase activity, WUE water use efficiency) in two F<sub>4</sub> populations (POP-CT and POP-LT) by single environment mapping with composite interval mapping (CIM) and joint analysis of all environments with mixed-linear-model-based composite interval mapping (MCIM). Green/red and sapphire/pink rectangular, circle, triangle, rhombus, hexagon, and pentagram represented identified QTLs for Pn, Gs, Ci, Tr, RuBP, and WUE under well-watered/drought-stressed environment in POP-CT and POP-LT with CIM, respectively. Yellow and blue rectangular, circle, triangle, rhombus, hexagon, and pentagram represented identified QTLs for Pn, Gs, Ci, Tr, RuBP, and WUE in POP-CT and POP-LT among all watering environments with MCIM, respectively.



## Figure 5

Epistasis of QTLs were identified for photosynthetic-related traits in POP-CT (a) and POP-LT (b) by joint analysis among all environments with mixed-linear-model-based composite interval mapping (MCIM). Black/brown dotted lines represented dominance-by-additive/dominance (DA/DD) epistatic interaction effects, respectively. Green/red and sapphire/pink rectangular, circle, triangle, rhombus, hexagon, and pentagram represented corresponding QTLs for net photosynthetic rate (Pn), stomatal conductance (Gs), intercellular CO<sub>2</sub> concentration (Ci), transpiration rate (Tr), ribulose 1,5-biphospate carboxylase activity (RuBP), and water use efficiency (WUE) under well-watered/drought-stressed environment in POP-CT and POP-LT, respectively

## Supplementary Files

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