



## Biochemical, gas exchange, and chlorophyll fluorescence analysis of maize genotypes under drought stress reveals important insights into their interaction and homeostasis

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### Abstract

Many studies have been conducted on maize to study the effect of drought on yield at the flowering stage, but understanding biochemical and photosynthetic response against drought at the seedling stage needs to be well established. Thus, to understand differential changes and interaction of biochemical and photosynthetic parameters at the seedling stage under drought, a greenhouse experiment with twelve maize genotypes under severe drought (30% field capacity) and irrigated (90–100% field capacity) conditions were performed. Drought differentially altered biochemical and photosynthetic parameters in all genotypes. A sharp increase in hydrogen peroxide, malondialdehyde (MDA), and total antioxidant capacity (TAOC) were seen and a positive association between  $H_2O_2$  and TAOC, and MDA and transpiration rate ( $E$ ) was observed under drought. Nonphotochemical quenching increased under drought to avoid the photosystem damage. PCA biplot analysis showed that reducing  $E$  and increasing photosynthetic efficiency would be a better drought adaptation mechanism in maize at the seedling stage.

**Keywords:** chlorophyll fluorescence; drought; gas exchange; interaction; maize.

### Highlights

- Stomatal and nonstomatal limitations of photosynthesis occur in maize under severe drought
- A strong positive association was found between  $H_2O_2$  and TAOC
- Reducing  $E$  and increasing  $F_v/F_m$  would be a better drought adaptation mechanism

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**Abbreviations:** APX – ascorbate peroxidase; ASH – ascorbate; ATP – adenosine triphosphate; CAT – catalase; Chl – chlorophyll;  $C_i/C_a$  – the ratio of intercellular to ambient  $CO_2$  concentration; DAS – days after sowing;  $E$  – transpiration rate;  $F_v/F_m$  – maximal quantum efficiency of PSII;  $g_{H_2O}$  – water vapor conductance; GPX – glutathione peroxidase; GSH – glutathione; GST – glutathione S-transferase; MDA – malondialdehyde; NPQ – nonphotochemical quenching;  $O_2^-$  – superoxide radical;  $^1O_2$  – singlet oxygen;  $^{\cdot}OH$  – hydroxyl radical; PETC – photosynthetic electron transport chain;  $P_N$  – net assimilation rate; ROS – reactive oxygen species; SOD – superoxide dismutase; TAOC – total antioxidant capacity; VPD – air to leaf vapor pressure deficit; VWC – volumetric water content;  $Y_{NPQ}$  – quantum yield of NPQ-related energy loss;  $\Phi_{PSII}$  – quantum yield of photosynthetic electron transport.

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## Introduction

Maize (*Zea mays* sp.) is widely used as food, feed, and biofuel around the world. Its yield is hugely limited by drought in various parts of the world. Though maize has higher productivity compared to other cereals, its productivity is greatly limited by biotic and abiotic stresses (Assefa and Ayalew 2019). Around 40% of Africa's maize-growing area faces occasional drought stress, resulting in yield losses of 10–25%, while 25% of the maize crop suffers frequent drought, with losses of up to half (Fisher *et al.* 2015). Fifty years of meteorological and annual yield data (1958–2007) of the United States of America estimated that drought caused 13% of the USA's maize and soybean yield variability (Zipper *et al.* 2016). Drought interferes with plant growth, nutrient accumulation, water relations, and photosynthesis, thus reducing the yield (Farooq *et al.* 2009, Praba *et al.* 2009). Accordingly, plants have developed well sensing and signal transduction systems to respond to water limitation and osmotic stress (Bhaskara *et al.* 2012). Oxidative signaling is one of the major network complexes that operates under water-limiting conditions despite regulating plant growth and development under normal conditions (Tsukagoshi *et al.* 2010, Noctor *et al.* 2014). It generally happens through the reactive oxygen species (ROS) network which includes, hydrogen peroxide ( $H_2O_2$ ), superoxide radical ( $O_2^-$ ), singlet oxygen ( $^1O_2$ ), and the hydroxyl radical ( $^{\bullet}OH$ ). Under ambient growth conditions, ROS are produced continuously in the plants (Jakob and Heber 1996), and usually  $^1O_2$  and  $^{\bullet}OH$  production is kept at a minimum, while  $H_2O_2$  and  $O_2^-$  production is kept at a higher rate (Noctor and Foyer 1998). They function as various signaling molecules and participate under diverse stimuli (Singh *et al.* 2019). Generally, due to the photosynthesis and respiration process, ROS generations are inevitable and are mainly produced in chloroplasts, mitochondria, and peroxisomes.

The leakage of electrons from the photosynthetic electron transport chain (PETC) leads to the formation of ROS, particularly singlet oxygen [by energy transfer from triplet excited state chlorophyll (Chl) to  $O_2$ ] and hydrogen peroxide (by reduction of  $O_2$  through Mehler reaction) (Mehler 1951, Asada and Takahashi 1987). Mainly these ROS are highly unstable causing significant damage to DNA, cell membranes, proteins, and lipids by unrestricted oxidation, if not controlled or scavenged properly (Richter and Schweizer 1997, Dat *et al.* 2000, Apel and Hirt 2004). To avoid the cytotoxic effect of ROS, the plant has developed various scavenging mechanisms, which mainly operated through enzymatic (superoxide dismutase, SOD; ascorbate peroxidase, APX; glutathione peroxidase, GPX; glutathione S-transferase, GST; catalase, CAT) and nonenzymatic (ascorbate, ASH; glutathione, GSH; alpha-tocopherol; carotenoids and flavonoids) scavenging systems (Mittler *et al.* 2004, Gill and Tuteja 2010). But during abiotic stress, especially during drought, the redox homeostasis balance breaks down due to increased and uncontrolled production of ROS (Sgherri *et al.* 1993, Smirnoff 1993, Boo and Jung 1999, Polle 2001), creating

an imbalance in ROS production and scavenging. Mostly during water stress, initial signaling from abscisic acid from roots to shoot (Zhang and Davies 1987, Davies and Zhang 1991) leads to the closing of stomata to reduce transpiration flux, which in turn declines the  $CO_2$  intake leading to higher leakage of electrons from PETC to  $O_2$  by the Mehler reaction (Smirnoff 1993, Dat *et al.* 2000). Further, the reduced  $CO_2$  fixation decreases the consumption of ATP and NADPH, which in turn reduces  $NADP^+$  (major electron acceptor in PSI) regeneration through the Calvin cycle. This provokes the over-reduction of the photosynthetic electron transport chain (Cruz de Carvalho 2008) and leakage of electrons to molecular oxygen thus producing higher amount of  $H_2O_2$  (Asada 1999), leading to drought-induced oxidative stress.

In addition to cell damage, drought-induced oxidative stress impairs the photosystem machinery of the plant by creating the disequilibrium between light capture and utilization (Mattos and Moretti 2015), hence reducing photosynthetic efficiency. Usually, under drought-induced reduced  $CO_2$  fixation, the rate of absorption of light energy by photosynthetic pigments exceeds the rate of its consumption in chloroplasts, hence the absorbed light energy accelerates the process of photoinhibition (inhibition of PSII repair). Further, during drought, the  $CO_2$  assimilation rate showed a decrease due to stomatal and nonstomatal limitations (Cossins and Chen 1997). Additionally, drought not only reduces stomatal conductance and diffusion of  $CO_2$  but also decreases ribulose-1,5-bisphosphate (RuBP) generation, ATP synthesis, and other vital metabolic processes (Cossins and Chen 1997). Chl fluorescence is one of the most prominent approaches used to study the photosynthetic efficiency of plants (Murchie and Lawson 2013).  $F_v/F_m$  (maximal quantum efficiency of PSII) ratio is a key parameter to detect the PSII photoinhibition induced by stress (Krause and Weis 1991).

Maize as a C<sub>4</sub> plant fixes  $CO_2$  more efficiently by reducing photorespiration but it is very sensitive to drought at various stages of development (Sinclair *et al.* 1990, Farré *et al.* 2000). Many studies have been done at the anthesis stage to understand the effect of drought on yield and other agronomic traits, but knowledge of the differential response of biochemical and photosynthetic traits and their interaction is limited and need to be well explored. This present study aims at studying how maize responds to drought at the seedling stage through biochemical and photosynthetic changes and how they interact with each other, which could help in selecting the best physiologically adapted genotype for drought. Based on these observed parameters HD-6 genotype relatively performed well under drought compared to others.

## Materials and methods

**Plant growth conditions and drought treatments:** Twelve genotypes were used in the present study (text table). Seeds were surface-sterilized with sodium hypochlorite (3%) for 10 min and sown in a 30 × 30 cm pot filled with well-mixed sandy loam soil. Each pot

was inoculated with 7 g (300 spores g<sup>-1</sup>) *Rhizophagus irregularis*, arbuscular mycorrhiza formulation powder. Fertilizers were applied according to soil nutrient analysis and plants were grown under optimum growth conditions in a greenhouse at the Kunming Institute of Botany, Yunnan, China from 2019 to 2020. The average day/night temperature of the greenhouse was 29.5/20°C and relative humidity was 55–80% throughout the growth period.

No.	Genotype	Source
1	GJ-3	Local cultivar
2	JH-19A-501-36	YAAS-CIMMYT
3	YS-18B-150-44	YAAS-CIMMYT
4	GJ-14	Local cultivar
5	JH-19A-501-23	YAAS-CIMMYT
6	GJ-2	Local cultivar
7	La-posta-seq	YAAS-CIMMYT
8	JL-118	Local cultivar
9	YS-18B-150-43	YAAS-CIMMYT
10	HD-6	Local cultivar
11	JH-18A-514-38	YAAS-CIMMYT
12	GJ-1	Local cultivar

Before the initiation of the experiment, a pre-soil analysis was done to find out a field capacity (FC) of the soil. Soil (of bulk density 1.45 g cm<sup>-3</sup>) used for the experiment was filled in the same pots used for the experiment and saturated fully. After 24 and 48 h of saturation, soil moisture was measured by an *SM150T* soil moisture sensor (*Delta-T Devices Ltd.*, United Kingdom). After 24 h, the volumetric water content (VWC) was averaged at 20.8%, and after 48 h, it was 18%. Soil moisture content after 2–3 d after irrigation or saturation has been generally considered as field capacity (Veihmeyer and Hendrickson 1931, Kirkham 2005). Thus 18% VWC was considered moisture at FC and 5–5.5% VWC (30% of FC) was regarded as severe drought stress. The experiment was arranged in a combined block design (CBD) containing two treatments, *i.e.*, control (90–100% FC) and drought (30% FC) with 15 plants per genotype per treatment. Each pot was sown with two seeds and thinned to one at 14 d after sowing (DAS). Drought treatment was initiated at 21 DAS and brought soil moisture content to 30% FC (13–15 d after stress initiation) and all observations and sampling were taken simultaneously. Drought treatment was monitored daily by *Delta 150* soil moisture meter. Leaf (top, the most fully expanded) samples from five biological replicates from all treatments were harvested and frozen immediately in liquid nitrogen and later stored in a –80°C freezer for biochemical analysis.

**Malondialdehyde (MDA) and reactive oxygen species assay:** Fresh leaf tissue were homogenized with phosphate buffer to prepare a 10%-aliquot solution and used for following bioassays in triplicate with required dilutions. MDA was measured to detect the rate of lipid peroxidation in maize leaves by the thiobarbituric acid reactive

substances (TBARS) assay method using the MDA assay kit (A003-3) obtained from *Nanjing Jiancheng Bioengineering Institute* (<http://www.njjcbio.com/>), China. In principle, thiobarbituric acid reacts with MDA to generate a red compound that has the maximum absorbance at 532 nm; the method was used with minor modifications and expressed as nmol ml<sup>-1</sup> of homogenate (Heath and Packer 1968, Dhindsa *et al.* 1981). Hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) content was estimated using commercial kit A064-1 manufactured and supplied by *Nanjing Jiancheng Bioengineering Institute*, China, based on the principle of its reaction with molybdate to form coordination complex and recording its absorbance at 405 nm in spectrophotometer plate reader. The amount of H<sub>2</sub>O<sub>2</sub> was calculated according to the user manual formula of the kit with little modifications and expressed as  $\mu\text{mol}(\text{H}_2\text{O}_2) 0.1 \text{ ml}^{-1}$  of homogenate.

**Total antioxidative capacity (TAOC):** Production of total enzymatic [superoxide dismutase (SOD), glutathione peroxidase (GPX), catalase (CAT), glutathione S-transferase (GST), *etc.*] and nonenzymatic (vitamins, amino acids, and metalloproteins) antioxidants were estimated by commercial A015 kit produced and supplied by *Nanjing Jiancheng Bioengineering Institute*, China. The kit is based on the principle that antioxidative compounds can reduce Fe<sup>3+</sup> to Fe<sup>2+</sup>, and Fe<sup>2+</sup> reacts with phenanthroline to produce a stable complex, which was measured at 520 nm to calculate TAOC and expressed as U 0.1 ml<sup>-1</sup> of homogenate.

**Photosynthesis and Chl fluorescence measurements:** Gas exchange and Chl fluorescence were measured in the youngest fully expanded leaf using a portable gas-exchange fluorescence system (GFS-3000 and PAM-Fluorometer 3056-FL, Heinz Walz GmbH, Germany). Gas exchange was measured between 9:00–11:00 h by setting an absolute CO<sub>2</sub> concentration of 400 ppm, relative humidity of 75%, cuvette temperature of 25°C, and PAR of 1,500  $\mu\text{mol}(\text{photon}) \text{ m}^{-2} \text{ s}^{-1}$ . A light-response curve was calculated for both drought and control plants and found constant at PAR of 1,500  $\mu\text{mol}(\text{photon}) \text{ m}^{-2} \text{ s}^{-1}$ . Assimilation rate ( $P_N$ ), transpiration rate ( $E$ ), water vapor conductance ( $g_{\text{H}_2\text{O}}$ ), and intercellular CO<sub>2</sub> concentration ( $C_i$ ) were calculated according to von Caemmerer and Farquhar (1981). Chl fluorescence parameters maximal quantum efficiency of PSII ( $F_v/F_m$ ), quantum yield of photosynthetic electron transport [ $\Phi_{\text{PSII}} = (F_m' - F)/F_m'$ ], nonphotochemical quenching [ $\text{NPQ} = (F_m - F_m')/F_m'$ ], and quantum yield of NPQ-related energy loss [ $Y_{\text{NPQ}} = (F/F_m') - (F/F_m)$ ] were calculated from the Chl fluorescence measurement according to Genty *et al.* (1989) and Baker (2008). All the Chl fluorescence measurements were done before dawn (dark-adapted) by exposing them to strong actinic light.

**Statistical analysis:** The means were compared between control and drought by two-way analysis of variance (ANOVA) and Tukey's post-hoc test, with  $p=0.05$  significance level. The positive relative or delta change

was calculated by subtracting values under control with drought or *vice versa* (whichever is applicable) and were divided by control values except for TAOC and were ranked from the lowest to the highest. Lower delta change suggests that lesser difference between control (C) and drought (T, treatment), and is considered a better performer for that respective trait. While for TAOC relative or delta change was calculated by subtracting values under drought with control and ranked from the highest to lowest value, indicating higher TAOC under drought is essential for a plant to scavenge ROS and avoid cellular damage. Pearson's correlation (two-tailed) and principal component analysis (PCA) biplot analysis for both genotypes and recorded parameters was performed in *Origin Pro 2022* (<https://www.originlab.com/>).

## Results

**Hydrogen peroxide:** The  $\text{H}_2\text{O}_2$  content drastically increased due to drought in all genotypes (Fig. 1A). Under control, GJ-2 showed the highest  $\text{H}_2\text{O}_2$  followed by GJ-1 and the least amount found in JH-18A-514-38, while in drought-treated samples, the highest  $\text{H}_2\text{O}_2$  was found in GJ-2 followed by GJ-3 while the least was found in YS-18B-150-43 and YS-18B-150-44. The positive delta change  $[(T - C)/C]$  was the lowest in GJ-1 followed by JH-19A-501-36 and highest in JH-18A-514-38 (Table 1). Genotypes were ranked according to a positive delta (from the lowest to the highest) for their comparison and interpretation. The delta change shows the relative decrease or increase of the value of the parameter in drought compared to its respective control. Further, under drought,  $\text{H}_2\text{O}_2$  showed a higher variation than that of control (Table 2). A summary of the mean and standard deviation (SD) of the measured parameters is presented in Table 2.

**Total antioxidants capacity:** The TAOC significantly increased in all genotypes under drought except JH-19A-501-23, JL-118, and GJ-1 (very marginal reduction was seen under drought) (Fig. 1B). Genotypes HD-6 and GJ-2 had the highest TAOC in control, whereas JL-118 was the lowest. Similarly HD-6 contained the highest TAOC in drought followed by GJ-2 and the lowest was seen in JL-118. Genotypes were ranked based on treatment-control (T-C) from the highest to lowest (Table 1). The highest TAOC content genotypes under drought may be considered good oxidative stress-balancing lines.

**Malondialdehyde:** Drought conditions significantly increased MDA content (Fig. 1C). Genotype JH-19A-501-23 contained the highest MDA followed by YS-18B-150-43 and the least in La-posta-seq in control, while under drought, GJ-3 had a higher MDA followed by YS-18B-150-43 and the least in La-posta-seq. The positive delta change  $[(T - C)/C]$  was lowest in JH-19A-501-23 followed by HD-6 and the highest in GJ-3 (Table 1). Genotypes were ranked according to positive delta (the lowest to the highest). Lower delta change is supposed to be more tolerant to drought.

**Net assimilation rate ( $P_N$ ):** The  $P_N$  was significantly reduced in all genotypes under drought (Fig. 2D). All genotypes significantly differed under control for  $P_N$ , but not in drought. GJ-14 had the highest  $P_N$  followed by YS-18B-150-43 and the lowest in La-posta-seq under control, while under drought, YS-18B-150-43 showed the highest  $P_N$  followed by YS-18B-150-44 and the lowest in GJ-2. However, HD-6 and JL-118 exhibited the lowest and highest positive delta change  $[(C - T)/C]$ , respectively (Table 1).

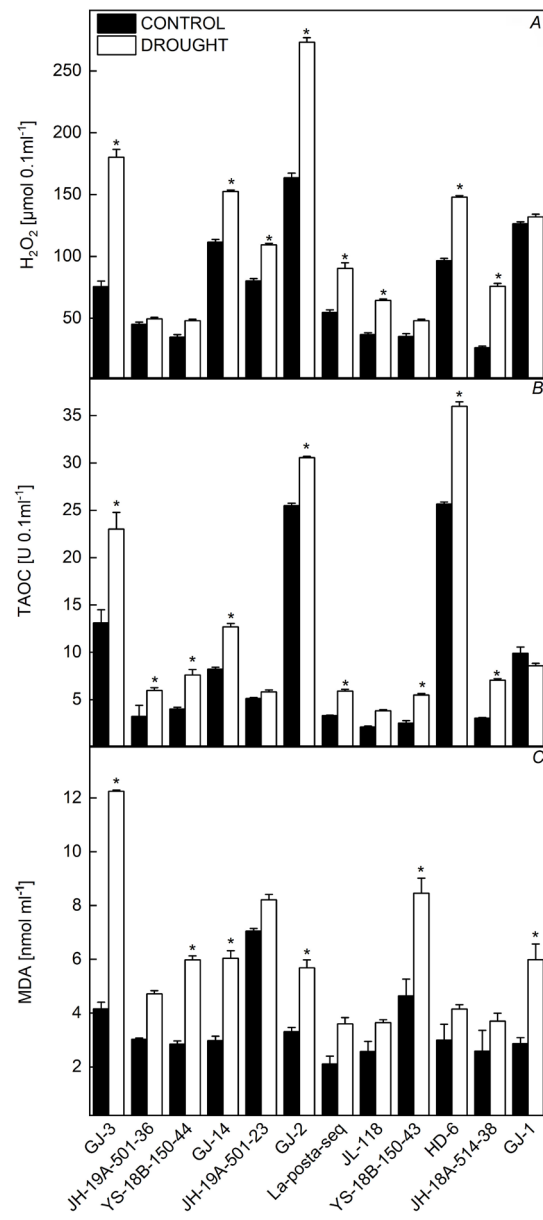


Fig. 1. Changes in (A) hydrogen peroxide ( $\text{H}_2\text{O}_2$ ), (B) total antioxidant capacity (TAOC), and (C) malondialdehyde (MDA) content in different maize genotype leaves in a greenhouse experiment. Values are means  $\pm$  SE,  $n = 5$ . \* indicates significant differences at  $P < 0.05$ .

Table 1. Ranking of genotypes based on their relative or delta change of measured parameters under drought (D) and control (C). All traits except TAOC are ranked from the lowest to the highest delta change values, while TAOC are ranked from the highest to lowest. Genotypes that showed lower delta change (less difference between control and drought) except for TAOC, were considered better performers for that respective trait. However, higher TAOC under drought compared to control is considered a better performer. HD-6 (bold) performed comparatively well under drought.

Rank	$H_2O_2$ (T - C)/C	MDA (T - C)/C	TAOC T - C	E (C - T)/C	VPD (T - C)/C	$g_{H2O}$ (C - T)/C	$P_N$ (C - T)/C	$C_i/C_a$ (T - C)/C	$F_v/F_m$ (C - T)/C	$\Phi_{PSII}$ (C - T)/C	NPQ (T - C)/C	$Y_{NPQ}$ (T - C)/C
1	GJ-1	JH-19A-501-23	<b>HD-6</b>	JH-19A-501-23	JH-19A-501-23	JH-19A-501-23	JH-19A-501-23	<b>HD-6</b>	<b>HD-6</b>	JH-19A-501-23	YS-18B-150-43	JH-19A-501-23
2	JH-19A-501-36	<b>HD-6</b>	GJ-3	GJ-3	JL-118	YS-18B-150-43	JH-18A-514-38	JH-19A-501-23	JH-19A-501-23	JH-19A-501-23	YS-18B-150-43	JH-19A-501-23
3	YS-18B-150-43	JL-118	GJ-2	JH-19A-501-36	JL-118	JH-19A-501-36	YS-18B-150-44	YS-18B-150-44	JH-19A-501-36	YS-18B-150-43	GJ-1	<b>HD-6</b>
4	JH-19A-501-23	JH-18A-514-38	GJ-14	JL-118	JH-19A-501-36	<b>HD-6</b>	YS-18B-150-43	La-posta-seq	YS-18B-150-43	JH-19A-501-36	GJ-14	GJ-1
5	GJ-14	JH-19A-501-36	JH-18A-514-38	GJ-14	JL-118	JH-19A-501-36	JH-19A-501-23	GJ-2	GJ-2	JH-18A-514-38	<b>HD-6</b>	JH-18A-514-38
6	YS-18B-150-44	GJ-2	YS-18B-150-44	YS-18B-150-43	GJ-1	YS-18B-150-44	JL-118	JH-19A-501-23	JH-19A-501-23	GJ-2	GJ-3	GJ-14
7	<b>HD-6</b>	La-posta-seq	YS-18B-150-43	<b>HD-6</b>	GJ-14	JH-18A-514-38	La-posta-seq	GJ-14	YS-18B-150-44	GJ-14	JH-18A-514-38	JH-19A-501-36
8	La-posta-seq	YS-18B-150-43	JH-19A-501-36	GJ-2	GJ-2	JH-19A-501-36	GJ-1	GJ-1	YS-18B-150-44	JH-19A-501-36	GJ-3	
9	GJ-2	GJ-14	La-posta-seq	GJ-14	GJ-3	YS-18B-150-44	GJ-1	JH-19A-501-36	GJ-14	GJ-1	YS-18B-150-44	GJ-2
10	JL-118	GJ-1	JL-118	GJ-1	YS-18B-150-43	La-posta-seq	GJ-14	JH-18A-514-38	JL-118	GJ-3	GJ-2	YS-18B-150-44
11	GJ-3	YS-18B-150-44	JH-19A-501-23	La-posta-seq	JH-18A-514-38	GJ-1	JH-18A-514-38	JL-118	GJ-3	JL-118	La-posta-seq	La-posta-seq
12	JH-18A-514-38	GJ-3	GJ-1	JH-18A-514-38	La-posta-seq	GJ-14	GJ-2	La-posta-seq	La-posta-seq	JL-118	JL-118	

**Air to leaf vapor pressure deficit (VPD):** The VPD was calculated according to [von Caemmerer and Farquhar \(1981\)](#) based on intercellular  $H_2O$  mole fraction within the leaf (ppm)( $w_i$ ) and  $H_2O$  mole fraction in the cuvette (ppm) ( $w_a$ ) and expressed as  $Pa\ kPa^{-1}$ . The formula for VPD calculated by default in *GFS-3000*,  $VPD = (w_i - w_a) / [1 - (w_i + w_a/2)]$ . VPD increased in all genotypes under drought compared to control. The variation for VPD was found to be higher in drought condition between genotypes compared to control ([Fig. 2B, Table 2](#)). Under control, JL-118 showed the highest VPD followed by JH-18A-514-38 and the lowest was in GJ-2, similarly under drought, JH-18A-514-38 had the highest VPD followed by La-posta-seq and the lowest was in GJ-2. The positive delta change  $[(T - C)/C]$  was the lowest in JH-19A-501-23, followed by HD-6, and the highest was in La-posta-seq ([Table 1](#)).

**The ratio of intercellular to ambient  $CO_2$  ( $C_i/C_a$ ):** The  $C_i/C_a$  ratio was used to evaluate stomatal acclimation ([Sage 1994](#)). All genotypes varied under both control and drought for  $C_i/C_a$  ratio ([Fig. 2E](#)). Genotype GJ-2 showed a higher  $C_i/C_a$  ratio followed by GJ-14 and the lowest in GJ-3 under control, while in drought, GJ-14 showed the highest followed by JH-19A-501-36 and the lowest in YS-18B-150-43. The lowest and highest positive delta change  $[(T - C)/C]$  was seen in HD-6 and GJ-3, respectively ([Table 1](#)).

**Transpiration rate (E):** Transpiration was reduced significantly under drought. Significant variation was found in transpiration between genotypes in both control and drought ([Fig. 2A](#)). Genotype GJ-14 showed the highest  $E$ , followed by JH-18A-514-38, and the lowest in GJ-1 in control, while under drought, JH-19A-501-23 had the highest  $E$ , followed by GJ-3, and the lowest was in GJ-1. The positive delta change  $[(C - T)/C]$  was the lowest in JH-19A-501-23, followed by GJ-3, and the highest was in JH-18A-514-38, and all the genotypes were ranked from the lowest to the highest ([Table 1](#)).

**Water vapor conductance ( $g_{H2O}$ ):** Significant variation was found for  $g_{H2O}$  between genotypes under both control and drought. It decreased significantly under drought compared to control ([Fig. 2C](#)). Genotype GJ-14 showed the highest  $g_{H2O}$  followed by GJ-2 and the lowest was seen in JH-18A-514-38 under control, while in drought, JH-19A-501-23 showed the highest  $g_{H2O}$  followed by JH-19A-501-36 and the lowest in JH-18A-514-38. Similarly, JH-19A-501-23 and GJ-14 exhibited the lowest and highest positive delta change  $[(C - T)/C]$ , respectively ([Table 1](#)).

**Maximal quantum efficiency of PSII ( $F_v/F_m$ ):** The  $F_v/F_m$  ratio differed considerably between control and drought ([Fig. 3A](#)). Generally,  $F_v/F_m$  ranges from 0 to 0.84. Genotype GJ-14 showed higher  $F_v/F_m$  followed by GJ-3 and the least by HD-6 under control, while in drought, it ranged from 0.638 (La-posta-seq) to 0.77 (YS-18B-150-43). The lowest and highest positive delta change  $[(C - T)/C]$

Table 2. Mean and standard deviation (SD) of the measured parameters under control and drought.  $C_i/C_a$  – the ratio of intercellular to ambient  $\text{CO}_2$  concentration;  $E$  – transpiration rate;  $F_v/F_m$  – maximal quantum efficiency of PSII;  $g_{\text{H}_2\text{O}}$  – water vapor conductance; MDA – malondialdehyde; NPQ – nonphotochemical quenching;  $P_N$  – net assimilation rate; TAOC – total antioxidant capacity; VPD – air to leaf vapor pressure deficit;  $\Phi_{\text{PSII}}$  – quantum yield of photosynthetic electron transport.

Parameter	Control		Drought	
	Mean	SD	Mean	SD
$\text{H}_2\text{O}_2$ [ $\mu\text{mol 0.1 ml}^{-1}$ ]	73.84	43.39	114.25	67.46
MDA [ $\text{nmol ml}^{-1}$ ]	3.43	1.33	6.04	2.55
TAOC [ $\text{U 0.1 ml}^{-1}$ ]	8.82	8.51	12.71	10.92
$E$ [ $\text{mmol}(\text{H}_2\text{O}) \text{ m}^{-2} \text{ s}^{-1}$ ]	2.05	0.39	0.85	0.39
VPD [ $\text{Pa kPa}^{-1}$ ]	22.49	5.57	27.14	8.88
$g_{\text{H}_2\text{O}}$ [ $\text{mol}(\text{H}_2\text{O}) \text{ m}^{-2} \text{ s}^{-1}$ ]	91.58	35.06	36.73	17.67
$P_N$ [ $\mu\text{mol m}^{-2} \text{ s}^{-1}$ ]	10.31	1.87	1.62	0.90
$C_i/C_a$	0.49	0.09	0.81	0.13
$F_v/F_m$	0.79	0.02	0.73	0.04
$\Phi_{\text{PSII}}$	0.78	0.02	0.70	0.05
NPQ	0.02	0.00	0.03	0.01

was observed in HD-6 and La-posta-seq, respectively (Table 1).

**Quantum yield of photosynthetic electron transport ( $\Phi_{\text{PSII}}$ ):** The  $\Phi_{\text{PSII}}$  was reduced in drought treatment in all genotypes (Fig. 3B). Genotype GJ-3 showed a higher yield followed by GJ-14 and the least by HD-6 under control, while in drought, it ranged from La-posta-seq (0.603) to YS-18B-150-43 (0.754). The positive delta change  $[(C - T)/C]$ , was the lowest in HD-6 followed by JH-19A-501-23 and the highest was observed in La-posta-seq (Table 1).

**Nonphotochemical quenching (NPQ):** The NPQ increased considerably in all genotypes under drought compared to control (Fig. 3C). NPQ was the lowest in JH-19A-501-36 followed by GJ-2 and the highest was in JH-19A-501-23 under control. Under drought, NPQ was the lowest in YS-18B-150-43 followed by JH-19A-501-36, and the highest was observed in JL-118. The lowest and highest positive delta change  $[(T - C)/C]$  was seen in JH-19A-501-23 and JL-118, respectively (Table 1).

**Quantum yield of NPQ-related energy loss ( $Y_{\text{NPQ}}$ ):** The  $Y_{\text{NPQ}}$  denotes the fraction of energy dissipated in the form of heat *via* the regulated nonphotochemical quenching mechanism. The  $Y_{\text{NPQ}}$  was higher in all genotypes under drought than that in control (Fig. 3D). Genotype YS-18B-501-36 showed the lowest  $Y_{\text{NPQ}}$  followed by GJ-2 and the highest was observed in JH-19A-501-23 under control. Under drought, genotype YS-18B-150-43 showed the lowest followed by YS-18B-501-36, and the highest was observed in JL-118. The positive delta change  $[(T - C)/C]$  was the lowest in YS-18B-150-43 and the highest in JL-118 (Table 1).

**Correlation and Principal Component Analysis (PCA):** Correlation was performed between traits measured under control (Table 3) and drought (Table 4) to check the pattern of the association.

Under control conditions, TAOC and  $\text{H}_2\text{O}_2$  were significantly correlated positively ( $r = 0.77^*$ ). Similarly,  $C_i/C_a$  and  $g_{\text{H}_2\text{O}}$  ( $r = 0.76^{**}$ ),  $F_v/F_m$  and  $P_N$  ( $r = 0.68^*$ ), and  $\Phi_{\text{PSII}}$  and  $F_v/F_m$  ( $r = 0.91^{**}$ ) were significantly associated positively. However,  $\text{H}_2\text{O}_2$  and VPD;  $C_i/C_a$  and VPD;  $g_{\text{H}_2\text{O}}$  and VPD were significantly negatively correlated. Under drought,  $\text{H}_2\text{O}_2$  and TAOC ( $r = 0.78^{**}$ ), MDA and  $E$  ( $r = 0.59^*$ ), and  $g_{\text{H}_2\text{O}}$  and  $E$  ( $r = 0.78^{**}$ ) were significantly positively correlated, and  $\Phi_{\text{PSII}}$  and NPQ were negatively correlated. Further PCA biplot analysis was performed separately for control (Fig. 4A) and drought (Fig. 4B). Three principal components for control covered more than 76% of the variation (Table 5) and its biplot showed that MDA and NPQ;  $\text{H}_2\text{O}_2$  and  $C_i/C_a$ ;  $\Phi_{\text{PSII}}$ ,  $F_v/F_m$ , and  $E$ ; and  $g_{\text{H}_2\text{O}}$  and  $P_N$  were grouped, while VPD spotted separately, which is mostly influenced by temperature and relative humidity. However, no specific clustering of genotypes was observed under control, suggesting genetic diversity of the genotypes used.

Under drought, three principal components covered 70% of the variation and there were four clusters, cluster 1 (NPQ, VPD), cluster 2 (MDA,  $g_{\text{H}_2\text{O}}$ ,  $C_i/C_a$ ), cluster 3 ( $F_v/F_m$ ,  $\Phi_{\text{PSII}}$ ), and  $P_N$  spotted separately (Fig. 4B). Further, more genotype clusters were seen in the drought biplot, where GJ-3 and JH-19A-501-23; JH-19A-501-36 and GJ-14; La-posta-seq and JL-118; and YS-18B-150-43 and JH-18A-514-38 were clustered, respectively, while HD-6 spotted separately towards  $F_v/F_m$  vector. Genotype La-posta-seq was spotted separately in the control biplot, while it clustered together with JL-118 in drought with higher NPQ and VPD. Genotype GJ-3 grouped with JH-19A-501-23 under drought towards  $E$  vector, but in control, it was in mid-VPD vector.

The NPQ and MDA showed higher variation in control compared to drought, while  $F_v/F_m$  and  $\Phi_{\text{PSII}}$  had greater variation in drought. A higher variation (mean sum of squares) in genotype  $\times$  environment interaction was observed in  $\text{H}_2\text{O}_2$ ,  $g_{\text{H}_2\text{O}}$ , VPD, TAOC, and MDA, suggesting that these traits are highly influenced by the drought (Table 6). GJ-2 maintained higher TAOC both in control and drought. Under control HD-6 showed higher  $E$ ,  $F_v/F_m$ , and  $\Phi_{\text{PSII}}$ , while under drought, HD-6 maintained higher photosynthetic efficiency and lower  $E$ , making it more drought tolerant. The lower photosynthetic efficiency of genotypes La-posta-seq and JL-118 could be due to higher NPQ and VPD under drought. YS-18B-150-44 was found to be the most stable genotype under both control and drought (near to origin in both biplots). Genotype GJ-14 was stable for both  $\text{H}_2\text{O}_2$  and  $C_i/C_a$  under both control and drought.

## Discussion

**Drought significantly increases membrane damage and total antioxidant capacity:** Hydrogen peroxide, though involved in various signaling processes (Singh *et al.*

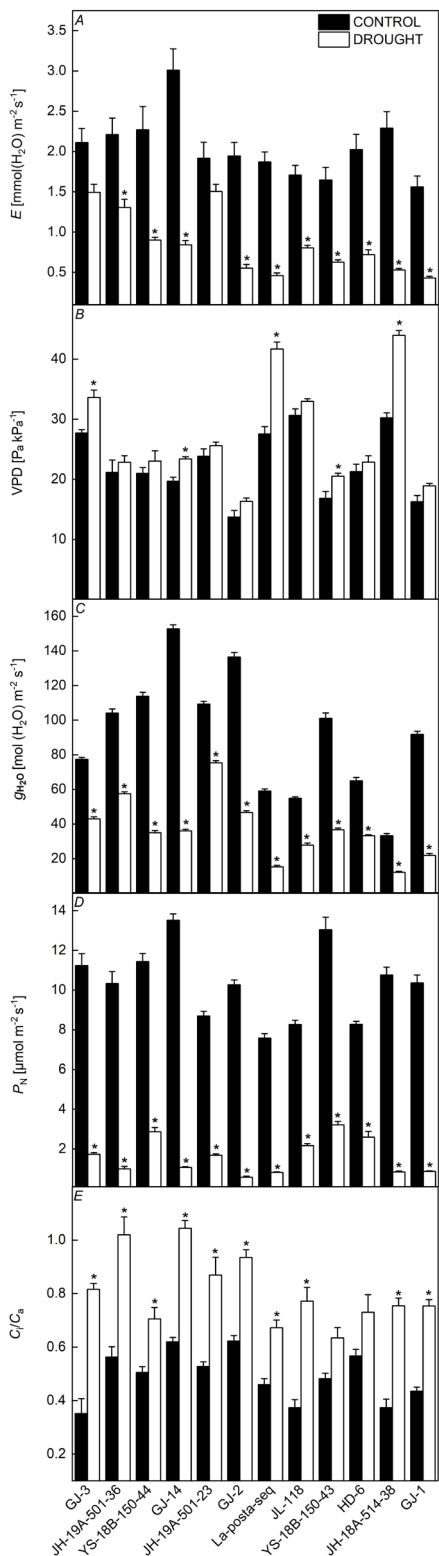


Fig. 2. Gas-exchange parameters of maize seedlings leaves under control and drought stress. (A) Transpiration rate ( $E$ ), (B) vapour pressure deficit (VPD), (C) water vapour conductance ( $g_{H_2O}$ ), (D) assimilation rate ( $P_N$ ), and (E) intercellular/ambient  $CO_2$  concentration ( $C/C_a$ ). Values represent means  $\pm$  SE,  $n = 5$ . \* indicates significant differences at  $P < 0.05$ .

2019), becomes toxic and causes damage to cells if it is not scavenged to maintain below damageable level (Dat *et al.* 2000).  $H_2O_2$  increased in all genotypes under drought suggesting stress signal induction and occurrence of drought-induced oxidative stress. It is also supported by a low correlation ( $r = 0.25$ ) between MDA and  $H_2O_2$  under drought suggesting  $H_2O_2$  is also involved in signal induction in addition to cell membrane damage (Smirnoff and Arnaud 2019). This is supported by GJ-1 and

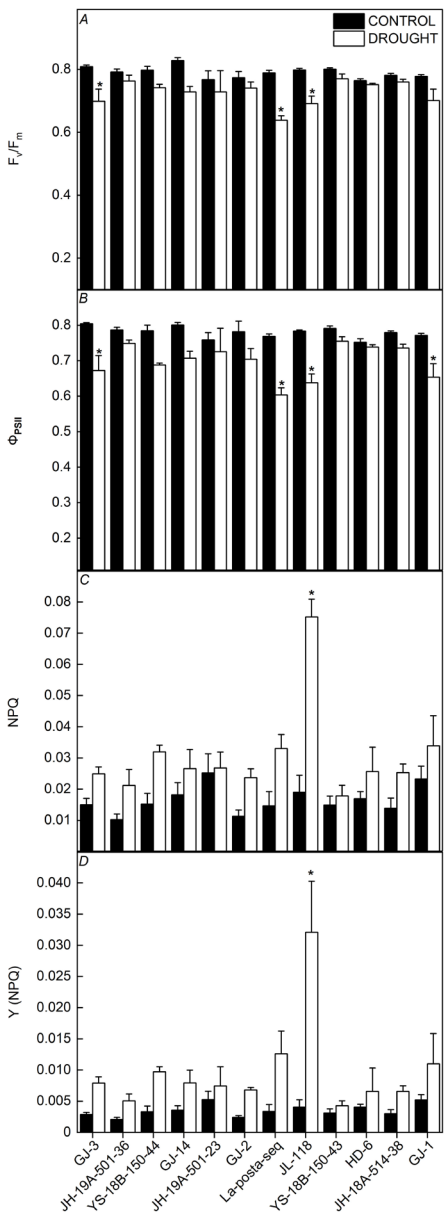


Fig. 3. Chlorophyll fluorescence parameters of maize seedlings leaves under control and drought stress. (A) Maximal quantum efficiency of photosystem II ( $F_v/F_m$ ), (B) quantum yield of photosynthetic electron transport ( $\Phi_{PSII}$ ), (C) nonphotochemical quenching (NPQ), and (D) quantum yield of NPQ-related energy loss ( $Y_{NPQ}$ ). Values represent means  $\pm$  SE,  $n = 5$ . \* indicates significant differences at  $P < 0.05$ .

Table 3. Pearson's correlation between recorded parameters under control conditions. \* Significant at  $P<0.05$  and \*\* significant at  $P<0.01$ .  $C_i/C_a$  – the ratio of intercellular to ambient  $\text{CO}_2$  concentration;  $E$  – transpiration rate;  $F_v/F_m$  – maximal quantum efficiency of PSII;  $g_{\text{H}_2\text{O}}$  – water vapor conductance; MDA – malondialdehyde; NPQ – nonphotochemical quenching;  $P_N$  – net assimilation rate; TAOC – total antioxidant capacity; VPD – air to leaf vapor pressure deficit;  $Y_{\text{NPQ}}$  – quantum yield of NPQ-related energy loss;  $\Phi_{\text{PSII}}$  – quantum yield of photosynthetic electron transport.

	$\text{H}_2\text{O}_2$	MDA	TAOC	$E$	VPD	$g_{\text{H}_2\text{O}}$	$P_N$	$C_i/C_a$	$F_v/F_m$	$\Phi_{\text{PSII}}$	NPQ	$Y_{\text{NPQ}}$
$\text{H}_2\text{O}_2$	1	0.067	0.768**	0.036	-0.633*	0.539	0.025	0.529	-0.023	-0.120	0.179	0.181
MDA	0.067	1	-0.035	-0.163	-0.131	0.291	0.069	0.108	-0.226	-0.136	0.463	0.353
TAOC	0.768**	-0.035	1	-0.001	-0.442	0.202	-0.133	0.434	-0.217	-0.246	-0.126	-0.057
$E$	0.036	-0.163	-0.001	1	0.057	0.395	0.483	0.395	0.528	0.418	-0.223	-0.322
VPD	-0.633*	-0.131	-0.442	0.057	1	-0.740**	-0.419	-0.697*	-0.045	-0.005	0.046	0.027
$g_{\text{H}_2\text{O}}$	0.539	0.291	0.202	0.395	-0.745**	1	0.549	0.762**	0.448	0.328	0.021	-0.087
$P_N$	0.025	0.069	-0.133	0.483	-0.419	0.549	1	0.170	0.682*	0.748**	-0.193	-0.326
$C_i/C_a$	0.529	0.108	0.434	0.395	-0.697*	0.762**	0.170	1	0.049	-0.145	-0.145	-0.137
$F_v/F_m$	-0.023	-0.226	-0.217	0.528	-0.045	0.448	0.682*	0.049	1	0.916**	-0.305	-0.459
$\Phi_{\text{PSII}}$	-0.120	-0.136	-0.246	0.418	-0.005	0.328	0.748**	-0.145	0.916**	1	-0.396	-0.564
NPQ	0.179	0.463	-0.126	-0.223	0.046	0.021	-0.193	-0.145	-0.305	-0.396	1	0.973**
$Y_{\text{NPQ}}$	0.181	0.353	-0.057	-0.322	0.027	-0.087	-0.326	-0.137	-0.459	-0.564	0.973**	1

Table 4. Pearson's correlation between recorded parameters under drought. \* Significant at  $P<0.05$  and \*\* significant at  $P<0.01$ .  $C_i/C_a$  – the ratio of intercellular to ambient  $\text{CO}_2$  concentration;  $E$  – transpiration rate;  $F_v/F_m$  – maximal quantum efficiency of PSII;  $g_{\text{H}_2\text{O}}$  – water vapor conductance; MDA – malondialdehyde; NPQ – nonphotochemical quenching;  $P_N$  – net assimilation rate; TAOC – total antioxidant capacity; VPD – air to leaf vapor pressure deficit;  $Y_{\text{NPQ}}$  – quantum yield of NPQ-related energy loss;  $\Phi_{\text{PSII}}$  – quantum yield of photosynthetic electron transport.

	$\text{H}_2\text{O}_2$	MDA	TAOC	$E$	VPD	$g_{\text{H}_2\text{O}}$	$P_N$	$C_i/C_a$	$F_v/F_m$	$\Phi_{\text{PSII}}$	NPQ	$Y_{\text{NPQ}}$
$\text{H}_2\text{O}_2$	1	0.250	0.782**	-0.050	-0.304	0.158	-0.428	0.383	-0.102	-0.070	-0.220	-0.205
MDA	0.250	1	0.136	0.595*	-0.198	0.463	0.254	0.037	0.055	0.135	-0.361	-0.335
TAOC	0.782**	0.136	1	-0.028	-0.308	0.089	-0.007	0.141	0.160	0.183	-0.291	-0.296
$E$	-0.050	0.595*	-0.028	1	-0.042	0.785**	0.149	0.426	0.122	0.249	-0.119	-0.126
VPD	-0.304	-0.198	-0.308	-0.042	1	-0.490	-0.209	-0.305	-0.451	-0.377	0.261	0.289
$g_{\text{H}_2\text{O}}$	0.158	0.463	0.089	0.785**	-0.490	1	0.096	0.533	0.330	0.451	-0.276	-0.289
$P_N$	-0.428	0.254	-0.007	0.149	-0.209	0.096	1	-0.533	0.310	0.254	0.103	0.086
$C_i/C_a$	0.383	0.037	0.141	0.426	-0.305	0.533	-0.533	1	0.210	0.268	-0.161	-0.171
$F_v/F_m$	-0.102	0.055	0.160	0.122	-0.451	0.330	0.310	0.210	1	0.949**	-0.487	-0.541
$\Phi_{\text{PSII}}$	-0.070	0.135	0.183	0.249	-0.377	0.451	0.254	0.268	0.949**	1	-0.606*	-0.649*
NPQ	-0.220	-0.361	-0.291	-0.119	0.261	-0.276	0.103	-0.161	-0.487	-0.606*	1	0.995**
$Y_{\text{NPQ}}$	-0.205	-0.335	-0.296	-0.126	0.289	-0.289	0.086	-0.171	-0.541	-0.649*	0.995**	1

JH-19A-501-36 which showed a minute increase in  $\text{H}_2\text{O}_2$  content in drought compared to control, thus ranking first and second respectively for  $\text{H}_2\text{O}_2$  production and scavenging. However, MDA was higher in GJ-1 under drought though it showed a marginal increase of  $\text{H}_2\text{O}_2$  under drought compared to control, while GJ-3 showed a significant increase in both  $\text{H}_2\text{O}_2$  and MDA under drought compared to control (Fig. 1A,C). An increase in  $\text{H}_2\text{O}_2$  content damages the cell membrane by reacting with lipids and proteins causing lipid peroxidation. MDA content indicates the level of lipid peroxidation and membrane damage (Porcel and Ruiz-Lozano 2004, Abid *et al.* 2018) but MDA content under drought is more positively associated with water loss (transpiration rate) ( $r = 0.595^*$ ) (Table 4), indicating stomatal closure is a drought-adaptive

mechanism (Cornic 2000). Likewise, a study on *Eugenia uniflora* L. also showed a strong correlation between MDA and  $E$  under water stress (Toscano *et al.* 2016).

Further, TAOC and  $\text{H}_2\text{O}_2$  had a significantly higher correlation both under control and drought suggesting that as  $\text{H}_2\text{O}_2$  increased, total antioxidants also increased to scavenge them. Likewise, Anjum *et al.* (2017) reported that drought stress increased  $\text{H}_2\text{O}_2$ ,  $\text{O}_2^-$ , and MDA compared to well-watered maize hybrids, and there was also increased activity of various enzymatic and nonenzymatic antioxidant activities under drought. Similarly, drought tolerance was associated with increased antioxidants and redox-regulating enzymes [catalase (CAT), ascorbate peroxidase (APX), glutathione reductase (GR), and glutathione peroxidase (GPX)] in maize (Avramova *et al.*

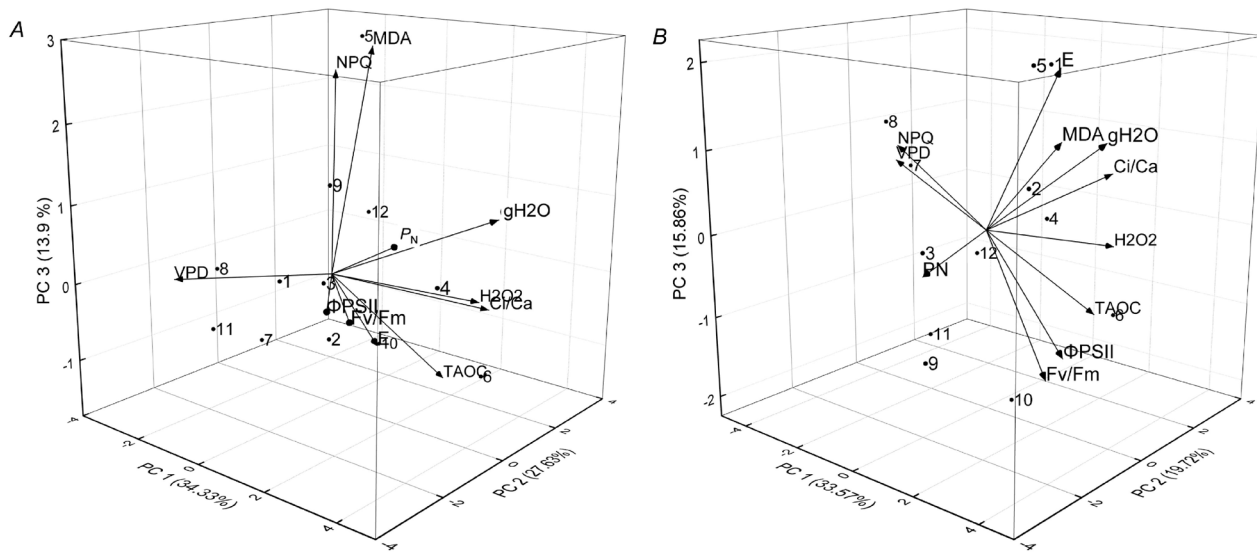


Fig. 4. Principal component (PC) biplot analysis of recorded parameters [hydrogen peroxide ( $H_2O_2$ ), total antioxidant capacity (TAOC), malondialdehyde (MDA) content, transpiration rate ( $E$ ), vapour pressure deficit (VPD), water vapour conductance ( $g_{H2O}$ ), assimilation rate ( $P_N$ ), intercellular/ambient  $CO_2$  concentration ( $C_i/C_a$ ), maximal quantum efficiency of photosystem II ( $F_v/F_m$ ), quantum yield of photosynthetic electron transport ( $\Phi_{PSII}$ ), and nonphotochemical quenching (NPQ)] and genotypes [1 – GJ-3, 2 – JH-19A-501-36, 3 – YS-18B-150-44, 4 – GJ-14, 5 – JH-19A-501-23, 6 – GJ-2, 7 – La-posta-seq, 8 – JL-118, 9 – YS-18B-150-43, 10 – HD-6, 11 – JH-18A-514-38, 12 – GJ-1] under control (A) and drought (B).

Table 5. Principal component (PC) analysis and percentage [%] variance coverage under control and drought.

PC	Control			Drought		
	Eigenvalue	Variance [%]	Cumulative [%]	Eigenvalue	Variance [%]	Cumulative [%]
1	3.7762	34.33	34.33	3.6932	33.57	33.57
2	3.0398	27.63	61.96	2.1691	19.72	53.29
3	1.5319	13.93	75.89	1.7450	15.86	69.16
4	0.8726	7.93	83.82	1.3644	12.40	81.56
5	0.7536	6.85	90.67	0.9394	8.54	90.10
6	0.5319	4.83	95.51	0.5860	5.33	95.43
7	0.3484	3.17	98.68	0.2693	2.45	97.88
8	0.0919	0.84	99.51	0.1521	1.38	99.26
9	0.0382	0.35	99.86	0.0583	0.53	99.79
10	0.0127	0.12	99.97	0.0206	0.19	99.98
11	0.0028	0.03	100.00	0.0027	0.02	100.00

Table 6. Mean sum of the square of all traits due to environment, genotype, and their interaction.  $C_i/C_a$  – the ratio of intercellular to ambient  $CO_2$  concentration;  $E$  – transpiration rate;  $F_v/F_m$  – maximal quantum efficiency of PSII;  $g_{H2O}$  – water vapor conductance; MDA – malondialdehyde; NPQ – nonphotochemical quenching;  $P_N$  – net assimilation rate; TAOC – total antioxidant capacity; VPD – air to leaf vapor pressure deficit;  $Y_{NPQ}$  – quantum yield of NPQ-related energy loss;  $\Phi_{PSII}$  – quantum yield of photosynthetic electron transport.

	$H_2O_2$	MDA	TAOC	$E$	$P_N$	$g_{H2O}$	VPD	$C_i/C_a$	$F_v/F_m$	$\Phi_{PSII}$	NPQ	$Y_{NPQ}$
Environment	32,431.29	134.459	294.100	25.974	1,385.66	54,645.43	383.507	1.866	0.089	0.153	0.004	$8.32 \times 10^{-4}$
Genotype	20,941.48	20.325	690.262	0.530	7.83	3,332.77	298.005	0.057	0.003	0.006	$7.30 \times 10^{-4}$	$1.72 \times 10^{-4}$
Interaction	2,113.90	6.006	18.983	0.339	6.13	1,287.62	30.120	0.018	0.004	0.005	$5.06 \times 10^{-4}$	$1.45 \times 10^{-4}$

2017). Concerning MDA and TAOC, HD-6 showed lower MDA and higher TAOC increase in drought from control. However, in the present study, no significant negative

correlation between MDA and TAOC was observed, which could be due to the operation of other membrane damaging substances other than ROS produced under drought or

disruption of the ROS-scavenging system under severe drought (Sharma *et al.* 2012). Based on the H<sub>2</sub>O<sub>2</sub> content, MDA, and total antioxidant content, HD-6 was observed to have a good balance of H<sub>2</sub>O<sub>2</sub> and lower cell membrane damage under drought. However, JH-19A-501-23 had the lowest MDA delta change, but showed less TAOC. Further, with respect to MDA and H<sub>2</sub>O<sub>2</sub> content, GJ-3 was found to be susceptible to drought stress.

**Severe drought stress imposes both stomatal and non-stomatal limitations on photosynthesis:** ROS are well-known causes of damage to photosystems of the plant (Khorobrykh *et al.* 2020). In the present study, the negative association of the assimilation rate with H<sub>2</sub>O<sub>2</sub> ( $r = -0.43$ ) and  $C_i/C_a$  ( $r = -0.53$ ) under drought (Table 4) could be due to disruption of D1 proteins of PSII by ROS (Mishra and Ghanotakis 1994, Miyao *et al.* 1995). The D1 protein is the main subunit of PSII (Barber *et al.* 1997) and is directly involved in photosynthetic electron transport (Edelman and Mattoo 2008). In HD-6 and JL-118,  $P_N$  is reduced by fewer units in drought compared to other genotypes from their respective control (Fig. 2D). The VPD, which is measured by the difference between water vapor pressure in the leaf and the water vapor pressure of the ambient air, gives an accurate idea of leaf water balance (Grossiord *et al.* 2020). Both genotypes HD-6 and JH-19A-501-23 showed relatively less delta change for VPD suggesting only a marginal increase of VPD in drought compared to control. An increase in VPD makes stomata smaller and reduces stomatal conductance to save water and limit photosynthesis (Grossiord *et al.* 2020). La-post-seq and JH-19A-514-38 showed significantly higher VPD and accordingly their  $P_N$  was the lowest among all genotypes under drought (Fig. 2B,D). Further, VPD had a negative correlation ( $r = -0.21$ ) with  $P_N$  under drought (Table 4). The photosynthetic rate is limited by both stomatal and nonstomatal limitations (Cossins and Chen 1997). During the initiation of drought or under mild to moderate drought stress, the decline in photosynthesis occurs mostly due to a reduction in the internal cellular CO<sub>2</sub> concentration by stomatal closure (Lawlor 2002). However nonstomatal limitations also reduce photosynthesis due to impaired metabolic processes, such as RuBP synthesis, ATP synthesis, and electron transfer (Cossins and Chen 1997). In the present study,  $C_i/C_a$  ratio significantly increased in all genotypes under drought stress, though there was a reduction in stomatal conductance. Similar increases in  $C_i/C_a$  ratio under severe drought were previously reported in rice (Ji *et al.* 2012), maize (Zhang *et al.* 2015), cowpea (Singh and Reddy 2011), and wheat (Martin and Ruiz-Torres 1992). Genotype HD-6 showed the lowest delta change with a marginal increase in internal CO<sub>2</sub> concentration under drought compared to control suggesting nonstomatal limitation was relatively lesser in HD-6 compared to others. The highest  $C_i/C_a$  delta change of GJ-3 could be due to pronounced production of H<sub>2</sub>O<sub>2</sub> and higher MDA content under drought compared to control (Table 1). Further,  $C_i/C_a$  showed a positive correlation ( $r = 0.38$ ) with H<sub>2</sub>O<sub>2</sub> and  $g_{H2O}$  ( $r = 0.53$ ) suggesting that

water loss increases drought stress and thus enhances H<sub>2</sub>O<sub>2</sub> production.

Concerning  $g_{H2O}$ , JH-19A-501-23 and GJ-3 showed lesser delta change. Drought stress usually reduces stomatal conductance and transpiration to save water, which reduces the CO<sub>2</sub> influx, thus reducing the final photosynthetic assimilation (Oukarroum *et al.* 2009, Pinheiro and Chaves 2011). The delta change of transpiration rate was also the lowest in JH-19A-501-23 and GJ-3 suggesting that under drought, the transpiration rate of JH-19A-501-23 did not compromise lot and its  $P_N$  [(C – T/C)] was also comparatively higher (Table 1). Overall, based on gas-exchange parameters, genotypes HD-6 and JH-19A-501-23 were found to be more tolerant than other genotypes.

**Drought disrupts photosynthesis and tolerant genotype (HD-6) shows comparatively higher PSII photochemical efficiency:** Chl fluorescence provides detailed information on the state of PSII and its response to environmental change (Murchie and Lawson 2013). The  $F_v/F_m$  ratio denotes maximal photochemical efficiency of PSII reaction centers (Butler 1978) and low  $F_v/F_m$  implies underutilization of light energy absorbed by PSII reaction centers (Fracheboud and Leipner 2003).

$F_v/F_m$  negatively correlated with VPD ( $r = -0.45$ ) suggesting high VPD may reduce photosynthetic efficiency under drought stress. A larger value of VPD was associated with a reduction in stomatal conductance (Dos Santos *et al.* 2017) and thus reduced photosynthesis (Shirke and Pathre 2004). Under drought, HD-6 showed only marginal reduction in  $F_v/F_m$  (0.751) compared to its respective control (0.764) (Fig. 3A, Table 1) implying relatively higher photosynthetic efficiency in HD-6 under drought. The delta change of  $F_v/F_m$  was also lowest for HD-6. However, in JH-19A-501-23,  $F_v/F_m$  ratio declined by 0.05 in drought compared to the control.

The  $\Phi_{PSII}$ , effective PSII quantum yield, denotes the fraction of absorbed energy used in photochemistry, which determines the efficiency of PSII (Tsai *et al.* 2019). It tells the proportion of the light used in PSII photochemistry (Murata 1992). It is affected by the rate of electron transport or the concentrations of electron acceptors, *e.g.*, NADP<sup>+</sup>, available at the acceptor side of PSI (Tsai *et al.* 2019).  $F_v/F_m$  showed a significantly high positive correlation ( $r = -0.95^{**}$ ) with  $\Phi_{PSII}$ . Genotypes HD-6 and JH-19A-501-23 showed a relatively low decline in the  $\Phi_{PSII}$  under drought compared to other genotypes (Fig. 3B, Table 1).

Nonphotochemical quenching (NPQ) is widely used to measure the nonradiative energy dissipation (Bilger and Björkman 1990). The surplus fraction of the absorbed light is dissipated as heat known as NPQ (Chen *et al.* 2019). Drought stress not only damages PSII but also the light-harvesting complex (Hura *et al.* 2007). NPQ is one of the strategies adopted by plants to mitigate damage by harmlessly quenching the excitation of chlorophyll within the light-harvesting antennae of PSII by converting excitation energy into thermal energy, which can then

be released (Kasajima *et al.* 2011), thus reducing the formation of free radicals. Likewise in the present study, under control conditions, NPQ positively correlated with MDA, but under drought, it was negatively correlated with  $H_2O_2$  ( $r = -0.22$ ) and MDA ( $r = -0.36$ ) (Table 5). Further, NPQ is negatively associated with  $\Phi_{PSII}$  ( $-0.606^*$ ) under drought, which could be due to NPQ-led downregulation of photosynthesis (Murchie and Lawson 2013) by competing with photochemistry for the absorbed energy (Bilger and Björkman 1990). In JH-19A-501-23, NPQ increased marginally under drought (1.06 times) compared to other genotypes (Fig. 3C), whereas in JL-118 under drought, NPQ increased around 4 times compared to control, implying comparatively less photosystem damage and downregulation of photosynthesis in JH-19A-501-23 under drought. However, in HD-6, under drought, NPQ increased 1.5 times compared to control. The  $Y_{NPQ}$  also decreased marginally under drought in YS-18B-150-43, JH-19A-501-23, and HD-6 compared to control.  $Y_{NPQ}$  denotes the fraction of energy dissipated in the form of heat *via* the regulated nonphotochemical quenching mechanism (Huang *et al.* 2012) and it also showed a negative correlation with  $\Phi_{PSII}$  ( $r = -0.65^*$ ).

Overall, based on biochemical, gas exchange, and chlorophyll fluorescence parameters, HD-6 performed well under drought followed by JH-19A-501-23.

**Principal component analysis and genotype  $\times$  environment interaction:** PCA for measured parameters under control revealed that  $H_2O_2$  and  $C_i/C_a$  clustered together, suggesting that internal  $CO_2$  influences  $H_2O_2$  accumulation. Higher  $CO_2$  concentration in soybean was reported to have accumulated higher  $H_2O_2$  in leaf tissues (Cheeseman 2006). Further,  $E$ ,  $F_v/F_m$ , and  $\Phi_{PSII}$  interact and influence  $CO_2$  assimilation and photosynthetic efficiency, while MDA accumulation could increase NPQ to remove excess light as heat. Previous studies have found that mutants lacking the capacity to induce NPQ are very sensitive to photoinhibition and drought stress (Cousins *et al.* 2002, Allorent *et al.* 2013).

However, under drought, clustering of MDA,  $g_{H2O}$ ,  $C_i/C_a$ , and  $E$  (Fig. 4B) suggests that higher stomatal conductance and transpiration increase membrane damage which finally leads to a collapse of photosystems (shown by the increase in  $C_i$ ). Further, VPD increases NPQ (grouped together) to remove excess heat and save the plant from photoinhibition, while under control, VPD is spotted alone in the control biplot. The assimilation rate was less influenced by other factors under drought suggesting the operation of other unmeasured processes. Furthermore, biplot analysis under control and drought showed that genotypes behaved differently in these two conditions. More elaborately, HD-6 had higher photosynthetic efficiency in drought, but under control, it was more aligned towards  $E$  (Fig. 4A), suggesting that increasing photosynthetic efficiency and reducing transpiration rate could be a better tolerance mechanism under drought. Supporting this, GJ-3, which was spotted near MDA and NPQ vectors in control, was shown to have a higher transpiration rate thus having reduced

photosynthetic efficiency and being relatively susceptible. A higher genotype  $\times$  environment interaction, the mean sum of the square of  $H_2O_2$ ,  $g_{H2O}$ , VPD, TAOC, and MDA (Table 6), suggests that these traits are highly influenced by the drought; thus, these traits may be important when selecting for drought-tolerant genotypes. Based on all measured parameters, HD-6 genotype performed well, with higher TAOC and the lower decrease in  $P_N$  and photosynthetic efficiency under drought compared to control.

**Conclusion:** Plants under drought stress responded through complex processes and interactions. Drought increased the generation of ROS and caused lipid peroxidation. Drought-tolerant genotypes balance the ROS by producing a higher amount of antioxidants to scavenge them thus reducing cell damage.  $CO_2$  assimilation rate was reduced by both stomatal and nonstomatal inhibition of photosynthesis under severe drought stress. An increase in transpiration rate was found to increase MDA content under drought. Likewise, MDA,  $E$ ,  $g_{H2O}$ , and  $C_i/C_a$  clustered together in PCA analysis showing their strong interactions. However, the assimilation rate associated with  $g_{H2O}$  under control conditions, but under drought, it was located separately in PCA, suggesting some other factors involved in increasing the assimilation rate under drought. Correlation analysis showed that assimilation rate was negatively associated with  $H_2O_2$  and  $C_i$  content under drought, implying that imbalance of ROS impairs and inhibits photosystems. Similarly, MDA was positively associated with NPQ in control, while under drought, MDA was strongly influenced by  $g_{H2O}$ , rather than NPQ. Thus looking for variation with higher stomatal adjustment, lower  $H_2O_2$  content and lesser  $C_i$  under severe drought could able to identify drought-tolerant donors in maize. Further, reducing transpiration rate and increasing photosynthetic efficiency ( $F_v/F_m$ ) under drought would be a better drought-adaptation mechanism for maize. Based on biochemical and photosynthetic parameters, HD-6 was observed to be more tolerant against severe drought stress and behaved differently than other genotypes. The holistic and multi-omics approach could reveal more information on the interaction of biochemical and photosynthetic parameters.

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