Photoprotection and optimization of sucrose usage contribute to faster recovery of photosynthesis after water deficit at high temperatures in wheat

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Abstract
Plants are increasingly exposed to events of elevated temperature and water deficit, which threaten crop productivity. Understanding the ability to rapidly recover from abiotic stress, restoring carbon assimilation and biomass production, is important to unravel crop climate resilience. This study compared the photosynthetic performance of two Triticum aestivum L. cultivars, Sokoll and Paragon, adapted to the climate of Mexico and UK, respectively, exposed to one week water deficit and high temperatures, in isolation or combination. Measurements included photosynthetic assimilation rate, stomatal conductance, in vitro activities of Rubisco (EC 4.1.1.39) and invertase (INV, EC 3.2.1.26), antioxidant capacity and chlorophyll a fluorescence. In both genotypes, under elevated temperatures and water deficit (WD38°C), the photosynthetic limitations were mainly due to stomatal restrictions and to a decrease in the electron transport rate. Chlorophyll a fluorescence parameters clearly indicate differences between the two genotypes in the photoprotection when subjected to WD38°C and showed faster recovery of Paragon after stress relief. The activity of the cytosolic invertase (CytINV) under these stress conditions was strongly related to the fast photosynthesis recovery of Paragon. Taken together, the results suggest that optimal sucrose export/utilization and increased photoprotection of the electron transport machinery are important components to limit yield fluctuations due to water shortage and elevated temperatures.
Abbreviations — A, net photosynthesis assimilation rate; cytINV, cytosolic invertase; ETR, electron transport rate; FRAP, ferric reducing antioxidant power; gs, stomatal conductance; LHCII, Light-harvesting complex II, LRWC, leaf relative water content; LWP, leaf water potential; NPQ, total non-photochemical quenching; PAR, Paragon; Qa, quinone A; Qb, quinone B; qN, non-photochemical quenching; qP, photochemical quenching; RCA, Rubisco activase; RH, relative humidity; RuBP-ribulose 1,5-biphosphate; SOK, Sokoll; SDW, soil dry weight; SFC, soil field capacity; SRWC, soil relative water content; TEAC, Trolox equivalents antioxidant capacity; TSP, Total soluble protein; vacINV, vacuolar invertase; Vi- Rubisco initial activity; Vt- Rubisco total activity; WD, water deficit; WD25°C, water deficit at 25°C; WD38°C, water deficit at 38°C; WW, well-watered; WW25°C, well-watered at 25°C; WW38°C, well-watered at 38°C.

Introduction

Global warming is a serious threat to crop production. Wheat is the world’s most harvested crop per area, however, wheat yield is below the average of the other major crops (e.g. maize and rice) being therefore only the second most-produced cereal grain, with 26% of the world share (FAOSTAT 2017). Around 40% of the global wheat yield fluctuations are explained by climatic variation, and heatwaves and drought are among the principal stressors (Deryng et al. 2014, Zampieri et al. 2017). Each degree-Celsius increase in global mean temperature reduces, on average, the global yield of wheat by 6% (Zhao et al. 2017). To improve wheat yield in a changing climate, and ensure food security for an increasing world population, it is essential to comprehend how wheat plants respond to fluctuations in temperature and water availability, and the mechanisms involved in fast recovery of plant growth upon relief from high temperatures and extended drought.

When subjected to high temperatures, plants usually use evaporative cooling to reduce leaf temperature (Carmo-Silva et al. 2012, Costa et al. 2013). However, in response to water shortage, higher plants close the stomata to limit water loss by transpiration. When both conditions are present, stomatal closure reduces transpiration and consequently the plant temperature rises and intercellular CO2 concentration decreases (Chaves et al. 2003, Carmo-Silva et al. 2012, Duque et al. 2013) . High temperatures and drought negatively affect photosynthetic CO2 fixation at different levels, depending on the stress intensity, decreasing biomass accumulation (Zandalinas et al. 2018, Lamaoui et al. 2018, Tricker et al. 2018, Raja et al. 2020). Even if high temperature increases the maximum rate (Vmax) of the primary carboxylation enzyme of C3 photosynthesis (Rubisco, EC 4.1.1.39), it also increases the inhibition of Rubisco by sugar phosphate derivatives and thus Rubisco activation state decreases (Salvucci and Crafts-Brandner, 2004a,b). The efficiency of Rubisco depends on the activity of Rubisco’s catalytic chaperone, Rubisco Activase (RCA), to promote the release of inhibitory sugar phosphates from active sites. However, RCA is extremely thermal sensitive and depends on the redox status and ADP/ATP ratio (Carmo-Silva et al. 2015). To improve plant tolerance to increased
temperatures, bioengineering approaches aiming to enhance Rubisco activity by increasing the thermotolerance of RCA have been suggested (Scafaro et al. 2016, Mueller-Cajar 2017, Shivhare and Mueller-Cajar 2017, Scafaro et al. 2019, Degen et al. 2020). Lower internal CO₂ concentration and high temperatures also reduce Rubisco specificity for CO₂ relative to O₂, resulting in an increase of photorespiration, which leads to the release of previous fixed CO₂ and higher demand for ATP (Walker et al. 2016).

Moreover, imbalances between CO₂ assimilation and the rate of light capture usually lead to an excess of energy in the system that can result in reactive oxygen species (ROS) formation and photoinhibition if the capacities of dissipation, scavenging and repairing are exceeded (Yamamoto 2016). Among the main energy dissipation mechanisms are the non-photochemical quenching (qN), generally compartmented in three major components, energy-dependent quenching, qE, state-transition quenching, qT, and photoinhibition quenching, ql), cyclic electron flow around photosystem I and chlororespiration (Rumeau et al. 2007, Ruban 2016, Wang and Fu 2016). ROS detoxification is generally conducted enzymatically and by the production of several antioxidant compounds (Mittler et al. 2004; Foyer 2018; Begum et al. 2019) When energy dissipation and ROS detoxification fails, oxidative damage occurs. Many studies reported the reduction of the electron transfer from water to NADP⁺, due to reversible and irreversible inhibition of photosystem II (PSII) caused by oxidative stress in face of elevated temperatures and/or drought. The main processes involved are the damage of the oxygen-evolving complex (Heckathorn et al. 1998, Tiwari et al. 2008, Chen et al. 2016), the degradation and aggregation of the D1 protein (Kamata et al. 2005, Komayama et al. 2007, Allakhverdiev et al. 2008, Takahashi and Murata 2008) and changes on the membrane fluidity (Gounaris et al. 1983, Aronsson et al. 2008, Yamamoto 2016a).

Therefore, when photosynthetic performance and plant growth are challenged by water shortage and elevated temperatures, optimization of sucrose export, uptake, and utilization, e.g. through adjustment of source – sink relations via invertase activity (INV, EC 3.2.1.26), can contribute to reducing yield fluctuations. Invertases mediate the hydrolytic cleavage of sucrose into hexose monomers and are involved in regulating carbohydrate partitioning, developmental processes, hormone responses and biotic and abiotic interactions (Roitsch and González 2004). Invertases localized in the vacuole (VacINVs) play a major role in the osmotic regulation (Nägele et al. 2010, Ruan 2014, Weiszmann et al. 2018), while cytosolic invertases (CytINVs) control sugar homeostasis and the maintenance of constant glucose levels to sustain cellular functions (Ruan et al. 2010, Lunn 2016, Figueroa and Lunn 2016).

The aims of the present study were to (1) characterise the photosynthetic limitations of two wheat genotypes, Paragon and Sokoll, adapted to distinct climate conditions, under water deficit and/or high temperature, and (2) to determine which factors are responsible for photosynthetic performance and recovery from high temperature in the absence or presence of water deficit. To test the hypothesis that the UK-adapted cultivar Paragon would be less resistant to heat stress and water deficit compared
to the Mexican-adapted cultivar Sokoll, the two genotypes were subject to water deficit and elevated temperatures, in isolation or in combination, and compared for net assimilation rate, stomatal conductance, Rubisco and invertase in vitro activities, antioxidant capacity and chlorophyll a fluorescence.

Materials and methods

Plant growth conditions

Two Triticum aestivum L. (wheat) genotypes were selected on the basis that these are adapted to distinct climate conditions: Paragon is a traditional UK spring wheat elite cultivar, while Sokoll is a synthetic-derived cultivar developed by the International Maize and Wheat Improvement Centre (CMMYT, Mexico). Plants of both genotypes were grown from seeds in a controlled environment chamber (Fitoclima 5000 EH, Aralab) in 1-L pots containing horticultural substrate (Compo Sana Universal, Compo Sana). Light was provided by fluorescent lamps (Osram Lumilux L 58W/840 cool white lamps) placed at specific distances from the plants to obtain an average photosynthetic photon flux density (PPFD) of 300 μmol m⁻² s⁻¹ at the top of the canopy, with a photoperiod of 16 h. Due to space constraints, temperature assays were performed in two consecutive experiments. After full germination, all plants were initially grown under a control temperature (25/18°C day/night), with 50% relative humidity (RH) for 21 days.

For experiments under control temperature, plants remained at 25/18°C (day/night) with 50% RH throughout the experiment. Three weeks post-germination plants were randomly assigned to two irrigation treatments: five plants per cultivar were maintained well-watered (WW; minimum 80% field capacity, WW25°C) throughout the experiment and five plants were subject to water deficit (WD, 30±5% field capacity, WD25°C) for 7 days. For experiments under elevated temperature, 21-day-old plants were also exposed to high temperatures (38/31°C day/night) with 60% RH and randomly assigned to the irrigation treatments: ten plants per cultivar were maintained WW (80±5% field capacity, WW38°C) and ten plants were subject to WD (30±5% field capacity, WD38°C) for 5 days. From the 10 plants allocated to WW38°C or WD38°C, 5 were randomly selected for recovery after 5 days of stress, re-watered and maintained at control temperatures for 7 days. WD was established by withholding watering and sustaining a minimum of 30±5% field capacity. The soil water content was determined gravimetrically by weighing the pots, and irrigation was provided to compensate evapotranspiration and keep the field capacity in the WW and WD pots. Leaf samples for biochemical analyses were collected at the end of the respective temperature and irrigation treatment, 5-7 h after the beginning of the photoperiod, frozen into liquid nitrogen and stored at -80°C.

Leaf and soil water status

Plant water status was estimated by leaf relative water content (LRWC) following the methodology described by Čatský (1960). Fresh leaf samples from the flag leaf (1-2 cm²) were collected, fresh weight
was immediately measured in an electronic scale (Sartorius BP221S), turgid weight (LTW) was determined after saturating samples by immersion in deionized water overnight, and dry weight (LDW) was measured after oven-drying samples at 70°C for 48 h. Soil relative water content (SRWC) was determined by following a similar procedure; although soil field capacity (SFC) was achieved by watering the pots to saturation and allowing water drainage for 2 hours, and dry weight (SDW) was measured after oven-drying samples at 110°C for 36 h. Leaf water potential was measured with a C-52 thermocouple chamber (Wescor), 20 mm² leaf discs were cut and equilibrated for 30 min in the chamber before the readings were recorded by a PSYPRO water potential datalogger (Wescor) in the psychrometric mode.

**Thermal imaging**

Thermal images were obtained using a thermal camera (Flir 50bx, FLIR Systems Inc.) with emissivity set at 0.95 and approximately 1 m distance from the plants. Before each set of measurements, background temperature was determined by measuring the temperature of a crumpled sheet of aluminium foil in a similar position to the leaves of interest with the emissivity set at 1.0 following the methodology described by Costa et al. (2013). Thermal images were analysed with the software FLIR Tools (FLIR Systems, Inc.). The temperature of each plant was determined from the temperature of five leaves using the function “area”. Visible images (RGB) were collected to complement the analysis of thermal images.

**Gas exchange and chlorophyll a fluorescence steady-state measurements**

Parallel measurements of photosynthetic gas exchange and chlorophyll a fluorescence were performed in a non-detached fully expanded leaf from each plant using a gas exchange system (IRGA LCpro+, ADC BioScientific) combined with a chlorophyll fluorescence imaging system (Imaging-PAM Chlorophyll Fluorometer M-series Mini version, Heinz Walz GmbH). Control air temperature was set to 25°C, PPFD at the leaf level set to 226 μmol m⁻² s⁻¹ and the CO₂ concentration in the leaf chamber set to 400 μmol CO₂ mol⁻¹ air allowing the leaf to reach steady-state assimilation rate (A) and stomatal conductance (gs). A and gs were calculated by the LCpro+ software according to von Caemmerer and Farquhar (1981). Chlorophyll a steady-state fluorescence was analysed using the Imaging Win analytical software (Heinz Walz GmbH). PSII effective quantum yield (ΦPSII) was obtained according to Genty et al. (1989), photochemical (qP) and non-photochemical (qN) quenching were calculated according to Oxborough and Baker (1997) and total non-photochemical fluorescence quenching (NPQ) was calculated using the Stern-Volmer approach (Krause and Jahns 2007). Electron transport rate (ETR) was then calculated as: $ETR = 0.5ΦPSII \times PPFD \times abs$. Absorptivity (abs) was measured for each leaf before the chlorophyll a fluorescence measurement.

**Chlorophyll a fluorescence induction**
The kinetics of the rapid fluorescence induction rise was recorded on fully expanded dark-adapted leaves (10 minutes) exposed to a saturating light pulse (3500 μmol m$^{-2}$ s$^{-1}$) for 1 second to obtain the OJIP Chl a fluorescence transient rise (Handy PEA, Hansatech Instruments). Fluorescence parameters derived from the extracted data, namely specific energy fluxes per QA-reducing PSII reaction center and photosynthetic performance indexes were calculated according to Strasser and collaborators (Strasser et al. 2004, Tsimilli-Michael and Strasser 2008) with the nomenclature presented in Stirbet and Govindjee (2011).

**Antioxidant capacity**

Antioxidant metabolites were extracted from frozen leaf samples (0.1-0.3 g FW) by homogenisation in pure methanol with 1.4 mm zirconium oxide beads (Precellys) in a tissue homogenizer (Precellys Evolution, Precellys) and then centrifuged at 20 000 g for 5 min. Trolox equivalents antioxidant capacity (TEAC) and ferric reducing antioxidant power (FRAP) were measured in the supernatant using a 96-well microtiter plate. TEAC was determined by the reaction of the sample supernatant and 2,2’-Azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS), solution 1:20 in phosphate buffer pH 7.4 (0.7-0.8 optical density). The reaction mixtures were incubated 6 min at room temperature before measuring absorbance at 734 nm (ELx808, BioTek Instruments, Inc.). 6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid (Trolox) standards (0-0.8 mM in 96% ethanol) were measured alongside the samples and used to prepare the respective calibration curve. FRAP was measured by the reaction of the sample supernatant with a solution consisting of 0.3 mM acetate buffer, 10 mM 2,4,6-tripyridyl-s-triazine (TPTZ) and 20 mM FeCl$_3$. The reaction mixtures were incubated 4 min at room temperature before measuring the absorbance at 593nm (ELx808, BioTek Instruments Inc.). FeSO$_4$ standards (0-1.0 mM) in ddH$_2$O were measured alongside the samples and used to prepare the respective calibration curves. Samples and standards were measured in triplicate alongside blanks containing no sample.

**Rubisco activity**

Rubisco was extracted from the leaves by grinding frozen samples (0.1-0.3 g FW) in a cold mortar with quartz sand, 1% (w/v) insoluble polyvinylpyrrolidone (PVP), ice-cold extraction medium (1/10 FW per mL) containing 50 mM Bicine-KOH pH 8.0, 1 mM ethylenediaminetetraacetic acid (EDTA), 5% (w/v) polyvinylpyrrolidone (PVP25000), 6% polyethylene glycol (PEG$_{4000}$), 10 mM 1,4-dithiothreitol (DTT), 50 mM β-mercaptoethanol and 1% (v/v) protease inhibitor cocktail for plant extracts (Sigma-Aldrich), adapted from Carmo-Silva et al. (2010). Leaf extracts were then centrifuged at 14 000 g and 4°C for 5 min. The supernatant was kept at 4°C and used immediately for measurement of Rubisco activities by the incorporation of $^{14}$CO$_2$ into acid-stable products at 25 and 38°C, following the protocol described in Parry et al. (1997) with modifications. The reaction mixture contained 100 mM Bicine-NaOH pH 8.2, 40 mM MgCl$_2$, 10 mM NaH$^{14}$CO$_3$ (7.4 kBq μmol$^{-1}$) and 0.4 mM ribulose 1,5-bisphosphate(Ru BP).
Rubisco initial activity (Vi) was determined by adding the supernatant to the mixture and stopping the reaction after 60-180s with 10 M HCOOH. Total activity (Vt) was measured after incubating the same volume of extract for 3 min with all the reaction mixture components except RuBP, to allow carbamylation of all the Rubisco available catalytic sites. The reaction was then started by adding RuBP and stopped as above. All measurements were carried out in triplicate and control reactions were quenched with HCOOH prior to the addition of RuBP. The mixtures were completely dried at 70°C overnight and the residues re-hydrated in 0.5 mL ddH₂O, then mixed with 5 mL scintillation cocktail (Ultima Gold, Perkin-Elmer). Radioactivity due to ¹⁴C incorporation in the acid-stable products was measured by liquid scintillation counting (LS7800, Beckman). The activation state of Rubisco was calculated as the ratio Vi / Vt × 100. Total soluble protein (TSP) content was determined according to the Bradford method (Bradford 1976) using BSA Fraction V as standard protein.

**Invertase activity**
Cytosolic invertase (CytInv) and vacuolar invertase (VacInv) were extracted from the leaves by grinding frozen samples (0.1-0.3 g FW) in a cold mortar with quartz sand, 1% (w/v) PVPP, ice-cold extraction medium containing 40 mM TRIS-HCl pH 7.6, 3 mM MgCl₂, 1 mM EDTA, 0.1 mM phenylmethylsulfonyl fluoride (PMSF), 1 mM benzamidine, 14 mM β-mercaptoethanol, 24 μM nicotinamide adenine dinucleotide phosphate (NADP⁺), according to Jammer et al. (2015), with modifications. Leaf extracts were then centrifuged at 20 000 g for 10 min at 4°C. The supernatant was kept at 4°C and dialysed overnight with 20 mM potassium phosphate buffer pH 7.4 at 4°C in a dark room. Extracts were aliquoted, frozen in liquid nitrogen and stored at -20°C. The activities were measured in thawed samples using 96-well microtiter plates. Reaction mixtures containing 10 mM sucrose and dialysed protein extract were incubated for 30 min at 37°C, cooled for 5 min on ice to stop the reaction, and then incubated for 30 min at room temperature with GOD-POD reagent (10 U mL⁻¹ of Glucose oxidase from *Aspergillus niger* (GOD), 0.8 U mL⁻¹ peroxidase from horseradish (POD) and 0.8 mg mL⁻¹ ABTS in 0.1 M potassium phosphate buffer (pH 7.0). The amount of liberated glucose was determined by measurement of absorbance at 405 nm at 30°C (ELx808, BioTek Instruments Inc.). Glucose standards (0-50 nmol) were measured alongside the samples and used to prepare the respective calibration curves. All measurements were carried out in triplicate alongside blanks containing no sucrose. TSP content was determined according to the Bradford method (Bradford 1976) using BSA Fraction V as standard protein.

**Statistical analysis**
The statistical significance of trait variation was tested by factorial ANOVA, with cultivars, irrigation and temperature regimes as fixed factors. Post-hoc comparison between treatments was performed with Duncan test (P < 0.05) using IBM SPSS Statistics, Version 25 (IBM, USA). Multivariate analysis was performed with MixOmics R package (Rohart et al. 2017) using Rstudio software.
Results

Leaf and soil water relations under drought and high temperatures

To characterise the leaf and soil water status of Sokoll and Paragon plants, leaf and soil relative water content (LRWC and SRWC, respectively) and leaf water potential (LWP) were estimated at the end of each experimental condition (Table 1). Well-watered (WW) plants presented leaf relative water content (LRWC) and leaf water potential (LWP) around or above 80% and -1 M Pa, respectively, suggesting good cellular hydration. On the other hand, water deficit (WD) conditions led to a decrease in LRWC and LWP values (lower than 70% and -1 MPa, respectively), revealing a reduction in hydration and a considerable driving force for water movement through the plant. Under WD25°C, Paragon presented higher LRWC than Sokoll, even though no significant differences were found for LWP and soil relative water content (SRWC), showing the capacity of this genotype to maintain cellular hydration under these conditions. The canopy temperature (Tcanopy) increased in both cultivars when subject to high temperatures. Under WW38°C, Tcanopy was significantly lower in Sokoll compared to Paragon, indicating the ability of Sokoll to avoid heat and maintain optimal cell temperature. No differences were observed between the genotypes when subjected to WD38℃, the observed LRWC under 50% and low LWP indicate severe drought stress, and Tcanopy was also highest in these plants.

Effects of drought and high temperature on photosynthesis

WD plants had significantly lower net photosynthesis assimilation rate (A), stomatal conductance (gs) and electron transport rate (ETR) compared to WW plants, except for Paragon at 25°C (Fig. 1A-C). Steady-state photosynthetic gas-exchanges were comparable for both genotypes under WW conditions. A strong positive relationship between A and gs was observed (r=0.914, P<0.0001 and r=0.974 P<0.0001, Paragon and Sokoll respectively, Table S1), suggesting a possible stomatal limitation to photosynthesis, and between A and ETR (r= 0.966, P<0.0001 and r=0.797, P<0.0001, Table S1), suggesting limitations at the photosystems level.

Effect of water deficit and high temperatures on Rubisco in vivo activities measured at control and high temperatures

To verify if the limitations in the carbon fixation found under stress conditions were a result of an imbalance in the Calvin-Benson-Bassham cycle, the in vivo Rubisco activity was assessed at the two growth temperatures. When Rubisco activity was measured at 25°C, the initial and total velocities decreased significantly under WD (WD25°C and WD38°C) and elevated temperatures (WW38°C) (Fig. 2A,B). However, the activation state of Rubisco remained largely unchanged between the various conditions (Fig. 2C). When Rubisco assays were performed at 38°C, activities were higher compared to measurements at 25°C, although the increase of initial velocity was higher than in total velocity (Fig. 2D,E). A significant difference was also observed between plants grown at 38°C under different
irrigation regimes. No significant differences were observed in Rubisco activation state when measured
at this temperature (Fig. 2F). The lack of differences in net photosynthetic assimilation rate of WW38°C
plants (Fig. 1A) would indicate that even the reduced level of Rubisco activity in these plants (~10 µmol
CO₂ m⁻² s⁻¹, Fig. S1D) is sufficient to support photosynthesis at the growth light levels (PPFD <300
µmol photons m⁻² s⁻¹).

**Effect of water deficit and high temperatures on the antioxidant capacity and chlorophyll a
fluorescence**

To analyse how both genotypes cope with possibly harmful consequences caused by energy excess
under stress, chlorophyll a fluorescence and two dissipation mechanisms, ROS scavenging and non-
photochemical dissipation, were quantified. A decrease of photochemical quenching (qP) was observed
in Sokoll WD25°C and in both genotypes at WD38°C (Fig. 3A-B). Under the same conditions, non-
photochemical quenching (qN, NPQ) increased (Fig. 3C-D). Moreover, the two genotypes showed an
increase in the antioxidant capacity (FRAP and TEAC) under drought at both temperatures (Fig. 3E,
F). In order to thoroughly understand how the different biochemical processes in the photosystems are
affected by stress conditions, the chlorophyll a kinetic parameters were correlated with the antioxidant
capacity and NPQ, and ETR (Fig. 4). A positive correlation was observed between the antioxidant
capacity and NPQ, as well as an inverse correlation to ETR. In all conditions, Sokoll showed a stronger
correlation between the number of electron carriers per electron transport chain (Sₘ) and ETR than
Paragon. The strength of the correlation between energy fluxes (JₑABS, JₑDI, JₑET2 and JₑRE1), ETR and NPQ
changed for both genotypes under WD (Fig. 4A,C). This was particularly the case in Paragon in
WD38°C (Figs 4C,S2 and Table S2), supported by the increase of JₑABS, JₑDI and JₑRE1 to control
conditions. In Sokoll the positive correlation between ETR and both electron transport fluxes (JₑET2 and
JₑRE1, Fig. 4C) indicated a decrease of electron transport rate on the entire flux until photosystem I.

**Recovery from high temperatures conditions**

Following 5 days of exposure to high temperatures and/or drought, wheat plants were allowed to recover
for 7 days (at 25°C and WW) and their photosynthetic performance was compared by measuring
chlorophyll a fluorescence, net photosynthetic assimilation and stomatal conductance. Even though no
differences were detected on the fraction of open PSII reaction centres (qP, Fig. 5A,B), a significant
increase on the non-photochemical quenching was observed relative to control (qN, NPQ, Fig. 5A,C,D).
The increase in NPQ was only accompanied by a decrease in the electron transport rate of Sokoll
recovering from WD38°C (Fig. 5E). Paragon presented higher LRWC and LWP when recovering from
WD38°C than Sokoll (Table 1), even though no significant differences were found, indicating a higher
capacity of this genotype to return to control cellular hydration and recover the driving force for water
movement through the plant. Slower recovery of Sokoll ETR and higher NPQ suggest that WD is
promoting photoinhibition in Sokoll. The photosynthetic assimilation rate and stomatal conductance
(Fig. 5F,G) increased in Paragon plants recovered after growing at 38°C in WW and WD conditions relative to control. However, in Sokoll, the photosynthetic assimilation rate decreased significantly in recovery from WD38°C and gs decreased when recovering from both conditions. All parameters reflecting the photosynthetic capacity revealed a better recovery from WD38°C in Paragon compared to Sokoll. Once again, results suggest that stomatal conductance impairment and recovery are a limiting factor for photosynthesis rate under water deficit and high temperature.

Invertase in vivo activities under water deficit and high temperatures

To verify if other sources of energy were used to cope with stress besides the direct usage of photoassimilates, the activity of invertases isoenzymes (located in the cytosol and vacuole) were measured. Results showed that the activity of vacINV was higher in Paragon for all the conditions compared to Sokoll (Fig. 6A). However, modulation of cytINV was observed according to different stress conditions (Fig. 6B): the cytINV activity increased in plants growing at 38°C with an interesting difference between WD38°C to WW38°C and WW25°C in Paragon. Even though the CytINV activity slightly increased, no significant differences were found for all conditions in Sokoll (Fig 6B). Overall, in Paragon, cytINV was negatively correlated to the assimilation rate ($r=-0.774$, $P<0.0001$, Table S1). Together with the previous results that showed a better recovery of this genotype after the combination of water deficit and high temperature, these data suggest that an increase of sucrose catabolism, when the production of photosynthetic assimilates decreases, improved wheat recovery from stress conditions.

Discussion

Two wheat cultivars, Paragon and Sokoll, were studied for their ability to withstand water deficit and high temperatures, in isolation or in combination. Paragon is a traditional UK spring wheat elite cultivar (Moore 2015), while Sokoll is a synthetic-derived cultivar developed by the International Maize and Wheat Improvement Centre (CIMMYT, Mexico), known to show good productivity under elevated temperatures (Solís Moya and Camacho Casas 2016). As these genotypes are adapted to distinct environmental conditions, it is of relevance to determine which factors are responsible for their photosynthetic performance. Therefore, the present study aimed to first characterise the photosynthetic limitations of the two genotypes under water deficit and/or high temperature and then to assess photosynthetic recovery from high temperature in the absence or presence of drought. To achieve this goal, Paragon and Sokoll were compared using several established parameters, namely net assimilation rate, stomatal conductance, Rubisco and invertase in vitro activities, antioxidant capacity and chlorophyll $a$ fluorescence.

Under increased temperatures a natural heat avoidance strategy of plants is to decrease leaf temperature through increased transpiration (Carmo-Silva et al. 2012, Zandalinas et al. 2018). Albeit at
25°C, both genotypes showed a mean leaf temperature slightly higher than the atmospheric temperature (Paragon = 26.87°C; Sokoll = 26.33°C), when subjected to 38°C both genotypes showed a decrease of leaf temperature relative to atmospheric temperature, which was statistically significant in Sokoll at WW38°C (Table 1). Additionally, both genotypes maintained similar photosynthetic assimilation and electron transport rates compared to control conditions (Fig. 1A,C). However, in vitro Rubisco activity decreased more than 10-fold (Fig. 2), in agreement with previous reports (Galmés et al. 2013, Perdomo et al. 2016, 2017). The maintenance of assimilation rates despite this abrupt decline in Rubisco activity can be explained by the increase in catalytic rate under increased temperature. When measured at 38°C, the initial activity was 5 times higher than when measured at 25°C (Fig. 2A,D) and showed rates comparable to the rates of photosynthesis in the same plants. In vivo, the Rubisco chaperone (RUBISCO ACTIVASE, RCA) helps to overcome possible dead-end inhibition of Rubisco by promoting ATP-dependent conformational changes at the closed sites of Rubisco (Feller, Crafts-Brandner and Salvucci, 1998, Crafts-Brandner and Salvucci, 2000, Salvucci and Crafts-Brandner, 2004) and may contribute to sustaining Rubisco activities at adequate levels to support carbon assimilation (Perdomo et al. 2017). Under our experimental conditions and without water restrictions, photosynthesis occurred at sufficient rates to supply carbon for cellular growth and metabolic energy.

Despite no direct impact of high temperatures was found on photosynthetic assimilation, stomatal conductance and electron transport rate, and in spite of the better performance of Paragon at WD25°C, no differences between genotypes were observed at WD38°C, since these parameters significantly decreased in both Paragon and Sokoll (Fig. 1A,C). These results illustrate that when combined, water deficit and high temperatures have a synergistic effect, both genotypes showed severe leaf dehydration (LRWC> 50%, Table 1) and a serious reduction of stomatal conductance (less than 15% of control values, Fig. 1B). Under such stress conditions, photosynthesis no longer provides a source of carbon and other mechanisms are required to enable plants’ intense reprogramming effort to acclimatise, survive and, mostly, to recover physiological functions after re-watering. Various stress conditions result in the coordinated regulation of both source - sink relations and direct defence responses (Roitsch 1999, Jan et al. 2019, Kosar et al. 2020). Notably, the activities of the different invertase isoenzymes are affected by drought and heat stress (Albacete et al. 2011). Paragon recovered faster from high temperatures and water deficit conditions (Fig. 5) presented higher activity of cytINV and slightly higher activity of vacINV (Fig. 6A,B). These results are suggesting that genotypes with high capacity to hydrolyse sucrose recover faster from episodes of high temperatures combined with drought and therefore reduce the impact of climate fluctuation in yield. Marques da Silva and Arrabaça (2004), in the C₄ grass Setaria sphacelata, found that the higher amount of soluble carbohydrates and the lower amount of starch in leaves exposed to long-term water deficit played a minor role on the osmoregulation against desiccation, suggesting that high availability of hexoses is mainly due to changes on the sucrose metabolism to support other cellular functions. Pinheiro and Chaves (2011) also suggested a connection between cytINV and ABA, sucrose, starch, and ROS metabolism in response
to acute drought stress. Higher activity of vacINV has been reported in maize leaves under water deprivation conditions (Pelleschi et al. 1997, Trouverie et al. 2003), although in sugarcane (Wang et al. 2017), cytINV was also shown to play a more prominent role than vacINV under abiotic stress. In barley, activities of both vacINV and cytINV were repressed after a heat stress episode (Antonio Cuesta-Seijo et al. 2019). In tomato, ectopic expression of cell wall invertases resulted in drought tolerance that was accompanied by also changes in cytINV and vacINV (Albacete et al. 2015). Barratt et al. (2009) demonstrated that cytINV may be the primary route by which carbon from sucrose is supplied to non-photosynthetic tissues in Arabidopsis, suggesting, in concordance to our results, that it would grant a source of carbon to feed cellular functions when photosynthesis is impaired. Secchi and Zwieniecki (2012, 2016) suggested that, under severe drought, high levels of sugar accumulation and invertase activity could prime the xylem for the accelerated restoration of xylem function upon return to hydrated conditions. The authors proposed that the reduction of stomatal conductance and embolism reduces the transpiration flow, subsequently changing the balance of carbohydrate fluxes in xylem instigating the accumulation of sucrose in the apoplast. That mechanism can trigger a cellular stress response promoting starch degradation, leading to the increase of cellular soluble sugar concentration and membrane sucrose gradient. The suggested model is in accordance to our results, Paragon showed high activity of invertases under severe drought (WD38°C, Fig. 6) and the resuming high osmotic level could help xylem embolism refilling and the recovery of transport. When water is delivered from roots, the fast recovery of transpiration could consequently help to explain the faster recovery of photosynthesis, leaf water potential and leaf hydration (Fig. 5 and Table 1). The observed evidence highlighted the role of sucrolytic enzymes in the supply of carbon from sucrose needed to the massive metabolic reorganization employed to tolerate stress, helping plants to recover faster and being less affected by heat and water deficit episodes.

In the present study, WD38°C affected the photochemical capacity in both genotypes, increasing NPQ and qN (Fig. 3B,C) and decreasing qP (Fig. 3A), followed by a decrease of ETR (Fig. 1C). Generally, in higher plants, qE is assumed as the major component of qN, as a short time adaptation to deal with the overproduction of ATP and NADPH and the accumulation of protons in the thylakoid lumen when CO₂ fixation decreases (Krause and Jahns 2007, Takahashi and Murata 2008). Generally, if the energy dissipation mechanisms (qE, qT) and ROS detoxification fail, oxidative damage occurs, leading to photoinhibition (Murata et al. 2007, Yamamoto 2016). The increase in the ROS scavenging activity was observed in both genotypes under WD38°C (Fig. 3E,F). In Paragon, an increase of the absorbed photon flux (J_ABS) was not followed by an increase in the maximum trapped flux (J_oTR) and the electron transport from QA to QB (J_oET2), probably because of the observed increase in the dissipated energy flux (J_DI) (Figs 4, S2 and Table S2), which avoid the overreduction of the electron transport chain. Additionally, the photochemical function of this genotype fully recovered upon stress release, as shown by the recovery of qP and ETR to values similar to control conditions (Fig. 5B,E). The increase in dissipated energy flux may be related to a photoprotective mechanism based on the aggregation and
detachment of the light-harvesting complex II (LHCII) from the reaction center of PSII (Ruban et al. 2012; Ruban 2016). In higher plants, LHCII aggregates are common sites of energy dissipation facilitated by PsbS (qE) or induced by redox-controlled LHCII phosphorylation (qT) (Minagawa 2011), active in plants under CO₂ starvation and heat stress (Šiffel and Vácha 1998, Šiffel and Braunová 1999, Tang et al. 2007). On the other hand, in Sokoll, the reduction of ETR highly correlates to the decrease of both electron transport fluxes (JoET2 and JoRE1, Fig. 4 WD38℃), and despite the full recovery of qP, NPQ levels remained at high levels and ETR stayed below control condition, indicating slower and limited recovery (Fig. 5). Chlorophyll fluorescence parameters clearly indicate differences in photoprotection when both genotypes were subjected to WD38℃ and faster recovery of Paragon after stress relief.

Modulation of the cytosolic invertase was observed and suggests a relationship between an increase of CytINV activity under stress and the recovery of photosynthesis upon high temperatures and water deficit conditions. Upon water shortage and elevated temperatures, when photosynthetic performance and growth priorities are altered, optimization of sucrose export and utilization in conjunction with increased photoprotection of the electron transport machinery could contribute to the recovery of photosynthetic capacity, and consequently to reduce yield fluctuations under climate change. The integration of cell physiological phenotyping via the semi-highthroughput determination of enzyme activity signatures (Jammer et al. 2015) with ecophysiological measurements proved to be a powerful holistic phenomics approach (Großkinsky et al. 2015).

Author contributions
P.M.P.C. planned and carried out the experiments, analysed and interpreted the results. E.C.S. and J.M.S. contributed to the interpretation of the results and supervised the research. A.B.S. and T.R. provided critical feedback. P.M.P.C. took the lead in writing the manuscript. All authors discussed the results and contributed to the final manuscript.

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Data availability statement

The data that support the findings of this study are available from the corresponding author upon request and data supporting findings of this study are available in the supplementary material of this article.

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Table 1. Leaf and soil water status, and canopy temperature of Paragon and Sokoll wheat plants exposed to a combination of heat stress and water deficit and recovery from heat stress conditions. Plants were grown for 3 weeks, then exposed to heat stress (38°C versus control, 25°C), water deficit (WD versus well-watered WW) and re-watered at control temperature (25°C) after heat stress conditions (RWW38°C and RWD38°C). Values are means ± SD (n = 5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, P<0.05). LRWC- leaf relative water content; LWP- leaf water potential; SRWC- soil relative water content; Tcanopy- canopy temperature.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Genotype</th>
<th>LRWC (% H2O)</th>
<th>LWP (MPa)</th>
<th>SRWC (% H2O)</th>
<th>Tcanopy (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25 °C WW</td>
<td>Paragon</td>
<td>90.11 ± 8.82 c</td>
<td>-0.50 ± 0.08 c</td>
<td>88.45 ± 5.84 c</td>
<td>26.87 ± 0.65 a</td>
</tr>
<tr>
<td></td>
<td>Sokoll</td>
<td>90.20 ± 1.73 c</td>
<td>-0.81 ± 0.12 bc</td>
<td>80.11 ± 4.88 b</td>
<td>26.33 ± 0.19 a</td>
</tr>
<tr>
<td>WD</td>
<td>Paragon</td>
<td>68.24 ± 12.45 b</td>
<td>-1.16 ± 0.16 ab</td>
<td>26.74 ± 4.84 a</td>
<td>28.79 ± 0.62 b</td>
</tr>
<tr>
<td></td>
<td>Sokoll</td>
<td>31.89 ± 8.87 a</td>
<td>-1.39 ± 0.10 a</td>
<td>29.12 ± 0.92 a</td>
<td>27.89 ± 1.10 b</td>
</tr>
<tr>
<td>38 °C WW</td>
<td>Paragon</td>
<td>78.60 ± 8.47 bc</td>
<td>-0.82 ± 0.06 bc</td>
<td>87.57 ± 2.11 c</td>
<td>35.04 ± 0.98 c</td>
</tr>
<tr>
<td></td>
<td>Sokoll</td>
<td>80.38 ± 4.74 bc</td>
<td>-0.77 ± 0.09 bc</td>
<td>75.02 ± 5.32 b</td>
<td>33.37 ± 0.40 d</td>
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<tr>
<td>WD</td>
<td>Paragon</td>
<td>39.60 ± 17.71 a</td>
<td>-1.30 ± 0.59 a</td>
<td>30.44 ± 1.69 a</td>
<td>36.95 ± 0.74 e</td>
</tr>
<tr>
<td></td>
<td>Sokoll</td>
<td>43.06 ± 26.64 a</td>
<td>-1.55 ± 0.58 a</td>
<td>28.42 ± 2.72 a</td>
<td>37.52 ± 0.47 e</td>
</tr>
<tr>
<td>Recovery</td>
<td>RWW 38 °C</td>
<td>Paragon</td>
<td>86.46 ± 1.36 c</td>
<td>-0.76 ± 0.03 bc</td>
<td>90.13 ± 5.25 c</td>
</tr>
<tr>
<td></td>
<td>Sokoll</td>
<td>94.91 ± 4.82 cd</td>
<td>-0.74 ± 0.05 bc</td>
<td>91.69 ± 6.14 c</td>
<td>25.58 ± 0.4 a</td>
</tr>
<tr>
<td>RWD</td>
<td>Paragon</td>
<td>90.83 ± 3.42 c</td>
<td>-0.72 ± 0.1 bc</td>
<td>88.96 ± 4.1 c</td>
<td>26.33 ± 0.44 a</td>
</tr>
<tr>
<td>38 °C</td>
<td>Sokoll</td>
<td>78.31 ± 21.18 bc</td>
<td>-0.98 ± 0.16 ab</td>
<td>89.3 ± 3.22 c</td>
<td>26.43 ± 0.21 a</td>
</tr>
</tbody>
</table>

Figure 1. Steady-state photosynthesis of Paragon (PAR) and Sokoll (SOK) wheat plants exposed to a combination of heat stress and water deficit. (A) Net CO2 assimilation, (B) stomatal conductance (gs) and (C) electron transport rate (ETR) were measured at growth light and ambient CO2 in fully expanded leaves of wheat 3-week-old plants under well-watered (WW) and water deficit (WD) conditions and exposed to control (25°C) and heat stress conditions (38°C). Values are means ± SD (n = 5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, P<0.05).
Figure 2. Effect of high temperature and drought on Rubisco activity (expressed by total soluble protein, TSP) and activation state in two wheat genotypes, Paragon (PAR) and Sokoll (SOK). (A-C) Rubisco initial (Vi) and total (Vt) activities and activation state were measured at 25°C and (D-F) 38°C in extracts of fully expanded leaves from 3-week-old wheat plants under well-watered (WW) and water deficit (WD) conditions and exposed to control (25°C) and heat stress conditions (38°C). Values are means ± SD (n = 4-5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, P<0.05).
**Figure 3.** Effect of high temperature and drought on chlorophyll $a$ fluorescence and the antioxidant scavenging capacity in two wheat genotypes, Paragon (PAR) and Sokoll (SOK). (A) Chlorophyll $a$ fluorescence imaging of the photochemical (qP) and non-photochemical (qN) quenching components in representative leaves. (B) Photochemical quenching (qP), (C) non-photochemical quenching (qN) (D) total non-photochemical quenching (NPQ), (E) ferric reducing antioxidant power (FRAP) and (F) trolox equivalents antioxidant capacity (TEAC) in fully expanded leaves of 3-week-old wheat plants under well-watered (WW) and water deficit (WD) conditions and exposed to control ($25^\circ C$) and heat stress conditions ($38^\circ C$). Values are means ± SD (n = 4-5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, $P<0.05$).
Figure 4. Heatmap representation of the correlation between chlorophyll a fluorescence kinetics (OJIP parameters) and antioxidant capacity or steady-state chlorophyll a fluorescence of two wheat genotypes, Paragon (PAR) and Sokoll (SOK), under different stresses. Canonical correlations were determined according to the effect of (A) water deficit (at 25°C, WD25°C), (B) high temperatures (well-watered, WW38°C), and (C) water deficit combined with high temperatures (WD38°C) relative to control plants (WW25°C). All parameters were measured in fully expanded leaves of 3-week-old plants. OJIP parameters included are: absorbed photon flux (J<sub>ABS</sub>); maximum trapped exciton flux (J<sub>oTR</sub>); dissipated energy flux (J<sub>DI</sub>); electron transport flux from QA to QB (J<sub>oET2</sub>); electron transport flux until PSI acceptors (J<sub>oET1</sub>); number of electron carriers per electron transport chain (S<sub>m</sub>); performance index for energy conservation from photons absorbed by PSII antenna to the reduction of QB (PI<sub>ABS</sub>) and until the reduction of PSI acceptors (PI<sub>TOTAL</sub>). Mean values ± SD (n = 5 biological replicates) are in supplementary data, Table S1. Steady-state chlorophyll a fluorescence parameters are non-photochemical quenching (NPQ) and electron transport rate (ETR). Antioxidant capacity was determined by trolox equivalents antioxidant capacity (TEAC) and ferric reducing antioxidant power (FRAP). Different colours denote positive (red) or negative (blue) correlations between variables (n=5 biological replicates).
Figure 5. Recovery of the photochemistry and stomatal function of two wheat genotypes, Paragon (PAR) and Sokoll (SOK), after exposure to high temperatures and water deficit. (A) Chlorophyll \( a \) fluorescence imaging of the photochemical (qP) and non-photochemical (qN) quenching components in representative leaves. (B) Photochemical quenching (qP), (C) non-photochemical quenching (qN), (D) total non-photochemical quenching (NPQ), (E) electron transport rate (ETR), (F) net photosynthetic assimilation rate (A), (G) stomatal conductance (gs). Measurements at growth PPFD in fully expanded leaves of 33-day-old wheat plants recovering for 7 days under well-watered (WW) conditions and 25°C after exposure to WW (RWW 38°C) or water deficit (RWD 38°C) conditions and high temperature (38°C) for 5 days. Values are means ± SD (n=5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, \( P<0.05 \)).
Figure 6. Effect of high temperature and water deficit on cytoplasmic and vacuolar invertases activities in two wheat genotypes, Paragon (PAR) and Sokoll (SOK). (A) Vacuolar Invertase (vacINV) and (B) cytoplasmic invertase (cytINV) activities were measured at 30°C in fully expanded leaves of 3-week-old wheat plants under well-watered (WW) and water deficit (WD) conditions and exposed to control (25°C) and high temperatures (38°C). Values are means ± SD (n=4-5 biological replicates). Different letters denote statistically significant differences between treatments (Duncan analysis, P<0.05).

Supplementary data

Fig.S1. Effect of high temperature and drought on Rubisco activity (expressed by leaf area) and activation state in two wheat genotypes, Paragon (PAR) and Sokoll (SOK).

Table S1. Pearson correlation matrix between net photosynthetic assimilation rate (A), stomatal conductance (gs), electron transport rate (ETR) and cytoplasmic invertase (cytINV) in two wheat genotypes, Paragon and Sokoll, under well-watered (WW) and water deficit (WD) conditions and exposed to control (25°C) and high temperatures (38°C).

Table S2. OJIP parameters of Paragon and Sokoll wheat plants exposed to a combination of heat stress and water deficit and recovered under well-watered conditions.

Fig. S2. Chlorophyll a fluorescence induction curves (OJIP curves) of Paragon and Sokoll wheat plants exposed to water deficit, heat stress, a combination of heat stress and water deficit and recovered under well-watered conditions.