

Moderate heat stress prevented the observed biomass and yield stimulation caused by elevated CO₂ in two well-watered wheat cultivars

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Abstract

Key message Heat stress (HS) under well-watered conditions was not detrimental to leaf photosynthesis or yield but modified the elevated CO_2 response of photosynthesis and yield in two contrasting wheat cultivars.

Abstract Climate change is increasing the frequency of extreme events such as heat waves, adversely affecting crop productivity. While positive impacts of elevated carbon dioxide (eCO₂) on crop productivity are evident, the interactive effects of eCO₂ and environmental stresses are still unclear. To investigate the interactive effects of elevated CO₂ and heat stress (HS), we grew two contrasting wheat cultivars, early-maturing Scout and high-tillering Yitpi, under non-limiting water and nutrients at ambient (aCO₂, 450 ppm) or elevated (eCO₂, 650 ppm) CO₂ and 22 °C in the glasshouse. Plants were exposed to two 3-day HS cycles at the vegetative (38.1 °C) and/or flowering (33.5 °C) stage. At aCO₂, both wheat cultivars showed similar responses of photosynthesis and mesophyll conductance to temperature and produced similar grain yield. Relative to aCO₂, eCO₂ enhanced photosynthesis rate and reduced stomatal conductance and maximal carboxylation rate (V_{cmax}). During HS, high temperature stimulated photosynthesis at eCO₂ in both cultivars, while eCO₂ stimulated photosynthesis in Scout. Electron transport rate (J_{max}) was unaffected by any treatment. eCO₂ equally enhanced biomass and grain yield of both cultivars in control, but not HS, plants. HS reduced biomass and yield of Scout at eCO₂. Yitpi, the cultivar with higher grain nitrogen, underwent a trade-off between grain yield and nitrogen. In conclusion, eCO₂ improved photosynthesis of control and HS wheat, and improved biomass and grain yield of control plants only. Under well-watered conditions, HS was not detrimental to photosynthesis or growth but precluded a yield response to eCO₂.

Keywords Wheat \cdot Elevated CO₂ \cdot Photosynthetic acclimation \cdot Temperature response \cdot Heat stress \cdot Grain yield

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Introduction

Ongoing climate change is threatening the production of agricultural crops including wheat (*Triticum aestivum*) (IPCC 2014; Asseng et al. 2015; Mishra et al. 2021). By the end of this century, atmospheric carbon dioxide concentration ([CO₂]) is expected to reach 700 ppm, increasing surface temperatures by 1.1-2.6 °C (IPCC 2014). For every degree of the temperature increase, global wheat production is predicted to decrease by 6-10% (Asseng et al. 2015; García et al. 2015). Crop models are important tools for assessing the impact of climate change (Asseng et al. 2013). However, they largely lack the ability to consider genotype-specific responses to elevated [CO₂] (eCO₂) and their interaction with other environmental conditions. Hence, it is important to better understand how plants respond to eCO₂ interactions with the environment. Photosynthesis, a fundamental

process driving crop growth and yield, can partially explain the interactive effects of eCO_2 with environmental stresses and provide a mechanistic basis for crop models (Yin and Struik 2009).

During photosynthesis, ribulose-1, 5-bisphosphate carboxylase/oxygenase (Rubisco) catalyzes the carboxylation and oxygenation of ribulose-1, 5-bisphosphate (RuBP). eCO_2 increases photosynthetic rates (A_{sat}) and reduces photorespiration and stomatal conductance (g_s) . Generally, higher photosynthetic rates enhance the growth and productivity of plants leading to increased leaf area, plant size and crop yield (Krenzer and Moss 1975; Sionit et al. 1981; Hocking and Meyer 1991; Mitchell et al. 1993; Kimball et al. 1995; Mulholland et al. 1998; Cardoso-Vilhena and Barnes 2001; Högy et al. 2009; Kimball 2016; Fitzgerald et al. 2016; Kimball 1983). Following long term CO₂ enrichment, photosynthetic capacity defined as the rubisco activity or electron transport rate under high irradiance per unit leaf area, may diminish due to lower amount of Rubisco (Nie et al. 1995; Rogers and Humphries 2000; Ainsworth et al. 2003) or reduced activation of Rubisco (Delgado et al. 1994).

Optimum temperature range for wheat growth is 17-23 °C, with a minimum of 0 °C and maximum of 37 °C (Porter and Gawith 1999). Global warming involves a gradual increase in mean temperature as well as increased frequency and intensity of heat waves. Heat can adversely affect crop growth and disrupt reproduction depending on the timing, intensity and duration (Sadras and Dreccer 2015). Higher daytime temperatures (below damaging level) increase photosynthesis up to an optimum temperature, above which photosynthesis decreases mainly due to higher photorespiration (Berry and Bjorkman 1980; Long 1991). High night time temperatures increase respiration and reduce overall photosynthetic carbon gain (Prasad et al. 2008). At the whole plant level, high temperatures accelerate growth (Fischer 1980) and shorten crop duration (Hatfield and Prueger 2015), hence reducing grain yield due to insufficient time to capture resources. Losses due to short crop duration are usually higher than benefits of growth stimulation at high temperature (Wardlaw and Moncur 1995).

The severity of the damage caused by abrupt temperature increases above the optimum range (termed heat stress, HS) depends on the magnitude and duration of HS as well as the developmental stage of the plant (Wahid et al., 2007). HS may reduce photosynthesis due to reduced chlorophyll content, impaired photosystem II and lower Rubisco activation (Berry and Bjorkman 1980; Eckardt and Portis 1997). Furthermore, HS can directly damage cells and increase grain abortion resulting in reduced growth, biomass and grain yield (Stone and Nicolas 1996, 1998; Wardlaw et al. 2002; Farooq et al. 2011). Around anthesis, HS (> 30 °C) reduces seed setting due to lower pollen viability, leading to

poor fertilization, abnormal ovary development and slower pollen growth (Balla et al. 2019).

The interactive effects of eCO₂ and HS on plant growth and yield can be positive, negative or neutral (Wang et al. 2008, 2011). Plants may acclimate to changes in growth temperature and shift the optimum temperature for photosynthesis, which can maximize the photosynthetic rate at the growth temperature (Yamori et al. 2014). Elevated CO₂ increases the temperature optimum of photosynthesis (Long 1991; Alonso et al. 2009) by reducing photorespiration and improving tolerance to photoinhibition (Hogan et al. 1991). In addition, differences in plasticity of photosynthetic parameters with respect to growth temperature have been found responsible for differences in photosynthetic temperature acclimation (Yamori et al. 2010) and respiratory temperature acclimation can generate apparent acclimation of photosynthetic processes (Way and Yamori 2014). The impact of HS on photosynthesis will depend on whether Rubisco, electron transport or end-product synthesis is limiting at eCO₂ (Sage and Kubien 2007). Enhanced growth and leaf level intrinsic water use efficiency (iWUE) by eCO₂ may help compensate for the negative impact of HS; conversely, heat-induced shortening of the grain-filling stage and grain abortion could limit the benefits of eCO_2 (Lobell and Gourdji 2012). In addition, decreased g_s under eCO₂ may limit transpirational cooling and therefore exacerbate HS. Thus, HS counteracts the positive effect of eCO_2 on yield components and may aggravate the negative effect of eCO₂ on grain quality due to the high sensitivity of wheat to temperature stress especially during anthesis and grainfilling stage (Wang and Liu 2021).

Many studies have investigated the response of wheat to eCO_2 in enclosures and in the field (Wang and Liu 2021). However, only a few studies have considered eCO2 interaction with temperature increases in wheat (Rawson 1992; Delgado et al. 1994; Morison and Lawlor 1999; Jauregui et al. 2015; Cai et al. 2016) and rarely with HS (Coleman et al. 1991; Wang et al. 2008). Studies considering HS have addressed mainly the biomass or yield aspects and not the physiological processes such as photosynthesis (Stone and Nicolas 1994, 1996, 1998). Interactive effects of eCO₂ and HS on photosynthesis have been reported in a limited number of studies (Wang et al. 2008, 2011; Macabuhay 2016; Macabuhay et al. 2018; Chavan et al. 2019). Macabuhay et al. (2018) studied interactive effects of eCO₂ and (experimentally imposed) heatwaves on wheat (cv Scout and Yitpi) grown in a dryland cropping system and concluded that eCO2 may moderate some effects of HS on grain yield but such effects strongly depend on seasonal conditions and timing of HS. In another glasshouse experiment on the interactive effects of severe HS (achieved by increasing RH to lower transpirational cooling) and eCO₂ in wheat (cv Scout), we found that eCO₂ mitigated the negative impacts of HS at anthesis on photosynthesis and biomass, but grain

yield was reduced by HS in both CO_2 treatments (Chavan et al. 2019). However, HS can occur throughout plant growth, including during vegetative, flowering or grain filling stages. Moreover, irrigation and relative humidity during HS plays an important role in how plants handle HS and plants may cope well with HS in well-watered conditions. In addition, different crop genotypes may respond variably to the interaction of eCO_2 with HS.

Here, we build on our previous work by comparing the interactive effects of eCO_2 and HS in two commercial wheat cultivars. In this study, we investigated the impact of HS during well-watered conditions at ambient RH. Scout and Yitpi have similar genetic background but distinct agronomic features. Scout is a mid-season maturity cultivar with very good early vigor that can produce leaf area early in the season. Scout has a putative water-use efficiency (WUE) gene, which has been identified using carbon isotope discrimination (Condon et al. 2004). Yitpi is a good early vigor, freely tillering, late flowering and long maturity cultivar (Seednet 2005; Pacificseeds 2009; Bahrami et al. 2017).

Although Scout is known to be a high yielding variety with very good grain quality and high reproductive sink (Pacific seeds 2009), we hypothesized that Yitpi will produce higher grain yield due to its ability to initiate more tillers and its longer time to flower and mature (Hypothesis 1). Fast growing plants with high sink capacity show a greater eCO₂-induced growth stimulation (Poorter 1993) and less photosynthetic acclimation (Delgado et al. 1994) compared to slow growing counterparts with low sink capacity. Consequently, we hypothesized that Yitpi will show greater photosynthetic, growth and yield response to eCO₂ due to its greater vegetative sink capacity (tillering) relative to Scout with restricted tillering (Hypothesis 2). The greater growth stimulation at eCO₂ may buffer Yitpi against HS damage compared to Scout. Thus, HS may decrease yield in Scout more than Yitpi and aCO₂ more than eCO₂ (Hypothesis 3). HS is more damaging at the reproductive relative to the vegetative developmental stage (Farooq et al. 2011). Hence, we expect less damage in plants exposed to HS at the vegetative stage relative to the flowering stage (Hypothesis 4).

To test these hypotheses, Scout and Yitpi were grown at controlled ambient or elevated CO_2 conditions and subjected to one or two heat stresses at the vegetative (HS1) and/or flowering (HS2) stage. Plant growth, biomass and leaf photosynthetic parameters were measured at different time points across the life cycle of the plants.

Materials and methods

Plant culture and treatments

The experiment was conducted in the glasshouse facility located at the Hawkesbury campus of Western Sydney University (WSU). Seeds of commercial winter wheat cultivars Scout and Yitpi were procured from Agriculture Victoria (Horsham). Cultivars were selected based on their use in the Australian Grains Free Air CO₂ Enrichment (AGFACE) project investigating climate change impacts on wheat growth and yield (Houshmandfar et al. 2017). For germination, 300 seeds of each cultivar were sterilized using 1.5% NaOCl₂ for 1 min followed by incubation in the dark at 28 °C for 48 h in petri plates. Sprouted seeds were planted in germination trays using seed raising and cutting mix (Scotts, Osmocote[®]) at ambient CO_2 (a CO_2 , 400 μ L L⁻¹), temperature (22/14 °C day/night), relative humidity (RH, 50-70%) and natural light (midday average 500 μ mol m⁻² s⁻¹) (Figure S1). The growth stages are denoted by decimal code (DC) according to (Zadoks et al. 1974) along with the time points here after. Two-week-old seedlings (DC12) were transplanted to individual cylindrical pots (15 cm diameter and 35 cm height) using sieved soil collected from local site. At transplanting stage (T0) pots were distributed into two aCO₂ (400 μ L L⁻¹) and two eCO_2 (650 µL L⁻¹) chambers (Figure S1B). Some plants were exposed to one or two HS cycles at the vegetative (HS1, 10 weeks after planting, WAP, DC 32) and/or the flowering (HS2, 15 WAP, DC 63) stages for 3 days with temperature ramp up from 14 °C night temperature (8 pm to 6 am) to 38 °C or 33 °C during mid-day (10 pm to 4 pm) respectively at 60% daytime RH (Figures S1, S2). The two HS cycles created four sets of heat treatments at each CO₂ concentration as follows: (1) Control-plants were not exposed to HS at any stage, (2) HS1-plants were exposed to HS at vegetative (DC32) stage only, (3) HS2—plants were exposed to HS at reproductive (DC63) stage only and (4) HS1 + 2—plants were exposed to both the heat stresses HS1 and HS2 (Fig. 1).

Thrive all-purpose fertilizer (Yates) was applied monthly throughout the experiment to maintain similar nutrient supply in all treatment combinations. Pots were regularly swapped between left and right benches as well as between front and back for randomization within chamber. Pots and treatments were also swapped between the two ambient and two elevated CO_2 chambers for randomization among chambers.

Fig. 1 Experimental design. Depicting plant growth plotted over 5-time points (T0 T1, T2 T3 and T4) across the wheat life cycle till maturity. The circles represent harvest of 10 plants at each time point. Green circles represent control plants grown at ambient or elevated CO₂ Upward directed red and brown arrows point to timing and duration of two heat stresses (HS), HS1 (vegetative stage) and HS2 (flowering stage) respectively. The red circles represent plants subjected to HS1 and brown circles represent plants subjected to HS2. Red and brown dotted circles represent plants subjected to both heat stresses. Downward small black arrows represent timing of single point gas exchange measurements. Thermometer symbol represents timing of temperature response measurements

Time point (at the end of week period)	Weeks After Planting (WAP)	Zadoks Growth Scale	Growth Period/Stage	
T0 (W ₀ - W ₂)	2	DC12	Transplanting: 2 leaves unfolded	
T1 (W ₂ -W ₇)	7	DC28	Tillering: Main shoot and 6/8 tillers, 6/8 leaves unfolded	
HS1	10	DC32	Vegetative stage	
T2 (W ₇ -W ₁₂)	12	DC35	Pre-anthesis: 4 th node detectable	
HS2	15	DC63	Flowering stage	
T3 (W ₁₂ -W ₁₇)	17	DC65	Anthesis: 50 % flowering	
T4 (W ₁₇ -W ₂₅)	25	DC90	Grain filling: Maturity	



Growth and biomass measurements

The full factorial experimental design included four chambers (two chambers for each CO_2 treatment) and five destructive harvests at time points T0 (2 WAP, DC12), T1 (6 WAP, DC28), T2 (10 WAP, DC35), T3 (17, DC65) and T4 (25 WAP, DC90) (Fig. 1). Ten plants per treatment per cultivar were measured and harvested at each time point except for T4 where plants exposed to only HS1 were not harvested. Morphological parameters were measured followed by determinations of root, shoot and leaf dry mass. Samples were dried for 48 h in the oven at 60 °C immediately after harvesting. Leaf area was measured at time point T1, T2 and T3 using a leaf area meter (LI-3100A, LI-COR, Lincoln, NE, USA). Plant height, leaf number,

tiller number and ear (grain inflorescence) number along with developmental stage information (booting (DC45), half-emerged (DC55) or fully emerged (DC60)) were recorded at time points T2 and T3).

Leaf gas exchange measurements

The youngest fully developed leaf (which was the flag leaf at T3) was used to measure gas exchange parameters. Steady state leaf gas exchange measurements were performed at time points T1, T2 and T3 using a portable open gas exchange system (LI-6400XT, LI-COR, Lincoln, USA) to measure light-saturated (photosynthetic photon flux density (PPFD)=1500 µmol m⁻² s⁻¹) photosynthetic rate (A_{sat}), stomatal conductance (g_s), ratio of intercellular to ambient

 $CO_2(C_i/C_a)$, leaf transpiration rate (E), dark respiration (R_d) and dark- and light-adapted chlorophyll fluorescence (Fv/Fm and Fv'/Fm', respectively). Dark adapted leaf measurements were conducted by switching off light for 15 min. Steady state leaf gas exchange measurements were also performed during (3rd day of HS) and after heat shock (next day after HS) along with recovery stage (5 days after HS). Plants were moved to a neighboring chamber with ambient CO₂ levels for short time (20-30 min for each plant) where air temperature was separately manipulated to achieve the desired leaf temperature. The LI-COR 6400-40 leaf chamber fluorometer (LCF) was used to measure gas exchange parameters at a PPFD of 1500 μ mol m⁻² s⁻¹ at two reference CO₂ (CO2R) concentrations (400 and 650 μ L L⁻¹) and two leaf temperatures (25 and 35 °C). Photosynthetic down regulation or acclimation was examined by comparing the measurements at common CO₂ (ambient and elevated CO₂ grown plants measured at 400 μ L L⁻¹ CO₂ partial pressure) and growth CO_2 (aCO₂ grown plants measured at 400 µL L⁻¹ CO₂ partial pressure and eCO₂ grown plants measured at 650 μ L L⁻¹ CO₂ partial pressure).

Dark respiration (R_d) was measured after a dark adaptation period of 15 min. Intrinsic water use efficiency (iWUE) was calculated as A_{sat} (µmol m⁻² s⁻¹)/ g_s (mol m⁻² s⁻¹). The response of A_{sat} to variations in sub-stomatal CO₂ mole fraction (*Ci*) (A-Ci response curve) was measured at T3 in 8 steps of CO₂ concentrations (50, 100, 230, 330, 420, 650, 1200 and 1800 µL L⁻¹) at leaf temperature of 25 °C. Measurements were taken around mid-day (from 10 am to 3 pm) on attached last fully expanded or flag leaves of the main stems. Before all leaf gas exchange measurements, the leaf was allowed to stabilize for 10–20 min until it reached a steady state of CO₂ uptake and stomatal conductance. Ten replicate plants per treatment were measured except for plants exposed only HS1 at T3 were not measured.

Mesophyll conductance and temperature response

Mesophyll conductance (g_m) was determined by concurrent gas exchange and stable carbon isotope measurements using portable gas exchange system (LI-6400-XT, LI-COR, Lincoln, NE, USA) connected to a tunable diode laser (TDL) (TGA100, Campbell Scientific, Utah, USA) for the two wheat cultivars grown at ambient atmospheric CO₂ levels. A_{sat} and ${}^{13}CO_2/{}^{12}CO_2$ carbon isotope discrimination were measured after T1 at five leaf temperatures (15, 20, 25, 30 and 35 °C) and saturating light (1500 µmol quanta m⁻² s⁻¹). Leaf temperature sequence started at 25 °C decreasing to 15 °C and then increased up to 35 °C. A-Ci response curves were measured at each leaf temperature. Dark respiration was measured by switching light off for 20 min at the end of each temperature curve.

Measurements were made inside a growth cabinet (Sanyo) to achieve desired leaf temperature. The photosynthetic carbon isotope discrimination (Δ) to determine g_m was measured as follows (Evans et al. 1986):

$$\Delta = \frac{1000\varepsilon \left(\delta^{13}C_{sam} - \delta^{13}C_{ref}\right)}{1000 + \delta^{13}C_{sam} - \varepsilon \left(\delta^{13}C_{sam} - \delta^{13}C_{ref}\right)}$$
(1)

where,

$$\varepsilon = \frac{C_{ref}}{C_{ref} - C_{sam}} \tag{2}$$

 C_{ref} and C_{sam} are the CO₂ concentrations of dry air entering and exiting the leaf chamber, respectively, measured by the TDl. g_m was calculated using correction for ternary effects (Farquhar and Cernusak 2012; Evans and Von Caemmerer 2013) following the next expression:

$$g_m = \frac{\frac{1+t}{1-t} \left(b - a_i - \frac{eR_d}{A + R_d} \right) \frac{A}{C_a}}{\left(\Delta_i - \Delta_o - \Delta_e - \Delta_f \right)}$$
(3)

where, Δ_i is the fractionation that would occur if the gm were infinite in the absence of any respiratory fractionation (e=0), Δ_0 is observed fractionation, Δ_e and Δ_f are fractionation of ¹³C due to respiration and photorespiration respectively (Evans and Von Caemmerer 2013).

$$\Delta_i = \frac{1}{1-t} a' \frac{1}{(1-t)} \left((1+t)b - a' \right) \frac{C_i}{C_a} \tag{4}$$

$$\Delta_e = \frac{1+t}{1-t} \left(\frac{eR_d}{(A+R_d)C_a} (C_i - \Gamma *) \right)$$
(5)

$$\Delta_f = \frac{1+t}{1-t} \left(f \frac{\Gamma *}{C_a} \right) \tag{6}$$

where,

$$t = \frac{\left(1 + a'\right)E}{2g_{ac}^t} \tag{7}$$

The constants used in the model were as follows: E denotes transpiration rate; g_{ac}^{t} is total conductance to CO₂ diffusion in the boundary layer ($ab = 2.9\%_{0}$) and in air ($a = 4.4\%_{0}$); a' is the combined fractionation of CO₂ across boundary layer and stomata; net fractionation caused by RuBP and PEP carboxylation ($b = 27.3\%_{0}$) (Evans et al. 1986); fractionation with respect to the average CO₂ composition associated with photorespiration ($f = 11.6\%_{0}$) (Lanigan et al. 2008) and we assumed null fractionation associated with mitochondrial respiration in light (e = 0).

Nitrogen and carbon estimation

Leaf discs were cut from the flag leaves used for gas exchange measurements at time points T2 and T3 which were then oven dried at 60 °C for 48 h. Leaf discs were processed for nitrogen (N) and carbon (C) content using an elemental analyzer (Dumas method). N and C were also estimated from other plant components including leaf, stem, root and grain harvested at T1, T3 and T4. Ground samples were processed for C & N with a CHN analyzer (LECO Tru-Mac CN-analyser, Leco corporation, USA) using an automated dry combustion method (Dumas method). Leaf N per unit area (N_{area}) was calculated as N (mmol g^{-1})×LMA (g m⁻²). Photosynthetic nitrogen use efficiency (PNUE) was calculated as A_{sat} (µmol m⁻² s⁻¹)/leaf N_{area} (mmol m⁻²). Protein content was estimated using N and multiplication factor of 5.7 (Mosse 1990; Bahrami et al. 2017). N utilization efficiency was calculated as grain yield per total plant N.

Statistical and temperature analysis

All data analyses and plotting were performed using R computer software (R Core Team 2022). The effect of treatments and their interactions was analyzed using linear modeling with 'anova' in R. Significance tests were performed with anova and post hoc Tukey test using the 'glht' function in the multcomp R package. Coefficient means were ranked using post-hoc Tukey test. The Farquhar-von Caemmerer-Berry (FvCB) photosynthesis model was fit to the A_{sat} response curves to C_i (A-Ci response curve) measured at T3 or chloroplastic CO₂ mole fraction (Cc), which was estimated from the g_m measuremed (A-Cc response curve) after T1. We used the plantecophys R package (Duursma 2015) to perform the fits, using measured g_m and R_d values, resulting in estimates of maximal carboxylation rate (V_{cmax}) and maximal electron transport rate (J_{max}) for D-ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) using measured R_d values. Temperature correction parameter (Tcorrect) was set to False while fitting A-C_i curves. Temperature response of V_{cmax} and J_{max} were calculated by standard Arrhenius and modified Arrhenius functions, respectively (Medlyn et al. 2002). Estimated V_{cmax} and J_{max} values at five leaf temperatures were then fit using nonlinear least square (nls) function in R to determine energy of activation for V_{cmax} (EaV) and J_{max} (EaJ) and entropy (Δ SJ). Temperature responses of V_{cmax} and R_d were fit using Arrhenius equation as follows,

$$f(Tk) = k_{25} \cdot \exp\left[\frac{E_a \cdot (Tk - 298)}{R \cdot 298 \cdot Tk}\right]$$
(8)

where, Ea is the activation energy (in J mol⁻¹) and k_{25} is the value of R_d or V_{cmax} at 25 °C. R is the universal gas constant

(8.314 J mol⁻¹ K⁻¹) and Tk is the leaf temperature in K. The activation energy term Ea describes the exponential rate of rise of enzyme activity with the increase in temperature. The temperature coefficient Q_{10} , a measure of the rate of change of a biological or chemical system as a consequence of increasing the temperature by 10 °C was also determined for *Rd* using the following equation:

$$R_d = R_{d25} \cdot Q_{10}^{[(T-25)/10]} \tag{9}$$

A peaked function (Harley et al. 1992) derived Arrhenius function was used to fit the temperature dependence of J_{max} , and is given by the following equation:

$$f(Tk) = k_{25} \cdot \exp\left[\frac{E_a \cdot (Tk - 298)}{R \cdot 298 \cdot Tk}\right] \left[\frac{1 + \exp\left(\frac{298 \cdot \Delta S - H_d}{298 \cdot R}\right)}{1 + \exp\left(\frac{Tk \cdot \Delta S - H_d}{Tk \cdot R}\right)}\right]$$
(10)

where, Ea is the activation energy and k_{25} is the J_{max} value at 25 °C, Hd is the deactivation energy and S is the entropy term. Hd and Δ S together describe the rate of decrease in the function above the optimum. Hd was set to constant 200 kJ mol⁻¹ to avoid over parametrization. The temperature optimum of J_{max} was derived from Eq. 10 (Medlyn et al. 2002) and written as follows:

$$T_{opt} = \frac{H_d}{\Delta S - R \cdot \ln\left[\frac{E_a}{(H_d - E_a)}\right]}$$
(11)

The temperature response of A_{sat} was fit using a simple parabola equation (Crous et al. 2013) to determine temperature optimum of photosynthesis:

$$A_{sat} = A_{opt} - b \cdot \left(T - T_{opt}\right)^2 \tag{12}$$

where, T is the leaf temperature of leaf gas exchange measurement for A_{sat} , T_{opt} represents the temperature optimum and A_{opt} is the corresponding A_{sat} at that temperature optimum. Steady state gas exchange parameters g_m , g_s , C_i and J_{max} to V_{cmax} ratio were fit using nls function with polynomial equation:

$$y = A + Bx + Cx^2 \tag{13}$$

Results

Two commercial wheat cultivars Scout and Yitpi were grown under aCO_2 or eCO_2 (daytime average of 450 or 650 µL L⁻¹, respectively; 65% RH and 22 °C), natural sunlight and well-watered conditions (Figure S1). Both aCO_2 and eCO_2 grown plants were exposed to two 3-day HS cycles at the vegetative (HS1, 10 WAP, DC32, daytime average of 38 °C) and flowering stage (HS2, 15 WAP, DC63, daytime average of 33.5 °C), while daytime RH was maintained at 60%. HS2 was lower in intensity relative to HS1 due to the cool winter conditions. Both HS cycles had similar overall effects on growth and yield parameters, refuting our fourth hypothesis that HS during the reproductive stage is more damaging. Hence, we mostly compare the control plants to those exposed to both heat stresses. Grain filling started 17

Photosynthetic temperature responses of the two wheat cultivars at aCO₂

A-C_i curves together with g_m were measured at five leaf temperatures to characterize the thermal photosynthetic responses of the two wheat cultivars grown at aCO₂ (Fig. 2;



Fig. 2 Temperature response of photosynthetic parameters: CO_2 assimilation rate (**a**), mesophyll conductance (**b**), stomatal conductance (**c**) and intercellular CO_2 (**d**), V_{cmax} (**e**), J_{max} (f), J_{max} / V_{cmax} (**g**) and dark respiration (**h**) over leaf temperatures (15, 20, 25, 30 and

35 °C) in plants grown at aCO_2 . Scout and Yitpi are depicted using circles with solid cultivars and triangles with broken cultivars respectively. Data in panels (**a–f**) and (**h**) are fit using nonlinear least square (nls) function in R

Table 1Summary of modelledparameters for temperatureresponse of photosynthesis

Parameter	Constant	Scout	Yitpi
$A_{sat} (\mu mol \ m^{-2} \ s^{-1})$	T _{opt} (°C)	23.4±1 a	23.4±0.7 a
	A _{opt}	24.6±1 a	$22 \pm 0.6 \text{ b}$
$g_m (\mathrm{mol} \;\mathrm{m}^{-2} \;\mathrm{s}^{-1} \;\mathrm{bar}^{-1})$	T_{opt} (°C)	27.9 ± 2.2 a	32.5±6.9 b
	g_m at 25 °C	0.31±0.01 a	$0.25 \pm 0.01 \text{ b}$
	g_m at T_{opt}	0.30 ± 0.01	0.25 ± 0.01
$V_{cmax} (\mu { m mol} { m m}^{-2} { m s}^{-1})$	V _{cmax} at 25 °C	192.7 ± 17.1 a	198.4 ± 17.7 a
	EaV (kJ mol ⁻¹)	43.3 <u>+</u> 8.74 a	46.4 ± 8.7 a
$J_{max} (\mu mol \ m^{-2} \ s^{-1})$	J _{max} at 25 °C	187.9±13.1 a	186.1±5.7 a
	T _{opt} (°C)	29.6±0.3 a	30.5±0.3 a
	J_{max} at T _{opt}	205.7 ± 10.2	215.4 ± 13.4
	EaJ (kJ mol ^{-1})	37.7 ± 13.2 a	41.1±5.8 a
	$\Delta SJ (J \text{ mol}^{-1} \text{ K}^{-1})$	648.3±5.3 a	647 ± 2.4 a
	Hd (kJ mol^{-1})	200	
$Rd \ (\mu mol \ m^{-2} \ s^{-1})$	<i>Rd</i> at 25 °C	1.25 ± 0.02 a	1.25 ± 0.02 a
	EaR (kJ mol ⁻¹)	30.9±1.6 a	33.2±1.7 a
	Q ₁₀	1.51 ± 0.03 a	1.56 ± 0.04 a

Summary of coefficients derived using nonlinear least square fitting of CO₂ assimilation rates and maximal rate of carboxylation (V_{cmax}) and maximal rate of RuBP regeneration (J_{max}) determined using A-C_i response curves and dark respiration measured at five leaf temperatures 15, 20, 25, 30 and 35 °C. Values are means with standard errors. Derived parameters include temperature optima (T_{opt}) of photosynthesis (A_{opt}); activation energy for carboxylation (EaV); activation energy (EaJ), entropy term (Δ SJ) and T_{opt} and corresponding value for J_{max} with deactivation energy (Hd) assumed constant; and activation energy (EaR) and temperature coefficient (Q₁₀) for dark respiration. Letters indicate significance of variation in means

Table 1). Overall, both cultivars had similar photosynthetic temperature responses. A_{sat} and g_s increased with leaf temperature up to an optimum (T_{opt}) around 23.4 °C and decreased thereafter, while C_i slowly decreased with temperature. Mesophyll conductance (g_m) increased up to 25 °C then plateaued (Fig. 2b). Scout had slightly higher A_{sat} , g_s and g_m than Yitpi at most leaf temperatures (Fig. 1a–d). R_d linearly increased with temperature, and both cultivars had similar Q₁₀ (Fig. 2h, Table 1). V_{cmax} and J_{max} exponentially increased with leaf temperature, but J_{max} declined above T_{opt} (30 °C) in both cultivars (Fig. 1e, f). There was no significant difference in V_{cmax} , J_{max} or their activation energies between the two wheat cultivars (Fig. 2e–g; Table 1). The ratio of J_{max}/V_{cmax} was similar for the two cultivars and linearly decreased with leaf temperature (Fig. 2g).

eCO₂ stimulated photosynthesis and reduced stomatal conductance in both wheat cultivars

Overall, the two wheat cultivars had similar A_{sat} , g_s , iWUE (A_{sat}/g_s), R_d , Fv/Fm, V_{cmax} and J_{max} measured under most conditions (Figs. 1, 2, 3, 4; Tables S1, S2, S3). Under control (non-HS) conditions, eCO₂ enhanced A_{sat} measured at growth CO₂ (A_{growth}) and 25°C and reduced g_s in both cultivars at T1, T2 and T3 (Figs. 3, S2; Tables S1, S2, S3). When measured at common CO₂ and 25 °C, eCO₂-grown plants had lower A_{sat} (- 12% at T2, p < 0.01) and g_s (- 10% at T2,

p < 0.001) relative to aCO₂. This photosynthetic downregulation was more persistent in Yitpi compared to Scout (Fig. 3; Tables S1, S2, S3).

High temperature during HS enhanced photosynthesis under eCO₂

The two HS cycles did not reduce A_{growth} measured at 25°C during or after HS (Fig. 3a–d; Tables S1, S2, S3). During both HS1 and HS2, eCO₂ stimulated A_{growth} measured at 25°C in Scout but not Yitpi. Relative to 25 °C, A_{growth} increased at 35 °C in Scout (10–14%) and Yitpi (12–18%) grown at eCO₂ but not at aCO₂. Immediately after the recovery from HS, A_{growth} was upregulated in eCO₂-grown Scout (Figs. 3a–d, S3). During both HS cycles, dark-adapted Fv/Fm measured at 25 °C tended to be lower in Yitpi grown at eCO₂ relative to aCO₂. In both cultivars, Fv/Fm decreased at 35 °C relative to 25 °C, indicating transient damage to PSII due to HS at both CO₂ treatments (Fig. 3e–h; Tables S1, S2, S3).

Following long-term recovery from HS1 and/or HS2, the eCO₂ stimulation of A_{growth} was still marginally apparent in all T3 plants, being the strongest in eCO₂-grown Yitpi (Fig. 5a, b; Tables S1, S2, S3). The reduction of g_s at eCO₂ was weak in all plants (Fig. 5c, d; Tables S1, S2, S3). Hence, iWUE was stimulated by eCO₂ in all treatments, while PNUE was unaffected (Fig. 5e–h; Tables S1, S2, S3). There



 aCO₂ grown meas at 400 — eCO₂ grown meas at 400 eCO₂ grown meas at 650 -

Fig. 3 Response of leaf gas exchange parameters to eCO₂ under non-HS conditions. Measurements were made at 25 °C before each harvest (T1, T2 and T3) for CO₂ assimilation rates (a, b) and stomatal conductance (c, d) in Scout (Circles) and Yitpi (Triangles). Plants were grown and measured at aCO₂ (blue solid cultivars), grown and

measured at eCO₂ (red solid cultivars), and grown at eCO₂ and measured at 400 μ L CO₂ L⁻¹ (red dashed cultivars). Statistical significance levels (t-test) for the growth condition within each cultivar are shown and they are: *=p<0.05; **=p<0.01: ***=p<0.001

was a good correlation between A_{growth} and g_s ($r^2 = 0.51$, p < 0.001) across all treatments (Fig. 6a).

 V_{cmax} and J_{max} were derived from A-C_i response curves measured at 25 °C during the recovery stage after HS2. For control and HS plants, growth at eCO₂ marginally reduced V_{cmax} in Scout (- 14%, p=0.09) and Yitpi (- 15%, p=0.06) but had no effect on J_{max} . HS had no effect on V_{cmax} or J_{max} in either cultivar (Fig. 5i–l; Tables S1, S2, S3). V_{cmax} and J_{max} correlated well ($r^2 = 0.75$, p < 0.001) across treatments (Fig. 6b).

Yitpi produced more tillers and grains than Scout

When compared at aCO₂, the two wheat cultivars differed in phenology and growth habit. Scout developed faster and flowered earlier than Yitpi. At T2, 43% of tillers had ears in Scout compared to 11% in Yitpi (Figure S3). At T2, Scout was 74% (p < 0.001) taller than Yitpi but at T3 both cultivars had similar height (Fig. 7i, j; Tables S4, S5). In contrast, Yitpi accumulated more biomass relative to Scout by producing more tillers. At T3, Yitpi had 42% (p < 0.005) more total plant biomass, 130% (p < 0.001)more tillers, 254% (p < 0.001) larger leaf area, 128%(p < 0.001) more leaves and 61% (p < 0.001) larger leaf size compared to Scout (Fig. 7; Tables S4, S5).

At the final harvest (T4), Yitpi had more plant biomass (84%, p < 0.001), tillers (88%, p < 0.001) and number of grains (54%, p < 0.001). Conversely, Scout had larger grain size (+31%, p < 0.001), a higher proportion (100%) of its tillers developed ears and more ears filled grains compared to higher tillering Yitpi (88%). Hence, both cultivars had relatively similar grain yield (g/plant) (Fig. 8a-f; Tables S4, S6). Higher (178%, p < 0.001) harvest index (HI) in Scout was due to early maturity and consequent leaf senescence leading to loss of biomass at final harvest (Tables S5). The final harvest was undertaken four weeks after all ears had matured on Scout to give ample time for grain filling in Yitpi (Fig. 8; Tables S4, S6).



Fig. 4 Response of photosynthesis and chlorophyll fluorescence to HS in Scout and Yitpi grown at aCO_2 or eCO_2 . CO_2 assimilation rates (**a-d**) and dark-adapted chlorophyll fluorescence, Fv/Fm (**e-h**) were measured at growth CO_2 and 25 °C in Scout (Circles) and Yitpi (Tri-

angles). Open and closed symbols represent control and HS plants, respectively. In addition, plants were measured at 35 $^{\circ}C$ (*) during both HS cycles

eCO₂ similarly stimulated wheat biomass and grain yield under non-HS conditions

The increase in plant biomass at eCO_2 depended on the growth stage (Figs. 6, 7; Tables S4, S5). However, the overall stimulation was not different between the two cultivars as evident from the non-significant eCO_2 x cultivar interaction at all harvests (Table S4). By T3 (anthesis), when

both cultivars were still within the exponential growth stage, eCO₂ stimulated plant biomass of Yitpi (+29%, p<0.001) and Scout (+9%, p<0.001) under control conditions. The number of tillers, total leaf area, mean leaf size or leaf mass area were not significantly affected by eCO₂ in either cultivar (Fig. 7; Tables S4, S5). eCO₂ increased allocation to stem relative to leaf biomass, particularly in Yitpi. Accordingly, there was a strong correlation across treatments between





Fig. 5 Response of photosynthetic parameters to eCO_2 and HS at anthesis (T3) in Scout and Yitpi. CO_2 assimilation rate (**a**, **b**), stomatal conductance (**c**, **d**), photosynthetic water use efficiency (**e**, **f**) and photosynthetic nitrogen use efficiency (**g**, **h**) were measured at growth CO_2 . V_{cmax} (**i**, **j**) and J_{max} (**k**, **l**) were derived from ACi curves measured at 25 °C. Cultivars indicate means and shaded region is

95% confidence interval. Data shown for control (not exposed to any heat stress) and plants exposed to both heat stress cycles (HS1+2). Statistical significance levels (t-test) for the growth condition within each cultivar are shown and they are: *=p<0.05; **=p<0.01: ***=p<0.001

stem and leaf biomass ($r^2 = 0.83$, p < 0.001) and between total biomass and leaf area ($r^2 = 0.83$, p < 0.001) in Scout but not in Yitpi. However, the two cultivars followed common relationship for root *versus* shoot biomass ($r^2 = 0.41$, p < 0.001) and leaf area *versus* leaf number ($r^2 = 0.82$, p < 0.001) across all treatments suggesting no effect of cultivar, eCO₂ or HS on these common allometric relationships (Figure S4; Table S5).

At the final harvest T4 (seed maturity), eCO_2 enhanced biomass and equally stimulated grain yield by increasing grain number in both cultivars (+64% in Scout and +50% in Yitpi) under control conditions only (Fig. 8a–d; Tables S4, S5, S6). Harvest index was not directly affected by any treatments but showed a significant interaction (p < 0.05) between CO₂ and cultivar, such that HI was higher in Yitpi under eCO₂ (Tables S5 and S6).

eCO₂ did not stimulate the grain yield of HS plants

At T3, moderate HS (34-38 °C) applied under wellwatered conditions and 60% RH during the vegetative (HS1 applied after T1) and flowering (HS2 applied after T2) stages had no significant impact on plant biomass of either wheat cultivar or CO_2 treatment. By T4, there were significant HS × CO_2 × cultivar interactions (p < 0.01) for biomass and grain yield. HS1 + 2 reduced the biomass and grain yield of eCO₂-grown Scout relative to aCO₂-grown counterparts. Unlike control plants, the biomass and yield of HS plants were not enhanced by eCO₂ (Fig. 8; Tables S4, S5).

eCO₂ reduced grain N in Yitpi but not in Scout

Neither eCO_2 or HS had a significant effect on flag leaf N content in either cultivar at T2 or T3, but eCO_2 reduced aggregate leaf N content (- 18%) at T3 in Yitpi only (Cultivar × CO_2 p < 0.05) (Table S7). Yitpi had higher grain N content (+ 26%) than Scout in control plants grown at a CO_2 (Fig. 8g, h; Table S6). In control plants, eCO_2 significantly reduced grain protein content in Yitpi (- 18%, p < 0.05) but not in Scout due to significant cultivar × CO_2 interaction (p < 0.01), while HS had no effect on protein content in either cultivar (Fig. 8g, h; Table S6).



Fig. 6 Relationships with leaf gas exchange and grain yield across treatments. CO_2 assimilation rate plotted as a function of stomatal conductance (**a**) (both a CO_2 and e CO_2 grown plants measured at 400 µL L⁻¹), J_{max} plotted as a function of V_{cmax} (**b**) and grain protein plotted as a function of yield (**c**) in Scout (Circles) and Yitpi (Triangles). Ambient and elevated CO_2 are depicted in blue and red, respectively. Control and heat stressed plants depicted using open and closed symbols. Panel a depicts data for control, HS1, HS2 and both heat stresses (HS1+2), while panels b and cinclude only control and HS1+2

Discussion

Two wheat cultivars with contrasting morphology and phenology, but similar photosynthesis and grain yield

The effects of future climate conditions, including eCO_2 , will depend on the environmental conditions (e.g., water and heat stress) and the crop's agronomic features. Here, we compared the interactive effects of eCO_2 and HS on two commercial wheat cultivars, Scout and Yitpi, with contrasting phenology and growth habit. Plants were grown under well-watered and fertilized conditions to remove any confounding effects of water or nutrient limitations on the eCO_2 or HS responses. RH was kept constant to minimize the negative impact of dry air during HS. Finally, we compared the effects of applying HS at the vegetative and flowering stages.

Free tillering Yitpi produced substantially more tillers, leaf area and biomass relative to the faster developing Scout. Accordingly, our first hypothesis predicted that Yitpi will have higher grain yield. The results only partially supported this hypothesis, because relative to Yitpi, Scout had higher harvest index (HI) due to its early maturing and senescing habit. While Yitpi initiated more tillers, a lower proportion of these tillers produced ears and filled grains. In contrast, Scout produced less tillers but flowered earlier which allowed enough time for all its tillers to produce ears and fill bigger grains by the final harvest. Hence, both cultivars had relatively similar yields due to bigger grain size in Scout and higher grain number in Yitpi. It is worth noting that some field trials have reported slightly higher grain yields in Scout than Yitpi (National variety trial report, GRDC, 2014). Our results are consistent with a previous study using different wheat cultivars with contrasting source-sink relationships which reported that the freely tillering cultivar "Silverstar" translated into more spikes while restricted tillering cultivar "H45" had more and heavier kernels per spike than "Silverstar" (Tausz-Posch et al. 2015). Thus, early vigor and maturity compared to high tillering capacity seem to be equally beneficial traits for high grain yield in the Australian environment.

The two wheat cultivars showed similar photosynthetic traits and response to temperature and eCO_2 . The temperature response of g_m as well as the values recorded at 25 °C (0.25–0.31 mol m⁻² s⁻¹ bar⁻¹) for Scout and Yitpi (Fig. 2b) were similar to what has been reported for wheat (0.39) and other crop species (cotton = 0.73, soybean = 0.49 and rice = 0.67) (Von Caemmerer and Evans 2015; Jahan et al. 2021). In contrast to our expectations that Scout would have higher WUE due to its selection based on a carbon isotope discrimination gene (Condon et al. 2004), both wheat cultivars showed similar iWUE under most measurement and growth conditions in this study (Fig. 5e, f; Table S2).

Elevated CO₂ stimulated photosynthesis but reduced photosynthetic capacity in both cultivars

Long term exposure to eCO_2 may reduce photosynthetic capacity due to lower amount of Rubisco in a process referred as 'acclimation' (Nie et al. 1995; Rogers and Humphries 2000; Ainsworth et al. 2003). Alternatively, eCO_2 may 'down-regulate' photosynthetic capacity by reducing

Fig. 7 Response of plant growth and morphological traits to elevated CO₂ and HS: Total dry mass (a, b), tillers or number of tillers (c, d), leaf area (e, f), leaf number (g, h) and height (i, j) were measured at different time points across the life cycle of wheat cultivars Scout (Circles) and Yitpi (Triangles). Ambient and elevated CO2 are depicted in blue and red color, respectively. Open symbols connected with solid cultivars and closed symbols connected with dashed cultivars represent control and HS plants, respectively. HS1 and HS2 depict the timing of HS applied at 10 and 15 weeks after planting respectively



Rubisco activation or other regulatory mechanisms without affecting Rubisco content (Delgado et al. 1994). In the current study, eCO₂ similarly increased A_{growth} (+21%) measured at growth CO₂ and reduced both A_{sat} (-12%) measured at common CO₂ (Fig. 3, T2) and V_{cmax} (Fig. 5i, j) in both cultivars. In contrast, flag leaf N (and possibly Rubisco) was not significantly affected in either cultivar (Table S6). Hence, the wheat cultivars have likely undergone a photosynthetic downregulation (less rubisco activation)—rather than acclimation (reduction in rubisco content)—in response to eCO₂ (Delgado et al. 1994; Leakey et al. 2009). These results partially countered our second hypothesis suggesting that Yitpi will show less photosynthetic acclimation due to its higher sink capacity. The interaction of eCO_2 with plant traits are complex. On the one hand, eCO_2 is expected to cause less feedback inhibition on photosynthesis in plants with high sink capacity (Ainsworth et al. 2004). On the other, fastgrowing plants show a proportionally larger response to eCO_2 (Poorter and Navas 2003). Hence, high tillering in Yitpi and fast development in Scout both led to a relatively small observed photosynthetic downregulation in response to growth at eCO_2 . This allowed a sustained photosynthetic stimulation, which in turn led to a significant biomass and yield enhancement by CO_2 enrichment in both wheat cultivars (Figs. 6, 7). Photosynthetic responses of wheat in current study are in agreement with earlier enclosure studies Fig. 8 Response of total plant dry mass and grain parameters to growth at eCO₂ and HS at maturity (T4): Total dry mass (a, b), grain dry mass (c, d), grain number (e, f) and grain nitrogen (g, h) were measured at the final harvest. Cultivars indicate means and shaded region is 95% confidence interval. Ambient and elevated CO2 are depicted in blue and red color respectively. Heat stress levels include plants not exposed to any heat stress (control) and both heat stresses (HS1+2). Statistical significance levels (t-test) for the growth condition within each cultivar is shown and they are: * = p < 0.05; ** = p < 0.01: *** = p < 0.001



which generally have higher response to eCO_2 than the FACE studies (Kimball et al. 1995, 1999; Hunsaker et al. 1996; Osborne et al. 1998; Long et al. 2006; Cai et al. 2016).

Elevated CO₂ stimulated grain yield similarly in both wheat cultivars

In disagreement with our second hypothesis, eCO_2 similarly stimulated plant biomass and grain yield in early-maturing Scout and high tillering Yitpi (Figs. 6, 7; Table S4, S5). In Scout, the biomass stimulation was

associated with increased tillering (one extra tiller per plant). In contrast, Yitpi produced many tillers at aCO₂ and the additional fixed carbon at eCO₂ was allocated to the existing tillers. At seed maturity, eCO₂ stimulated grain yield similarly in both cultivars as a result of the trade-off between grain yield components (Dias de Oliveira et al. 2015). In particular, eCO₂ stimulated grain number in both cultivars, while grain size increased in Scout only (Fig. 8; Table S6). Generally, eCO₂ stimulates grain yield by increasing the number of tillers and consequently, ears per plant (Zhang et al. 2010; Bennett et al. 2012) as a result of increased carbon supply due photosynthetic stimulation by eCO_2 , which has also been reported in FACE studies (Högy et al. 2009; Tausz-Posch et al. 2015; Fitzgerald et al. 2016). However, in our study, the increase in grain yield at eCO_2 was mainly due to the increase in the number of grains per ear. In line with our results, Tausz-Posch et al. (2015) reported comparable grain yield stimulation by eCO_2 in two different wheat cultivars with contrasting source-sink relationships. Moreover, grain yield of twenty wheat cultivars that differed in tillering propensity, water soluble carbohydrate accumulation, early vigor and transpiration efficiency responded similarly to eCO_2 in glasshouse settings (Ziska et al. 2004; Bourgault et al. 2013).

Elevated CO₂ reduced grain N in Yitpi only

Overall, there is a negative relationship between grain yield and quality (Taub et al. 2008; Pleijel and Uddling 2012). Hence, increased grain yield at eCO_2 results in lower grain N and hence protein content (Seneweera and Conroy 1997; Bahrami et al. 2017). In our study, eCO₂ reduced grain N in Yitpi under control conditions. Scout was characterized by having larger grains which accumulated less N than Yitpi. Moreover, eCO₂ reduced total leaf N (-18%) at T3 and grain N (-17%) at T4 in Yitpi but not in Scout. This is consistent with the results from FACE study with same cultivars which reported - 14% reduction in N content by eCO₂ in above ground dry mass in Yitpi but not in Scout under well-watered conditions (Bahrami et al. 2017). The higher biomass accumulation in free tillering Yitpi may have exhausted the nutrient supply or nitrogen uptake may have slowed down (Carreras Navarro et al. 2020), such that further biomass stimulation by eCO₂ lead to a significant dilution in N content (Taub and Wang 2008).

Wheat cultivars with early vigour such as Scout have greater root biomass accumulation as well as greater early N uptake which may have avoided a negative effect of eCO₂ on leaf and grain N (Liao et al. 2004; Bahrami et al. 2017). Accordingly, Scout maintained a higher N utilization efficiency (grain yield per total plant N) relative to Yitpi under all treatments (Table S6). Increased grain yield is strongly associated with higher grain number per unit area (Zhang et al. 2010; Bennett et al. 2012) which dilutes the amount of N translocated per grain. Quality deterioration due to lower protein via reduced N is of critical concern in future high CO2 climate considering that even additional supply of N does not prevent N dilution in grain under eCO_2 (Tausz et al. 2017). In addition, eCO_2 has strong detrimental effect on other nutrient availability and remobilization from leaves to grains (Tcherkez et al. 2020).

HS had little effects on wheat photosynthesis or yield at aCO₂

One of the key finding of this study was that the application of HS events (HS1, HS2 or HS1+2) was not detrimental to aCO₂-grown wheat plants (Figs. 5, 7; Tables S4, S5, S6). Thus, our hypothesis that HS will reduce photosynthesis, biomass and yield at aCO₂ was rejected. This finding is in contrast to previously reported studies where HS reduced the grain yield and negatively affected the growth and development in wheat (Stone and Nicolas 1996, 1998; Farooq et al. 2011; Coleman et al. 1991). During heat waves in the field, the vapor pressure deficit (VPD) increases and soil moisture decreases leading to lower stomatal conductance and consequently lower transpiration rate. Thus, plants are unable to cool down and leaf temperatures rise beyond optimum levels causing damage. The negligible effect of HS in our study could be explained by the ability of well-watered plants to maintain leaf temperature below damaging levels due to transpirational cooling (Perera et al. 2019; Deva et al. 2020) even with air temperatures reaching up to 38 °C. At moderate (~60%) relative humidity, there is sufficient water vapour gradient to sustain high transpiration rates when soil water is available, as was the case in our experiment. In most cases, g_s was not significantly affected (Tables S1 and S2), and even slightly higher at T3 in HS-pants relative to the control (Fig. 5c, d) Well-watered crops can maintain grainfilling rate, duration and size under HS (Dupont et al. 2006). and high temperatures can increase crop yields if not exceeding critical optimum growth temperature (Welch et al. 2010). Also, in the current study, the night temperatures were not increased during HS which favors plant growth by reducing respiratory losses (Prasad et al. 2008).

In particular, HS did not elicit a direct negative impact on photosynthesis or chlorophyll fluorescence in either cultivar or CO₂ treatment. During HS, high temperature transiently reduced maximum efficiency of PSII (Fv/Fm) in both cultivars and CO₂ treatments (Fig. 4e-h). However, unchanged Fv/Fm measured at 25 °C confirmed that photosynthesis did not suffer long-term damage during or after HS. Moreover, HS was not severe enough to negatively affect A_{growth} measured at 25 °C. These results are corroborated by the insensitivity of V_{cmax} and J_{max} to HS (Fig. 5i–l), but contrast with previously reported studies where HS reduced photosynthesis in wheat at the vegetative (Wang et al. 2008) and the flowering (Chavan et al. 2019; Balla et al. 2019) stages. HS lowers membrane thermostability by inducing reactive oxygen species (ROS) and altering the membrane protein structures, which lead to changes in the fluidity of the thylakoid membrane and separation of light harvesting complex from the photosystems (Wahid et al. 2007; Poudel 2020). We were unable to measure leaf temperatures in the current study, but we speculate that, in well-watered wheat plants

growing at moderate RH, leaf temperatures might not have increased beyond damaging levels to the membranes during the HS events.

Repeated HS may result in priming which involves preexposure of plants to a stimulating factor such as HS (Wang et al. 2017) and enable plants to cope better with later HS events (Balla et al. 2021). However, there was no difference between HS applied at the vegetative (HS1) and/or flowering stage (HS2) in rejection of our fourth hypothesis, and this may additionally be due to the short term duration of the two HS cycles (3 days each). Hence, our study demonstrated the benign effect that HS has on crop yield when separated from water stress and plants are able to transpire.

HS precluded an eCO₂ response in biomass and grain yield

In our study, the impact of HS depended on the wheat cultivar and growth CO_2 (Tables S1 and S4). Elevated CO_2 and temperature interactions can be complex, dynamic and difficult to generalize as they can go in any direction depending on plant traits and other environmental conditions (Rawson 1992). Plant development is generally accelerated by increased temperature; eCO_2 can accelerate it further in some instances or may have neutral or even retarding effects in other cases (Rawson 1992).

While eCO₂ stimulated wheat biomass and grain yield under control (non-HS) conditions, HS precluded a yield response to eCO₂ in Yitpi and reduced biomass and yield in eCO₂-grown Scout relative to aCO₂-grown counterparts (Fig. 8). These results are in contrast with previous studies that reported similar wheat yield reduction at ambient or elevated CO_2 in response to severe (Chavan et al. 2019) or moderate HS (Zhang et al. 2018). The results also partially refuted our third hypothesis that HS may decrease yield more at aCO_2 than eCO_2 , while partially agreeing that HS will have a more negative impact on Scout relative to Yitpi, albeit for different reasons than what we originally suggested. The negative effect of HS on Scout biomass and grain yield at eCO₂ occurred despite the eCO₂ stimulation of A_{growth} under HS (T3, Fig. 5). However, over the long term, A_{growth} was stimulated in eCO₂-grown Yitpi and not Scout (Fig. 5).

Lack of a biomass stimulation despite high photosynthetic rates during HS under eCO_2 could be due to the short duration of HS (3 days), which may not have been long enough to stimulate biomass gain. In addition, nutrient limitation at eCO_2 may have restricted the eCO_2 growth response. Typically, eCO_2 studies show reduced N content in wheat and other crops (Taub and Wang 2008; Leakey et al. 2009; Bahrami et al. 2017). Hence, the wheat plants may have exhausted available nutrients due to increased demand by growing sinks at eCO_2 , which may limited further stimulation by high temperature. HS may be more damaging at eCO_2 due to reduce transpirational cooling as a result of reduced g_s at eCO_2 , leading to higher leaf temperatures. However, A_{growth} increased in response to high temperature (35 °C) under eCO_2 but not under aCO_2 during HS (Fig. 4a–d), which refutes the suggestion of HS-damage to photosynthesis.

Higher g_s during HS at moderate RH in well-watered conditions may increase A_{growth} by increasing C_i in both aCO₂ and eCO₂ grown plants. Furthermore, lower photorespiration under eCO₂ allows additional increase in A_{growth} with temperature when measured at 35 °C relative to 25 °C (Long 1991). Under aCO₂, photorespiration increases with temperature reducing A_{growth} measured at 35 °C relative to 25 °C. Our results also point to a shift in T_{ont} of photosynthesis (~24 °C at aCO₂) to higher temperatures for plants grown at eCO₂ (Sage and Kubien 2007). This would come about as a result of lower photorespiration at eCO_2 as well as the slight upregulation of photosynthetic rates observed in eCO2-grown Scout at the recovery stage of HS (Fig. 4a, c). However, at T3, A_{growth} was similar between aCO_2 and eCO_2 grown plants (Fig. 5a) indicating the short-term nature of this photosynthetic upregulation.

Conclusions

The two wheat cultivars, Scout and Yitpi differed in growth and development but produced similar grain yield. Under control conditions, eCO2 stimulated biomass and yield similarly in both cultivars. HS was not damaging to photosynthesis, growth, biomass or grain yield under wellwatered and moderate RH conditions. However, HS interacted with eCO_2 , leading to similar or lower biomass and grain yield at eCO₂ relative to both aCO₂ in plants exposed to HS. This interactive effect precluded the positive effects of eCO₂ in HS-plants. eCO₂ improved photosynthetic rates in control and HS plants. Also, high temperature stimulated photosynthesis under eCO₂ but not under aCO₂ during HS which suggests increased optimum temperature of photosynthesis at eCO_2 . We speculate that in the current study, HS plants were able to cool down using high transpiration which helped to maintain lower leaf temperatures despite high air temperatures during HS. The current study provides important insights into the effect of short-term moderate temperature increases in well-watered conditions under future elevated CO2, potential role of transpirational cooling during HS and interactions between HS and eCO₂ which will be useful in breeding cultivars for future climate and improving crop model accuracy to predict crop performance in future high CO₂ environment with frequent heat waves.

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Author contributions All authors conceived the project. SGC maintained the plants and collected the data. SGC and RAD analysed the data. SGC and OG prepared the manuscript with input from other co-authors.

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Data availability All data supporting the findings of this study are available within the paper and within its supplementary materials published online. Reuse of the data is permitted after obtaining permission from the corresponding author.

Code availability Software application and codes used are all publicly available.

Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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References

- Ainsworth EA, Davey PA, Hymus GJ, Osborne CP, Rogers A, Blum H, Nösberger J, Long SP (2003) Is stimulation of leaf photosynthesis by elevated carbon dioxide concentration maintained in the long term? A test with *Lolium perenne* grown for 10 years at two nitrogen fertilization levels under free air CO2 enrichment (FACE). Plant Cell Environ 26:705–714. https://doi.org/ 10.1046/j.1365-3040.2003.01007.x
- Ainsworth EA, Rogers A, Nelson R, Long SP (2004) Testing the "source–sink" hypothesis of down-regulation of photosynthesis in elevated [CO2] in the field with single gene substitutions in

glycine max. Agric for Meteorol 122:85–94. https://doi.org/10. 1016/j.agrformet.2003.09.002

- Alonso A, Pérez P, Martínez-Carrasco R (2009) Growth in elevated CO2 enhances temperature response of photosynthesis in wheat. Physiol Plant 135:109–120. https://doi.org/10.1111/j.1399-3054. 2008.01177.x
- Asseng S, Ewert F, Rosenzweig C, Jones JW, Hatfield JL, Ruane AC, Boote KJ, Thorburn PJ, Rötter RP, Cammarano D, Brisson N, Basso B, Martre P, Aggarwal PK, Angulo C, Bertuzzi P, Biernath C, Challinor AJ, Doltra J, Gayler S, Goldberg R, Grant R, Heng L, Hooker J, Hunt LA, Ingwersen J, Izaurralde RC, Kersebaum KC, Müller C, Naresh Kumar S, Nendel C, O'Leary G, Olesen JE, Osborne TM, Palosuo T, Priesack E, Ripoche D, Semenov MA, Shcherbak I, Steduto P, Stöckle C, Stratonovitch P, Streck T, Supit I, Tao F, Travasso M, Waha K, Wallach D, White JW, Williams JR, Wolf J (2013) Uncertainty in simulating wheat yields under climate change. Nat Clim Change 3:827–832. https://doi. org/10.1038/nclimate1916
- Asseng S, Ewert F, Martre P, Rötter RP, Lobell DB, Cammarano D, Kimball BA, Ottman MJ, Wall GW, White JW, Reynolds MP, Alderman PD, Prasad PVV, Aggarwal PK, Anothai J, Basso B, Biernath C, Challinor AJ, De Sanctis G, Doltra J, Fereres E, Garcia-Vila M, Gayler S, Hoogenboom G, Hunt LA, Izaurralde RC, Jabloun M, Jones CD, Kersebaum KC, Koehler A-K, Müller C, Naresh Kumar S, Nendel C, O'Leary G, Olesen JE, Palosuo T, Priesack E, Eyshi Rezaei E, Ruane AC, Semenov MA, Shcherbak I, Stöckle C, Stratonovitch P, Streck T, Supit I, Tao F, Thorburn PJ, Waha K, Wang E, Wallach D, Wolf J, Zhao Z, Zhu Y (2015) Rising temperatures reduce global wheat production. Nat Clim Change 5:143–147. https://doi.org/10. 1038/nclimate2470
- Bahrami H, De Kok LJ, Armstrong R, Fitzgerald GJ, Bourgault M, Henty S, Tausz M, Tausz-Posch S (2017) The proportion of nitrate in leaf nitrogen, but not changes in root growth, are associated with decreased grain protein in wheat under elevated [CO2]. J Plant Physiol 216:44–51. https://doi.org/10.1016/j. jplph.2017.05.011
- Balla K, Karsai I, Bónis P, Kiss T, Berki Z, Horváth Á, Mayer M, Bencze S, Veisz O (2019) Heat stress responses in a large set of winter wheat cultivars (*Triticum aestivum* L.) depend on the timing and duration of stress. PLoS ONE 14:e0222639. https:// doi.org/10.1371/journal.pone.0222639
- Balla K, Karsai I, Kiss T, Horváth Á, Berki Z, Cseh A, Bónis P, Árendás T, Veisz O (2021) Single versus repeated heat stress in wheat: what are the consequences in different developmental phases? PLoS ONE 16:e0252070. https://doi.org/10.1371/journ al.pone.0252070
- Bennett D, Izanloo A, Reynolds M, Kuchel H, Langridge P, Schnurbusch T (2012) Genetic dissection of grain yield and physical grain quality in bread wheat (*Triticum aestivum* L.) under waterlimited environments. Theor Appl Genet 125:255–271. https:// doi.org/10.1007/s00122-012-1831-9
- Berry J, Bjorkman O (1980) Photosynthetic response and adaptation to temperature in higher plants. Annu Rev Plant Physiol 31:491– 543. https://doi.org/10.1146/annurev.pp.31.060180.002423
- Bourgault M, Dreccer MF, James AT, Chapman SC (2013) Genotypic variability in the response to elevated CO2 of wheat lines differing in adaptive traits. Funct Plant Biol 40:172–184
- Cai C, Yin X, He S, Jiang W, Si C, Struik PC, Luo W, Li G, Xie Y, Xiong Y, Pan G (2016) Responses of wheat and rice to factorial combinations of ambient and elevated CO2 and temperature in FACE experiments. Glob Change Biol 22:856–874. https://doi. org/10.1111/gcb.13065
- Cardoso-Vilhena J, Barnes J (2001) Does nitrogen supply affect the response of wheat (*Triticum aestivum* cv. Hanno) to the

combination of elevated CO2 and O3? J Exp Bot 52:1901–1911. https://doi.org/10.1093/jexbot/52.362.1901

- Carreras Navarro E, Lam SK, Trębicki P (2020) Elevated carbon dioxide and nitrogen impact wheat and its aphid pest. Front Plant Sci. https://doi.org/10.3389/fpls.2020.605337
- Chavan SG, Duursma RA, Tausz M, Ghannoum O (2019) Elevated CO2 alleviates the negative impact of heat stress on wheat physiology but not on grain yield. J Exp Bot. https://doi.org/10.1093/ jxb/erz386
- Coleman JS, Rochefort L, Bazzaz FA, Woodward FI (1991) Atmospheric CO2, plant nitrogen status and the susceptibility of plants to an acute increase in temperature. Plant Cell Environ 14:667–674. https://doi.org/10.1111/j.1365-3040.1991.tb01539.x
- Condon AG, Richards RA, Rebetzke GJ, Farquhar GD (2004) Breeding for high water-use efficiency. J Exp Bot 55:2447–2460. https:// doi.org/10.1093/jxb/erh277
- Crous KY, Quentin AG, Lin Y-S, Medlyn BE, Williams DG, Barton CVM, Ellsworth DS (2013) Photosynthesis of temperate *Eucalyptus globulus* trees outside their native range has limited adjustment to elevated CO2 and climate warming. Glob Change Biol 19:3790–3807. https://doi.org/10.1111/gcb.12314
- Delgado E, Mitchell RAC, Parry MAJ, Driscoll SP, Mitchell VJ, Lawlor DW (1994) Interacting effects of CO2 concentration, temperature and nitrogen supply on the photosynthesis and composition of winter wheat leaves. Plant Cell Environ 17:1205–1213. https://doi.org/10.1111/j.1365-3040.1994. tb02018.x
- Deva CR, Urban MO, Challinor AJ, Falloon P, Svitákova L (2020) Enhanced leaf cooling is a pathway to heat tolerance in common bean. Front Plant Sci 11:19. https://doi.org/10.3389/fpls. 2020.00019
- Dias de Oliveira EA, Siddique KHM, Bramley H, Stefanova K, Palta JA (2015) Response of wheat restricted-tillering and vigorous growth traits to variables of climate change. Glob Change Biol 21:857–873. https://doi.org/10.1111/gcb.12769
- Dupont FM, Hurkman WJ, Vensel WH, Tanaka C, Kothari KM, Chung OK, Altenbach SB (2006) Protein accumulation and composition in wheat grains: effects of mineral nutrients and high temperature. Eur J Agron 25:96–107. https://doi.org/10.1016/j.eja. 2006.04.003
- Duursma RA (2015) Plantecophys—an R package for analysing and modelling leaf gas exchange data. PLoS ONE 10:e0143346. https://doi.org/10.1371/journal.pone.0143346
- Eckardt NA, Portis AR (1997) Heat denaturation profiles of ribulose-1,5-bisphosphate carboxylase/oxygenase (rubisco) and rubisco activase and the inability of rubisco activase to restore activity of heat-denatured rubisco. Plant Physiol 113:243–248
- Evans J, Von Caemmerer S (2013) Temperature response of carbon isotope discrimination and mesophyll conductance in tobacco. Plant Cell Environ 36:745–756. https://doi.org/10.1111/j.1365-3040.2012.02591.x
- Evans J, Sharkey T, Berry J, Farquhar G (1986) Carbon isotope discrimination measured concurrently with gas exchange to investigate CO2 diffusion in leaves of higher plants. Funct Plant Biol 13:281–292
- Farooq M, Bramley H, Palta JA, Siddique KHM (2011) Heat stress in wheat during reproductive and grain-filling phases. Crit Rev Plant Sci 30:491–507. https://doi.org/10.1080/07352689.2011. 615687
- Farquhar GD, Cernusak LA (2012) Ternary effects on the gas exchange of isotopologues of carbon dioxide. Plant Cell Environ 35:1221– 1231. https://doi.org/10.1111/j.1365-3040.2012.02484.x
- Fischer RA (1980) Influence of water stress on crop yield in semiarid regions
- Fitzgerald GJ, Tausz M, O'Leary G, Mollah MR, Tausz-Posch S, Seneweera S, Mock I, Löw M, Partington DL, McNeil D,

Norton RM (2016) Elevated atmospheric [CO2] can dramatically increase wheat yields in semi-arid environments and buffer against heat waves. Glob Change Biol 22:2269–2284. https://doi. org/10.1111/gcb.13263

- García GA, Dreccer MF, Miralles DJ, Serrago RA (2015) High night temperatures during grain number determination reduce wheat and barley grain yield: a field study. Glob Change Biol 21:4153– 4164. https://doi.org/10.1111/gcb.13009
- Harley PC, Thomas RB, Reynolds JF, Strain BR (1992) Modelling photosynthesis of cotton grown in elevated CO2. Plant Cell Environ 15:271–282. https://doi.org/10.1111/j.1365-3040.1992.tb00974.x
- Hatfield JL, Prueger JH (2015) Temperature extremes: Effect on plant growth and development. Weather Clim Extrem 10(Part A):4–10. https://doi.org/10.1016/j.wace.2015.08.001
- Hocking PJ, Meyer CP (1991) Effects of CO2 enrichment and nitrogen stress on growth and partitioning of dry matter and nitrogen in wheat and maize. Aust J Plant Physiol (australia) 18:339
- Hogan KP, Smith AP, Ziska LH (1991) Potential effects of elevated CO2 and changes in temperature on tropical plants. Plant Cell Environ 14:763–778. https://doi.org/10.1111/j.1365-3040.1991. tb01441.x
- Högy P, Wieser H, Köhler P, Schwadorf K, Breuer J, Franzaring J, Muntifering R, Fangmeier A (2009) Effects of elevated CO2 on grain yield and quality of wheat: results from a 3-year free-air CO2 enrichment experiment. Plant Biol 11:60–69. https://doi. org/10.1111/j.1438-8677.2009.00230.x
- Houshmandfar A, Fitzgerald GJ, O'Leary G, Tausz-Posch S, Fletcher A, Tausz M (2017) The relationship between transpiration and nutrient uptake in wheat changes under elevated atmospheric CO2. Physiol Plant. https://doi.org/10.1111/ppl.12676
- Hunsaker DJ, Kimball BA, Pinter PJ, LaMorte RL, Wall GW (1996) Carbon dioxide enrichment and irrigation effects on wheat evapotranspiration and water use efficiency. Trans ASAE (USA) 39:1345
- IPCC (2014) Climate change 2014: synthesis report. Contributions of Working Group I, Working Group II and Working Group III, special report on renewable energy sources and climate change mitigation and special report on managing the risks of extreme events and disasters to advance climate change adaptation to the fifth assessment report (AR5). IPCC, Geneva, Switzerland. p 80
- Jahan E, Thomson PC, Tissue DT (2021) Mesophyll conductance in two cultivars of wheat grown in glacial to super-elevated CO2 concentrations. J Exp Bot 72:7191–7202. https://doi.org/10. 1093/jxb/erab320
- Jauregui I, Aroca R, Garnica M, Zamarreño ÁM, García-Mina JM, Serret MD, Parry M, Irigoyen JJ, Aranjuelo I (2015) Nitrogen assimilation and transpiration: key processes conditioning responsiveness of wheat to elevated [CO2] and temperature. Physiol Plantarum 155:338–354. https://doi.org/10.1111/ppl. 12345
- Kimball BA (1983) Carbon dioxide and agricultural yield: an assemblage and analysis of 430 prior observations. Agron J 75:779–788. https://doi.org/10.2134/agronj1983.000219620075000 50014x
- Kimball BA (2016) Crop responses to elevated CO2 and interactions with H2O, N, and temperature. Curr Opin Plant Biol 31:36–43. https://doi.org/10.1016/j.pbi.2016.03.006
- Kimball BA, Pinter PJ Jr, Garcia RL, La Morte RL, Wall GW, Hunsaker DJ, Wechsung G, Wechsung F, Kartschall T (1995) Productivity and water use of wheat under free-air CO2 enrichment. Glob Change Biol 1:429–442
- Kimball BA, LaMorte RL, Pinter PJ, Wall GW, Hunsaker DJ, Adamsen FJ, Leavitt SW, Thompson TL, Matthias AD, Brooks TJ (1999) Free-air CO2 enrichment and soil nitrogen effects on energy balance and evapotranspiration of wheat. Water Resour Res 35:1179–1190. https://doi.org/10.1029/1998WR900115

- Krenzer EG, Moss DN (1975) Carbon dioxide enrichment effects upon yield and yield components in wheat. Crop Sci 15:71–74. https:// doi.org/10.2135/cropsci1975.0011183X001500010020x
- Lanigan GJ, Betson N, Griffiths H, Seibt U (2008) Carbon Isotope fractionation during photorespiration and carboxylation in senecio. Plant Physiol 148:2013–2020. https://doi.org/10.1104/pp. 108.130153
- Leakey ADB, Ainsworth EA, Bernacchi CJ, Rogers A, Long SP, Ort DR (2009) Elevated CO2 effects on plant carbon, nitrogen, and water relations: six important lessons from FACE. J Exp Bot 60:2859–2876. https://doi.org/10.1093/jxb/erp096
- Liao M, Fillery IRP, Palta JA (2004) Early vigorous growth is a major factor influencing nitrogen uptake in wheat. Funct Plant Biol 31:121–129. https://doi.org/10.1071/fp03060
- Lobell DB, Gourdji SM (2012) The influence of climate change on global crop productivity. Plant Physiol 160:1686–1697. https:// doi.org/10.1104/pp.112.208298
- Long SP (1991) Modification of the response of photosynthetic productivity to rising temperature by atmospheric CO2 concentrations: has its importance been underestimated? Plant Cell Environ 14:729–739. https://doi.org/10.1111/j.1365-3040.1991.tb01439.x
- Long SP, Ainsworth EA, Leakey ADB, Nösberger J, Ort DR (2006) Food for thought: lower-than-expected crop yield stimulation with rising CO2 concentrations. Science 312:1918–1921. https:// doi.org/10.1126/science.1114722
- Macabuhay AA (2016) Physiological and biochemical responses of wheat to combined heat stress and elevated CO2 during grain-filling
- Macabuhay A, Houshmandfar A, Nuttall J, Fitzgerald GJ, Tausz M, Tausz-Posch S (2018) Can elevated CO2 buffer the effects of heat waves on wheat in a dryland cropping system? Environ Exp Bot 155:578–588. https://doi.org/10.1016/j.envexpbot.2018.07.029
- Medlyn BE, Dreyer E, Ellsworth D, Forstreuter M, Harley PC, Kirschbaum MUF, Le Roux X, Montpied P, Strassemeyer J, Walcroft A, Wang K, Loustau D (2002) Temperature response of parameters of a biochemically based model of photosynthesis. II. A review of experimental data. Plant Cell Environ 25:1167–1179. https:// doi.org/10.1046/j.1365-3040.2002.00891.x
- Mishra D, Shekhar S, Chakraborty S, Chakraborty N (2021) High temperature stress responses and wheat: impacts and alleviation strategies. Environ Exp Bot. https://doi.org/10.1016/j.envexpbot. 2021.104589
- Mitchell RAC, Mitchell VJ, Driscoll SP, Franklin J, Lawlor DW (1993) Effects of increased CO2 concentration and temperature on growth and yield of winter wheat at two levels of nitrogen application. Plant Cell Environ 16:521–529. https://doi.org/10. 1111/j.1365-3040.1993.tb00899.x
- Morison JIL, Lawlor DW (1999) Interactions between increasing CO2 concentration and temperature on plant growth. Plant Cell Environ 22:659–682. https://doi.org/10.1046/j.1365-3040.1999. 00443.x
- Mosse J (1990) Nitrogen-to-protein conversion factor for ten cereals and six legumes or oilseeds. A reappraisal of its definition and determination. Variation according to species and to seed protein content. J Agric Food Chem 38:18–24. https://doi.org/10.1021/ jf00091a004
- Mulholland BJ, Craigon J, Black CR, Colls JJ, Atherton J, Landon G (1998) Growth, light interception and yield responses of spring wheat (*Triticum aestivum* L.) grown under elevated CO2 and O3 in open-top chambers. Glob Change Biol 4:121–130. https://doi. org/10.1046/j.1365-2486.1998.00112.x
- Nie GY, Long SP, Garcia RL, Kimball BA, Lamorte RL, Pinter PJ, Wall GW, Webber AN (1995) Effects of free-air CO2 enrichment on the development of the photosynthetic apparatus in wheat, as

indicated by changes in leaf proteins. Plant Cell Environ 18:855–864. https://doi.org/10.1111/j.1365-3040.1995.tb00594.x

- Osborne CP, Roche JL, Garcia RL, Kimball BA, Wall GW, Pinter PJ, Morte RLL, Hendrey GR, Long SP (1998) Does leaf position within a canopy affect acclimation of photosynthesis to elevated CO2? Analysis of a wheat crop under free-air CO2 enrichment. Plant Physiol 117:1037–1045. https://doi.org/10.1104/pp.117.3. 1037
- Pacificseeds (2009) LongReach scout wheat.https://www.pacificsee ds.com.au/images/Icons/Products/Wheat/SNSWVICSA/Scout VICSA.pdf. Accessed 15 Jan 2017
- Perera RS, Cullen BR, Eckard RJ (2019) Using leaf temperature to improve simulation of heat and drought stresses in a biophysical model. Plants (basel) 9:8. https://doi.org/10.3390/plants9010008
- Pleijel H, Uddling J (2012) Yield vs. quality trade-offs for wheat in response to carbon dioxide and ozone. Glob Change Biol 18:596–605. https://doi.org/10.1111/j.1365-2486.2011.2489.x
- Poorter H (1993) Interspecific variation in the growth response of plants to an elevated ambient CO2 concentration. Vegetatio 104–105:77–97. https://doi.org/10.1007/BF00048146
- Poorter H, Navas M-L (2003) Plant growth and competition at elevated CO2: on winners, losers and functional groups. New Phytol 157:175–198. https://doi.org/10.1046/j.1469-8137.2003.00680.x
- Porter JR, Gawith M (1999) Temperatures and the growth and development of wheat: a review. Eur J Agron 10:23–36. https://doi.org/ 10.1016/S1161-0301(98)00047-1
- Poudel PB (2020) Heat stress effects and tolerance in wheat: a review. J Biol Today's World 9:6
- Prasad PVV, Pisipati SR, Ristic Z, Bukovnik U, Fritz AK (2008) Impact of nighttime temperature on physiology and growth of spring wheat. Crop Sci 48:2372–2380. https://doi.org/10.2135/ cropsci2007.12.0717
- R Core Team (2022) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/
- Rawson H (1992) Plant responses to temperature under conditions of elevated CO2. Aust J Bot 40:473–490
- Rogers A, Humphries SW (2000) A mechanistic evaluation of photosynthetic acclimation at elevated CO2. Glob Change Biol 6:1005–1011. https://doi.org/10.1046/j.1365-2486.2000.00375.x
- Sadras V, Dreccer MF (2015) Adaptation of wheat, barley, canola, field pea and chickpea to the thermal environments of Australia. Crop Pasture Sci 66:1137–1150. https://doi.org/10.1071/CP15129
- Sage RF, Kubien DS (2007) The temperature response of C3 and C4 photosynthesis. Plant Cell Environ 30:1086–1106. https://doi.org/10.1111/j.1365-3040.2007.01682.x
- Seednet (2005) Yitpi. http://www.seednet.com.au/documents/Yitpi Factsheet2005.pdf. Accessed 15 Jan 2017
- Seneweera SP, Conroy JP (1997) Growth, grain yield and quality of rice (*Oryza sativa* L.) in response to elevated CO2 and phosphorus nutrition. Soil Sci Plant Nutr 43:1131–1136. https://doi.org/ 10.1080/00380768.1997.11863730
- Sionit N, Mortensen DA, Strain BR, Hellmers H (1981) Growth response of wheat to CO2 enrichment and different levels of mineral nutrition. Agron J 73:1023–1027. https://doi.org/10. 2134/agronj1981.00021962007300060027x
- Stone P, Nicolas M (1994) Wheat cultivars vary widely in their responses of grain yield and quality to short periods of postanthesis heat stress. Funct Plant Biol 21:887–900
- Stone PJ, Nicolas M (1996) Effect of timing of heat stress during grain filling on two wheat varieties differing in heat tolerance. II. Fractional protein accumulation. Funct Plant Biol 23:739–749
- Stone PJ, Nicolas ME (1998) The effect of duration of heat stress during grain filling on two wheat varieties differing in heat

tolerance: grain growth and fractional protein accumulation. Funct Plant Biol 25:13–20

- Taub DR, Wang X (2008) Why are nitrogen concentrations in plant tissues lower under elevated CO2? A critical examination of the hypotheses. J Integr Plant Biol 50:1365–1374. https://doi.org/10. 1111/j.1744-7909.2008.00754.x
- Taub DR, Miller B, Allen H (2008) Effects of elevated CO2 on the protein concentration of food crops: a meta-analysis. Glob Change Biol 14:565–575. https://doi.org/10.1111/j.1365-2486.2007. 01511.x
- Tausz M, Norton RM, Tausz-Posch S, Löw M, Seneweera S, O'Leary G, Armstrong R, Fitzgerald GJ (2017) Can additional N fertiliser ameliorate the elevated CO2-induced depression in grain and tissue N concentrations of wheat on a high soil N background? J Agron Crop Sci 203:574–583. https://doi.org/10.1111/jac.12209
- Tausz-Posch S, Dempsey RW, Seneweera S, Norton RM, Fitzgerald G, Tausz M (2015) Does a freely tillering wheat cultivar benefit more from elevated CO2 than a restricted tillering cultivar in a water-limited environment? Eur J Agron 64:21–28. https://doi.org/10.1016/j.eja.2014.12.009
- Tcherkez G, Ben Mariem S, Larraya L, García-Mina JM, Zamarreño AM, Paradela A, Cui J, Badeck F-W, Meza D, Rizza F, Bunce J, Han X, Tausz-Posch S, Cattivelli L, Fangmeier A, Aranjuelo I (2020) Elevated CO2 has concurrent effects on leaf and grain metabolism but minimal effects on yield in wheat. J Exp Bot 71:5990–6003. https://doi.org/10.1093/jxb/eraa330
- Von Caemmerer S, Evans JR (2015) Temperature responses of mesophyll conductance differ greatly between species. Plant Cell Environ 38:629–637. https://doi.org/10.1111/pce.12449
- Wahid A, Gelani S, Ashraf M, Foolad MR (2007) Heat tolerance in plants: an overview. Environ Exp Bot 61:199–223. https://doi. org/10.1016/j.envexpbot.2007.05.011
- Wang X, Liu F (2021) Effects of elevated CO2 and heat on wheat grain quality. Plants 10:1027. https://doi.org/10.3390/plants10051027
- Wang D, Heckathorn SA, Barua D, Joshi P, Hamilton EW, LaCroix JJ (2008) Effects of elevated CO2 on the tolerance of photosynthesis to acute heat stress in C3, C4, and CAM species. Am J Bot 95:165–176. https://doi.org/10.3732/ajb.95.2.165
- Wang D, Heckathorn SA, Wang X, Philpott SM (2011) A meta-analysis of plant physiological and growth responses to temperature and elevated CO2. Oecologia 169:1–13. https://doi.org/10.1007/ s00442-011-2172-0
- Wang X, Liu F, Jiang D (2017) Priming: a promising strategy for crop production in response to future climate. J Integr Agric 16:2709– 2716. https://doi.org/10.1016/S2095-3119(17)61786-6
- Wardlaw IF, Moncur L (1995) The response of wheat to high temperature following anthesis. I. The rate and duration of kernel filling. Funct Plant Biol 22:391–397. https://doi.org/10.1071/pp9950391

- Wardlaw IF, Blumenthal C, Larroque O, Wrigley CW (2002) Contrasting effects of chronic heat stress and heat shock on kernel weight and flour quality in wheat. Funct Plant Biol 29:25–34
- Way DA, Yamori W (2014) Thermal acclimation of photosynthesis: on the importance of adjusting our definitions and accounting for thermal acclimation of respiration. Photosynth Res 119:89–100. https://doi.org/10.1007/s11120-013-9873-7
- Welch JR, Vincent JR, Auffhammer M, Moya PF, Dobermann A, Dawe D (2010) Rice yields in tropical/subtropical Asia exhibit large but opposing sensitivities to minimum and maximum temperatures. PNAS 107:14562–14567. https://doi.org/10.1073/pnas.10012 22107
- Yamori W, Noguchi K, Hikosaka K, Terashima I (2010) Phenotypic plasticity in photosynthetic temperature acclimation among crop species with different cold tolerances. Plant Physiol 152:388– 399. https://doi.org/10.1104/pp.109.145862
- Yamori W, Hikosaka K, Way DA (2014) Temperature response of photosynthesis in C3, C4, and CAM plants: temperature acclimation and temperature adaptation. Photosynth Res 119:101–117. https://doi.org/10.1007/s11120-013-9874-6
- Yin X, Struik PC (2009) C3 and C4 photosynthesis models: an overview from the perspective of crop modelling. NJAS 57:27–38. https://doi.org/10.1016/j.njas.2009.07.001
- Zadoks JC, Chang TT, Konzak CF (1974) A decimal code for the growth stages of cereals. Weed Res 14:415–421. https://doi.org/ 10.1111/j.1365-3180.1974.tb01084.x
- Zhang H, Turner NC, Simpson N, Poole ML (2010) Growing-season rainfall, ear number and the water-limited potential yield of wheat in south-western Australia. Crop Pasture Sci 61:296–303. https://doi.org/10.1071/CP09288
- Zhang X, Högy P, Wu X, Schmid I, Wang X, Schulze WX, Jiang D, Fangmeier A (2018) Physiological and proteomic evidence for the interactive effects of post-anthesis heat stress and elevated CO2 on wheat. Proteomics 18:1800262. https://doi.org/10.1002/ pmic.201800262
- Ziska LH, Morris CF, Goins EW (2004) Quantitative and qualitative evaluation of selected wheat varieties released since 1903 to increasing atmospheric carbon dioxide: can yield sensitivity to carbon dioxide be a factor in wheat performance? Glob Change Biol 10:1810–1819. https://doi.org/10.1111/j.1365-2486.2004. 00840.x

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