- 1 Effect of leaf temperature on estimating physiological traits of wheat leaves from
- 2 hyperspectral reflectance
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15 Abstract

A growing number of leaf traits can be predicted from hyperspectral reflectance data. These 16 include structural and compositional traits, such as leaf mass per area, nitrogen and 17 chlorophyll content, but also physiological traits such a Rubisco carboxylation activity, 18 electron transport rate and respiration rate. Since physiological traits vary with leaf 19 temperature, how does this impact on predictions made from reflectance measurements? 20 We investigated this with two wheat varieties, by repeatedly measuring each leaf through a 21 sequence of temperatures imposed by varying the air temperature in a growth room. The 22 23 function predicting Rubisco capacity normalised to 25 °C predicted the same value, regardless of leaf temperatures ranging from 20 to 35°C. Leaf temperature affected none of 24 the predicted traits: V_{cmax25}, J, chlorophyll content, LMA, N content per unit leaf area or 25 26 V_{cmax25}/N . However, as others have derived models to predict Rubisco activity that includes 27 variation associated with leaf temperature, we discuss whether these functions may include a temperature signal within the reflectance spectra. 28

Keywords: Leaf temperature, hyperspectral reflectance, Rubisco carboxylation activity,
 electron transport rate, leaf dry mass per area, chlorophyll content, leaf nitrogen, *Triticum aestivum*

32 Introduction

Plant breeders continually strive to improve crop yield. For cereals, there has been a recognition that future increases could benefit from improving photosynthesis (Parry *et al.*, 2011; Reynolds *et al.*, 2009). Crop growth is not simply related to a measurement of photosynthetic rate of a particular leaf under one condition. Instead, photosynthesis

integrated over a day with contributions from all the leaves in the canopy drives crop 37 growth. Subsequent conversion into biomass and the partitioning into harvested grains 38 39 determines yield. All these processes combined pose a major challenge on how to 40 meaningfully measure photosynthesis with the goal of improving yield. However, there are 41 a few examples that have compared historical sequences of cultivars and observed 42 correlations between leaf photosynthetic rate and wheat yield (Beche et al., 2014; Fischer et al., 1998; Gaju et al., 2016; Yao et al., 2019). It has also been found that radiation use 43 44 efficiency (above ground biomass produced per unit of intercepted photosynthetically active radiation) has been increasing over time with changing wheat varieties in both the UK 45 (Shearman et al., 2005) and Australia (Sadras et al., 2012). Interestingly, both studies found 46 the same rate of increase (0.012 g MJ 1 y 1). 47

48 It is possible to survey photosynthetic properties between wheat genotypes (Driever et al., 2014; Silva-Pérez et al., 2019), but detailed phenotyping is time-consuming which 49 limits the number of genotypes that can be sampled. A promising alternative is to predict 50 photosynthetic traits from leaf reflectance spectra. Serbin et al. (2012) derived models 51 predicting nitrogen concentration, leaf dry mass per unit area, maximum Rubisco 52 53 carboxylase activity (V_{cmax}) and photosynthetic electron transport rate (J) from hyperspectral reflectance measured on leaves of Populus tremuloides and P. deltoides. Leaf temperature 54 55 varied between 20 and 30 °C, depending on the glasshouse regime, which strongly influenced V_{cmax} . A single model was presented that applied to both species, regardless of 56 57 leaf temperature. It was argued that this implied that the derived V_{cmax} was not being predicted indirectly from another trait such as nitrogen (Serbin et al., 2012). The 58 hyperspectral reflectance approach was also successfully used to predict V_{cmax} for Glycine 59 max measured between 26 and 34°C (Ainsworth et al., 2014) and Nicotiana tabacum 60 (Meacham-Hensold et al., 2019) and Zea mays (Yendrek et al., 2017) measured at various 61 62 temperatures in the field.

In order to be able to make useful comparisons of Rubisco capacity between plants 63 which may differ in their leaf temperature during sampling, one needs to know both V_{cmax} 64 and leaf temperature. Alternatively, one could use the temperature responses of the 65 Rubisco enzyme kinetic parameters (Bernacchi et al., 2003; Silva-Perez et al., 2017) to 66 convert gas exchange estimates of V_{cmax} to a common temperature, e.g. 25°C (V_{cmax25}), 67 which are then used to build a model from reflectance data. This has been done for a group 68 69 of 37 broadleaf tree species (Dechant et al., 2017), wheat (Silva-Perez et al., 2018) and 21 tropical tree species from Panama and Brazil (Wu et al., 2019). Heckmann et al. (2017) also 70 presented predictions of V_{cmax} for Brassica, Moricandia and Z. mays from reflectance 71 72 spectra, but measured only at 25°C. While Dechant et al. (2017) normalised their gas exchange to 25° C, the reflectance spectra were collected at prevailing leaf temperatures. It 73 is not known whether the prediction of V_{cmax25} from leaf reflectance is insensitive to the 74 temperature of the leaf during the reflectance measurement. We, therefore, set out to 75 specifically assess whether leaf trait predictions from hyperspectral reflectance varied with 76 77 leaf temperature by repeatedly measuring the same leaf sequentially through a range of 78 temperatures in two wheat cultivars. We hypothesized that leaf temperature would not 79 affect predicted values of leaf traits obtained using leaf hyperspectral reflectance.

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81 Materials and Methods

82 Plant material and growth conditions

Expt 1 Two spring wheat genotypes (*Triticum aestivum* Kukri and Seri) were grown in a naturally lit greenhouse (day/night temperatures set at 25/15 °C) at the Australian National University in Canberra during Sep-Nov 2018. Three seeds were sown in well-drained 3.5-litre pots filled with commercial potting mix, containing basal fertilizer Osmocote (Scotts). Pots were laid out according to randomized block design with six replicates and blocks representing the replications. After emergence, seedlings were thinned down to one plant per pot. Plants were watered daily until the end of the experiment.

Temperature treatment was given in a controlled environmental chamber, with day/night 90 temperatures set at 25/15 °C and irradiance set to 500 μ mol photons m⁻² s⁻¹. All the 91 measurements were made seven days after anthesis. Plants were moved to the chamber 92 one day before the actual measurements so that plants could acclimatize to the chamber's 93 94 environmental conditions. The next day, measurements were made at a chamber temperature of 15, 25, 35 and 15 $^{\circ}$ C, in the described sequence. After achieving the desired 95 chamber temperature, plants were acclimatized at least 1 hr before the measurements 96 97 were made.

Expt 2 Two spring wheat genotypes (Kukri and Seri) and one triticale (Hawkeye) were grown
in a greenhouse with temperature set to 20/15 °C (day/night). Seeds were sown on multiple
days in March 2018 and each genotype was sown separately in shallow tray with raising mix.
After germination, seedlings were transplanted into 5L pots filled commercial potting mix,
containing basal fertilizer Osmocote (Scotts). Five-six weeks after sowing, half of the plants
were transferred to an adjacent greenhouse room set at 32/20 °C, where they grew for onetwo more weeks before gas exchange measurements were made.

105 *Hyperspectral reflectance measurements (Expt 1)*

Hyperspectral reflectance spectra were measured with a FieldSpec[®]4 (Analytical Spectral 106 107 Devices, Boulder, CO, USA) full range spectroradiometer (350-2500 nm) attached to a leaf 108 clip (Analytical Spectral Devices, Boulder, CO, USA) with a fibre optic cable. Leaf clip had an 109 internal calibrated light source and two external panels i.e. a white panel to calibrate the 110 instrument and a black panel for taking measurements. A mask containing a black circular gasket was also attached to leaf clip, which was used to reduce the leaf-clip aperture to an 111 oval area (1.15 x 1.4 cm = 1.264 cm²) suitable for a wheat leaf (Silva Perez ref). For each 112 113 temperature, one reflectance measurement was made at the same place of the flag leaf of each plant by putting the leaf vertically to the leaf probe as explained elsewhere (Silva-Perez 114 115 et al., 2018).

Leaf reflectance spectra were processed according to Silva-Perez *et al.* (2018). A 'jump' correction associated with a change in the detectors at 1000 and 1800 nm was applied before the traits were predicted.

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119 *MultispeQ measurements (Expt 1)*

Linear electron transport (LET) and relative chlorophyll content (SPAD units) measurements were carried out using a handheld MultispeQ (Beta) device linked to the PhotosynQ platform (<u>www.photosynq.org</u>) (Kuhlgert *et al.*, 2016). Relative chlorophyll content (SPAD units) was estimated by measuring the transmittance of red (650 nm) and infrared (940 nm) light. LET was estimated from the measurements of quantum yield of photosystem II (III) via pulse-amplitude modulation (PAM) fluorometry at photosynthetically active radiation (PAR) of 1000 µmol photons m⁻² s⁻¹ (Kuhlgert *et al.*, 2016).

127 Gas-exchange measurements (Expt 2)

Gas exchange was measured on the most recently fully expanded leaves with a LI-6400XT 128 129 Portable Photosynthesis system (LI-COR Biosciences Inc., Lincoln, NE, USA) on plants placed inside a controlled environment cabinet (Thermoline Science Model-TRIL/SL). The air flow 130 rate was 500 μ mol s¹ with a PPFD of 1800 μ mol m² s¹ supplied by the LED light. Gas 131 exchange was measured at leaf temperatures of 15, 25 and 35° C. At each temperature, CO₂ 132 response curves were measured in $21\% O_2$ using inlet CO_2 concentrations of 400, 50, 100, 133 150, 250, 400, 600, 800, 1000, 400 μ mol mol⁻¹. Subsequently, the air was changed to 2% O₂ 134 with a CO_2 concentration in the leaf chamber of 380 µmol mol⁻¹, the flow reduced to 200 135 μ mol s⁻¹ and measurement continued for 60 minutes with concurrent sampling for carbon 136 isotope discrimination to determine mesophyll conductance (Evans and von Caemmerer, 137 2013). Maximum Rubisco carboxylase activity (V_{cmax}) was calculated from CO₂ response 138 curves using kinetic constants derived from wheat (Silva-Pérez et al., 2017). 139

140 Statistical analysis

141 Data were subjected to analysis of variance using various packages in R (R coreR, 2013).

142 Means were compared for significant differences using Tukey's multiple comparison tests at 143 5% probability level.

144 Results

145 The consequence of using leaf reflectance spectra collected from leaves with varying 146 temperatures to predict leaf traits was investigated with two wheat varieties.

The predicted value of V_{cmax25} was not affected by the leaf temperature when 147 reflectance spectra were measured, for either cultivar (Fig. 1A). Upon returning the growth 148 cabinet to 15°C, the V_{cmax25} values were not significantly different from the initial values. By 149 contrast, V_{cmax} values derived from gas exchange increased fourfold between 15 and 35°C 150 (Fig. 1B). Values for wheat grown under cool or hot conditions showed a difference at 35°C. 151 with the cool grown plants falling further below the theoretical line (consistent with an E_a of 152 63kJ mol⁻¹) than plants from the hot treatment. The values for plants from the hot 153 154 treatment superimpose the previously published data from Silva-Perez et al. (2017).

Predicted values for the rate of electron transport, J, were also independent of the leaf temperature when reflectance spectra were measured (Fig. 2A). In this case, the model was built from data collected under a PPFD of 1800 μ mol m⁻² s⁻¹ and leaf temperatures

158 mainly at 25°C but ranging up to 32°C. However, in contrast to V_{cmax}, J is less sensitive to leaf temperature, increasing by 30% between 15 and 25°C and then plateauing (Fig. 2B). The 159 temperature responses of J were comparable to the previously published data from Silva-160 Perez et al. (2017). Growth temperature shifted the temperature response. For plants 161 162 grown under hot conditions, J was less than that from plants grown under cool conditions at 15°C but greater at 35°C. We also measured J using a MultispeQ instrument following the 163 collection of leaf reflectance spectra at each temperature (denoted LET, Fig. 2A). This was 164 measured under a PPFD of 1000 μ mol m⁻² s⁻¹, similar to the irradiance in the growth cabinet. 165 As it was collected rapidly, it does not represent the steady state. However, it also indicated 166 that the rate of electron transport was similar between 20 and 30°C, then declined slightly 167 168 at 35°C.

169 Predicted values of chlorophyll content were insensitive to the leaf temperature when reflectance spectra were measured (Fig. 3). A similar result was observed for 170 chlorophyll content estimated with the MultispeQ. The absolute values obtained with the 171 MultispeQ were about 20% greater than that predicted from reflectance. The chlorophyll 172 173 content values are predicted from reflectance using a model built on measurements using 174 SPAD-502 chlorophyll meter (Minolta Camera Co., Ltd, Japan) whereas the MultispeQ uses 175 relative transmissions of red (650 nm) and infrared (940 nm) light. Additionally, MultispeQ 176 has two in built differences from Minolta SPAD; 1) MultispeQ takes a series of transmittance measurements over a range of increasing light intensities, and 2) MultispeQ also averages 177 values over a larger leaf area ($\sim 1 \text{ cm}^2$) (Kuhlgert et al. 2016). Nevertheless, additional 178 calibration comparisons were not made as the focus was on temperature. Values predicted 179 for three other leaf traits, leaf dry mass per unit leaf area (LMA), nitrogen content and 180 Rubisco carboxylation capacity normalised to 25 °C per unit leaf nitrogen (V_{cmax25}/N), were 181 also independent of the leaf temperature when reflectance spectra were measured (Fig. 4). 182 A statistical comparison of the effects of temperature treatments on various measured and 183 predicted leaf traits is also provided separately (Supp. Table 1). 184

The spectral response of correlations between single wavelength reflectance values and leaf temperature is shown superimposed on the reflectance spectrum (Fig. 5). Clear correlations were observed on the long wavelength shoulders of the two water absorption bands with peaks at 1531 and 2038nm and a third peak in the red edge at 720nm. By sequentially adding weighted reflectance values from single wavelengths, we were able to predict leaf temperature remarkably well with just four wavelengths (Fig. 6).

191 Discussion

Leaf hyperspectral reflectance is an optical signal that can provide information remotely and rapidly. With appropriate calibration data obtained from other methods, predictive models can be built for a range of leaf traits. The method has potential for use as a high-throughput tool for phenotyping photosynthetic traits at the leaf and canopy scale. While one might expect leaf reflectance to enable prediction of the amount of a substance e.g. leaf dry mass or nitrogen per unit area, it is harder to understand how physiological processes such as rates of reactions could contribute to reflectance. Thus, with Rubisco

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199 being such a major constituent of leaf protein, a reflectance model could arise from a signal 200 associated with leaf protein or nitrogen, as argued by Dechant et al. (2017). If this was the 201 case, then changing leaf temperature would alter Rubisco activity but not Rubisco content. 202 However, Serbin et al. (2012) successfully included variation in leaf temperature to derive a 203 model predicting Rubisco activity, demonstrating that the model was not fundamentally 204 associated with another constituent such as nitrogen. In contrast to gas exchange 205 measurements, where leaf temperature is measured directly to enable the calculation of 206 stomatal conductance, leaf temperature is not generally measured directly during the collection of hyperspectral reflectance. Consequently, it is necessary to consider how leaf 207 208 temperature affects the estimation of physiological traits, such as V_{cmax} and J, using leaf 209 hyperspectral reflectance.

210 Parameters independent of temperature

211 A leaf structural property that has been widely reported is leaf dry mass per unit leaf 212 area as it is easy to measure and relates to lifespan and other traits (Wright et al., 2004). 213 Robust predictions of LMA can be made from hyperspectral reflectance data (Ecarnot et al., 214 2013; Serbin et al., 2012; Silva-Perez et al., 2018). As LMA is a leaf property that would not 215 change in response to short term changes in temperature, models predicting LMA from 216 hyperspectral reflectance should also be insensitive to the temperature of the leaf during measurement. This was found to be true for LMA (Fig. 4A) as well as for other leaf 217 218 constituents, chlorophyll (Fig. 3) and nitrogen (Fig. 4B).

219 Photosynthesis is a process involving many constituents, but has been successfully modelled in C_3 plants by considering the properties of Rubisco (Farguhar *et al.*, 1980). 220 221 Knowing the amount of Rubisco per unit leaf area, its properties and a few assumptions, it is 222 possible to predict photosynthetic responses to irradiance, atmospheric CO_2 and temperature. The amount of Rubisco is unlikely to vary significantly during short term 223 224 changes in leaf temperature, but the carboxylase activity is strongly temperature dependent 225 (Badger and Collatz, 1977; Bernacchi et al., 2001; Sharwood et al., 2016). Therefore, models 226 using reflectance to predict Rubisco content or activity normalised to a fixed temperature 227 should be independent of leaf temperature. Indeed, this was what we observed (Fig. 1A).

228 Parameters that vary with temperature

229 Enzyme activities vary with temperature which can be described with the Arrhenius 230 equation through the energy of activation term E_a. V_{cmax} is the product of Rubisco content 231 and catalytic rate. It seems possible that models predicting V_{cmax} from reflectance 232 (Ainsworth et al., 2014; Serbin et al., 2012) may contain two components, one that is 233 independent of temperature (representing Rubisco protein content) and another that varies 234 with leaf temperature. Immediately prior to placing the leaf into the clip for measuring leaf 235 reflectance, Ainsworth et al. (2014) measured leaf temperature with an infrared thermometer, but they only report the relationship between predicted V_{cmax} and leaf 236 237 temperature. We, therefore, looked at our reflectance spectra collected at different leaf temperatures to see if we could predict leaf temperature. Predicted leaf temperature, using 238 239 reflectance of just four wavelengths, clearly correlated with measured leaf temperature,

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with an r^2 of 0.91 (Fig. 6). It is unlikely that this equation has general application as it arose from only two genotypes measured under one environment and it is known that the power and generality of models predicting dark respiration (Coast *et al.*, 2019) and V_{cmax25} (Wu *et al.*, 2019) improved as the diversity of calibration data increased. However, the point is that leaf temperature apparently can be extracted from reflectance spectra which could explain how models can predict reaction rates from reflectance.

246 The Arrhenius equation predicts an exponential increase in rate with increasing temperature, whereas models calculating parameters from reflectance sum linear 247 248 weightings of each reflectance at each wavelength and would, therefore, have linear 249 responses to temperature. The difference between an exponential and a linear relationship 250 may not be very noticeable over a narrow temperature range. In the case of V_{cmax} , values 251 derived from gas exchange deviated below the Arrhenius function at 35°C (Fig. 1B), such 252 that a linear function would fit the data well between 15 and 35°C. There are also 253 indications that a single value for E_a may not be appropriate across the temperature range 254 from 10 to 40°C. Sharwood et al. (2016) found it necessary to use lower values for E_a at leaf 255 temperatures above 25°C. The rate of electron transport, J, also varies with temperature 256 (Bernacchi et al., 2003; June et al., 2004; Medlyn et al., 2002) but reaches a maximum 257 around 30°C before decreasing again. As a result, the change in J between 20 and 35°C is less pronounced than for V_{cmax}. It is therefore uncertain whether models predicting J from 258 259 reflectance (Dechant et al., 2017; Serbin et al., 2012; Silva-Perez et al., 2018) would contain 260 components that vary with temperature. In the case of J for wheat (Silva-Perez et al., 2018), the reflectance model was built with data that varied little in leaf temperature and the 261 262 predicted values of J were found to be unaffected by leaf temperature when reflectance 263 was measured (Fig. 2A). However, given that J did not vary greatly over this temperature 264 range (Fig. 2), this may not be a very rigorous test. By contrast, as Serbin et al. (2012) 265 deliberately used variation in leaf temperature to generate a broader spread in J to build 266 their reflectance model – testing their function with multiple spectra obtained from a leaf 267 measured over a range of temperatures could be informative.

268 Conclusion

269 Leaf temperature varying between 20 to 35°C during the measurement of leaf 270 reflectance did not affect predicted values of leaf traits (V_{cmax25} , chlorophyll and nitrogen 271 contents per unit area, LMA and V_{cmax}/N), for wheat. It was possible to extract leaf temperature from reflectance spectra which may explain how models that predict rates that 272 273 vary with temperature (V_{cmax}, J, dark respiration) could arise. Models predicting traits that 274 vary with leaf temperature should be tested using multiple measurements from each leaf covering a range of temperature. Reflectance appears to have the potential to predict leaf 275 276 temperature, but to construct a robust model would require calibration with a broader set 277 of experiments.

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284 Supplementary material

285 Table S1. Effects of leaf temperature on leaf physiological traits

Supplementary. Table 1. Effects of leaf temperature on leaf physiological traits measured using MulitispeQ (leaf temperature, LET, relative chlorophyll) or predicted from leaf hyperspectral reflectance (V_{cmax25} , J, chlorophyll content, LMA, nitrogen content, V_{cmax25}/N) in leaves of two wheat genotypes exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C, in a growth chamber (Expt 1). Symbols represent the mean ± SE of six different leaves from six different plants. Sequential measurements were made on each leaf, seven days after anthesis.

	Ambient Air Temperature	Measured traits using MultispeQ			Predicted traits using leaf hyperspectral reflectance					
Genotype		Leaf Temperature (°C)	LET (μmol e ⁻ m ⁻² s ⁻¹)	Relative Chlorophyll (SPAD units)	V _{cmax25} (μmol CO ₂ s ⁻¹ g ⁻¹ (N))	J (μmol e ⁻ m ⁻² s ⁻¹)	Chlorophyll content (SPAD units)	LMA (g m ⁻²)	Nitrogen content (g m ⁻²)	V_{cmax25}/N (µmol CO ₂ s ⁻¹ g ⁻¹ (N))
Kukri	15 °C	19.6 + 0.4	142.9 ± 7.0	59.8 ± 0.7	162.5ª ± 1.7	196.2 ^{ªb} ± 3.6	51.8 ^{ab} ± 0.6	54.0 ± 1.2	2.81 ± 0.07	59.6 ^b ± 1.2
	25 °C	26.2 + 0.2	145.2 ± 9.2	59.9 ± 1.0	154.2 ^b ± 1.8	184.9 ^b ± 3.7	52.6ª ± 0.4	57.4 ± 0.8	2.90 ± 0.05	59.7 ^b ± 0.9
	35 °C	33.4 + 0.3	127.0 ± 4.3	60.6 ± 0.6	161.3 ^ª ± 2.2	196.3 ^{ªb} ± 4.5	50.8 ^b ± 0.4	56.3 ± 1.1	2.82 ± 0.06	64.1 ^ª ± 0.6
	15 °C	19.5 + 0.3	145.4 ± 8.1	59.2 ± 1.0	165.2° ± 2.6	203.8 ^ª ± 5.0	52.7° ± 0.3	55.7 ± 1.1	2.89 ± 0.05	$60.6^{b} \pm 1.0^{c}$
	LSD (5%)		n.s.	n.s.	6.2**	12.5*	1.3*	n.s.	n.s.	2.8*
Seri	15 °C	19.7 + 0.3	144.2ª ± 5.2	62.7 ± 0.8	173.9 ± 2.4	209.2 ± 4.0	53.7 ± 0.4	56.9 ^b ± 0.7	3.05 ± 0.05	60.1 ^b ± 0.8
	25 °C	27.3 + 0.1	148.6ª ± 6.1	63.3 ± 0.8	169.5 ± 2.2	203.5 ± 3.7	53.8 ± 0.2	$58.5^{ab} \pm 0.6$	3.08 ± 0.05	61.0 ^b ± 0.7
	35 °C	34.9 + 0.4	124.9 ^b ± 3.2	64.5 ± 0.8	169.4 ± 2.2	203.7 ± 4.3	53.5 ± 0.4	$60.4^{a} \pm 0.7$	3.05 ± 0.05	61.0 ^b ± 0.7 65.7 ^ª ± 0.7
	15 °C	19.6 + 0.2	145.1°±6.5	62.8 ± 0.7	168.4 ± 2.2	208.5 ± 3.7	53.0 ± 0.3	57.9 ^b ± 0.9	2.96 ± 0.04	$61.5^{b} \pm 0.5$
	LSD (5%)		15.9*	n.s.	n.s.	n.s.	n.s.	2.1*	n.s.	2.04***

*** = P <0.001, ** = P < 0.01, * = P < 0.05, n.s. = non-significant



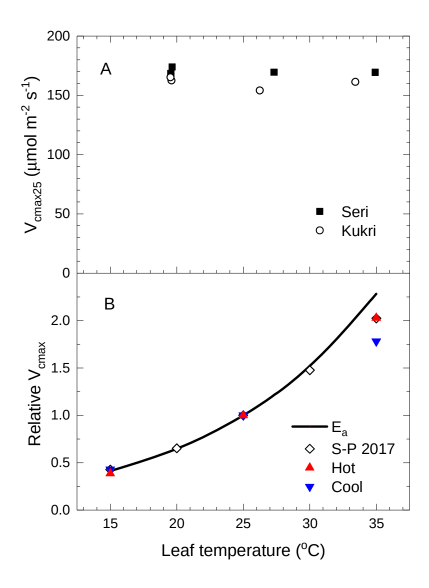


Fig. 1. Effects of leaf temperature on Rubisco activity in wheat. A. The maximum rate of carboxylation by Rubisco normalised to 25 °C (V_{cmax25}) predicted from leaf hyperspectral reflectance measurements at different leaf temperatures. Seven days after anthesis, two wheat genotypes were exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C, in a growth chamber (Expt 1). Symbols represent the mean ± SE of six different leaves from six different plants. B. Temperature response of V_{cmax} normalised to 1 at 25 °C derived from gas exchange measurements (symbols) or modelled ($E_a 63$ kJ mol⁻¹). Data from plants grown under hot (20/15 °C) or cool (32/20 °C) conditions (Expt 2), or from Silva-Perez et al. (2017).

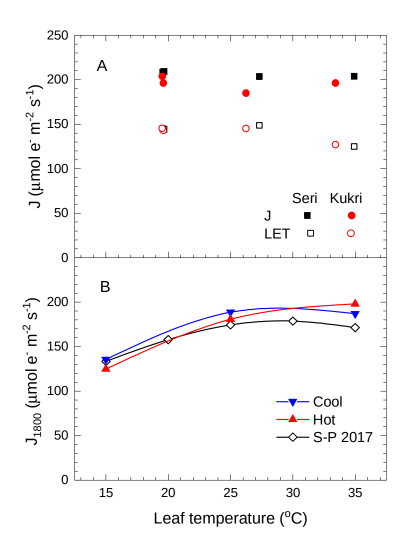


Fig. 2. Effects of leaf temperature on electron transport rate in wheat. A. Rate of electron transport (J) predicted from leaf reflectance under a PPFD of 1800 μ mol m⁻² s⁻¹ (solid symbols) and linear electron transport (LET) measured using MultispeQ under a PPFD of 1000 μ mol m⁻² s⁻¹ (open symbols) at each leaf temperature. Seven days after anthesis, two wheat genotypes were exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C, in a growth chamber (Expt 1). Symbols represent the mean ± SE of six different leaves from six different plants. . B. Rates of electron transport calculated from gas exchange measurements made under a PPFD of 1800 μ mol m⁻² s⁻¹ from plants grown under cool (20/15 °C) or hot (32/20 °C) conditions (Expt 2), or from Silva Perez et al. (2017).

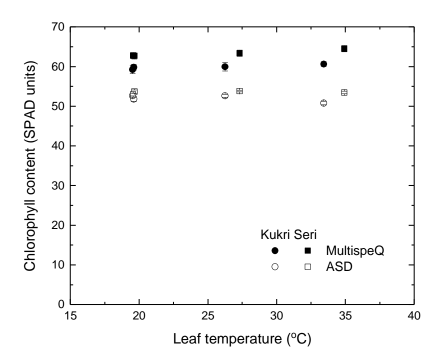


Fig. 3. Effects of leaf temperature on estimated chlorophyll content (SPAD units) in wheat. Seven days after anthesis, two wheat genotypes were exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C, in a growth chamber (Expt 1). Symbols represent the mean \pm SE of six different leaves from six different plants. Leaf chlorophyll content was estimated using two different devices i.e. direct measurements with MultispeQ or predicted from leaf hyperspectral reflectance measurements made with an ASD FieldSpec Spectroradiometer.

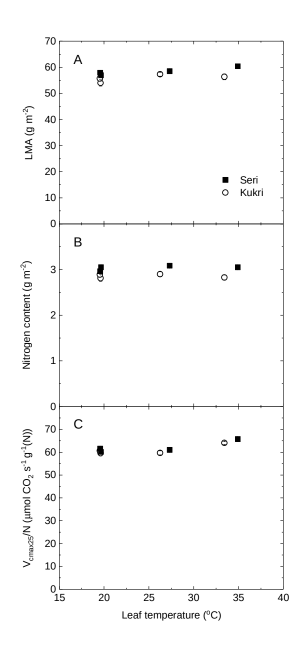


Fig. 4. Effects of leaf temperature on parameters predicted from leaf hyperspectral reflectance in wheat. A. Leaf dry mass per unit leaf area (LMA). B. Nitrogen content per unit leaf area. C. Rubisco carboxylation capacity normalised to 25 °C per unit leaf nitrogen (V_{cmax25} /N). Seven days after anthesis, two wheat genotypes were exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C, in a growth chamber (Expt 1). Symbols represent the mean ± SE of six different leaves from six different plants.

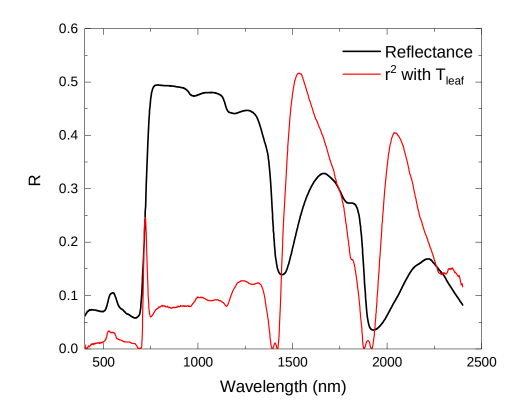


Fig. 5. Reflectance spectrum from a wheat leaf measured at 27°C (black thick line) with the spectral response of the correlation coefficient between leaf temperature and leaf reflectance (red thin line) superimposed.

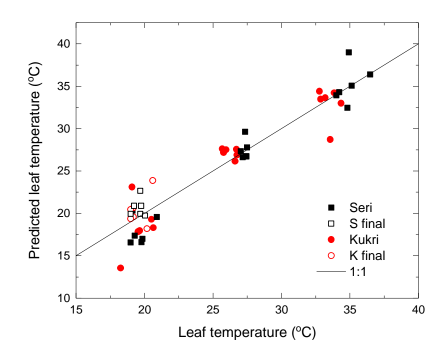


Fig. 6. Relationship between leaf temperature predicted from reflectance and measured using MultispeQ in wheat. Two wheat genotypes were exposed to a sequence of ambient air temperatures i.e. 15, 25, 35 and 15 °C which resulted in the leaf temperature increasing from 20 to 35°C (solid symbols) then decreased back to 20°C (open symbols) (Expt 1). Measurements were made on six different leaves from six different plants. The predicted leaf temperature was calculated as: $T_{leaf} = 1071.4 R_{1531} - 945.65 R_{1400} - 176.3 R_{1507} + 196.8 R_{697}$. The fitted regression equation (not shown) was y = 1.015x - 0.4, r² = 0.91.

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