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High C3 photosynthetic capacity and high intrinsic water use efficiency underlies the high productivity of the bioenergy grass *Arundo donax*

Richard J. Webster¹, Steven M. Driever², Johannes Kromdijk⁴, Justin McGrath⁴, Andrew D. B. Leakey^{3,4}, Katharina Siebke⁵, Tanvir Demetriades-Shah⁶, Steve Bonnage⁷, Tony Peloe⁸, Tracy Lawson⁹ & Stephen P. Long^{3,4}

Arundo donax has attracted interest as a potential bioenergy crop due to a high apparent productivity. It uses C3 photosynthesis yet appears competitive with C4 grass biomass feedstock's and grows in warm conditions where C4 species might be expected to be that productive. Despite this there has been no systematic study of leaf photosynthetic properties. This study determines photosynthetic and photorespiratory parameters for leaves in a natural stand of *A. donax* growing in southern Portugal. We hypothesise that *A. donax* has a high photosynthetic potential in high and low light, stomatal limitation to be small and intrinsic water use efficiency unusually low. High photosynthetic rates in *A. donax* resulted from a high capacity for both maximum Rubisco ($V_{c,max}$ 117 µmol CO₂ m⁻² s⁻¹) and ribulose-1:5-bisphosphate limited carboxylation rate (J_{max} 213 µmol CO₂ m⁻² s⁻¹) under light-saturated conditions. Maximum quantum yield for light-limited CO₂ assimilation was also high relative to other C3 species. Photorespiratory losses were similar to other C3 species under the conditions of measurement (25%), while stomatal limitation was high (0.25) resulting in a high intrinsic water use efficiency. Overall the photosynthetic capacity of *A. donax* is high compared to other C3 species, and comparable to C4 bioenergy grasses.

Giant reed (*Arundo donax* L.) has attracted interest as a potential bioenergy crop, due to a high apparent productivity and suitability as an accepted feedstock for cellulosic fuel production¹. It is found throughout the Mediterranean climate zones of the world and has been cultivated in Asia, southern Europe and North Africa for over a thousand years. It is a rhizomatous perennial herbaceous grass that produces stems of ca. 2–3 cm diameter that may grow to heights of 3–6 m. It is found in moist grasslands and wetlands although it is able to thrive across a wide range of soil types, while also being tolerant of mild drought and salinity, and is tolerant of water-logged soils^{2–4}. In climates with warm winters it is evergreen but dies back in climates with freezing winter temperatures. Although there are related species within the genus, molecular analysis suggests that *A. donax* is monophyletic in origin and its spread has not involved hybridisation with the related species. These analyses also suggest that it originated in west Asia and then spread to the Mediterranean region⁵. The high productivity of *A. donax* is achieved in warm growing conditions despite the fact that it uses C3 photosynthesis, which might be expected to be a competitive disadvantage compared to C4 species under such growing conditions.

¹Institute of Biological Environmental and Rural Sciences, Aberystwyth University, Aberystwyth, U.K. ²Centre for Crop Systems Analysis, Wageningen University, Wageningen, the Netherlands. ³Department of Plant Biology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA. ⁴Carl R. Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA. ⁵Heinz Walz GmbH, Effeltrich, Germany. ⁶LI-COR Biosciences UK Ltd., Cambridge, UK. ⁷ADC BioScientific Ltd., Hoddesdon, UK. ⁸Delta-T Devices Ltd., Cambridge, UK. ⁹School of Biological Sciences, University of Essex, Wivenhoe Park, Colchester, U.K. Correspondence and requests for materials should be addressed to R.J.W. (email: rcw@aber.ac.uk)

Side-by-side trials in central Italy over 11 years showed an average yield of above-ground dry biomass of 28.7 t ha⁻¹ yr⁻¹ for the highly productive C4 perennial grass *Miscanthus x giganteus* Deu et Greef, but 37.7 t ha⁻¹ yr⁻¹ for A. donax⁶. Although this appears a relatively low yield for M. x giganteus under irrigated conditions in the Mediterranean⁷, the reported yield for A. donax is high by any measure. A. donax in central and southern Italy showed average dry matter yield over 3 years of up to 74 t ha⁻¹ yr⁻¹, which exceeds the highest yields reported for *M. x giganteus*⁸. Similarly, in a multi-year study in Alabama *A. donax* achieved an average yield of $35.5 \text{ t ha}^{-1} \text{ yr}^{-1}$ compared to 23.5 t ha⁻¹ yr⁻¹ for switchgrass (Panicum virgatum L.)⁹. In a comparison with C4 sweet sorghum (Sorghum bicolor L.) in northern Italy, light penetration into an A. donax canopy studied over a three year period was found to be 60% higher and the efficiency of conversion of intercepted radiation into biomass was ca. 60% higher. This might be explained by its more effective distribution of light through the canopy, if it is coupled with a similar efficiency of leaf photosynthesis^{10,11}. However, yields may be considerably lower on marginal land. On a sandy loam with limited nutrient availability A. donax required three years to accumulate 20 t ha⁻¹, and build a rhizome mass of 16 t ha⁻¹⁴, by contrast to *M. x giganteus* which has proved significantly more productive on marginal sites¹². While M. x giganteus was found particularly vulnerable to damage by water-logging of soils during autumn and winter, A. donax appeared to thrive in these situations. This tolerance of water-logging also allows A. donax to spread along watercourses from rhizome fragments where M. x giganteus cannot¹³. Its ability to survive anaerobic soil may explain its ability to tolerate high levels of reduced ions of heavy metals¹⁴. A. donax is clearly a resilient, but also very productive species. However, there is little fundamental knowledge available on how it is able to realise this productivity. Understanding this demands knowledge of leaf photosynthesis as well as canopy microclimate.

Leaf photosynthetic rates of CO₂ uptake (A) measured in A. donax growing on an estuary in S. Africa were between 20 and 37 μ mol m⁻² s^{-1 15}. By combining modulated chlorophyll fluorescence and gas exchange, photorespiration was estimated to decrease A by 30%. Although water potential declined to -2.1 MPa around noon on sunny days, stomatal conductance (g_s) and A changed little. Such low leaf water potential would normally be expected to at least cause partial stomatal closure in most species¹⁵. The rates of stomatal conductance reported are at the upper end of those found in other C3 species but not much greater than those of the productive C4 bioenergy grasses M. x giganteus and P. virgatum, as well as productive modern cultivars of maize (Zea mays L.)^{16,17}.

In summary, *A. donax* achieves and exceeds the productivities normally associated with C4 perennial grasses in warm climates. For a plant which forms a high leaf area index (LAI) monocultures in nature and as a crop, it would be expected to have a high photosynthetic capacity at both high and low light intensity under warm conditions. The high photosynthetic capacity is needed in the context of a dense canopy where shaded leaves need to achieve high efficiency, as well as those in full sun, and where all leaves need to be efficient under the low light conditions of dawn, dusk and cloudy days. Yet it is a C3 species, growing in warm conditions where only a C4 species might be expected to be photosynthetically efficient¹⁸. The limited published data suggests that *A. donax* may indeed have high light-saturated leaf photosynthetic rates, which may also be supported by high leaf conductances. High leaf conductances will allow a higher CO₂ concentration at ribulose-1:5-bisphosphate carboxylase/ oxygenase (Rubisco), so increasing the rate of carboxylation and decreasing photorespiration, but at the expense of water loss and water use efficiency¹⁹.

To better define the photosynthetic capacity of *A. donax* and understand how this may explain its high productivity this study aimed to define key *in vivo* properties. These were: J_{max} , the maximum whole chain electron transport rate supporting ribulose-1:5-bisphosphate (RuBP) regeneration, $V_{c,max}$ the maximum rate of carboxylation that can be supported by Rubisco, the light-saturated rate of CO₂ assimilation (A_{sat}) and the maximum quantum yield of CO₂ assimilation ($\Phi_{CO2,max}$) defined by the initial slope of the response of *A* to absorbed photon flux (αI). Concurrent measurement of water vapour flux and modulated chlorophyll fluorescence were used to determine g_s , stomatal limitation to CO₂ uptake (I), intercellular CO₂ concentration (c_i), and the whole chain electron transport rate (J). These parameters were determined for leaves in a natural stand of *A. donax* growing in a dry valley in S. Portugal. These are used to test the hypotheses that *A. donax* has a high photosynthetic potential in both high and low light, that stomatal limitation may be unusually small and thus intrinsic leaf water use efficiency poor. The *in vivo* measures: J_{max} , $V_{c,max}$, A_{sat} and $\Phi_{CO2,max}$, are key to parameterization of the steady-state biochemical model of C3 photosynthesis²⁰ which underlies most models of crop and ecosystem productivity²¹. This study therefore also serves as a key resource for parameterizing production models of this emerging crop.

Materials and Methods

Field location. The study was undertaken on a naturally occurring *A. donax* stand located at Quinta de São Pedro - Centro de estudos (Armadas, 2815–786 Sobreda, Portugal 38°38′40.6″N 9°11′34.5″W, altitude 85 m). A mixed 5 acre site with a disturbed dry valley community of annuals, patches of native macchia and scrub, pine woodland, introduced Acacia woodland and a few planted ornamentals on a Cambisol soils²².

Meteorological assessment. Relative humidity, air temperature and irradiance were measured over the sampling period using a Weather Station (WS-GP2 with GP2 Data Logger, Delta-T Devices Ltd, Cambridge, UK) (Fig. S1). Relative Humidity and air temperature were used to calculate vapour pressure deficit (VPD), as a virtual channel within the GP2. Soil temperature and soil water content were measured within the stand over the sampling period using soil moisture sensors (SM300 with GP2 Data Logger, Delta-T Devices Ltd, Cambridge, UK).

Stomatal Conductance and *In situ* **Chlorophyll Fluorescence.** Measurements of stomatal conductance were made with a diffusion porometer (AP4, Delta-T Devices Ltd, Cambridge, UK). All measurements of conductance and photosynthesis were made on the most recently expanded leaf, as judged by ligule emergences. Readings were taken on several leaves, locating the porometer at three points along each leaf: 1) the base just above the ligule, 2) the midpoint between ligule to tip; and 3) the area just below the leaf tip. Both the adaxial and abaxial surfaces were measured at these points at midday and mid-afternoon following the procedures of Monteith²³, and Potter²⁴. Modulated chlorophyll fluorescence was monitored to estimate the operating efficiency of PSII $(F_q'/F_m')^{25,26}$ on fully emerged leaves at midday and late afternoon using a fluorometer (MONITORING-PAM, Heinz Walz GmbH, Effeltrich, Germany) (Fig. S1).

Photosynthetic intercellular-CO₂ **response curves.** Leaves were sampled pre-dawn from the A. donax stand. Leaves were excised just above the ligule, and immediately re-cut under water and then kept in cool low-light conditions (20 °C and 20 μ mol m⁻² s⁻²) until use. The response of net leaf CO₂ uptake rate (A) to external CO_2 concentration (c_a) was assessed on five different leaves. Leaves were placed in the leaf cuvette of a portable open gas exchange system (LCPro-SD with 6.25 cm² cuvette; ADC BioScientific Ltd. Herts, England; or LI6400XT with 6 cm² cuvette; LI-COR Biosciences, Lincoln, Nebraska, USA). Air temperature was controlled at 25 °C, leaf temperature 27 °C and VPD at 0.96 kPa. To fully induce photosynthesis before measurements commenced, leaves were first allowed to equilibrate at a photon flux (I) of $500 \,\mu\text{mol}\,\text{m}^{-2}\,\text{s}^{-1}$ and reference c_a of $400 \,\mu\text{mol}\,\text{mol}^{-1}$ until A had reached a stable value, the I was then increase to 1500 μ mol m⁻² s⁻¹. When A had reached a stable value at $I = 1500 \,\mu$ mol m⁻² s⁻¹ c_a was then changed to the following levels in sequence, 300, 250, 200, 150, 100, 50, 400, 600, 900, 1200 and 1500 μ mol mol⁻¹. The leaf remained at each level until a stable A could be determined. Potential Rubisco carboxylation ($V_{c,max}$) and electron transport through photosystem II (J_{max}) were determined from the responses of A_{sat} to c_i . For each leaf values of A located above the transition between Rubisco-limited and RuBP- or electron transport-limited photosynthesis in the A/c_i response were used to solve for J_{max} using the equations representing RuBP regeneration-limited A²⁰. Values were temperature adjusted to 25 °C using the temperature response equations of Bernacchi et al. (2001)²⁷ for the Rubisco-limited, and Bernacchi et al. (2003)²⁸ for the RuBP regeneration limited portions of the A vs c_i curves. Stomatal limitation (l) at the current ambient c_a of 400 µmol mol⁻¹ was derived using the method outlined in Long and Bernacchi (2003)²⁹.

Photosynthetic light response curves. Sampling and measurements of *A* vs. *I* were performed in parallel with *A* vs. c_i curves. Leaves were placed in the leaf cuvette of an integrated open gas exchange system and modulated chlorophyll fluorometer (GFS-3000FL, LED-Array/PAM-Fluorometer 3055-FL and 3080-O2 Oxygen sensor. Heinz Walz GmbH, Effeltrich, Germany). Leaves were adapted to an incident photon flux of 1500 µmol m⁻² s⁻¹ prior to measurements with the [CO₂] at 400 µmol mol⁻¹ and leaf temperature controlled at 25 °C. The VPD of the air entering the gas exchange system was maintained between 0.8 and 1.1 kPa. Photosynthetic light response curves were then obtained at ambient (21%) and low (2%) oxygen concentrations. Photon flux was varied in a step-wise manner, either starting from the photon flux at which photosynthesis had been induced, after a steady-state *A* was obtained or alternating between different photon fluxes, in each case waiting for a steady-state to be obtained. Photon flux levels were 2000, 1500, 1000, 700, 500, 350, 250, 150, 75, 45 µmol m⁻² s⁻¹. At each light level, once a new steady state was reached gas exchange rates were recorded. Simultaneously, F_q'/F_m' was determined via modulated chlorophyll fluorescence and application of a saturating flash as previously described^{25,26,30}.

The light response of CO_2 assimilation rate is described by a four-parameter non-rectangular hyperbola, according to Marshall and Biscoe³¹:

$$A = \frac{\Phi_{\max} I + A_{\operatorname{sat}} - \sqrt{\left[\left(\Phi_{\max} I + A_{\operatorname{sat}}\right)^2 - 4\theta \Phi_{\max} I A_{\operatorname{sat}}\right]}}{2\theta} - R_d \tag{1}$$

where A is the CO₂ assimilation rate (μ mol m⁻² s⁻¹), Φ_{max} is the apparent maximum quantum yield (mol CO₂ (mol photons)⁻¹), *I* is the incident photon flux (μ mol m⁻² s⁻¹), *A*_{sat} is the CO₂ assimilation rate at saturating photon flux (μ mol m⁻² s⁻¹), θ is the curve convexity (dimensionless) and *R*_d is mitochondrial respiration in the light (μ mol m⁻² s⁻¹). The light compensation point (LCP) was determined as the incident photon flux (*I*) where *A* = 0 as predicted from the fitted curve.

The quantum yield of CO₂ assimilation, Φ_{CO2} , was determined according to Genty³⁰:

$$\Phi_{CO_2} = \frac{A + R_d}{\alpha I} \tag{2}$$

where A is corrected for respiratory loss (R_d) and divided by the absorbed photon flux (αI), with α as the leaf absorbance. The method used to determine absorbance under the chamber lighting conditions is described later.

The parameters Φ_{CO2} and F_q'/F_m' have been shown to be linearly related under conditions where photorespiration is suppressed^{30,32,33}. Under ambient (21%) and low (2%) oxygen atmosphere, the relationship can be used to assess the possible operation of electron acceptors other than CO₂, e.g. photorespiration or active oxygen production, which are manifest as an increased F_q'/F_m' relative to its corresponding Φ_{CO2}^{34} .

The relation of the quantum yield of CO₂ assimilation (Φ_{CO2}) after Valentini³⁵ to the operating efficiency of PSII (F_q'/F_m') under non-photorespiratory conditions can be described as a linear relationship³⁰, where *b* is the y axis intercept and *k* is the slope of the relationship between F_q'/F_m' and Φ_{CO2} :

$$\frac{F'}{F'm} = k * \Phi_{\rm CO_2} + b \tag{3}$$

Assuming 4 electrons are necessary for the fixation of a CO₂ molecule, this can be expressed as:

$$\frac{F'q}{F'm} = \frac{1}{4}k * \Phi_{e-} + b$$
(4)

where Φ_{e-} is the quantum yield of total electron flow, which can be rewritten as:

$$\Phi_{e-} = \frac{4\left(\frac{F'_q}{F'_m} - b\right)}{k} \tag{5}$$

This relationship is assumed to hold in the presence and in the absence of photorespiration, where in the presence of photorespiration it can be used to calculate the total electron flow, which is the sum of electron flow to the reactions of carboxylation (J_C) and oxygenation (J_O)³⁵:

$$J_T = I * \Phi_{e^-} = J_C + J_O \tag{6}$$

 $J_{\rm C}$ can be calculated by assuming 4 electrons are needed per CO₂ for carboxylation:

$$J_C = 4\left(A + R_d + R_p\right) \tag{7}$$

where R_p is the amount of CO₂ released by photorespiration. J_O can be calculated by assuming that 8 electrons are needed per CO₂ released in photorespiration:

$$O = 8 * R_p \tag{8}$$

Combining the equations for J_T , J_C and J_O then gives:

$$R_{p} = \frac{J_{T} - 4(A + R_{d})}{12}$$

$$J_{C} = \frac{1}{3}[J_{T} + 8(A + R_{d})]$$

$$J_{O} = \frac{2}{3}[J_{T} - 4(A + R_{d})]$$
(9)

Leaf transmission and reflectance. Immediately following completion of the gas exchange measurements leaf absorbance (α) was measured from 400 to 700 nm, with a dual-channel spectrometer and integrating spheres (SpectroClip-JAZ-TR, Ocean Optics, Oxford, UK). Six spectral measurements of leaf transmittance and reflectance were made per leaf. The average transmittance (τ) and reflectance (R) for each leaf was used to determine $\alpha = (1 - R - \tau)$. Total absorbance for the PAR spectrum was calculated. Total absorbed light was also calculated by combining the percentage of actinic light emitted by the blue (470 nm) and red (640 nm) LEDs in the leaf gas exchange chamber with α for the peak wavelength of the two LED types³⁶. The apparent maximum quantum yield (Φ_{max}) was then recalculated to give the maximum absolute quantum yield, i.e. the net number of CO₂ molecules absorbed by the leaf per photon absorbed:

$$\Phi_{CO_2,\max} = \frac{\Phi_{\max}}{\alpha} \tag{10}$$

Intrinsic leaf water use efficiency. Intrinsic leaf water use efficiency (LWUE) was calculated from gas exchange of CO₂ and H₂O as the ratio of CO₂ assimilation (*A*) over stomatal conductance (g_s) under 21% O₂ at photon fluxes of 1000 and 2000 µmol m⁻² s⁻¹. Contrary to the water use efficiency calculated from *A* over transpiration (*E*)³⁷, *A*/ g_s is considered to be more realistic and comparable between studies, as it is not influenced by changes in leaf to air VPD in the leaf chamber³⁸.

Results

The light response of *A. donax* for *A* at high and low O₂ showed an increase of *A* when O₂ was lowered, compared to that at 21% O₂, indicating the reduction of *A* in air due to photorespiration (Fig. 1a). At both O₂ concentrations, *A* did not reach a plateau, even at high photon flux. Both the light response of g_s and c_i were lowered when O₂ concentration was reduced (Fig. 1b). The response of g_s to light largely scaled with *A*, while c_i reached a plateau above a photon flux of 600 µmol m⁻² s⁻¹. From the response of *A* to light, parameters were derived (Table 1). The CO₂ assimilation rate at saturating photon flux (A_{sat}) was remarkably high at both ambient ($30.19 + /-0.32 µmol m^{-2} s^{-1}$) and low O₂ concentration ($34.86 + /-1.2 µmol m^{-2} s^{-1}$) whereas the predicted values of *A* at infinite photon flux density as described by the fitted hyperbola were $38.11 + /-0.56 µmol m^{-2} s^{-1}$ at ambient O₂ and $43.45 + /-2.08 µmol m^{-2} s^{-1}$ at low O₂ concentration. The maximum quantum yield of CO₂ assimilation ($\Phi_{CO2,max}$) at ambient and low O₂ level was found to be 0.056 (+/-0.0029) and 0.072 (+/-0.0045) respectively, mol CO₂ (mol absorbed photons)^{α 1}. The intrinsic leaf water use efficiency (A/g_s) (LWUE) was $62.9 (+/-2.38) µmol mol^{-1}$ at sub-saturating photon flux (1000 µmol m⁻² s⁻¹) and $66.0 (+/-1.97) µmol mol^{-1}$ at near-saturating photon flux (2000 µmol m⁻² s⁻¹).

Under 21% O₂, the electron flow to oxygenation reactions of Rubisco (J_0) and the rate of CO₂ production by photorespiration (R_p) increased with photon flux, until a plateau was reached at a photon flux of 1000 µmol m⁻² s⁻¹



Figure 1. (a) Light response of assimilation rate for fully emerged *A. donax* leaves measured at 21% oxygen (closed symbols) and 2% oxygen (open symbols). (b) c_i (triangle) and g_s (square) at a c_a of 400 µmol mol⁻¹, leaf temperature of 25 °C and VPD of between 0.8 and 1.1 kPa (values are means and standard errors, for 5 different plants).

Photosynthetic light response curves	21% Oxygen	2% Oxygen	
$A_{\rm sat} (\mu { m mol} { m CO}_2 { m m}^{-2} { m s}^{-1})$	38.11 (0.56)	43.45 (2.08)	
$\Phi_{\rm max}$ (mol CO ₂ mol photons ⁻¹)	0.050 (0.003)	0.064 (0.004)	
$\Phi_{\rm CO2,max}$ (mol CO ₂ mol photons ⁻¹)	0.056 (0.003)	0.072 (0.004)	
$R_{\rm d} (\mu { m mol} { m CO}_2 { m m}^{-2} { m s}^{-1})$	1.471 (0.15)	1.581 (0.16)	
θ(-)	0.543 (0.06)	0.601 (0.08)	
LCP (μ mol m ⁻² s ⁻¹)	29.88 (2.14)	24.89 (1.63)	



(Fig. 2.). However, the electron flow to carboxylation ($J_{\rm C}$) increased further with increasing photon flux and did not appear to reach a plateau (Fig. 2). It was estimated that *ca*. 25% of electron flow is accounted for by $J_{\rm O}$ under saturating light intensity. The ratio of CO₂ production by photorespiration, $R_{\rm p}$, to assimilation ($A_{\rm sat}$) was 0.216.

Total leaf absorbance from 400 nm to 700 nm was 0.87 (+/- 4.0×10^{-3} , n = 6). Total leaf absorbance at the peak emitting wavelengths of red (640 nm) and blue (470 nm) was 0.89 (S.E. +/- 4.47×10^{-3} , n = 6) and used to calculate F_q'/F_m' against the quantum yield of CO₂ assimilation (Φ_{CO2}) on an absorbed light basis. The relation between F_q'/F_m' and Φ_{CO2} under high and low O₂ showed significant positive correlations (Fig. 3). The difference







Figure 3. Relationship between operating efficiency of PSII (F_q'/F_m') and quantum yield of CO₂ assimilation (Φ_{CO2}) for *A. donax* calculated from light response of assimilation rate, measured at 21% oxygen (closed circle) and 2% oxygen (open circle) and both at a c_a of 400 µmol mol⁻¹, leaf temperature of 25 °C and VPD of between 0.8 and 1.1 kPa. Values are corrected for leaf absorbance, 0.89 ($\pm 4.47 \times 10^{-3}$), (values are means and standard errors, for 5 different plants).

in slope of the relation, representing the apparent electron demand to assimilate one CO_2 , at ambient and low O_2 concentration can be attributed to the presence or absence of photorespiration.

The response of *A* to c_i for *A*. *donax* leaves at saturating light intensity was determined at ambient O₂ concentration (Fig. 4). Subsequently, parameters were derived to describe the A/c_i response and the stomatal limitation (Table 2). The maximum velocity of carboxylation by Rubisco ($V_{c,max}$), was 117.8 µmol m⁻² s⁻¹ and the maximum rate of electron transport (J_{max}) was 213.7 µmol m⁻² s⁻¹. The CO₂ compensation point, Γ , was 44.2 µmol mol⁻¹ and the limitation to *A* imposed by stomata and leaf boundary layer, *l*, was 0.25 (Table 2).





Photosynthetic intercellular-CO ₂ response	21% Oxygen
$V_{ m c,max}$ (µmol CO ₂ m ⁻² s ⁻¹)	117.8 (8.8)
$V_{ m c,max25}$ (µmol CO ₂ m ⁻² s ⁻¹)	95.0 (6.9)
$J_{\rm max} (\mu { m mol} { m CO}_2 { m m}^{-2} { m s}^{-1})$	213.7 (21.2)
$J_{\rm max25}$ (µmol CO ₂ m ⁻² s ⁻¹)	184.0 (18.1)
Γ (µmol mol ⁻¹)	44.2 (0.95)
l(-)	0.25 (0.05)

Table 2. Parameters derived from photosynthetic intracellular $CO_2(c_i)$ response curves; replication and conditions of measurement as given in Fig. 4. mean, (SE), n = 5.

Time	Leaf Surface	Stem	Middle	Tip	$F_{\rm q}'/F_{\rm m}'$
12:00	Abaxial	399 (61)	538 (72)	355 (25)	
12:00	Adaxial	293 (30)	422 (35)	326 (64)	0.63 (0.03)
16:00	Abaxial	231 (24)	373 (77)	341 (87)	
16:00	Adaxial	110 (10)	210 (18)	160 (28)	0.51 (0.08)

Table 3. Stomatal conductance (mmol m⁻² s⁻¹) at mid and late afternoon, on abaxial and adaxial leaf surfaces with measurements along the leaf surface from the stem to the leaf tip. (Values are means and standard errors for 4 or 5 individual plants). The operating efficiency of PSII (F_q'/F_m') at mid and late afternoon, on the middle adaxial leaf surfaces.

Soil water content between 20 cm and 40 cm soil depth averaged 0.14 m³ m⁻³ (+/- 0.01) across the sampling area over the sampling period, with an average soil temperature of 21 °C (+/- 0.9). Stomatal conductance of *A. donax*, measured on ad- and abaxial surfaces of leaves on three different positions along the leaf blade, were highest at noon (12:00) and declined in the afternoon (16:00, Table 3). The highest stomatal conductance was observed in the middle of the leaf at both time points and both surfaces. The greatest decline in stomatal conductance between noon and afternoon was observed in the middle of the leaf, on both adaxial and abaxial surfaces (Table 3). Even though there was a great decline in stomatal conductance, the operating efficiency of PSII (F_q'/F_m') at noon and in the afternoon in the field did not decrease as much (Table 3). This implies that stomatal conductance was still not greatly limiting for photosynthesis during the afternoon.

Discussion

Light-saturated capacity for photosynthetic CO₂ uptake is determined by the maximum rate of carboxylation $(V_{c,max})$ and the maximum rate of electron transport, (J_{max}) , depending on the inter-cellular CO₂ concentration $(c_i)^{20}$. Averaged across 109 C3 species, $V_{c,max}$ was 64µmol m⁻² s⁻¹ and J_{max} 134µmol m⁻² s⁻¹ ³⁹, this compares to values found here for *A. donax* of 117.8 and 213.7 µmol CO₂ m⁻² s⁻¹, respectively, which are almost double the C3 average. These values are for a natural and unfertilized stand. Typically in C3 crops these values will increase with nitrogen status⁴⁰. However, the values found here for *A. donax* are high even compared to fertilized or N-fixing productive crops: *Phalaris arundinacae* (83.9 and 150.6 µmol m⁻² s⁻¹)⁴¹, *Triticum aestivum* (59 and 139 µmol m⁻² s⁻¹)⁴⁰, *Oryza sativa* (91 and 190 µmol m⁻² s^{-1 42}, *Helianthus annuus* (29 and Heinz walz µmol m⁻² s⁻¹)⁴³, *Glycine max* (83 and 160 µmol m⁻² s⁻¹)⁴⁴, and *Phaseolus vulgaris* (88 and 178 µmol m⁻² s⁻¹)⁴⁵. This highlights the high light-saturated maximum photosynthetic capacity of *A. donax*. Compared to other tall grass species considered as bioenergy sources, *A. donax* shows high *A* in full sunlight. Figure 1 shows an average *A* of 30.2 µmol m⁻² s⁻¹ at 25 °C, this compares to values at this temperature of 28 µmol m⁻² s⁻¹ for *M. x giganteus*⁴⁶, 24 µmol m⁻² s⁻¹ in field grown stands of *Panicum virgatum* (switchgrass)¹⁶, and 22 µmol m⁻² s⁻¹ for the C3 bioenergy grass, *Phalaris arundinacea* (Reed canary grass)⁴⁷.

Is this *A* achieved by lower losses to photorespiration in *A. donax*? If we assume "average" Rubisco kinetic properties as described by Brooks and Farquhar⁴⁸ for spinach, then the CO₂ photosynthetic compensation point (Γ^*) would be 42.5 µmol mol⁻¹ at 25 °C. Using this value, with the measured dark respiratory rate (R_d) of 2 µmol m⁻² s⁻¹, average *A* of 30.19 µmol m⁻² s⁻¹ and c_i of 255 µmol m⁻² s⁻¹ at a photon flux of 2000 µmol m⁻² s⁻¹ (Fig. 1), the photorespiratory rate of CO₂ release (R_p) would be 6.3 µmol m⁻² s⁻¹, following the equations of Farquhar *et al.*²⁰. This calculated rate, based on properties of spinach, is very close to the rate derived here for *A. donax* by combining fluorescence and gas exchange measurements (Fig. 2) via the equations of Valentini *et al.*³⁵. This suggests that as a proportion of net photosynthesis, photorespiratory loss of carbon is just as great in *A. donax* as in other C3 species.

Is the rate of CO₂ assimilation in limiting light high in *A. donax*? Light limited photosynthesis is determined by the efficiency with which the leaf can absorb incident light (α), and the maximum absolute quantum yield of CO₂ assimilation ($\Phi_{CO2,max}$), i.e. the initial slope of the response of *A* to *I*, corrected for absorptance (α). The value is therefore the maximum ratio of net absorbed CO₂ molecules to absorbed photons. The $\Phi_{CO2,max}$ determined for *A. donax* here in normal air at 25 °C was 0.056 mol mol⁻¹. Osborne and Garrett⁴⁹ similarly measured $\Phi_{CO2,max}$ by combining gas exchange and integrating sphere measurements across a range of C3 herbage grasses and cereals, covering different ploidy levels and cultivars in normal air and at 25 °C. They reported an average $\Phi_{CO2,max}$ of 0.051, with a range of 0.047 to 0.055 across these C3 grasses. So this does place the values here for *A. donax* at the upper end of this range and 8% higher than the average. The realized efficiency of light limited photosynthesis will be the product of $\Phi_{CO2,max}$ and the absorptance of the leaf (α). In surveying a wide range of healthy leaves of C3 species from tropical to polar habitats, Long *et al.*⁵⁰, found an average α of 0.80, and a range across species from 0.65 to 0.90, similarly measured in an integrating sphere. By comparison, the α for *A. donax* of 0.89 therefore appears at the upper end of this range and 14% higher above the average. The product of the indicated above average $\Phi_{CO2,max}$ and α therefore represents a 24% increase in CO₂ uptake per unit incident light, under light-limiting conditions.

Does A. donax achieve high photosynthetic rates by minimizing stomatal limitation? The stomatal limitation imposed on photosynthesis, l, for A. donax found in this study (0.25 or 25%) was higher than for other C3 plants, which varied from 0.137 to 0.21751-54. This indicates that A. donax does not achieve its high photosynthetic rates through a high stomatal conductance. Indeed c_i/c_a at full sunlight (2000 μ mol m⁻² s⁻¹) as a measure of the balance between stomatal conductance and assimilation was 0.64. This is almost 11% less than the average c_i/c_a of 0.72 for a range of C3 species¹⁹. A lower c_i/c_a requires a lower stomatal conductance relative to the photosynthetic rate, indicating a higher leaf level water use efficiency for a given leaf-air water vapour pressure deficit than the average for C3 species. So, while stomatal conductance may appear high (Fig. 1b), the low values of l and c_i/c_a in fact show that conductance is low relative to the high rates of leaf CO₂ uptakes for a C3 plant. Nevertheless its high light-saturated and light-limited photosynthetic rates can still only be achieved at the expense of considerable transpiration. The relatively high stomatal conductance maintained into the late afternoon (Table 3) suggests that the plant may be able to tap deep water resources even in its native Mediterranean in the late summer. The LWUE of A. donax in this study (62.9 to $66.0 \,\mu$ mol mol⁻¹) was higher than generally found for herbaceous species (43 µmol mol⁻¹) and more similar to LWUE found for evergreen shrubs and deciduous trees (64 and 66 µmol mol⁻¹, respectively)⁵⁵. However, the LWUE of A. donax was still much lower than for C4 species such as *Miscanthus* and switchgrass (115 and 107 μ mol mol⁻¹)¹⁶. The high productivity of *A. donax* does resemble and exceeds that of C4 plants, however its LWUE is much lower and clearly C3-like. This supports the notion that, as discussed above, A. donax is able to achieve its high photosynthetic rates with substantial transpiration, but is still more efficient than most C3 species. This is interesting when considering the diversity of habitat distribution that A. donax is found in, which ranges from very wet loam to relatively dry sandy soils.

Overall this study has found that the photosynthetic capacity of *A. donax* in full sunlight is high compared to other C3 species, and comparable to C4 bioenergy grasses. This is not the result of lower photorespiratory rates, but rather a high capacity for both RuBP-limited and RuBP-saturated photosynthesis, as evidenced by values of both $V_{c,max}$ and J_{max} that are near double the average for C3 species. High photosynthetic rates were not achieved through a high stomatal conductance, in fact stomatal limitation was found to be greater not less than in other C3 species. Light-limited photosynthesis, which will determine carbon uptake during periods of low light flux, around dawn and dusk, and during heavy cloud, as well as in the lower canopy may be aided by relatively high maximum quantum yields of CO₂ assimilation and high leaf absorptances.

References

- EPA. Supplemental Final Rule for New Qualifying Renewable Fuels under the RFS Program. Environ. Prot. Agency Off. Transp. Air Qual. (2013).
- Lewandowski, I., Scurlock, J. M. O., Lindvall, E. & Christou, M. The development and current status of perennial rhizomatous grasses as energy crops in the US and Europe. *Biomass and Bioenergy* 25, 335–361 (2003).
- 3. Pilu, R. Giant reed (Arundo donax L.): A weed plant or a promising energy crop? African J. Biotechnol. 11, 9163–9174 (2012).
- 4. Di Nasso, N., Roncucci, N. & Bonari, E. Seasonal Dynamics of Aboveground and Belowground Biomass and Nutrient Accumulation
- and Remobilization in Giant Reed (*Arundo donax* L.): A Three-Year Study on Marginal Land. *BioEnergy Res.* 6, 725–736 (2013).
 Mariani, C. *et al.* Origin, diffusion and reproduction of the giant reed (*Arundo donax* L.): a promising weedy energy crop. *Ann. Appl. Biol.* 157, 191–202 (2010).
- Angelini, L. G., Ceccarini, L., Nassi o Di Nasso, N. & Bonari, E. Comparison of Arundo donax L. and Miscanthus x giganteus in a long-term field experiment in Central Italy: Analysis of productive characteristics and energy balance. Biomass and Bioenergy 33, 635–643 (2009).
- Heaton, E. A quantitative review comparing the yields of two candidate C₄ perennial biomass crops in relation to nitrogen, temperature and water. *Biomass and Bioenergy* 27, 21–30 (2004).
- Borin, M. et al. Biomass production and N balance of giant reed (Arundo donax L.) under high water and N input in Mediterranean environments. Eur. J. Agron. 51, 117–119 (2013).
- 9. Ping, H., Bransby, D. I. & van Santen, E. Long-term biomass yields of giant reed, mimosa and switchgrass in Alabama. *Biofuels, Bioprod. Biorefining* **8**, 59–66 (2014).
- 10. Ceotto, E. et al. Comparing solar radiation interception and use efficiency for the energy crops giant reed (Arundo donax L.) and sweet sorghum (Sorghum bicolor L. Moench). F. Crop. Res. 149, 159–166 (2013).
- 11. Drewry, D. T., Kumar, P. & Long, S. P. Simultaneous improvement in productivity, water use, and albedo through crop structural modification. *Glob. Chang. Biol.* 20, 1955–67 (2014).
- Arundale, R. et al. Yields of Miscanthus × giganteus and Panicum virgatum decline with stand age in the Midwestern USA. Glob. Chang. Biol. - Bioenergy 6, 1–13 (2014).
- Mann, J. J., Kyser, G. B., Barney, J. N. & DiTomaso, J. M. Assessment of Aboveground and Belowground Vegetative Fragments as Propagules in the Bioenergy Crops Arundo donax and Miscanthus × giganteus. bioenerg 6, 688–698 (2013).
- 14. Papazoglou, E. G., Karantounias, G. A., Vemmos, S. N. & Bouranis, D. L. Photosynthesis and growth responses of giant reed (*Arundo donax* L.) to the heavy metals Cd and Ni. In *Environ. Int.* **31**, 243–249 (2005).
- Rossa, B., Tuffers, A. V., Naidoo, G. & Willert, D. J. Von. Arundo donax L. (Poaceae) a C3 Species with Unusually High Photosynthetic Capacity. Bot. Acta 111, 216–221 (1998).
- Dohleman, F. G., Heaton, E. A., Leakey, A. D. B. & Long, S. P. Does greater leaf-level photosynthesis explain the larger solar energy conversion efficiency of Miscanthus relative to switchgrass? *Plant. Cell Environ.* 32, 1525–37 (2009).
- 17. Dohleman, F. G. & Long, S. P. More productive than maize in the Midwest. How does Miscanthus do it? *Plant Physiol*. Preview, 109.139162 (2009).
- 18. Long, S. P. & Spence, A. K. Toward cool C4 crops. Annu. Rev. Plant Biol. 64, 701-22 (2013).
- Long, S. P., Ainsworth, E. A., Rogers, A. & Ort, D. R. Rising Atmospheric Carbon Dioxide: Plants FACE the Future. Annu. Rev. Plant Biol. 55, 591–628 (2004).
- Farquhar, G. D., Caemmerer, S., Von & Berry, J. A. A Biochemical Model of Photosynthetic CO₂ Assimilation in Leaves of C₃ Species. *Planta* 149, 78–90 (1980).
- Wang, D. et al. A physiological and biophysical model of coppice willow (Salix spp.) production yields for the contiguous USA in current and future climate scenarios. Plant. Cell Environ. 38, 1850–1865 (2015).
- 22. ISRIC. (2014). at<www.isric.org>.
- 23. Monteith, J. L., Campbell, G. S. & Potter, E. A. Theory and performance of a dynamic diffusion porometer. Agric. For. Meteorol. 44, 27–38 (1988).
- Potter, L., Bingham, M. G., Baker, M. G. & Long, S. P. The potential of two perennial C₄ grasses and a perennial C₄ sedge as lignocellulosic fuel crops in N. W. Europe crop establishment and yields in E. England. Ann. Bot. 76, 513–520 (1995).
- Baker, N. R. Chlorophyll Fluorescence: A Probe of Photosynthesis *In Vivo. Annu. Rev. Plant Biol.* 59, 89–113 (2008).
 Murchie, E. H. & Lawson, T. Chlorophyll fluorescence analysis: a guide to good practice and understanding some new applications.
- J. Exp. Bot. 64, 3983–98 (2013).
 17. Bernacchi, C. J., Singsaas, E. L., Pimentel, C., Portis Jr., A. R. & Long, S. P. Improved temperature response functions for models of Rubisco-limited photosynthesis. *Plant, Cell Environ.* 24, 253–259 (2001).
- Bernacchi, C. J., Pimentel, C. & Long, S. P. In vivo temperature response functions of parameters required to model RuBP-limited photosynthesis. Plant. Cell Environ. 26, 1419–1430 (2003).
- Long, S. P. & Bernacchi, C. J. Gas exchange measurements, what can they tell us about the underlying limitations to photosynthesis? Procedures and sources of error. J. Exp. Bot. 54, 2393–2401 (2003).
- Genty, B., Briantais, J.-M. & Baker, N. R. The relationship between the quantum yield of photosynthetic electron transport and quenching of chlorophyll fluorescence. *Biochim. Biophys. Acta* 990, 87–92 (1989).
- 31. Marshall, B. & Biscoe, P. V. A Model for C₃ Leaves Describing the Dependence of Net Photosynthesis on Irradiance. J. Exp. Bot. 31, 29–39 (1980).
- Genty, B., Harbinson, J. & Baker, N. Relative quantum efficiencies of the two photosystems of leaves in photorespiratory and nonrespiratory conditions. *Plant Physiol. Biochem.* 28, 1–10 (1990).
- Edwards, G. E. & Baker, N. R. Can CO₂ assimilation in maize leaves be predicted accurately from chlorophyll fluorescence analysis? Photosynth. Res. 37, 89–102 (1993).
- 34. Lawson, T. *et al.* C3 photosynthesis in the desert plant *Rhazya stricta* is fully functional at high temperatures and light intensities. *New Phytol.* **201**, 862–873 (2014).
- 35. Valentini, R., Epron, D., Angelis, P. D. E., Matteucci, G. & Dreyer, E. *In situ* estimation of net CO₂ assimilation, photosynthetic electron flow and photorespiration in Turkey oak (*Q. cerris* L.) leaves: diurnal cycles under different levels of water supply. *Plant Cell Environ.* 18, 631–640 (1995).
- Naidu, S. L. & Long, S. P. Potential mechanisms of low temperature tolerance of C₄ photosynthesis in *Miscanthus x giganteus*: an *in vivo* analysis. *Planta* 220, 145–155 (2004).
- 37. Penman, H. L. & Schofield, R. K. Some physical aspects of assimilation and transpiration. in Symp. Soc. Exp. Biol 5, 115–129 (1951).
- Bacon, M. Water use efficiency in plant biology. (John Wiley & Sons, 2009).
 Wullschleger, S. D. Biochemical Limitations to Carbon Assimilation in C₃ Plants—A Retrospective Analysis of the A/C_i Curves from 109 Species. J. Exp. Bot. 44, 907–920 (1993).
- Evans, J. R. & Farquhar, G. D. In Model. Crop Photosynth. Biochem. to canopy (Boote, K. & Loomis, R.) 1–15 (American Society of Agronomy, Inc, 1991).
- He, Z., Bentley, L. P. & Holaday, A. S. Greater seasonal carbon gain across a broad temperature range contributes to the invasive potential of Phalaris arundinacea (Poaceae; reed canary grass) over the native sedge Carex stricta (Cyperaceae). Am. J. Bot. 98, 20–30 (2011).

- 42. Ziska, L. H. & Teramura, A. H. CO₂ Enhancement of Growth and Photosynthesis in Rice (*Oryza sativa*): Modification by Increased Ultraviolet-B Radiation. *Plant Physiol.* **99**, 473–481 (1992).
- Siebke, K. *et al.* Control of photosynthesis in leaves as revealed by rapid gas exchange and measurements of the assimilatory force FA. *Planta* 182, 513–522 (1990).
- Harley, P. C. & Sharkey, T. D. An improved model of C3 photosynthesis at high CO₂: Reversed O₂ sensitivity explained by lack of glycerate reentry into the chloroplast. *Photosynth. Res.* 27, 169–178 (1991).
- Sage, R. F., Sharkey, T. D. & Seemann, J. R. Acclimation of Photosynthesis to Elevated CO₂ in Five C3 Species. *Plant Physiol.* 89, 590–596 (1989).
- Naidu, S. L., Moose, S. P., Al-Shoaibi, A. K., Raines, C. A. & Long, S. P. Cold Tolerance of C4 photosynthesis in Miscanthus x giganteus: Adaptation in Amounts and Sequence of C4 Photosynthetic Enzymes. 132, 1688–1697 (2003).
- Ge, Z. M. et al. Acclimation of photosynthesis in a boreal grass (*Phalaris arundinacea* L.) under different temperature, CO₂, and soil water regimes. *Photosynthetica* 50, 141–151 (2012).
- Brooks, A. & Farquhar, G. D. Effect of temperature on the CO₂/O₂ specificity of ribulose-1,5-bisphosphate carboxylase/oxygenase and the rate of respiration in the light. *Planta* 165, 397–406 (1985).
- Osborne, B. A. & Garrett, M. K. Quantum yields for CO₂ uptake in some diploid and tetraploid plant species. *Plant. Cell Environ.* 6, 135–144 (1983).
- Long, S. P., Postl, W. F. & Bolhàr-Nordenkampf, H. R. Quantum yields for uptake of carbon-dioxide in C-3 vascular plants of contrasting habitats and taxonomic groupings. *Planta* 189, 226–234 (1993).
- Maroco, J. P., Rodrigues, M. L., Lopes, C. & Chaves, M. M. Limitations to leaf photosynthesis in field-grown grapevine under drought - metabolic and modelling approaches. *Funct. Plant Biol.* 29, 451 (2002).
- Bernacchi, C. J., Morgan, P. B., Ort, D. R. & Long, S. P. The growth of soybean under free air [CO₂] enrichment (FACE) stimulates photosynthesis while decreasing *in vivo* Rubisco capacity. *Planta* 220, 434–446 (2005).
- Ripley, B. S., Gilbert, M. E., Ibrahim, D. G. & Osborne, C. P. Drought constraints on C₄ photosynthesis: Stomatal and metabolic limitations in C₃ and C₄ subspecies of *Alloteropsis semialata. J. Exp. Bot.* 58, 1351–1363 (2007).
- Hu, L., Wang, Z. & Huang, B. Diffusion limitations and metabolic factors associated with inhibition and recovery of photosynthesis from drought stress in a C₃ perennial grass species. *Physiol. Plant.* **139**, 93–106 (2010).
- Medrano, H., Flexas, J. & Galmés, J. Variability in water use efficiency at the leaf level among Mediterranean plants with different growth forms. *Plant Soil* 317, 17–29 (2009).

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Author Contributions

R.W. and S.D. conceived and designed the study. R.W., S.D., S.L., J.K., J.M., A.L., K.S., T.D., T.L., T.P. and S.B. all contributed to data collection and analysis. R.W. and S.D. produced the final analysis. R.W., S.D. and S.L. primarily led manuscript preparation. All authors reviewed the final manuscript.

Additional Information

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