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# **RESEARCH PAPER**

# A meta-analysis of mesophyll conductance to CO<sub>2</sub> in relation to major abiotic stresses in poplar species

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# Abstract

Mesophyll conductance ( $g_m$ ) determines the diffusion of CO<sub>2</sub> from the substomatal cavities to the site of carboxylation in the chloroplasts and represents a critical component of the diffusive limitation of photosynthesis. In this study, we evaluated the average effect sizes of different environmental constraints on  $g_m$  in *Populus* spp., a forest tree model. We collected raw data of 815 A-C<sub>i</sub> response curves from 26 datasets to estimate  $g_m$ , using a single curve-fitting method to alleviate method-related bias. We performed a meta-analysis to assess the effects of different abiotic stresses on  $g_m$ . We found a significant increase in  $g_m$  from the bottom to the top of the canopy that was concomitant with the increase of maximum rate of carboxylation and light-saturated photosynthetic rate ( $A_{max}$ ).  $g_m$  was positively associated with increases in soil moisture and nutrient availability, but was insensitive to increasing soil copper concentration and did not vary with atmospheric CO<sub>2</sub> concentration. Our results showed that  $g_m$  was strongly related to  $A_{max}$  and to a lesser extent to stomatal conductance ( $g_s$ ). Moreover, a negative exponential relationship was obtained between  $g_m$  and specific leaf area, which may be used to scale-up  $g_m$  within the canopy.

Keywords: Abiotic stress, A–C<sub>i</sub> curve, mesophyll conductance, meta-analysis, photosynthesis, poplar.

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## Introduction

Carbon assimilation in plants is importantly determined by the diffusion efficiency of CO<sub>2</sub> from the atmosphere to the site of carboxylation. The rate of CO2 diffusion is affected by two main diffusion limitations. The first limitation controls the  $CO_2$  flux from the atmosphere to the sub-stomatal cavities through the stomata and is characterized by stomatal conductance  $(g_s)$ . The second limitation determines the diffusion of  $CO_2$  from the substomatal cavities to the sites of carboxylation in the chloroplasts and is characterized by mesophyll conductance  $(g_m)$ .  $g_m$  is composed of gaseous and liquid phase resistances (Flexas et al., 2008; Evans et al., 2009; Niinemets et al., 2009). CO<sub>2</sub> diffusion inside the leaves is complex, facing a series of structural barriers coupled with biochemical regulation. It has been shown that  $g_m$  is typically limited by liquid phase conductance both in species with soft mesophytic leaves and in species with tough xerophytic leaves (Tosens et al., 2012a, b; Tomás et al., 2013). The liquid phase is a multicomponent pathway that involves the mesophyll cell wall thickness and porosity, the plasmalemma, the chloroplast envelope, the chloroplast thickness, and the mesophyll surface area exposed to intercellular air spaces per unit of leaf area (Evans et al., 2009; Tosens et al., 2012b; Tomás et al., 2013). After extensive study during the past two decades,  $g_m$  is now widely accepted as a critical limiting factor to photosynthesis, which has to be considered in characterizing plant carbon gain potentials and responses to future climate change (Evans et al., 2009; Niinemets et al., 2009, 2011; Flexas et al., 2016).

Mesophyll conductance has been shown to respond to environmental stress and may govern functional plasticity of photosynthesis and plant fitness under limited resources (Galle et al., 2009; Barbour et al., 2010; Buckley and Warren, 2014; Théroux Rancourt et al., 2015; Flexas et al., 2016; Shrestha et al., 2018). However, recent findings on the response of  $g_m$  to abiotic stress are conflicting and inconclusive, demonstrating the complex nature of g<sub>m</sub> variation (Flexas et al., 2008; Niinemets et al., 2009; Zhou et al., 2014; Shrestha et al., 2018). This suggests that the environmental and species-specific responses (and consequently the level of acclimation) of  $g_m$  to growth conditions should be considered in predicting plant performance in the field. Among the contrasting environmental responses, growth temperature may (Warren, 2008; Silim et al., 2010) or may not (Dillaway and Kruger, 2010; Benomar et al., 2018) affect  $g_{\rm m}$ . Similarly, the increase in soil nitrogen may (Warren, 2004; Shrestha et al., 2018; Xu et al., 2020; Zhu et al., 2020) or may not (Bown et al., 2009) stimulate  $g_m$ . The magnitude of decrease in  $g_{\rm m}$  under water stress and low light differed among studies (Warren et al., 2003; Niinemets et al., 2006; Montpied et al., 2009; Bögelein et al., 2012; Tosens et al., 2012a; Zhou et al., 2014; Peguero-Pina et al., 2015; Théroux Rancourt et al., 2015). These discrepancies among studies result in part from (i) the absolute changes in structural and biochemical traits controlling  $g_m$ , as well as from changes in the relative

contribution of these traits (Marchi *et al.*, 2008; Tomás *et al.*, 2013), and (ii) the level of coordination between  $g_m$ ,  $g_s$ , and leaf specific hydraulic conductivity ( $K_L$ ) (Flexas *et al.*, 2013; Théroux-Rancourt *et al.*, 2014; Xiong *et al.*, 2017). Given the complex interplay between different factors controlling  $g_m$ , it is important to examine its acclimation at the genus and species level to gain a general insight into the mechanistic basis of changes in  $g_m$ .

Five methods exist to estimate  $g_m$ : (i) chlorophyll fluorescence coupled to gas exchange (Harley *et al.*, 1992), (ii) carbon isotope discrimination coupled to gas exchange (initially developed by Evans *et al.*, 1986), (iii) oxygen isotope discrimination (Barbour *et al.*, 2016), (iv)  $A-C_i$  curve fitting (Ethier and Livingston, 2004; Sharkey *et al.*, 2007), and (v) 1D modeling of  $g_m$  from leaf structural characteristics (Evans *et al.*, 2009; Tosens *et al.*, 2012b; Tomás *et al.*, 2013). All of these methods are based on specific assumptions and each one has its limitations (Flexas *et al.*, 2013; Tosens and Laanisto, 2018). The standard deviation of the estimate of  $g_m$  may vary from 10% to 40%, which may limit our understanding of  $g_m$  acclimation to growth conditions, particularly when the variation between treatments or studies is less than the error of estimates (Sun *et al.*, 2014a).

Populus spp., model crops in forestry characterized by high yield potential, have been the subject of numerous studies to understand the physiological response to environmental factors but research is still necessary to make assessment of effects sizes and to make generalizations (Larocque et al., 2013). A general understanding of the CO<sub>2</sub> pathway through mesophyll and how it is affected by environmental factors would be beneficial in the effort to (i) accurately predict canopy photosynthesis under different environmental conditions, particularly under warmer and drier climate, and improve global carbon assimilation models, and (ii) effectively select more resilient and productive cultivars for wood and bioenergy. In poplar plantations, organic amendments like biosolids and pig slurry are used to increase growth rate at a low cost (Paniagua et al., 2016). These amendments are rich in copper, the effect of which on photosynthetic activity, growth, and nutrient uptake has been welldocumented in Populus spp. (Tognetti et al., 2004; Borghi et al., 2008; Pietrini et al., 2017). In addition, poplar is a good candidate for environmental use in phytofiltration of contaminated water in agriculture lands, where copper is a major contaminant due to the large use of copper sulfate as a fungicide and in weed control (Fischerová et al., 2006; Marmiroli et al., 2011).

Substantial data of  $A-C_i$  response curves in the literature have been used to estimate photosynthetic parameters, not to estimate  $g_m$ , and such compiled dataset would provide a basis to make such assessments of the response of  $g_m$  to the environment. In this study, we compiled 815  $A-C_i$  response curves from 26 datasets of different poplar species and hybrids (Table 1). Published  $A-C_i$  curve-fitting approaches differ broadly regarding the rectangularity of the hyperbola, segmentations of

# Table 1. List of dataset sources used in the meta-analysis

Author	<i>Populus</i> species or hybrid parents	Number of genotypes	Treatment	Provenance of plant material	Growth Environ- ment	Number of curves
Attia <i>et al.</i> (2015)	P. balsamifera L. P. simonii Carrière P. balsamifera L. × P. simonii Carrière	3	N/A	Canada	Growth chamber	15
Benomar <i>et al.</i> (https:// doi.org/10.5061/ dryad.9cnp5hqhp)	P. maximowiczii A. Henry × P. balsamifera L.	2	Water stress	Canada	Growth chamber	12
Benomar (2012)	P. maximowiczii A. Henry × P. balsamifera L. P. balsamifera L. × P. trichocarpa Torr. & A. Gray	2	Spacing and canopy level	Canada	Plantation	52
Benomar <i>et al.</i> (2019)	P. maximowiczii A. Henry × P. balsamifera L. P. maximowiczii A. Henry × P. nigra L.	2	Temperature and nitrogen	Canada	Growth chamber	23
Borghi <i>et al.</i> (2007)	P. × euramericana (P. deltoides W. Bartram × P. nigra L.) (clone Adda)	1	Copper	Italy	Growth chamber	21
Borghi <i>et al.</i> (2008)	P. alba L. P. × Canadensis (P. nigra L. × P. deltoides W. Bartram)	2	Copper	Italy	Growth chamber	18
Calfapietra et al. (2005)	<i>P.</i> × euramericana ( <i>P. deltoides</i> W. Bartram × <i>P. nigra</i> L.)	1	Nitrogen and atmospheric CO <sub>2</sub> and canopy level	Italy	Plantation	60
Castagna <i>et al.</i> (2015)	P. × canadensis (P. nigra L. × P. deltoides W. Bartram) P. deltoides W. Bartram × P. maximowiczii A. Henry	2	Ozone and cadmium soil contamination	Italy	Greenhouse	16
Di Baccio <i>et al.</i> (2009)	P. × euramericana (P. deltoides W. Bartram × P. nigra L.) (clone i-214)	1	Zinc soil contam- ination	Italy	Growth chamber	12
Elferjani <i>et al.</i> (2016)	P. trichocarpa Torr. & A. Gray × P. balsamifera L. (clone 747215) P. balsamifera L. × P. maximowiczii A. Henry (clones 915004 and 915005) P. maximowiczii A. Henry × P. balsamifera L. (clone 915319)	4	Latitudinal gra- dient	Canada	Plantation	24
Li <i>et al.</i> (2013)	P. euphratica Oliv.	1	Ground water availability	China	In field under shelter (lysim- eter)	9
Merilo <i>et al.</i> (2010)	P. nigra L. P. alba L.	2	Atmospheric CO <sub>2</sub> (FACE) and nitrogen and canopy level	Italy	Plantation	104
Niinemets <i>et al.</i> (1998)	P. tremula L.	1	Canopy level	Estonia	Natural forest stands	14
Ripullone <i>et al.</i> (2003)	P. × euramericana (P. deltoides W. Bartram × <i>P. nigra</i> L.) (clone i-214)	1	Nitrogen	Italy	Greenhouse	14
Ryan <i>et al.</i> (2009)	<i>P. deltoides</i> W. Bartram × <i>P. trichocarpa</i> Torr. & A. Gray	2	Ozone	United Kingdom	Greenhouse	118

# Table 1. Continued

Author	<i>Populus</i> species or hybrid parents	Number of genotypes	Treatment	Provenance of plant material	Growth Environ- ment	Number of curves
Silim <i>et al.</i> (2010)	P. balsamifera L.	1	Habitat and growth tempera- ture	Canada	Greenhouse	30
Soolanayakanahally <i>et al.</i> (2009)	P. balsamifera L.	1	Latitudinal gra- dient	Canada	Greenhouse	72
Théroux-Rancourt et al. (https:// doi.org/10.5061/ dryad.7sqv9s4s0)	P. deltoides W. Bartram × P. nigra L. (clone 3570) P. maximowiczii A. Henry × (P. deltoides W. Bartram × P. trichocarpa Torr. & A. Gray) (clones 505372 and 505508) P. maximowiczii A. Henry × P. trichocarpa Torr. & A. Gray (clone 750361) P. maximowiczii A. Henry × P. balsamifera L. (clones 915302, 915313, 915318) (P. deltoides W. Bartram × P.	8	Water stress	Canada	Greenhouse	38
Théroux-Rancourt <i>et al.</i> (2014)	nigra L.) × <i>P. trichocarpa</i> Torr. & A. Gray (clone 915508) Assiniboine: [( <i>P.</i> × 'Walker': <i>P. deltoides</i> W. Bartram × <i>P.</i> × potrowol/insp. P. J. Sobrod. ov	2	N/A	Canada	Greenhouse and growth	3
	Regel) × male parent unknown] Okanese [( $P$ :×'Walker') × $P$ :× petrowskiana R. I. Schrod. ex Regel]				Chamber	
Théroux-Rancourt <i>et al.</i> (2015)	<ul> <li>(P. maximowiczii A. Henry)×</li> <li>(P. deltoidesW. Bartram ×</li> <li>P. trichocarpa Torr. &amp; A. Gray)</li> <li>P. maximowiczii A. Henry ×</li> <li>P. balsamifera L.</li> <li>'Walker' [P. deltoides W. Bartram</li> <li>× (P. laurifolia Ledeb. × P. nigra</li> <li>L.)] × P. deltoides W. Bartram</li> <li>'Walker' × P. petrowskyana Schr.</li> <li>P. balsamifera L.</li> </ul>	5	N/A	Canada	Greenhouse and growth chamber	12
Tissue and Lewis (2010)	P. deltoides W. Bartram	1	Phosphorous and atmospheric CO <sub>2</sub>	Australia	Growth chamber	76
Tognetti <i>et al.</i> (https:// doi.org/10.5061/dryad. w3r2280qq)	<i>P. × euramericana (P. nigra</i> L. × <i>P. deltoides</i> W. Bartram) (clone i-214)		Zinc soil contam- ination	Italy	Greenhouse	24
Tognetti <i>et al.</i> (2004)	P. deltoides W. Bartram × P. maximowiczii A. Henry P. × euramericana (P. deltoides W. Bartram × P. nigra L.) (clone i-214)	2	Heavy metals	Italy	Greenhouse	24
Tosens <i>et al.</i> (2012a)	<i>P. tremula</i> L.	1	Light and water stress	Estonia	Growth chamber	8
Velikova <i>et al</i> . (2011)	P. nigra L.	20	Nickel soil con- tamination	Italy	Growth chamber (climate chamber)	16
Xu <i>et al.</i> (2020)	<i>P. × euramericana (P. deltoides</i> W. Bartram <i>× P. nigra</i> L.) (cv. '74/76')	1	Nitrogen and ozone	China	Growth chamber	6

the model of photosynthesis, and determination of the transition value of CO<sub>2</sub> from carboxylation to electron transport (Harley et al., 1992; Ethier and Livingston, 2004; Manter and Kerrigan, 2004; Dubois et al., 2007; Sharkey et al., 2007; Pons et al., 2009; Gu et al., 2010). These approaches led to different fitted values (Miao et al., 2009; Sun et al., 2014a). Although  $A-C_{i}$  curve fitting is unreliable for species with large  $g_{m}$ , it can provide results similar to those obtained from direct measurements for species with medium to low  $g_m$  (Niinemets et al., 2005, 2006; Warren, 2006; Qiu et al., 2017; Xu et al., 2020). Using the compiled  $A-C_i$  response curves, we performed curve fitting using a single method (Ethier and Livingston, 2004) to alleviate the fitting method bias and to obtain uniformed estimates of  $g_m$ , maximum rate of carboxylation ( $V_{cmax}$ ) and rate of electron transport (J). We further collected related variables like leaf nitrogen content, stomatal conductance, and specific leaf area (SLA) when data were available. Our main goal was to find trends in the response of mesophyll conductance to prevalent abiotic stressors and to examine the relationship between  $g_m$  and other leaf traits. We believe that a meta-analytical approach to analyse the accumulated data on the diffusion of  $CO_2$  through the mesophyll diffusion pathway in relation to other photosynthesis-related traits provides key insights into the different controls on mesophyll conductance and into the environmental plasticity of mesophyll conductance. We aim to contribute to the efforts of improving poplar photosynthetic efficiency in poplar breeding programs, and to improve modelling of global carbon assimilation of biomass and bioenergy crops under climate change.

# Materials and methods

### Data collection

Data were collected by a web search in Web of Science, Scopus, and Google Scholar using the following key words: ('*Populus*' or 'poplar' or 'hybrid poplar' or 'aspen') and (' $V_{\rm cmax}$ ' or 'maximum rate of electron transport ( $J_{\rm max}$ )' or 'mesophyll conductance'). At this step, the abstract of every item was checked to confirm the paper is actually about  $g_{\rm m}$ . Then, we looked at the 'Materials and methods' section of selected papers where  $A-C_{\rm i}$  response curves of *Populus* spp. were measured.

To get raw data of  $A-C_i$  response curves, we contacted the corresponding authors or co-authors of the targeted studies by e-mail and via ResearchGate. We obtained 23 datasets from published studies and three datasets from Benhomar, Tognetti and Théroux-Rancourt studies (Table 1; datasets available at Dryad Digital Repository). Collectively, they provided a total of 815  $A-C_i$  response curves.

The total data of 72 genotypes were collected from measurements on plants growing in plantations (five studies), or under controlled conditions (greenhouse or growth chamber set-ups; 21 studies) with optimal and stressful conditions (Table 1). After compiling all  $A-C_i$ curves, the quality of the data was assessed based on the following criteria: (i) only curves with at least two points in the saturation region (*J* region) were retained; (ii) only fitted curves with *P*-value <0.05 using the method of Ethier and Livingston (2004) were retained, and consequently 65 curves that did not meet these conditions were removed; and (iii) based on the literature,  $g_m$  values in *Populus* spp. using at least two methods simultaneously never exceeded 1 mol m<sup>-2</sup> s<sup>-1</sup> (Singsaas et al., 2004; Flexas et al., 2008; Velikova et al., 2011; Tosens *et al.*, 2012*a*; Théroux-Rancourt *et al.*, 2014; Momayyezi and Guy, 2017; Xu *et al.*, 2020). Then,  $g_m$  values >1 mol m<sup>-2</sup> s<sup>-1</sup> were considered as non-available data (94 entries), and  $V_{cmax}$  and J values were retained for further analyses.

### Data subsets

To examine the effect of a given abiotic factor on  $g_{\rm m}$ , we estimated that a minimum of three studies is necessary to have reliable conclusions, regardless of the genotype used, except copper for which only two studies were examined because they had been conducted under the same experimental conditions. Then, we could come up with subsets of data that focused on the same variable and performed analyses on them separately (identified in the column 'Treatment' in Table 1). Our first goal was to examine the effect of variations in these factors on  $g_{\rm m}$ , light-saturated photosynthetic rate ( $A_{\rm max}$ ),  $g_{\rm s}$ , J,  $V_{\rm cmax}$ , and in a second step, the relationships between  $g_{\rm m}$  and other photosynthetic characteristics ( $A_{\rm max}$ ,  $g_{\rm s}$ , J,  $V_{\rm cmax}$ ). The data subsets included the following environmental factors:

- Canopy level: four studies addressed the photosynthetic activity of leaves at the bottom, middle and top of trees (Niinemets *et al.*, 1998; Calfapietra *et al.*, 2005; Merilo *et al.*, 2010; Benomar, 2012).
- Atmospheric CO<sub>2</sub>: we examined the response of trees to elevated atmospheric CO<sub>2</sub> from the studies of Calfapietra *et al.* (2005), Merilo *et al.* (2010) and Tissue and Lewis (2010). We considered 370 ppm as the control treatment in the three studies, while the elevated CO<sub>2</sub> was 550 ppm of CO<sub>2</sub> for the studies of Calfapietra *et al.* (2005) and Merilo *et al.* (2010), and 700 ppm for the study of Tissue and Lewis (2010).
- Copper (Cu) stress: datasets from the studies of Borghi *et al.* (2007) and Borghi *et al.* (2008) were used to examine the response of poplar trees to contamination of the substrate with Cu. Treatments were assigned to three levels of Cu: 0 (0–0.4 μM), 20 (20–25 μM) and 75 (75–100 μM).
- Soil nitrogen (N) content: high vs. low soil N content treatments were reported in four studies: Ripullone et al. (2003), Calfapietra et al. (2005), Benomar et al. (2018) and Xu et al. (2020). In the study of Merilo et al. (2010), the authors showed that no effect of nitrogen fertilization was observed due to high background nutrient availability in the plantation site.
- Soil moisture: water status of trees was assessed and data from four studies were classified into two treatments: control (optimal watering) vs. water deficit (Li et al., 2013; Tosens et al., 2012a; Théroux-Rancourt (data available at Dryad Digital Repository: https://doi. org/10.5061/dryad.7sqv9s4s0); Benomar (data available at Dryad Digital Repository: https://doi.org/10.5061/dryad.9cnp5hqhp).

For Xu *et al.* (2020), we extracted data from the article (means and standard errors) and generated three replicates assuming a normal distribution using the SURVEYSELECT procedure of SAS (version 9.4; SAS Institute, Cary, NC, USA). The reason is that the authors used the same curve fitting approach (Ethier and Livingston, 2004) the we used in this meta-analysis study (Table 1).

For studies with two or more investigated factors, we considered the different levels of the factor of interest and the control level of the rest of the factors to avoid between-factor interaction effects on the results. For example, in Calfapietra *et al.* (2005), trees were subject to different levels of N and CO<sub>2</sub>; when we focused on the effect of N, we selected trees exposed to ambient CO<sub>2</sub> only (control).

### Curve analysis

Mesophyll conductance and photosynthetic capacity variables,  $V_{\text{cmax}}$  and J, were estimated by fitting  $A-C_i$  curve with the non-rectangular hyperbola version (Ethier and Livingston, 2004) of the biochemical model of

 $C_3$  plants (Farquhar *et al.*, 1980). This method was calibrated for low  $g_m$  species (<0.3 mol m<sup>-2</sup> s<sup>-1</sup>) and its accuracy is similar to estimates using the chlorophyll fluorescence method and online carbon <sup>13</sup>C isotope discrimination (Niinemets *et al.*, 2005, 2006; Ethier *et al.*, 2006; Tomás *et al.*, 2013; Qiu *et al.*, 2017; Xu *et al.*, 2020). The model was fitted using non-linear regression techniques (Proc NLIN, SAS) following Dubois *et al.* (2007) and Sun *et al.* (2014a).

Briefly, the net assimilation rate  $(A_n)$  is given as:

$$A_{\rm n} = \min \left\{ A_{\rm c}, \ A_{\rm j} \right\} \tag{1}$$

with

$$A_{\rm c} = V_{\rm cmax} \frac{\left(C_{\rm c} - \Gamma^*\right)}{C_{\rm c} + K_{\rm c} \left(1 + \frac{O}{K_{\rm o}}\right)} - R_{\rm day} \tag{2}$$

$$A_{\rm j} = J \; \frac{C_{\rm c} - \Gamma^*}{4 \left(C_{\rm c} + 2\Gamma^*\right)} - R_{\rm day} \tag{3}$$

$$C_{\rm c} = C_{\rm i} - \frac{A_{\rm n}}{g_{\rm m}} \tag{4}$$

where  $A_c$  is the Rubisco-limited rate of CO<sub>2</sub> assimilation (µmol m<sup>-2</sup> s<sup>-1</sup>),  $A_j$  is the RuBP-limited rate of CO<sub>2</sub> assimilation (µmol m<sup>-2</sup> s<sup>-1</sup>),  $V_{cmax}$  is the maximum rate of carboxylation (µmol m<sup>-2</sup> s<sup>-1</sup>), O is the partial atmospheric pressure of O<sub>2</sub> (mmol mol<sup>-1</sup>).  $\Gamma^*$  is the CO<sub>2</sub> compensation point in the absence of mitochondrial respiration,  $R_{day}$  is mitochondrial respiration in the light (µmol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>),  $C_c$  is the chloroplast CO<sub>2</sub> (µmol mol<sup>-1</sup>),  $C_i$  is the intercellular air space concentration of CO<sub>2</sub> (µmol mol<sup>-1</sup>),  $K_c$  (µmol mol<sup>-1</sup>) and  $K_o$  (mmol mol<sup>-1</sup>) are the Michaelis–Menten constants of Rubisco for CO<sub>2</sub> and O<sub>2</sub>, respectively, and J is the rate of electron transport (µmol m<sup>-2</sup> s<sup>-1</sup>). The values at 25 °C used for  $K_c$ ,  $K_o$ , and  $\Gamma^*$  were 272 µmol mol<sup>-1</sup>, 166 mmol mol<sup>-1</sup> and 37.4 µmol mol<sup>-1</sup>, respectively (Sharkey *et al.*, 2007), and their temperature dependencies were as in Sharkey *et al.* (2007).

In four datasets, measurements were carried out under a temperature that was different from the reference (25 °C). In this case,  $V_{\rm cmax}$  and J were normalized to 25 °C using the model of Kattge and Knorr (2007), which integrates the acclimation to growth temperature. However, the actual values of  $V_{\rm cmax}$  and J were more often significant compared with normalized values, and this was true using both ANOVA and regression analyses.

### Quantitative limitations analysis

The stomatal conductance  $(L_s)$ , mesophyll conductance  $(L_m)$ , and biochemical  $(L_b)$  relative limitations to photosynthesis were estimated following Grassi and Magnani (2005) as modified by Tomás *et al.* (2013):

$$L_{\rm s} = \frac{\left(\frac{g_{\rm tot}}{g_{\rm sc}}\right) \frac{\partial A_{\rm c}}{\partial A_{\rm c}}}{g_{\rm tot} + \frac{\partial A_{\rm c}}{\partial A_{\rm c}}} \tag{5}$$

$$L_{\rm m} = \frac{\left(\frac{g_{\rm tot}}{g_{\rm m}}\right) \frac{\partial A_c}{\partial A_c}}{g_{\rm tot} + \frac{\partial A_c}{\partial A_c}} \tag{6}$$

$$L_{\rm b} = \frac{g_{\rm tot}}{g_{\rm tot} + \frac{\partial A_c}{\partial C_c}} \tag{7}$$

where  $g_{tot}$  is the total CO<sub>2</sub> conductance and  $g_{sc}$  is the stomatal conductance to CO<sub>2</sub> ( $g_{sc}=g_{sw}/1.6$ ).

$$g_{\rm tot} = \frac{1}{\frac{1}{g_{\rm sc}} + \frac{1}{g_{\rm m}}}$$
(8)

$$\frac{\partial A_{\rm c}}{\partial C_{\rm c}} = \frac{V_{\rm cmax} \left( \Gamma^* + k_{\rm c} \left( 1 + {}^{\rm O} \middle/_{k_{\rm o}} \right) \right)}{\left( C_{\rm c} + k_{\rm c} \left( 1 + {}^{\rm O} \middle/_{k_{\rm o}} \right) \right)^2} \tag{9}$$

where  $\frac{\partial A_c}{\partial C_c}$  is the first derivative of  $A_c$  with respect to  $C_c$ .

Factors for which  $A_{\text{max}}$  changed significantly (canopy level, soil nitrogen, and soil moisture), the absolute contribution of stomatal conductance limitation ( $S_L$ ), mesophyll conductance (MC<sub>L</sub>), and biochemical photosynthetic capacity limitation ( $B_L$ ) to observed change of  $A_{\text{max}}$  were estimated following Grassi and Magnani (2005):

$$\frac{\mathrm{d}A_{\max}}{A_{\max}} = S_{\mathrm{L}} + \mathrm{MC}_{\mathrm{L}} + B_{\mathrm{L}} = L_{\mathrm{s}}\frac{\mathrm{d}g_{\mathrm{sc}}}{g_{\mathrm{sc}}} + L_{\mathrm{m}}\frac{\mathrm{d}g_{\mathrm{m}}}{g_{\mathrm{m}}} + L_{\mathrm{b}}\frac{\mathrm{d}V_{\mathrm{cmax}}}{V_{\mathrm{cmax}}}$$
(10)

where  $\frac{dA_{\text{max}}}{A_{\text{max}}}$  is the difference of  $A_{\text{max}}$  between the reference and the other treatments (within each factor) divided by  $A_{\text{max}}$  of the reference.

### Statistical analyses

Data analysis assessing the effect of the environmental factors on  $g_m$  and the relationship between  $g_m$  and the other traits were carried out using SAS version 9.4.

For studies that focused on nitrogen, soil moisture,  $CO_2$ , canopy level and copper, the effect of treatments on light-saturated photosynthetic rate ( $A_{max}$ ),  $g_m$ , and  $g_{sw}$  was assessed separately for each response variable, using mixed model analyses of variance of the primary data (Riley *et al.*, 2010; Mengersen *et al.*, 2013). 'Treatment' was the fixed effect while 'study' and 'genotype' nested within study were the random effects. The number of replicates was not necessarily balanced across treatments. The assumptions of normality of the residuals and homogeneity of variance were verified, and a log-transformation was made when necessary.

### Results

The number of studies on mesophyll conductance has rapidly increased since 2000, and more remarkably since 2013 (Fig. 1A), suggesting a growing interest among plant ecophysiologists in understanding the role of  $g_m$  in photosynthesis. This pattern was very similar to the increase of publication number on mesophyll conductance in *Populus* spp. (Fig. 1B).

### Canopy level

Light-saturated photosynthetic rate at an ambient CO<sub>2</sub> concentration (380–400 µmol mol<sup>-1</sup>),  $A_{\text{max}}$ , significantly increased from 7.19±0.44 µmol m<sup>-2</sup> s<sup>-1</sup> on average at the bottom leaves to 13.15±0.45 µmol m<sup>-2</sup> s<sup>-1</sup> at the mid-canopy, to 16.29±0.53 µmol m<sup>-2</sup> s<sup>-1</sup> at the upper canopy (Fig. 2A; Supplementary Table S1). Similar to  $A_{\text{max}}$ ,  $g_{\text{m}}$  had an ascending pattern, from the bottom (0.12±0.01 mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>) to the top of the canopy (0.24±0.02 mol m<sup>-2</sup> s<sup>-1</sup>) (Fig. 2C). Stomatal conductance ( $g_{\text{sw}}$ ) was the lowest at the bottom canopy (0.17±0.01 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) and then increased to 0.36±0.02 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup> at the mid and upper canopy (Fig. 2B). The  $g_{\text{m}}/g_{\text{sc}}$  ratio was significantly greater at the upper



**Fig. 1.** Cumulative number of published studies for mesophyll conductance ( $g_m$ ) between the years 2000 and 2020 (A), and cumulative number of published studies for mesophyll conductance ( $g_m$ ) in *Populus* spp. between the years 2001 and 2020 (B). The number of publications was determined using keywords (e.g.  $g_m$  and *Populus*) through database search available at the Web of Science Core Collection (https://clarivate.com/webofsciencegroup/solutions/web-of-science-core-collection/).

canopy (1.17±0.11), compared with the mid-canopy leaves (0.88±0.09) and was not different everywhere else (Fig. 2D).  $V_{\rm cmax}$  increased similarly to  $A_{\rm max}$  and  $g_{\rm m}$  from the bottom to the top of the canopy (Fig. 2E); however, SLA had an opposite trend (Fig. 2F).

### Atmospheric CO<sub>2</sub>

Increased atmospheric CO<sub>2</sub> had no effect on average  $A_{\text{max}}$  (14.43±0.60 µmol m<sup>-2</sup> s<sup>-1</sup>),  $g_{\text{m}}$  (0.21±0.02 mol m<sup>-2</sup> s<sup>-1</sup>) and  $g_{\text{m}}/g_{\text{sc}}$  (1.09±0.11) (Fig. 3A–C). However, average  $g_{\text{sw}}$  was higher (0.40±0.03 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) under 'Ambient', compared with 'Elevated' CO<sub>2</sub> (0.32±0.02 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) (Fig. 3B).

### Copper stress

 $A_{\text{max}}$  was not affected when soil Cu concentration increased from 0 to 20 or 75  $\mu$ M (9.67±0.95  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) (Fig. 4A). It should be noted that at the highest Cu level (75  $\mu$ M),  $A_{\text{max}}$  ranged from 4 to 15 µmol m<sup>-2</sup> s<sup>-1</sup>. Average  $g_{sw}$  significantly decreased under medium (20 µM, 0.17±0.02 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) and high Cu treatment (75 µM, 0.18±0.03 mol m<sup>-2</sup> s<sup>-1</sup>), compared with control treatment (Fig. 4B). Increasing Cu concentration in the soil did not affect  $g_m$  and the  $g_m/g_{sc}$  ratio (Fig. 4C, D).

### Soil nitrogen

 $A_{\rm max}$  was significantly greater (16.07±0.61 µmol m<sup>-2</sup> s<sup>-1</sup>) under high soil nitrogen (HN, 250 kg N ha<sup>-1</sup> y<sup>-1</sup> in field study or 20 mM for pot study) compared with low nitrogen treatment (LN, 12.93±0.65 µmol m<sup>-2</sup> s<sup>-1</sup>) (Fig. 5A). A high supply of nitrogen increased  $g_{\rm sw}$  (from 0.29±0.03 in LN to 0.36±0.03 mol m<sup>-2</sup> s<sup>-1</sup> in HN) and  $g_{\rm m}$  (from 0.19±0.02 to 0.23±0.02 to mol m<sup>-2</sup> s<sup>-1</sup>), but had no effect on the  $g_{\rm m}/g_{\rm sc}$  ratio (1.38±0.16 on average) (Fig. 5B–D).

### Soil moisture

Average  $A_{\text{max}}$  decreased by drought (range of leaf predawn water potential under water deficit,  $\Psi_{\text{leaf}} = -0.7$  to -0.8, soil water content=10%), dropping from  $17.13\pm0.71 \text{ }\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  to  $14.62\pm0.91 \text{ }\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  on average with the minimum value (3.83  $\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$ ) much lower than in watered trees (8.90  $\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$ ) much lower than in watered trees (8.90  $\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$ ) much lower than in watered trees (8.90  $\mu\text{mol }\text{m}^{-2}\text{ }\text{s}^{-1}$ ) (Fig. 6A). As expected, soil moisture deficit markedly altered  $g_{\text{sw}}$ , decreasing its average value from  $0.33\pm0.02 \text{ mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  in control trees to  $0.20\pm0.03 \text{ mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  under drought conditions (Fig. 6B). Drought had the same effect on  $g_{\text{m}}$ , but to a lesser extent than  $g_{\text{sw}}$ .  $g_{\text{m}}$  decreased from  $0.27\pm0.02 \text{ mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  to  $0.19\pm0.02 \text{ mol }\text{m}^{-2}\text{ }\text{s}^{-1}$  under soil moisture deficit (Fig. 6C). In addition, the  $g_{\text{m}}/g_{\text{sc}}$  ratio increased by 37% when plants were subject to drought (Fig. 6D).

### Quantitative limitations

In general, photosynthetic rate was mostly limited by CO<sub>2</sub> diffusion (up to 75%): stomatal limitation ( $L_{\rm s}$ ) and mesophyll limitation  $(L_{\rm m})$  (Table 2). Biochemical limitation  $(L_{\rm b})$  of photosynthesis rate was relatively low. Higher atmospheric CO<sub>2</sub> decreased biochemical limitation (16.62% to 14.73%) and increased mesophyll limitation (from 41.80% to 44.28%) while stomatal limitation remained unchanged. Within the canopy, stomatal and biochemical limitations were the greatest in the upper (47.79%) and the middle (16.05%) layers of the canopy, respectively (Table 2). The mesophyll conductance limitation was higher at the middle (50.47%) and the bottom (50.85%)than at the upper part of the canopy (40.51%). The decrease of  $A_{\rm max}$  (58.04 %) from the top, as a reference, to the bottom of the canopy (calculated with Equation 10) was mostly caused by mesophyll (absolute limitation=29.48%), followed by stomatal (19.16%) limitation and to a lesser extent by  $V_{\text{cmax}}$  (5.11%). At the middle of the canopy, the decrease of  $A_{\text{max}}$  (21.58%) was mostly due to  $g_{\rm m}$  (13.15%) and to a lesser extent to  $V_{\rm cmax}$ 



**Fig. 2.** Effect of the leaf position in the canopy (Bot, bottom; Mid, middle; Upp, upper) on light-saturated photosynthetic rate ( $A_{max}$ , A), stomatal conductance ( $g_{sw}$ , B), mesophyll conductance ( $g_m$ , C),  $g_m/g_{sc}$  ratio (D), maximum rate of carboxylation ( $V_{cmax}$ , E), and specific leaf area (*SLA*, F). For  $g_m/g_{sc}$  ratio,  $g_{sw}$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>). The horizontal line inside the boxes marks the median for the observations, the box ends indicate the upper (third) to lower (first) quartiles of the value ranges, and the whiskers indicate the highest and lowest observations. Means having the same letters are not significantly different at  $\alpha$ =0.05 (number of studies=4, number of genotypes=6).



**Fig. 3.** Effect of the atmospheric CO<sub>2</sub> concentration on light-saturated photosynthetic rate ( $A_{max}$ , A), stomatal conductance ( $g_{sw}$ , B), mesophyll conductance ( $g_{m}$ , C), and  $g_m/g_{sc}$  ratio (D). For  $g_m/g_{sc}$  ratio,  $g_{sw}$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>). The horizontal line inside the boxes marks the median for the observations, the box ends indicate the upper (third) to lower (first) quartiles of the value ranges, and the whiskers indicate the highest and lowest observations. Means having the same letters are not significantly different at  $\alpha$ =0.05 (number of studies=3, number of genotypes=4).

(4.74%) while the contribution of  $g_{sc}$  was marginal (2.76%). The copper stress resulted in an increase of the stomatal limitation and a decrease in mesophyll and biochemical limitations. Change in soil nitrogen did not affect the status of the

limitations. The decrease of  $A_{\text{max}}$  (20.11 %) under low soil nitrogen was mostly caused by  $g_{\text{sc}}$  (9.38%) and  $g_{\text{m}}$  (7.58%) and to a lesser extent by  $V_{\text{cmax}}$  (1.42%). Water stress increased stomatal limitation and decreased biochemical limitation but had



**Fig. 4.** Effect of the soil copper (Cu) concentration on light-saturated photosynthetic rate ( $A_{max}$ , A), stomatal conductance ( $g_{sw}$ , B), mesophyll conductance ( $g_m$ , C), and  $g_m/g_{sc}$  ratio (D). For  $g_m/g_s$  ratio,  $g_s$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>). The horizontal line inside the boxes marks the median for the observations, the box ends indicate the upper (third) to lower (first) quartiles of the value ranges, and the whiskers indicate the highest and lowest observations. Means having the same letters are not significantly different at  $\alpha$ =0.05 (number of studies=2, number of genotypes=3).



**Fig. 5.** Effect of the soil nitrogen content (HN, high nitrogen; LN, low nitrogen) on light-saturated photosynthetic rate ( $A_{max}$ , A), stomatal conductance ( $g_{sw}$ , B), mesophyll conductance ( $g_m$ , C), and  $g_m/g_{sc}$  ratio (D). For  $g_m/g_{sc}$  ratio,  $g_{sw}$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>). The horizontal line inside the boxes marks the median for the observations, the box ends indicate the upper (third) to lower (first) quartiles of the value ranges, and the whiskers indicate the highest and lowest observations. Means having the same letters are not significantly different at  $\alpha$ =0.05 (number of studies=5, number of genotypes=7).

no effect on mesophyll limitation (Table 2). Therefore, the observed decrease in  $A_{\text{max}}$  (21.02%) under water deficit was mainly due to stomatal (15.22%) and mesophyll limitation (7.64%).

# Relationship between $CO_2$ diffusion and photosynthetic activity

 $A_{\text{max}}$  was strongly correlated to both  $g_{\text{sw}}$  and  $g_{\text{m}}$  (P=0.001) and to  $V_{\text{cmax}}$  (P=0.001) over all the studies (Fig. 7A–C). Based

on the collected data,  $g_m$  was significantly correlated to  $g_{sw}$  (P=0.04). However, the relationship was not linear.  $g_m$  was the highest (0.4–0.5 mmol m<sup>-2</sup> s<sup>-1</sup>) when  $g_{sw}$  values were intermediate (0.2–0.4 mol m<sup>-2</sup> s<sup>-1</sup>), and lowest at high  $g_{sw}$  values (Fig. 7E).

We found a significant negative exponential relationship between SLA and  $g_m$  (*P*=0.001) (Fig. 7G) based on the collected data from studies that measured SLA (*n*=12). Leaf nitrogen content reported by three studies showed a significant correlation between  $g_m$  and N content per area ( $N_{area}$ )



**Fig. 6.** Effect of the soil moisture on light-saturated photosynthetic rate ( $A_{max}$ , A), stomatal conductance ( $g_{sw}$ , B), mesophyll conductance ( $g_m$ , C), and  $g_m/g_{sc}$  ratio (D). For  $g_m/g_{sc}$  ratio,  $g_{sw}$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>). The horizontal line inside the boxes marks the median for the observations, the box ends indicate the upper (third) to lower (first) quartiles of the value ranges, and the whiskers indicate the highest and lowest observations. Means having the same letters are not significantly different at α=0.05 (number of studies=4, number of genotypes=13).

Table 2.	The relative limitation (%)	of stomatal	conductance,	mesophyll	conductance,	and biochemistry	/ to photosynthesis	(for each
treatment,	the sum of relative limita	tions is 100°	%)					

Factor	Treatment	Stomatal limitation ( $L_{s}$ )	Mesophyll limitation ( <i>L</i> <sub>m</sub> )	Biochemical limitation (L <sub>b</sub> )
Canopy level	Bottom	37.71 (2.11)	50.86 (2.68)	11.43 (1.22)
	Middle	33.46 (1.99)	50.48 (2.4)	16.06 (1.34)
	Upper	45.97 (2.18)	40.52 (1.98)	13.69 (1.09)
Atmospheric CO <sub>2</sub>	Ambient	41.57 (2.3)	41.8 (2.27)	16.62 (1.43)
	Elevated	40.97 (1.87)	44.29 (2.04)	14.74 (1.21)
Copper stress	0 μM Cu	42.42 (6.78)	31.17 (7.38)	26.43 (7.1)
	20 µM Cu	57.88 (4.96)	25.16 (6.7)	16.96 (2.71)
	75 µM Cu	59.13 (7.58)	28.22 (6.92)	12.65 (1.29)
Soil nitrogen	High nitrogen	45.09 (2.49)	39.31 (2.07)	15.6 (1.46)
	Low nitrogen	46.02 (2.69)	39.27 (2.37)	14.71 (1.67)
Soil moisture	Control	42.47 (2.29)	38.04 (1.97)	19.49 (1.09)
	Drought	48.8 (2.78)	38.74 (2.11)	12.46 (1.25)

Data are expressed as means (SD).

(Fig. 7F).  $g_{\rm m}$  increased with  $N_{\rm area}$  until a saturation point (~0.25 mol m<sup>-2</sup> s<sup>-1</sup>).

# Discussion

### Canopy level

The scaling up of photosynthesis from leaves to the canopy and stands (using the model of Farquhar *et al.* (1980)) requires a deep understanding of within-canopy variations in leaf morpho-physiology and the main drivers of foliage acclimation to the dynamic gradient of environmental conditions (light, temperature, vapor pressure deficit (VPD) and soil moisture) (Niinemets *et al.*, 2006; Buckley and Warren, 2014; Niinemets *et al.*, 2015). Unfortunately, pieces of knowledge regarding the variation of  $g_m$  within the canopy and its mechanistic basis are scarce, in particular for *Populus* spp. This situation may explain why most global carbon cycle models remain ' $g_m$ -lacking', with possible consequences, such as overestimation of the fertilization effect of CO<sub>2</sub> on global gross primary production and underestimation of water-use efficiency (WUE) and canopy gross photosynthesis under future climate (Sun *et al.*, 2014b; Knauer *et al.*, 2019).

The steep and parallel increase of  $g_m$ ,  $A_{max}$ , and  $V_{cmax}$  from the bottom to the top of the canopy found here for *Populus* spp. is in agreement with the findings of Niinemets *et al.* (2006) for *Quercus ilex* L., Montpied *et al.* (2009) for *Fagus sylvatica* L., and Warren *et al.* (2003) for *Pseudotsuga menziesii* (Mirbel) Franco. A decrease of  $g_m$  from the bottom to the top of the canopy was also reported (Bögelein *et al.*, 2012; Cano *et al.*, 2013). We



**Fig. 7.** Relationship between light-saturated photosynthetic rate ( $A_{max}$ ), stomatal conductance ( $g_{sw}$ ), mesophyll conductance ( $g_m$ ), maximum rate of carboxylation ( $V_{cmax}$ ), electron transport rate (J),  $g_m/g_{sc}$  ratio, specific leaf area (SLA) and per area leaf nitrogen concentration ( $N_{area}$ ). For  $g_m/g_{sc}$  ratio,  $g_{sw}$  for water (mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>) was divided by 1.6 to obtain  $g_{sc}$  (mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>).

observed a greater  $g_m$  limitation under shade conditions (mid and bottom of canopy), which may lead to an overestimation of canopy photosynthesis. Overall, our results highlight the need to incorporate the acclimation of  $g_m$  to light conditions along the canopy in process-based models.

We observed a significant inverse relationship between  $g_m$ and SLA, comparable to previous studies (Niinemets et al., 2006; Montpied et al., 2009; Tosens et al., 2012b). This suggests that the increase in leaf thickness (lower SLA), e.g. in developing leaves and in leaves grown under higher light, may be associated with increased  $g_m$  (Tosens *et al.*, 2012*b*). Contrary to this, a positive relationship between  $g_m$  and SLA was demonstrated across Solanum species (Muir et al., 2014), reflecting the effect of increased leaf density and mesophyll cell wall thickness on  $g_{\rm m}$ . These lines of evidence collectively demonstrate the complex nature of the relationship between SLA and  $g_{\rm m}$ , reflecting the circumstance that SLA is an inverse of the product of leaf thickness and density, which can respond differently to environmental drivers (Niinemets, 1999; Poorter et al., 2009). The profile of  $g_m$  within the canopy observed here may be partially attributable to the morphological acclimation of Populus spp. foliage to light availability within the canopy. Moreover, this inverse relationship between SLA and  $g_{\rm m}$  was used as an empirical model to estimate a maximum attainable  $g_m$  at different canopy layers for C3 plants and was implemented in the Community Land Model (CLM.4.5) (Sun et al., 2014b; Knauer et al., 2019).

The change in morphological traits and their role in the acclimation of  $g_m$  to a vertical gradient of environmental conditions within the canopy need additional investigation. For instance, shade acclimation of leaf morphology is associated with a lower surface area of chloroplasts exposed to intercellular air spaces  $(S_c/S)$  and thicker chloroplasts (Hanba et al., 2002; Niinemets et al., 2006; Tosens et al., 2012b; Peguero-Pina et al., 2015). Species-specific leaf development patterns (i.e. evergreen sclerophyllous versus deciduous broadleaves) affect limitations to gas diffusion, thus determining the carbon balance of leaves (Marchi et al., 2007). However, light acclimation may be species-specific and altered by water, soil nitrogen, and leaf ontogeny (Niinemets et al., 2006; Tazoe et al., 2009; Peguero-Pina et al., 2015; Shrestha et al., 2018). It is still unclear whether the  $g_{\rm m}$  profile within the canopy is the result of the change in SLA.

Our results showing higher  $g_{sw}$  and  $g_m/g_{sc}$  at the top of the canopy are in disagreement with the findings of Montpied *et al.* (2009) and Bögelein *et al.* (2012), suggesting a speciesand environment-specific gradient of  $g_m/g_s$ . Temperature and VPD responses of  $g_m$  and  $g_s$  are different (Cano *et al.*, 2013), resulting in different diurnal patterns of  $g_m$  and  $g_s$ . Then, the gradient of  $g_m/g_s$  ratio along the canopy may drive the WUE at the canopy level and the midday depression of photosynthetic rate regardless of the level of isohydry of clones (Cano *et al.*, 2013; Buckley and Warren, 2014; Stangl *et al.*, 2019).

### Atmospheric CO<sub>2</sub>

The response of photosynthetic capacity and diffusion of  $CO_2$ to free-air CO2 enrichment considerably differed between species and experimental set-ups. The decrease in  $A_{\text{max}}$  and g<sub>sw</sub> in response to elevated CO<sub>2</sub> shown in our meta-analysis is in agreement with numerous studies on Populus spp. and other species (Ainsworth and Rogers, 2007; Medlyn et al., 2013; DaMatta et al., 2016), but is in disagreement with the findings of some other studies, e.g. Sigurdsson et al. (2001) and Uddling et al. (2009). For  $g_m$ , the effect of growth CO<sub>2</sub> changed among studies and some species having an intrinsic low  $g_m$  are more likely to respond to elevated CO<sub>2</sub> than species with high intrinsic g<sub>m</sub> (Niinemets et al., 2011). However, several studies have reported that  $g_m$  may decrease or be unresponsive to CO<sub>2</sub> enrichment (Singsaas et al., 2004; Zhu et al., 2012; Kitao et al., 2015; Mizokami et al., 2019). This suggests that the increase of  $A_{\text{max}}$  under elevated CO<sub>2</sub> cannot be attributed solely to  $g_m$  variation (Singsaas *et al.*, 2004). The absence of  $g_m$ response to elevated CO<sub>2</sub> complicates the research on mechanisms underlying this variation. Unlike  $g_m$ , researchers have proposed some hypotheses such as least-cost theory, nitrogen limitation, and resources investment to explain the decrease of  $A_{\text{max}}$ ,  $V_{\text{cmax}}$ , and  $g_{\text{s}}$  under elevated CO<sub>2</sub> (Leakey *et al.*, 2009; Smith and Keenan, 2020).

### Copper stress

Similar to our findings, gm remained unchanged in the herbaceous plant Silene paradoxa L., exposed to high Cu concentration, although g<sub>s</sub> decreased significantly (Bazihizina et al., 2015). In other cases of exposure to other heavy metals, like nickel (Ni), Velikova et al. (2011) reported a significant decrease in chloroplast CO2 content and mesophyll conductance in black poplar (P. nigra L.) exposed to 200 µM Ni under a hydroponic set-up (compared with control of 30 µM Ni). This reduction of  $g_{\rm m}$  might be attributed to an alteration of leaf structure by toxic effects of high concentrations of heavy metals in mesophyll cells (Velikova et al., 2011). Hermle et al. (2007), reported an acceleration of senescence and necrosis of mesophyll cells in P. tremula L. leaves exposed to Cu, Zn, Cd, and Pb at 640, 3000, 10, and 90 mg kg<sup>-1</sup> soil, respectively, and a decrease of chloroplast size from the early stages of exposure. The study of Hermle et al. (2007) also reported the thickening of cell walls and change of their chemical composition in damaged mesophyll cells, which might have affected permeability of cell walls and diffusion of CO<sub>2</sub> through them. Mercury (Hg; HgCl<sub>2</sub> form) altered CO<sub>2</sub> diffusion through aquaporins, a membrane channel of CO<sub>2</sub> diffusion, in faba bean (Vicia faba L.) (Terashima and Ono, 2002) and significantly reduced  $g_m$  in *P. trichocarpa* Torr. & Gray. HgCl<sub>2</sub> may also decrease  $g_m$  indirectly by disrupting carbonic anhydrase activity, as reported by Momayyezi and Guy (2018), who demonstrated that carbonic

anhydrase activity is strongly associated with  $g_m$  variation in *P. trichocarpa* Torr. & Gray (Momayyezi and Guy, 2017).

### Soil nitrogen

The increase of  $A_{\text{max}}$  by the enhancement of  $V_{\text{cmax}}$  in response to more available soil nitrogen has been established in the literature. However, the possible contribution of  $g_m$  to this augmentation remains unexplored for several species. Our results showed a concomitant increase of  $g_m$  with a higher supply of N. A positive correlation between the level of expression of aquaporin genes (plasma membrane intrinsic proteins and tonoplast intrinsic proteins) and  $g_m$  has been reported (Hanba et al., 2004; Flexas et al., 2006; Kaldenhoff et al., 2008; Perez-Martin et al., 2014), although it is still unclear whether this is a direct effect or a pleiotropic effect reflecting simultaneous increase in  $A_{\text{max}}$ ,  $g_{\text{m}}$ , and  $g_{\text{s}}$  (Flexas *et al.*, 2012). Recent studies have demonstrated that an increase in  $g_m$  has coincided with an increase in the amount of aquaporins after fertilization (Miyazawa et al., 2008b; Zhu et al., 2020). The biochemical limitation to photosynthesis was relatively low (16%) and the absolute contribution of this limitation to the decrease in  $A_{\rm max}$  under low nitrogen was much lower again (1.5%). This suggests that the limitations to photosynthesis resulting from low soil nitrogen are more attributable to CO<sub>2</sub> diffusion for Populus spp.

### Soil moisture

Although many studies showed a decline of  $g_m$  in response to soil water deficit (Flexas et al., 2009; Galle et al., 2009; Tosens et al., 2012a), it remains unclear if this limitation is happening within the mesophyll environment or occurs as a result of a stomatal limitation, which decreases intercellular  $CO_2(C_i)$ . Ma et al. (2021) reported that, across a broad range of species,  $g_m$  and  $g_{\rm s}$  decline concomitantly, which has the effect of keeping the  $g_{\rm m}/g_{\rm sc}$  ratio constant for all species and between well-watered and water-stressed plants, but with variation between plant functional types. We report similar  $g_m/g_{sc}$  ratios within our soil moisture dataset. However, reports in poplar have shown that this concomitant decline is not present all the time or within the full range of g<sub>m</sub> and g<sub>s</sub> values observed. Théroux Rancourt et al. (2015) showed that, in hybrid poplar,  $g_m$  remained unchanged (~0.3 mol m<sup>-2</sup> s<sup>-1</sup>) following soil drying until  $\Psi_{\text{leaf}} \approx -1.2$  MPa, after which  $g_{\rm m}$  decreased significantly. Our results showed that although  $g_m/g_s$  increased under water deficit conditions, stomatal conductance was, in absolute term, the most important limitation to  $A_{\text{max}}$ , as reported elsewhere (Cano *et al.*, 2013). In a trial on Quercus robur L. and Fraxinus angustifolia Vahl grown in the field, Grassi and Magnani (2005) reported a concomitant decrease of both  $g_s$  and  $g_m$  in a dry year ( $\Psi_{soil} \approx -1.7$  MPa), compared with a wetter year ( $\Psi_{soil} \approx -0.2$  MPa). In *P. tremula* L.,  $g_{\rm m}$  significantly declined when  $\Psi_{\rm leaf}$  of saplings dropped from -0.3 to -0.7 MPa due to applied osmotic stress (Tosens *et al.*, 2012*a*). Simultaneously, drought stress induced a decrease in SLA accompanied with an increase in the cell wall thickness and a decrease in the chloroplast surface area exposed to the intercellular air space per unit leaf area (Tosens *et al.*, 2012*a*). Other studies have shown that biochemical changes induced by drought stress could decrease  $CO_2$  diffusion to carboxylation sites in the chloroplast (Miyazawa *et al.*, 2008a).

Adaptation to the local environment might be a key driver of  $g_m$  variation among taxa, similarly to other morphophysiological traits. Interspecific and intraspecific differences in  $g_m$  from mesic versus xeric environments (*Quercus* spp. and *Eucalyptus* spp.) were reported by Zhou *et al.* (2014). Their study showed that  $g_m$ , as well as  $g_s$ ,  $V_{cmax}$ , and J of species from drier regions was less sensitive to water deficit, which maintains transpiration and photosynthesis activity at higher rates under drought, compared with species from the mesic environment. Marchi *et al.* (2008) observed that structural protection of mesophyll cells had a priority over functional efficiency of photochemical mechanisms in *Olea europaea* L. (evergreen sclerophyllous) but not in *Prunus persica* L. (deciduous broadleaf), depending on age-related variation in mesophyll anatomy.

### Conclusion and future directions

The present review shows that  $g_m$  in *Populus* spp. varies predictably along light gradients and that it responds to changes in soil moisture and nutrient availability, but is not affected by metal concentration and increasing atmospheric CO<sub>2</sub> concentration. Although metabolic processes noticeably influence the response of  $g_m$  to environmental changes, physical constraints through leaf development and ageing need to be considered in scaling photosynthesis from leaf to canopy, and in breeding programs for high WUE. Because fast-growing Populus spp. trees are important players in combating climate change, mitigating carbon emissions to some extent, comparisons of genotypes with different adaptations to changing environments and breeding for novel genotype-climate associations are urgently needed. This study shows that the variability of  $g_{\rm m}$  in different experimental conditions offers a potential indicator for improving Populus spp. productivity and resilience. However, more research is yet needed, also combined with anatomical studies, to better understand the sources of variation of CO<sub>2</sub> diffusion through the mesophyll and their consequences on carbon assimilation and growth.

Moreover, determination of the efficiency and optimal age for early selection of fast-growing poplar clones require an understanding of the genetic control and age-based genetic correlations for traits related to  $g_m$  and growth. For that, a detailed evaluation of the genotypic control of the variances and clonal heritability of  $g_m$  is needed. Finally, the identification of molecular bases of the regulation of  $g_m$  is necessary to further refine a multi-criteria early selection approach of poplar clones dedicated to the future forestry

capable of ensuring better productivity and increased resistance to environmental stresses (frost, drought, water logging, heavy metals, heat waves, etc.).

# Supplementary data

The following supplementary data are available at JXB online.

Table S1. Analysis of variance of the effect of different factors on photosynthesis-related traits.

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# **Author contributions**

RE and LB conceived the study, developed methodology, carried out data curation, formal analysis, investigation, data visualization, wrote original draft, and reviewed and edited the manuscript. MM contributed to data curation and visualization, and to reviewing and editing the manuscript. RT and UN contributed to investigation, methodology, and to reviewing and editing the manuscript. MM carried out funding acquisition and contributed to reviewing and editing the manuscript. RTS, GTR, TT, FR. SBG, MSL, and CC contributed to reviewing and editing the manuscript.

# **Conflict of interest**

The authors declare that there is no conflict of interest.

# Data availability

Soil moisture data from Théroux-Rancourt are available at Dryad Digital Repository (https://doi.org/10.5061/dryad.7sqv9s4s0); Benomar data are available at Dryad Digital Repository (https://doi.org/10.5061/ dryad.9cnp5hqhp);Tognetti data are available at Dryad Digital Repository (https://doi.org/10.5061/dryad.w3r2280qq).All other datasets generated for this study are available from the corresponding author upon request.

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