Leaf temperature and its dependence on atmospheric CO2 and leaf size

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There is general concern that the rapid increase in atmospheric CO2 concentration will lead to reduced stomatal conductance and subsequent increases in leaf temperature. Such an increase in leaf temperature is expected to adversely impact a plethora of processes connected to leaf metabolism and microbial/fungal communities on leaves. A model is proposed that combines the leaf energy balance with leaf gas exchange and photosynthesis to explore such issues. The model represents a hybrid ecological/physiological approach described by systems of equations based on steady-state leaf-gas exchange theories and leaf energy/radiation balance, equilibrium thermodynamics within the leaf, stomatal data, and atmospheric CO2 concentration. The model allows separating air from leaf temperatures thereby permitting exploration of the dependence of leaf cooling or heating for any combination of environmental conditions (e.g., wind velocity, atmospheric humidity, and atmospheric CO2 level), anatomic leaf properties (e.g., leaf size), and physiologic quantities (e.g., assimilation rate and transpiration rate). The model permits to distinguish whether leaf cooling or heating is to be expected if these parameters are varied. Based on model calculations, it is shown that leaf temperature is far more impacted by leaf size or wind speed than reduction in stomatal conductance caused by elevated atmospheric CO2. The model results are consistent with measurements of leaf cooling and heating.

KEYWORDS
atmospheric CO2, gas exchange, leaf cooling, leaf energy exchange, leaf heating, leaf size

1 INTRODUCTION

Plants are affected by changes in atmospheric CO2 levels directly and indirectly. Being the substrate of photosynthesis, atmospheric CO2 affects plants directly through leaf gas exchange that is driven by the gas concentration difference between leaf interior and exterior. A rise in exterior CO2 concentration, termed $C_a$, throughout the rest of the text, does not necessarily lead to a proportional rise in photosynthesis, because the assimilation rate is affected by temperature, saturation kinetics of the involved biochemical apparatus, and leaf conductance (which are partially interdependent). The overall leaf conductance, $g$, offering the diffusional pathway connecting CO2 source (atmosphere) and sink (mesophyll), consists of a mesophyll conductance, a stomatal conductance $g_s$, and an aerodynamic conductance. By and large, $g_s$ is presumed to be the most restrictive though the remaining two conductances can be restrictive under certain conditions, such as low wind speed or soil moisture stress states. It can be generally observed that $g_s$ responds inversely to $C_a$ to match atmospheric supply with sink demand, by decreasing stomatal aperture and/or number of stomata (Ainsworth and Rogers, 2007; De Boer et al., 2011; Franks et al., 2013, 2014; Lambersma et al., 2011; Leakey et al., 2009; Wagner et al., 1996). Because the stomatal pathway allows water vapour to escape
from the leaf interior, photosynthesis and transpiration are coupled through \( g_s \). Stomatal conductance \( g_s \) is regulated by two mechanisms: the guard cells that can vary stomatal aperture on short time scales (of the order of minutes to tens of minutes), whereas the maximum value of \( g_s \) is limited by stomatal density and size and thus changes on much longer time scales (e.g., from one growing season to the next and/or even longer time scales defined by evolutionary adaptation).

Additionally, increases in global atmospheric CO\(_2\) concentrations influence photosynthesis indirectly, via increases in global air temperatures. Temperature controls numerous leaf metabolic processes thereby exerting controls on photosynthesis. Moreover, both leaf assimilation rate, \( A \), and leaf transpiration rate, \( E \), depend on temperature: A shows an optimum value, \( A_{\text{opt}} \), at the related temperature \( T_{\text{opt}} \), whereas the saturation vapour pressure increase monotonically with increasing temperature as described by the Clausius–Clapeyron equation. Processes at the leaf-level are, however, not dependent on air temperature \( T_a \) but on leaf temperature \( T_l \). The \( T_l \) can be different from \( T_a \) due to several processes that exchange energy between the leaf surface and the atmosphere (De Boeck, Van De Velde, De Groote, and Nijs, 2016; Jones, 2013; Smith, 1978; Nobel, 2005). Leaf energy exchange is affected by gas exchange because transpiration within the leaf consumes heat that is then "exported" to the atmosphere as latent heat (causing "evaporative cooling"). This process contributes substantially to the energy budget of the leaf (Gates, 1968; Jones, 1999, 2013).

Under elevated \( C_a \) (and otherwise identical conditions), a decrease in \( g_s \) is thus expected to reduce evaporative cooling thereby enhancing the risk for plants to reach a critically high \( T_l \) when \( T_a \) is also high. Rising \( T_l \) due to elevated \( C_a \) in the future was suggested to be particularly detrimental for tropical vegetation (De Boeck, Van De Velde, De Groote, and Nijs, 2016). Higher \( T_l \) due to less evaporative cooling would also affect not only plants. For example, for insects living close to the leaf surface, any changes in average \( T_l \) would mean a substantial altering of their typical habitat conditions (Pincebourde and Casas, 2006; Pincebourde and Woods, 2012; Zavala, Nabity, and Delucia, 2013).

The importance of evaporative cooling depends on leaf size because the thickness of the leaf boundary layer increases with the size of the object (Schuepp, 1993). Increases in boundary layer thickness particularly impedes conductive and convective cooling (heat transfer by heat conduction and air currents, respectively) as well as gas exchange. Under identical wind conditions, small leaves experience a thinner boundary layer than larger leaves. This difference is due to the fact that as air encounters a leaf surface, a boundary layer is initiated at this intersection point and begins to develop and grow on the leaf surface along the local wind direction. Smaller leaf dimensions prevent this growing boundary layer to become too thick resulting in average boundary layer thicknesses that are smaller when compared to larger leaves for the same incident wind speed. The thinner boundary layer makes conductive and convective cooling more effective. Hence, small leaves, in effect, are less dependent on transpiration for a cooling mechanism (Gates, 1968; Huang, Chu, Hsieh, Palmroth, and Katul, 2015). Elevated \( C_a \), particularly high levels of 1,000 ppm and more, are anticipated for the end of this century in worst case scenarios (Nakicenovic et al., 2000), appears especially problematic for large leaves with respect to leaf overheating. In fact, high \( C_a \) conditions of the past were suggested to have suppressed the evolution of megaphylls during early stages of land plant evolution (Beerling, Osborne, and Chaloner, 2001). The absolute area of a leaf, however, is less crucial for boundary layer thickness than leaf shape along the predominant wind direction. This is because the thickness of the boundary layer does not depend on the total area but rather on the "characteristic dimension", \( l_C \) (Nobel, 2005; Vogel, 2009), over which the boundary layer develops. Leaves with the same area can thus show different heat exchange characteristics depending on leaf shape (Roth-Nebelsick, 2001; Vogel, 1968). For leaves with \( l_C \leq 2cm \), heat transfer is dominated by convection (Gates, 1968).

The interrelation between \( g_s, A, E, T_w, T_l, l_C \), and wind speed, \( v_w \), are complex and mutually intertwined, leading to counteracting and sometimes counter-intuitive effects. As an illustration, consider the following situation: Supposing that \( T_l \) and vapour pressure difference (VPD) between the leaf interior (assumed saturated) and the atmosphere change slowly. A sudden drop in \( g_s \) will lead to a lower transpiration rate \( E \), meaning that fewer water molecules are vaporized, producing less evaporative cooling. Consequently, the leaf temperature \( T_l \) increases, leading to a higher saturation vapour pressure of water molecules within the leaf. This, in turn, leads to a higher VPD between leaf interior and the atmosphere (assuming that atmospheric temperature and humidity remain constant), thereby enhancing transpiration. Since transpiration rate \( E \) is proportional to the product of leaf conductance \( g_s \) and to VPD, it is not possible to predict which of the two rivaling effects—a decrease in \( g_s \) due to stomatal closure or an increase in VPD because of elevated leaf temperature—dominates in the end, unless a quantitative theory encompassing both effects is at hand. Even in this example, another complication arises because \( g_s \) is related to assimilation rate \( A \), which in turn varies with \( T_l \), as already mentioned.

Finally, variation of \( v_w \) can generate other counter-intuitive effects: Increasing wind speed reduces the thickness of the boundary layer, thereby increasing leaf conductance \( g_s \). Assuming as above that \( T_l \) and VPD react sluggishly, transpiration rate increases whereupon evaporative cooling is intensified. Thus, \( T_l \) begins to drop, water vapour saturation pressure within the leaf and VPD follow, and \( E \) decreases somewhat. Again, \( E \) is affected by two competing mechanisms, and a quantitative theory is required to assess the net effect. Observations reviewed elsewhere (Grace, 1988; Huang et al., 2015) show that increased wind speed can actually decrease transpiration rate for certain environmental conditions. In fact, higher wind speed can increase water use efficiency (WUE = \( A/E \)) just because of stomatal closure under these circumstances (Schymanski & Or, 2016).

Taking all these relations together, it appears difficult to predict how changing \( C_a \) might affect \( T_l \) in isolation. Because \( T_l \) is an essential ecophysiological quantity, it is, however, desirable to evaluate possible mutual interrelations with \( C_a \) and such an analysis cannot ignore the role of \( I_C \). These issues motivate the development of the mathematical model to be described here. In particular, the focus is on the effect of changing \( C_a \) on \( T_l \) using a newly proposed model that connects
optimal leaf-gas exchange and leaf heat exchange by using explicit interrelations between $g$, $A$, $E$, $T_a$, $T_s$, $l_c$, and $v_w$.

To achieve this target model while maintaining tractability, the so-called “reduced order model” of leaf-gas exchange derived elsewhere (Konrad, Katul, Roth-Nebelsick, & Grein, 2017) is coupled to established formulations describing the simultaneous leaf energy and radiation budgets (Jones, 2013; Nobel, 2005). The modelled parameters are $g$, $A$, $E$, and $T_s$ explored here under light saturation conditions and imposed environmental conditions prescribed in the atmosphere outside the leaf boundary layer (labelled as “free stream variables”). These free stream variables include $v_w$, $C_a$, and $T_s$ for different leaf sizes $l_c$. The model is tested using gas exchange measurements and literature data.

## 2 | THE MODEL

The target model combines the leaf energy balance with leaf gas exchange for CO$_2$ and water vapour as well as biochemical demand for CO$_2$. Hence, the target model parameters are those associated with the aforementioned processes. For tractability, a number of simplifications are necessary in the target model. Steady-state conditions are assumed so that the atmospheric supply and biochemical demand for CO$_2$ are in balance. Also, the leaf interior is assumed to be saturated so that within the sub-stomatal cavity, the relative humidity of air is 100%. This assumption is reasonable provided leaf water potential does not drop below $-5$ MPa (Farquhar & Raschke, 1978, but see Cemusak et al., 2018). The leaf interior is also assumed to be in thermal equilibrium with the leaf surface so that the leaf surface temperature can be used to approximate the saturation and actual vapour pressure within the sub-stomatal cavity using the Clausius–Clapeyron equilibrium equation. Minor temperature differences between abaxial and adaxial leaf side can arise (Rockwell, Holbrook, & Stroock, 2014) that can, however, be neglected for the purposes here. Furthermore, mesophyll resistance is not particularly considered in the model though it can be added if known. It is also assumed that the conditions just outside the leaf boundary layer or the free stream are quasi-stationary and spatially uniform so that the main spatial gradients are those associated with processes within the leaf boundary layer. Free convection is not included, which becomes relevant at extremely low wind speeds. The model also considers leaf-level processes in isolation and hence caution must be exercised when extrapolating its conclusions to larger spatial scales. Any physiological differences between sunlit and shaded leaves are assumed to be entirely captured by parameters in the biochemical demand equation for C$_3$ plants. Perhaps most restrictive is the assumption that $\kappa = C_i/C_a$ is constant and does not change with environmental conditions (e.g., VPD), where $C_i$ is the internal CO$_2$ concentration. The model does not consider pro-

### 2.1 | The gas exchange formulation

The gas exchange formulation is centred around the carbon economy of the leaf: The atmospheric supply of CO$_2$ to the leaf is given by Fick’s diffusion law, or its approximate integrated version, as

$$A = g(C_a - C_i). \quad (1)$$

When atmospheric supply is matched to biochemical demand (i.e., when every CO$_2$ molecule that collides with the leaf surface and enters through the guard cells into the sub-stomatal cavity is eventually assimilated), then $A$ can also be described by the Farquhar photosynthesis model for C$_3$ plants given as (Farquhar, Von Caemmerer, and Berry, 1980, 2001)

$$A = \frac{C_i - \Gamma}{C_i + K} - R_d. \quad (2)$$

thus offering another relation between $A$ and $C_i$ at a prescribed $C_a$. The parameters associated with the photosynthetic model in Equation (2) are as follows: $q$ is carboxylation limited by Rubisco or RuBP regeneration rate, $K$ is a parameter containing Michaelis–Menten constants of carboxylation and oxygenation, $\Gamma$ is the CO$_2$ compensation point, and $R_d$ is the mitochondrial respiration rate. The dependence of the biochemical parameters $q$, $\Gamma$, $K$, and $R_d$ on $T_s$ is based on biochemical considerations derived elsewhere (Bernacchi, Pimentel, and Long, 2003) and are not repeated here. When only a balance between demand and supply for CO$_2$ is imposed, the outcome of this balance results in one equation with two unknowns: $g$ and $C_i$. Hence, the gas exchange formulation remains mathematically "unclosed" and requires additional equations. Denoting the ratio of CO$_2$ concentration within the leaf and the atmosphere as

$$\kappa = \frac{C_i}{C_a}. \quad (3)$$

Equations (3) and (2) can be used to eliminate $C_i$ and $A$ from Equation (1) to obtain an explicit expression for $g$ given as

$$g = \frac{q(xC_a - \Gamma)}{(xC_a + K)(1 - x)C_a} - \frac{R_d}{(1 - x)C_a}. \quad (4)$$

A number of points can be made about the mathematical character of this expression when noting that $\kappa$ is a constrained quantity robust to changes in $C_a$ and to a leading order, may be treated as a constant. Some support for the insensitivity of $\kappa$ to $C_a$ has been documented in a number of studies reviewed elsewhere (Katul, Ellsworth, and Lai, 2000; Katul, Manzoni, Palmroth, and Oren, 2010). To begin with, increasing $C_a$ while maintaining $\kappa$ approximately constant means that the "reduction term" that includes $R_d$ (the second term on the right-hand side of (4)) becomes negligible and has minor impact on $g$ except for small $C_a$. More significant is the first term on the right-hand side of (4) and its dependence on $C_a$. While the numerator is
linear in \( C_a \), the denominator is quadratic in \( C_a \), meaning that increasing \( C_a \) must lead to a reduction in \( g \) for high \( C_a \) (and conversely for very low \( C_a \)). Employing expression (3), the assimilation rate becomes

\[
A = \frac{k C_a - \Gamma}{\alpha C_2 + K} - R_d. \tag{5}
\]

Different from the \( g \) expression in (4), the numerator and denominator of the right-hand side of (5) are both linear in \( C_a \), meaning that leaf photosynthesis (and by extrapolation the capacity of the biosphere to absorb atmospheric CO\(_2\) when maximal leaf area is attained) are expected to “saturate” at \( q - R_d \) when \( C_a \) is very large. Fick’s Law of diffusion for water vapour also leads to

\[
E = g \alpha (w_l - w_a) = \frac{q (\kappa C_a - \Gamma) - R_d (\kappa C_a + K)}{(\kappa C_a + K)(1 - \kappa) C_a} \alpha (w_l - w_a), \tag{6}
\]

where \( \alpha = D_{h2o}/D_{CO2} \) denotes the ratio of the molecular diffusional coefficients of water vapour and CO\(_2\) in air and \( w_l \) is the water vapour concentration within the leaf. The model featured in (1) through (6) assumes that all kinetic parameters as well as \( w_l \) depend on \( T_l \) and not the commonly available (or projected) \( T_p \). Figure 1a, b, and c illustrates the results from Equations (4), (5) and (6) for \( g, A \) and \( E \) as functions of \( C_a \) and \( T_l \) momentarily assuming isotothermal conditions (\( T_l = T_s \)). The calculations assume that \( \kappa \) is constant determined from isotopic measurements though it can be allowed to vary. To accommodate such variations in \( \kappa \), an additional equation (and model assumptions) are needed. One common approach to arrive at such an expression is to assume that leaves maximize \( A \) subject to some constraints such as water availability per unit leaf area in the rooting zone. Such optimization theories show that \( \kappa \) weakly varies with VPD with a sub-unity exponent and with several schemes predicting approximate scaling relations of the form \( k \propto \sqrt{\text{VPD}} \) (Hari, Mäkela, Berninger, and Pohja, 1999; Katul, Palmroth, and Oren, 2009; Katul et al., 2010; Medlyn et al., 2011; Prentice, Dong, Gleason, Maire, and Wright, 2014). This scaling appears to be robust to the precise assumptions made about the constraints imposed on the optimization as discussed elsewhere (Dewar et al., 2018). As expected from such analysis, increasing \( C_a \) increases \( A \) at all temperatures (in a non-linear manner) though the interplay between \( C_a \) and \( T_p \) is far more complex for the remaining two variables (\( g \) and \( E \)). To allow for differences between \( T_l \) and \( T_p \), additional expressions are needed to describe the combined leaf energy and radiation balances.

### 2.2 The leaf energy exchange equations

To allow for leaf and air temperature differences in the target model, a conventional leaf energy balance formulation (Nobel, 2005; Jones, 2013) is introduced to express \( T_l \) in terms of \( T_p \) and \( g \). For steady-state conditions, the leaf energy balance equation is given by

\[
\text{energy into leaf} = \text{energy out of leaf} + \left[ \text{energy consumed by leaf metabolism} \right]. \tag{7}
\]

Since the energy consumed by leaf metabolism is negligible compared to the amounts of energy flowing into and out of the leaf, it is ignored in what follows. Also, it is assumed that leaves have high thermal inertia (or volumetric heat capacity), so that any imbalance between the left hand side and the right hand side in unsteady conditions do not alter \( T_l \) appreciably over the time scales considered here (i.e., changes in the environmental conditions external to the leaf). Keeping only the terms that quantify the energy exchange processes leads to (Nobel, 2005)

\[
a(1 + r)S_c \frac{1}{h_f} \sin \gamma + a_r \sigma \left( T_{aw}^4 + T_{sky}^4 \right) \\
\approx 2c_r \sigma \left( T_l^4 - T_s^4 \right) + \frac{2K_w}{d_{ist}}(T_l - T_s) + 2H_{vap}[h_l(T_l) - w_3]d_3(T_l). \tag{8}
\]

The terms on the left-hand side quantify the absorption of radiation coming directly from the sun (shortwave radiation), from the closer surroundings (such as other plants) and of diffuse radiation from the sky (longwave radiation). The right-hand side quantifies radiation emitted by the leaf (first term) and heat exchange via conduction and convection (second term) and evaporative cooling by transpiration. Notice that the last term on the right-hand side can be written, due to (6), as \( H_{vap}E \), where \( H_{vap} \) is the latent heat of vaporization. Table 1 presents the definitions and typical values of the other parameters in Equation (8). The conduction/convection term depends on the thickness \( d_{ist} \) of the laminar boundary layer attached to the leaf surface and on the free stream wind velocity \( v_w \) as well as the characteristic leaf length \( l_c \). In principle, \( v_w \) experiences turbulent fluctuations, and it is often replaced by its time-averaged value. In this case, the turbulent intensity can also play a role in determining \( d_{ist} \) because the laminar boundary layer may be intermittently disturbed by the outer turbulent state and may not attain its full steady-state value. However, such effects are usually absorbed in empirical constants of expressions such as the formula of Nobel, 2005 that is suitable for flat surfaces, such as angiosperm leaves and is given by

\[
d_{ist} = 4 \times 10^{-3} (m/\sqrt{s}) \left( \frac{l_c}{v_w} \right). \tag{9}
\]

(m and \( s \) denote the units meter and second, respectively). As expected, \( d_{ist} \) increases with increasing \( l_c \) and is reduced with increasing \( v_w \). As earlier noted, the leaf internal humidity \( w_i \) close to its saturation value \( w_{sat}(T_l) \) that only depends on leaf temperature \( T_l \). In a closed system at thermal equilibrium, the Clausius–Clapeyron equation (Reif, 1974) may be used and it is given as

\[
w_i(T_l) = w_{sat}(T_l) = \frac{u}{T_l} e^{-\frac{\Gamma}{T_l^2}}. \tag{10}
\]
**TABLE 1**  The model parameters together with their dimensions

| Quantity [unit] | Explanation | Value |
|-----------------|-------------|-------|
| **Physiologic parameters** | | |
| $A$ [μmol/m²/s] | Assimilation rate | Calculated |
| $g$ [m/s] | Leaf conductance | Calculated |
| $E$ [mmol/m²/s] | Transpiration rate | Calculated |
| $C_l$ [mol/m³] | Leaf internal CO₂ | Calculated |
| $T_l$ [°C] | Leaf temperature | Calculated |
| $w_l$ [mol/m³] | Leaf internal humidity | ≈ saturated |
| $l_c$ [mm] | Characteristic leaf length | Varied |
| $κ$ [-] | $κ = C_l/C_a$ | 0.768 |
| $T_{l,c}$ [°C] | Temperature separating the regimes of leaf warming and leaf cooling | Calculated |
| $T_{l,∞}$ [°C] | Special case of temperature $T_{l,c}$ for vanishing transpiration | Calculated |
| **Environmental parameters** | | |
| $C_a$ [mol/m³] | Atmospheric CO₂ | Varied |
| $T_a$ [°C] | Air temperature | Varied |
| $w_a$ [mol/m³] | Leaf external humidity | Varied |
| $w_{sat}$ [mol/m³] | Saturation value of humidity (cf. expression (10)) | Calculated |
| $w_{rel}$ | Relative atmospheric humidity, $w_{rel} = w_a/w_{sat}$ | Varied |
| $v_w$ [m/s] | Wind velocity | Varied |
| $d_b$ [mm] | Thickness of boundary layer (cf. expression (9)) | Calculated |
| $S_0$ [J/m²/s] | Solar constant | 1.366 |
| $σ$ [J/m²/s/K⁴] | Stefan-Boltzmann constant | $5.67 \times 10^{-8}$ |
| $K_a$ [J/m²/K] | Coefficient of thermal conductivity of air at 20°C | $2.55 \times 10^{-2}$ |
| $H_{vap}$ [J/mol] | Vapourisation heat of water | $44.1 \times 10^3$ |
| $γ$ | Angle of sun above horizon | 45° |
| $τ_a$ [-] | Atmospheric transmittance for moderate clear sky at moderate elevation | ≈ 0.7 |
| $α$ [-] | Absorptance of leaf for global radiation | ≈ 0.60 |
| $r$ [-] | Reflectance of the surroundings for global radiation | ≈ 0.20 |
| $α_{IR}$ [-] | Leaf absorptivity for infrared radiation | ≈ 0.96 |
| $ε_{IR}$ [-] | Leaf emissivity for infrared radiation | ≈ 0.96 |
| $T_{surf}$ [°C] | Temperature of the surroundings | ≈ 15 |
| $T_{sky}$ [°C] | Radiation temperature of the clear sky | ≈ −20 |
| $D_{CO₂}$ [m²/s] | Diffusion constant of CO₂ at $T = 25°C$ | $1.55 \times 10^{-5}$ |
| $D_{H₂O}$ [m²/s] | Diffusion constant of water vapour at $T = 25°C$ | $2.49 \times 10^{-5}$ |
| $σ$ | $σ = D_{H₂O}/D_{CO₂}$ | 1.6 |

**Biochemical parameters from Quercus petraea**

| Quantity [μmol/m²/s] | Explanation | Value |
|-----------------------|-------------|-------|
| $q$ | Carboxylation limited by Rubisco or RuBP at $T = 25°C$ | 58 |
| $K$ [μmol/m³] | Contains Michaelis-Menten constants | 6,926 |
| $Γ$ [μmol/m³] | CO₂ compensation point | 1,584 |
| $R_{o}$ [μmol/m²/s] | Mitochondrial respiration rate | 1.0 |

Note: Specific values of parameters designated as "varied" are given in the figure captions.
with \( u = 2.035 \times 10^{10} \text{mol/m}^3 \) and \( v = 5.306 (T_l \text{ in Kelvin}) \). This equation must be viewed as approximate because the sub-stomatal cavity is not closed to mass exchange (i.e., \( E > 0 \)). However, even for such a system, corrections to the Clausius–Clapeyron equation are small when steady-state conditions are attained. Notice that the radiation and evaporation terms in (8) are always positive, because these two mechanisms only allow the emission of energy from the leaf, whereas the processes of conduction/convection can—depending on the sign of the difference \( T_l - T_a \)—decrease or increase the leaf’s energy content.

Figures 2 and 3 illustrate the various components of the leaf energy balance Equation (8) as they change with \( T_a \): The black line denotes the left-hand side, that is, the radiative energy absorbed by the leaf. The three coloured curves represent the right-hand side, namely emitted radiation (red), conduction/convection processes (green), and evaporation (blue). Their sum equals—for any given \( T_a \)—
the black line. The conduction/convection is the only term that can be bidirectional and is one reason why results from leaf energy balance calculations can be counter-intuitive on some occasions. Comparison of the sub-figures of Figures 2 and 3 suggests that radiation is almost unaffected by variations of air humidity and leaf size, whereas conduction/convection and evaporation are strongly influenced, which is what one would expect from the physics of this situation.

2.3 | The target model: Combining the leaf gas and energy exchange formulations

If the quantities appearing on the left hand side of Equation (8), the species specific photosynthetic parameters defining the Farquhar biochemical demand model, and the free stream state parameters \( C_a \), \( v_w \), \( T_a \), and \( w_s \) (or, alternatively, VPD = \( w_l - w_s \)) are provided, Equations (4) and (8) (together with (9) and (10)) can be viewed as a system of two algebraic equations for the two unknowns \( g \) and \( T_l \).

In principle, the solution procedure is straightforward: Insertion of (4) into (8) (to eliminate \( g \)) produces an intricate equation for \( T_l \) that admits no closed-form expression. However, the solution simplifies considerably upon the substitution of a canonical form

\[
T_l = T_a (1 + \xi),
\]

followed by a Taylor series expansion with respect to \( \xi \) around \( \xi = 0 \). Since the difference between \( T_l \) and \( T_a \) is small with respect to \( T_a \) \( \text{(in K)} \), (implying \( \xi \ll 1 \)), it is justified to discard from the expansion higher order terms beyond quadratic \( (\xi^2) \). Solving the resulting quadratic equation for \( \xi \) and inserting the outcome into (11) yields \( T_l \). Insertion of \( T_l \) in expression (4) for \( g \) completes the solution process. It is to be noted that \( 1/g \), the reciprocal of leaf conductance \( g \), combines in series both stomatal and aerodynamic resistances according to

\[
\frac{1}{g} = \frac{1}{g_s} + \frac{1}{g_{bl}}.
\]

Because \( g_{bl} = \frac{D_{CO2}}{a_{BL}} \), it is possible to use (4) to separate \( g_s \) from \( g \). Figure 4 features the resistances \( 1/g_s \) (solid lines) and \( 1/g_{bl} \) (dashed lines) that are related to stomatal conductance and boundary layer conductance. According to (12), the sum of related curves equals the

![FIGURE 2](https://example.com/figure2.png)

The individual components impacting the steady-state leaf energy balance as a function of air temperature \( T_a \) and characteristic leaf length \( l_c \). The black lines indicate the total heat absorbed by the leaf, and the coloured lines indicate the mechanisms by which the heat is distributed: radiation (red lines), conduction, and convection (green lines) and evaporation (blue lines). Input values: wind speed \( v_w = 1 \text{ m/s} \), \( C_a = 400 \text{ \mu mol/mol} \). Other input values are as given in Table 1 (biochemical parameters were borrowed from *Quercus petraea*). [Colour figure can be viewed at wileyonlinelibrary.com]
total leaf resistance $1/g$, the reciprocal of leaf conductance. As shown in Figure 4, for small leaves and high wind speeds (Figure 4d), stomatal resistance is much higher than boundary layer resistance; if leaf size increases and wind speed decreases, both resistances become comparable (Figure 4c). The behaviour of the boundary layer resistance is a direct consequence of expression (9) for the thickness of the boundary layer whereas stomatal resistance includes all the radiation, energy, and leaf gas exchange processes.

3 | APPLICATION OF THE MODEL TO LEAF HEATING AND COOLING

The target model is now applied to *Quercus petraea* to illustrate the mechanisms of leaf heating and cooling. This species is chosen because of its wide-spread distribution (and economical value) within Europe, and its well-studied physiological and radiative properties. The *Q. petraea* biochemical parameters used are given in Table 1. Specifically, the effect of complementing the leaf gas exchange equations of Section 2.1 with the energy exchange equations of Section 2.2 is depicted by Figure 1. The upper row of sub-figures follows from the gas exchange equations alone (ignoring leaf energy exchange by setting $T_l = T_a$), the lower row of sub-figures results from coupling both models thereby distinguishing $T_l$ from $T_a$ with all kinetic parameters driven by $T_l$. If leaf energy exchange is taken into account, the maxima of the curves representing $g$ and $A$ are shifted to higher values of $T_a$, but the general structure of the family of curves—especially the highly different $g$- and $A$-values for identical $T_a$- but different $C_a$-values—remains the same. Much more affected by the inclusion of energy exchange is the transpiration rate $E$ that shows (a) a much smaller gradient with respect to $T_a$ and (b) a lesser dependence on $C_a$.

As already stated, the photosynthetic parameters $q$, $\Gamma$, $K$, and $R_d$ of the model of Farquhar et al., 1980; see expression (2)) depend on $T_l$ according to the relations of Bernacchi et al. (2003). The interplay of these individual dependencies results in the overall temperature dependence of the assimilation rate $A$ shown in Figure 1b, e, and h that exhibits a maximum of $A$ at some ($C_a$-dependent) optimum temperature $T_{opt}$. Obviously, elevated $C_a$ shifts $T_{opt}$ towards higher values. This effect is also observed experimentally as discussed elsewhere (Duursma et al., 2014; Eamus, Duff, and Berryman, 1995; Ghannoum...
et al., 2010; Quentin, Crous, Barton, and Ellsworth, 2015; Reef et al., 2016). Notice that the maxima of \(g(T_a)\) and \(A(T_a)\) are located at the same temperature, \(T_{opt}\). This is not a coincidence, and it directly follows from Equations (1) and (3) that imply

\[
g = A / (1 - \kappa C_a).
\]

Since it is assumed that \(\kappa = \text{const.}\) and since \(C_a\) does not depend on local leaf or air temperature here, \(g\) inherits its temperature dependence from \(A\) described above and depicted in Figure 1.

### 3.1 | Mechanisms of leaf heating and cooling

Figures 5 and 6 are now used as a starting point to unravel the interplay of the various contributions to leaf heating and cooling. The coloured curves feature the dependency of leaf temperature on air temperature for different leaf sizes and several values of free stream air relative humidity. The diagonal thin black lines indicate equality of leaf and air temperature, that is, \(T_l = T_a\). Thus, segments of the coloured curves located above the thin black lines indicate leaf heating, coloured segments below these lines indicate leaf cooling. The dashed black lines result if evaporation is inhibited in (8) by setting \(g = 0\).

The coloured lines in Figures 5 and 6 are related to different atmospheric \(C_a\) levels. They originate when stomata are open and transpiration, causing evaporative cooling, contributes to the energy exchange between leaf and air. That is, the transpiring leaves make use of all three energy exchange mechanisms to regulate leaf temperature. Since the mechanisms of conduction and convection can transport heat into and away from the leaf, they can cool or heat the leaf depending on the sign of \(T_l - T_a\). Evaporation, which is based on heat consuming vaporization, in contrast, can transport heat only away from the leaf, that is, cooling it. Figures 5 and 6 suggest that evaporative cooling can have a considerable impact on the temperature \(T_i\), \(i(C_a)\), which is defined by the intersection of the coloured lines and

\[
\text{FIGURE 4} \quad \text{Impact of air temperature } T_a, \text{atmospheric CO}_2, \text{and characteristic leaf length } l_c \text{ on stomatal resistance } 1/g_s \text{ (solid lines) and boundary layer resistance } 1/g_{bl} \text{ (dashed lines). The sum of these curves sets the total leaf resistance related to leaf conductance and amounts to } 1/g = 1/g_s + 1/g_{bl}. \text{ Free stream values: relative humidity } w_{rel} = 0.6. \text{ Other input values are as given in Table 1 (biochemical parameters were borrowed from Quercus petraea) [Colour figure can be viewed at wileyonlinelibrary.com]}
\]
the thin black lines, separating the regimes of leaf warming and leaf cooling: If, for instance, $C_a$ is in the range $180 \mu\text{mol/mol} \ldots 1000 \mu\text{mol/mol}$ and characteristic leaf length and humidity have the values $l_c = 30 \text{ mm}$ and $w_{\text{rel}} = 0.6$, the related $T_{i,c}$-values are in the range $21.6^\circ\text{C} \ldots 23.9^\circ\text{C}$. This is about 15 K lower than $T_{i,\infty} \approx 37.9^\circ\text{C}$, the equivalent of $T_{i,c}$ for the case of vanishing transpiration derived from (4) for $C_a \rightarrow \infty$ (explaining the nomenclature). This case is illustrated by Figures 5e and 6e.

Inspection of Figures 5 and 6 also reveals how $T_{i,c}$ depends on air humidity, leaf size, and wind speed:

1. The horizontal sequences a–c, d–f, and g–i in Figures 5 and 6 show that $T_{i,c}$ reacts sensitively to increasing air humidity: If $w_{\text{rel}}$ is increased from the value 0.3 to 0.9, $T_{i,c}$ moves from about $22^\circ\text{C}$ to $30.6^\circ\text{C} \ldots 32.9^\circ\text{C}$ (for $l_c = 30 \text{ mm}$). The reason is obvious: Increasing $w_{\text{rel}}$ lowers VPD that impedes transpiration and evaporative cooling.

2. Increasing leaf size $l_c$ (vertical sequences a–g, b–h, and c–i in Figures 5 and 6) does not change the temperature $T_{i,c}$. However, the amount of both leaf warming and cooling increases with $l_c$ for a given air temperature $T_a$. In Figures 5 and 6, this is clearly visible at the points where both the coloured lines

---

**FIGURE 5** Impact of air temperature $T_a$, atmospheric CO$_2$, air humidity $w_{\text{rel}}$ and characteristic leaf length $l_c$ on leaf temperature $T_l$. The diagonal thin black lines indicate equality of leaf and air temperature, that is, $T_l = T_a$. Thus, segments of the coloured curves located above the thin black lines indicate leaf heating, and coloured segments below this line indicate leaf cooling. The dashed black lines result if evaporation is inhibited in (8) by setting $g = 0$. The free stream values: wind speed $v_w = 1 \text{ m/s}$. Other input values are as given in Table 1 (biochemical parameters were borrowed from *Quercus petraea*) [Colour figure can be viewed at wileyonlinelibrary.com]
and the dashed black line (representing the special case \( g = 0 \) of vanishing transpiration) intersect the \( T_\ell \)-axis.

3. Comparison of Figures 5 and 6 shows that a reduction of wind speed does not change the temperature \( T_{\ell,c} \) that separates the regimes of leaf warming and leaf cooling. It enhances, however, both leaf warming and cooling.

The main results of this section are not limited to \( Q. \) petraea. In the Appendix, we outline how they can be derived from the species-independent model equations of Section 2.

3.2 Effect of atmospheric CO2 on leaf temperature

In Figures 5 and 6, the curves related to different \( C_a \)-values lie almost on top of one another (with the exception of the green curve, related to subambient \( C_a \), which deviates somewhat from the general trend), indicating that the impact of \( C_a \) on \( T_\ell \) is nearly irrelevant. This can be understood from the structure of Equation (8). It depends on \( C_a \) solely via the term \( g \) (by relation (4)), whereas \( T_{\ell} \) appears in several terms of which the emission term (containing \( T_4^\ell \)) and the expression \( w_l(T_\ell) \)
FIGURE 7  Leaf conductance $g$ (left column), assimilation rate $A$ (centre column), and transpiration rate $E$ (right column) as a function of characteristic leaf length $l_c$ for different values of air temperature $T_a =$ (5°C, 15°C, 25°C, 35°C; from top row to bottom row). Wind velocity is $v_w = 1$ m/s, relative atmospheric humidity is $w_{rel} = 0.6$. Other input values are as given in Table 1 (biochemical parameters were borrowed from *Quercus petraea*). [Colour figure can be viewed at wileyonlinelibrary.com]
FIGURE 8  Leaf conductance $g$ (left column), assimilation rate $A$ (centre column), and transpiration rate $E$ (right column) as a function of characteristic leaf length $l_c$ for different values of air temperature $T_a = (5^\circ C, 15^\circ C, 25^\circ C, 35^\circ C)$ (from top row to bottom row). Wind velocity is $v_w = 0.1 \text{ m/s}$, relative atmospheric humidity is $w_{rel} = 0.6$. Other input values are as given in Table 1 (biochemical parameters were borrowed from Quercus petraea) [Colour figure can be viewed at wileyonlinelibrary.com]

(containing an exponential involving $T_e$, according to the Clausius-Clapeyron Equation (10)) are especially notable: The fourth power and the exponential effectuate that even large variations in $C_a$ can be balanced by small variations in $T_e$, provided that all other variables are kept constant and Equation (8) is to be valid.)
Leaf temperature $T_l$ as function of characteristic leaf length $l_c$ for various combinations of air temperature $T_a = (5\, ^\circ C, 15\, ^\circ C, 25\, ^\circ C, 35\, ^\circ C)$ and relative atmospheric humidity $w_{rel} = (0.3, 0.6, \text{and } 0.9)$. Wind velocity is $v_w = 1\, m/s$. Other input values are as given in Table 1 (biochemical parameters were borrowed from Quercus petraea) [Colour figure can be viewed at wileyonlinelibrary.com]
FIGURE 10  Leaf temperature $T_l$ as function of characteristic leaf length $l_c$ for various combinations of air temperature $T_a = (5\,^\circ C, 15\,^\circ C, 25\,^\circ C, 35\,^\circ C)$ and relative atmospheric humidity $w_{rel} = (0.3, 0.6, \text{and} 0.9)$. Wind velocity is $v_w = 0.1\,\text{m/s}$. Other input values are as given in Table 1 (biochemical parameters were borrowed from *Quercus petraea*) [Colour figure can be viewed at wileyonlinelibrary.com]
Effect of leaf size and wind speed on assimilation rate, transpiration rate, and leaf temperature

Figures 7–10 illustrate the impact of $l_c$ on leaf conductance, assimilation rate, transpiration rate, and leaf temperature if free stream wind speed is varied. All curves share a common feature: Their slopes are steepest for small $l_c$ and decrease smoothly with increasing $l_c$, such that the curves flatten out more and more. The reason for this behaviour is the way in which $l_c$ enters Equation (8): $l_c$ acts on the conduction/convection term in Equation (8) through the boundary layer thickness that is related to leaf size by $d_b \propto \sqrt{l_c}$, according to (9). This square root dependence explains, at least partly, that size changes in large leaves have much less impact on the dependent variables than size changes in small leaves. Although there is no clear-cut threshold separating both regimes, a value of roughly $l_c \approx 30$ mm seems to be reasonable for which further increases in $l_c$ have minor impacts on $g$, $A$, and $E$.

The most interesting feature of leaf size is its interaction with air humidity and air temperature: Leaves may experience heating (for $T_a \leq 20$ °C) or cooling (for $T_a \geq 20$ °C) depending on $T_{opt}$. Both effects are much more pronounced for small leaves than for large leaves. Warming occurs also for high humidities (i.e., small VPD), but cooling is in this case very limited due to low transpiration. Ecologically, this temperature compensatory feature makes sense: the warming is beneficial for assimilation, the cooling reduces the hazard of overheating.

Incidentally, Figures 9 and 10 corroborate the nearly negligible dependence of leaf temperature on atmospheric CO$_2$ already encountered for a fixed leaf size in Figures 5 and 6 for arbitrary values of $l_c$.

4 | DISCUSSION

There is persistent interest in the ecological role of leaf size and shape, and the selective pressures that underlie the evolution of leaf architecture (Givnish, 1987; Givnish and Vermeij, 1976; Leigh, Septant, Close, and Nicotra, 2017; Nicotra et al., 2011; Niinemets et al., 2007; Peppe et al., 2011). While possible functional aspects of leaf size and shape are broad, comprising, for example, self-shading or herbivory (Brown, Lawton, and Grubb, 1991; Falster and Westoby, 2003; Givnish, 1984), it is physically inevitable that the potential thickness of the boundary layer is dictated by them, with consequences for leaf temperature $T_l$ and transpiration rate $E$.

The model results reported here corroborate various observations and findings on leaf temperature under various environmental conditions (Michaletz et al., 2015; Gates, Hiesey, Milner, and Nobs, 1964; Linacre, 1967; Helliker and Richter, 2008; Pincebourde and Woods, 2012). The proposed model extends these approaches by combining leaf-gas exchange (Section 2.1) with leaf energy balance (Section 2.2), allowing reconstruction and prediction of leaf cooling and heating in environments of different atmospheric CO$_2$ concentrations.

Whereas the results of this study support the ecological relevance of the characteristic leaf size $l_c$, no evidence could be found that $C_a$ exerts a selective pressure on $l_c$ via $g$, and evaporative cooling, at least on the time scales analysed here. A decrease in $g$ in response to rising $C_a$ has been observed and is predicted by Equation (4) under conditions already discussed. This decrease in $g$ has prompted anticipations that leaf temperatures would increase under elevated $C_a$ as a consequence of reduced evaporative cooling (Beerling and Berner, 2005; Paschalis, Katul, Fatichi, Palmroth, and Way, 2017; Voelker et al., 2016). Figures 5 and 6, however, show that leaf temperature is more robust against variations in atmospheric CO$_2$ concentration than against variations in leaf size and atmospheric humidity.

As illustrated by the model results, a combination of physical and physiological mechanisms counteracts $C_a$-induced decrease of evaporative cooling, particularly under conditions of high air temperature. These are highlighted and discussed below.

1. Physiologically, photosynthesis is stimulated strongest when $T_l$ is close to $T_{opt}$ where assimilation rate $A$ and—in view of relation (13)—leaf conductance $g$ are maximal. Figure 1 illustrates that $T_{opt}$ is displaced to higher values when $C_a$ increases. Thus, rising $C_a$ accompanied by increasing $T_{opt, \langle T_{opt} \rangle}$ should promote increasing A and should lead to increased stomatal opening. The increased stomatal opening leads to increased evaporative cooling, and particularly so for high $C_a$ provided $\kappa$ is kept constant. Higher $g$ for both elevated $C_a$ and $T_{opt, \langle T_{opt} \rangle}$ compared to ambient $T_a$ is actually supported by various experiments (Wang, Heckathorn, Wang, and Philpott, 2012). Slightly higher $E$ under elevated $C_a$ and high VPD has also been reported by whole-tree-flux measurements (Duursma et al., 2014).

2. Physically, leaf heating increases the saturation vapour pressure of the leaf internal air. When assuming that air humidity outside the leaf (i.e., the free stream value) is not changing drastically (even under elevated $C_a$), it is expected that evaporation and thus evaporative cooling will be enhanced (Campbell and Norman, 1998).

When assessing the anticipated decrease in $g$ as a response to rising $C_a$, one should keep in mind that the decisive quantity representing evaporative cooling in Equation (8) is transpiration $E$. This quantity is formed from the product of $g$ and $(w(T_l) - w_a)$, according to (6). Comparison of Figure 1d and f shows that $E$ responds to variations in $C_a$ and $T_f$ mildly, in contrast to $g$ that exhibits a much more pronounced dependence on both $C_a$ and $T_f$. The reason for that is the factor $(w(T_l) - w_a)$: (a) Its steep, positive slope with respect to $T_l$ attenuates the negative slope exhibited by $g$ for $T_{opt, \langle T_{opt} \rangle}$ (as illustrated by Figure 1f) and (b) it dominates the product $E = \alpha(w(T_l) - w_a)g$ with respect to temperature and "squeezes" the separated curves of Figures 1d,g into the compact bands of Figures 1f,i). This closeness of the curves related to different $C_a$-values implies that rising $C_a$ will not cause a tremendous increase of leaf temperature, if any.

As pointed out in Section 3.1, evaporative cooling is not the only mechanism that provides leaf cooling. The combination of radiation emission and heat conductance/convection contributes also to a leaf cooling effect. The work here shows that this cooling effect comes...
into operation for $T_a > T_i, \infty$ ($T_i, \infty$ has been defined in 3.1: It measures the expected leaf temperature in the absence of evaporative cooling). As typical value for $T_i, \infty$, we found above $T_i, \infty \approx 37.9^{\circ}C$. Thus, this mechanism intensifies leaf cooling if temperatures have already risen to values that are dangerous for vital leaf functions.

The combination of physical and physiological mechanisms leads to the circumstance that values of $T_a$ appear to be weakly affected by elevated $C_a$, regardless of leaf size (see Figures 9 and 10). The anticipated phenomenon of leaves heating up under elevated $C_a$ may therefore be largely non-existent.

These results are in contrast to expectations that periods of elevated $C_a$ generate necessarily rising leaf temperatures (Cernusak et al., 2013; Paschalis et al., 2017) as well as for the past (Haworth et al., 2014; Lee, Upchurch, Murchie, and Lomax, 2015; McElwain, Beerling, and Woodward, 1999). For the past, $C_a$ is usually expected to represent a crucial element for the evolution of leaf shape and size under conditions of high $C_a$ levels (Beerling and Royer, 2011; Lee et al., 2015; McElwain et al., 1999). Atmospheric CO2 increased and decreased repeatedly during land plant evolution, with levels often close to or even higher than 1,000 ppm (Anagnostou et al., 2016; Beerling and Royer, 2011; Franks et al., 2014; Montañez et al., 2007; Steinhorsdottir and Vajda, 2015). Under such conditions, both smaller leaf size and dissected leaves were suggested as appropriate strategies to avoid excessive leaf temperatures under periods of high $C_a$ (Haworth et al., 2014; Lee et al., 2015; McElwain et al., 1999). According to the results of the present study, elevated $C_a$ does not have the potential to act upon leaf evolution via $T_a$. This also means that palaeoclimate proxy approaches that include leaf size, such as CLAMP, are not affected by $C_a$. During periods of supposedly high CO2, large and entire leaves did in fact not disappear. For example, angiosperm leaf size even increased from the early to the late Palaeocene-Eocene Thermal Maximum, a period of both high CO2, large and entire leaves did in fact not disappear (Haworth et al., 2014; Lee et al., 2015; McElwain et al., 1999). Atmospheric CO2 increased and decreased repeatedly during land plant evolution, with levels often close to or even higher than 1,000 ppm (Anagnostou et al., 2016; Beerling and Royer, 2011; Franks et al., 2014; Montañez et al., 2007; Steinhorsdottir and Vajda, 2015). Under such conditions, both smaller leaf size and dissected leaves were suggested as appropriate strategies to avoid excessive leaf temperatures under periods of high $C_a$ (Haworth et al., 2014; Lee et al., 2015; McElwain et al., 1999). According to the results of the present study, elevated $C_a$ does not have the potential to act upon leaf evolution via $T_a$. This also means that palaeoclimate proxy approaches that include leaf size, such as CLAMP, are not affected by $C_a$. During periods of supposedly high CO2, large and entire leaves did in fact not disappear. For example, angiosperm leaf size even increased from the early to the late Palaeocene-Eocene Thermal Maximum, a period of both high CO2, large and entire leaves did in fact not disappear (Haworth et al., 2014; Lee et al., 2015; McElwain et al., 1999). Atmospheric CO2 increased and decreased repeatedly during land plant evolution, with levels often close to or even higher than 1,000 ppm (Anagnostou et al., 2016; Beerling and Royer, 2011; Franks et al., 2014; Montañez et al., 2007; Steinhorsdottir and Vajda, 2015). Under such conditions, both smaller leaf size and dissected leaves were suggested as appropriate strategies to avoid excessive leaf temperatures under periods of high $C_a$ (Haworth et al., 2014; Lee et al., 2015; McElwain et al., 1999).

Leaf warming by high $I_c$ is potentially highest under low air temperature $T_a$ with further enhancement by simultaneously high $w_{rel}$ (see upper two rows of Figures 9 and 10). Large leaves may thus be particularly beneficial in cooler climate zones during the morning hours by stimulating assimilation rate $A$, as was already suggested by Okajima, Taneda, Noguchi, and Terashima (2012), and in warm and hot climate zones during the afternoon hours by promoting leaf cooling to avoid leaf damage. These functional aspects emphasize the need for knowing leaf temperature when evaluating gas exchange under different environmental conditions (Michaletz et al., 2015).

The relations between leaf temperature, atmospheric CO2 and leaf size provided by the model here imply the following: Leaf temperature depends on leaf size, but is almost independent of atmospheric CO2 despite reductions in leaf conductance with increased atmospheric CO2.

**CONFLICT OF INTEREST**

All authors declare no conflict of interest.

**ACKNOWLEDGEMENTS**

This manuscript is the extended version of a talk given at the Special Session Cenozoic plant diversity: Gradients in time and space and their impact on early humans (ROCEEH/NECLIME) of the EPPC 2018 in Dublin. The authors would like to thank the organizers of this NECLIME session (Angela A. Bruch, Alexandra-Jane Henrot, Louis François, Natalia Rudaya, and Torsten Utescher) for the invitation to contribute to this Special Issue. This study was supported by a grant of the Volkswagen Foundation to A. R. N. within the programme "Research in Museums" (Refs. 87139, 87160, 87160-1). The authors would also like to thank the reviewers for their constructive comments.

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**How to cite this article:** Konrad W, Katul G, Roth-Nebelsick A. Leaf temperature and its dependence on atmospheric CO2 and leaf size. *Geological Journal, 2021*;56:866–885. [https://doi.org/10.1002/gj.3757](https://doi.org/10.1002/gj.3757)

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**APPENDIX A**

To motivate the results of Section 3.1, Equation (8) is considered in greater detail.

The special case of vanishing transpiration

First, the special case when transpiration (and thus evaporation) does not occur (e.g., when \( C_{w} \) is very large) is presented. This means that stomata are closed (\( g = 0 \)), implying that leaf evaporation and the last term in Equation (8) vanish. In Figures 5 and 6, this case is represented by the dashed black lines. These lines and the solid black lines, defined by \( T_{l} = T_{w} \), intersect at the temperature

\[
T_{l}=\frac{\sqrt{1+r^2}\left(\frac{1}{\sin\gamma}-\frac{r_{n}^{2}/\sin\gamma}{\pi}+a_{IR}\left(\frac{T_{w}^{4}+a_{IR}^{4}}{2\pi a_{IR}^{4}}\right)\right)}{2\pi a_{IR}^{4}}
\]

which can be calculated from insertion of \( g = 0 \) and \( T_{w} = T_{l} \) into Equation (8). All expressions under the square-root sign are necessarily positive (including \( \sin\gamma \)) and the angle \( \gamma \) denotes the height of the sun above the horizon is restricted to the range \( 0 < \gamma < 90\)°; this guarantees the existence of \( T_{l} \). Notice that \( T_{l} \) only depends on the radiative properties of the leaf and can therefore serve as a reference temperature that characterizes the radiative interactions of the leaf energy balance. The \( T_{l} \) is independent of \( w_{at} \) and \( l_{c} \), which
explains why it is located at the same position \((T_i = \approx 37.9{^\circ}C)\) in all sub-figures of Figures 5 and 6.

To draw further conclusions, the slope of the dashed black \((g = 0)\) line is calculated. Taking the total derivative of \((8)\) (assuming \(g = 0\)),

\[
0 = 8\varepsilon\sigma T_l^4 \frac{dT_l}{dT_a} + \frac{2K_a}{\partial_T} (dT_l - dT_a) \quad (A2)
\]

and then solving for \(dT_l/dT_a\), we obtain

\[
\frac{dT_l}{dT_a}_{|g = 0} = \frac{K_a}{K_a + 4d_0\varepsilon\sigma T_l^4} \quad (A3)
\]

If evaporation vanishes \((g = 0)\) the two results \((A1)\) and \((A3)\) allow the following conclusions to be drawn:

1. For \(T_o < T_{i,\infty}\) leaves are warmer than the surrounding air, whereas for \(T_o > T_{i,\infty}\) leaves are cooler than the air. Notice that the processes of radiation emission and heat conduction/convection alone produce this heating/cooling phenomenon, and evaporative cooling is not involved by definition.

2. Because all quantities in expression \((A3)\) are positive and the numerator is smaller than the denominator, the slope of the dashed black line is positive and restricted to the \((0)\) interval \(0...1\), implying that for "turned-off" leaf evaporation, \(T_i\) rises whenever \(T_o\) rises but by a smaller amount than \(T_o\). In other words, the combination of radiative, conductive, and convective energy exchange mechanisms between leaves and atmosphere mitigates the impact of \(T_o\)-variations on \(T_i\).

Including transpiration

If evaporation from the leaf interior is included \((i.e., \ g > \ 0)\), the leaf temperature (and thus heating and cooling) is described by the coloured lines in Figures 5 and 6. They run completely below the dashed black lines related to \(g = 0\), they exhibit smaller slopes than these, and they intersect the solid black line (defined by \(T_o = T_i\)) at lower values than \(T_{i,\infty}\), implying that the temperature \(T_i, (C_a)\) separating the leaf heating and cooling regimes drops. The reason for all these, in terms of physics, has already been stated; since vaporization consumes heat, evaporation can transport heat only away from the leaf, that is, evaporation can only cool the leaves. In contrast, the mechanisms of conduction and convection can transport heat both ways, depending on transport direction, they can cool or heat the leaf.

It is instructive to relate several features depicted in Figures 5 and 6 to Equation \((8)\).

1. Especially Figure 5 suggests quite clearly that the coloured lines approach for low temperatures the dashed black lines signifying vanishing evaporation. This makes sense: At low temperatures, assimilation yield is low, as illustrated by Figures 1b, e, and h, and plants prefer to close stomata to avoid water loss. Upon setting \(g = 0\), Equation \((8)\) reduces to the equation defining the dashed black lines.

2. With increasing leaf temperature, leaf conductance \(g\) and transpiration rate \(E\) increase (see Figures 1a, c) until \(g\) reaches its maximum value at \(T_i = T_{\text{opt}}\); the maximum of \(E(T_i)\) is located at a slightly higher temperature \(T_i = T_{\text{max}}^E\) (which can be traced to the positive slope of the factor \(w(T_i)\) in relation \((6)\)). At \(T_i = T_{\text{max}}^E\) the slopes of the coloured lines coincide with the slope of the dashed black lines. This can be understood as follows: If the assumption \(g = 0\) is dropped, relation \((8)\) implies the unrestricted version of relation \((A3)\)

\[
\frac{dT_l}{dT_a} = \frac{K_a}{K_a + 4d_0\varepsilon\sigma T_l^4 + d_0H_vap\frac{dT_l}{dT_a}} \quad (A4)
\]

Clearly, for \(\partial E(T_i)/\partial T_i = 0\) which indicates the maximum of \(E(T_i)\), this expression reduces to \((A3)\). Thus, maximum leaf temperature reduction caused by evaporation alone occurs at \(T_i = T_{\text{max}}^E\).

If leaf temperature increases beyond \(T_i = T_{\text{max}}^E\) the coloured lines first approach and finally terminate at the dashed black lines. The green line in Figure 1a illustrates that this happens when the stomata close; physiologically, this is due to high leaf temperature because then \(\text{(a) assimilation has declined already (as can be concluded from Figure 1b) and \(b)\) keeping water within the plant may have become vitally important.}

Relaxing the condition \(g = 0\) to \(g > 0\) by including evaporation as energy exchange mechanism requires a reformulation of the conclusions that were drawn at the end of section Appendix A.1:

1. For \(T_o < T_{i,\infty}\) leaves are warmer than the surrounding air, whereas for \(T_o > T_{i,\infty}\) leaves are cooler than the air.

2. For leaf temperature \(T_i < T_{\text{max}}^E\), all quantities in expression \((A4)\) are positive and the numerator is smaller than the denominator, the slope of the dashed black line is positive and restricted to the \((0)\) interval \(0...1\). This guarantees that for \(T_i < T_{\text{max}}^E\), leaf evaporation \(T_i\) rises whenever \(T_o\) rises but by a smaller amount than \(T_o\). For \(T_i > T_{\text{max}}^E\), expression \(\partial E(T_i)/\partial T_i\) in the denominator of \((A4)\) becomes negative. But if \(T_i\) obeys the condition \(T_i < \sqrt[4]{(H_vap/4\varepsilon\sigma)(\partial E(T_i)/\partial T_i)}\), the conclusion \(dT_i < dT_o\) still applies, that is, rising air temperature is mitigated.