Effects of exogenous ABA on photosynthesis and stomatal conductance of cut twigs from oak seedlings

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Introduction

Abscisic acid (ABA) plays a major role in plant-water relations. It has been shown to promote stomatal closure in many species including trees (Johnson, 1987), and there is growing evidence that it could be a root-produced effector for water stress reactions (Zhang et al., 1987). These assertions are based on studies with cut twigs supplied with exogenous ABA and on measured increases of ABA concentrations in xylem sap.

However, many questions remain open to discussion: are the concentrations of exogenous ABA necessary to promote a sensible reaction of the same magnitude as those of free ABA measured in the xylem during water stress? Is the rapid stomatal closure promoted by exogenous ABA the direct cause of the observed decline in net photosynthesis (Downton et al., 1988) or is there some direct effect of ABA on mesophyll photosynthesis (Raschke and Hedrich, 1985)? Do forest trees display the same responses to ABA as other species?

We have therefore, as a preliminary to a detailed survey of the role of ABA in reactions of oak species to water deficits, tested the reactions of cut twig photosynthesis to exogenous ABA. The effects of shoot removal on gas exchange were assessed prior to use of this technique with ABA.

Materials and Methods

Plant material

3 yr old seedlings, grown on a sand-peat soil (50/50, v/v) in 8 l pots, were transferred into a climate chamber (February) to accelerate bud break prior to measurements conducted during March and April 1988. Species: Quercus robur Fig. 1.L., Q. petraea L. (seeds collected near Nancy), and Q. pubescens L. (Avignon).

Gas exchange measurements

These were made in an open flow chamber. Twig transpiration was estimated using a bypass flow (300 l-h⁻¹), and net CO₂ assimilation was calculated from CO₂ reduction in the main flow (60 l-h⁻¹). Chamber volume was 9 l; time lags between apparent assimilation and transpiration appeared during rapid rate changes. Steady state calculations were therefore only conducted after stabilization to avoid artifacts.

Climate in the chamber

Photosynthetic photon flux density: about 600 ± 20 μmol·m⁻²·s⁻¹; temperature: 24°C;
molar fraction of CO₂ in the chamber (cₐ): 350 ± 5 μmol-mol⁻¹; leaf to air difference in water vapor molar fraction (dω): about 12–15 mmol-mol⁻¹, depending upon leaf temperature and stomatal conductance. Leaf water potential (ψ₇) was monitored in the chamber with a Wes-cor in situ leaf micropsychrometer.

**Gas exchange parameters**

Net CO₂ assimilation (A), transpiration (E), stomatal conductance for CO₂ (g) and mesophyll CO₂ molar fraction (cᵢ) were calculated according to von Caemmerer and Farquhar (1981). Results are presented either as time evolution of A, g and ψ₇, or as A vs cᵢ graphs.

**Twig removal**

Twigs bearing 3–4 leaves were enclosed in the chamber and gas exchange parameters determined after at least 2 h of equilibration. Thereafter, twigs were detached and their cut end immediately plunged into a nutrient solution. Gas exchange parameters and leaf water potential were monitored for at least 4 h after cutting.

**ABA application**

(+/-)2-cis-4-trans-Abscisic acid (Aldrich Chemie) was dissolved in the nutrient solution at 3 concentrations: 10⁻⁴, 10⁻⁵ and 10⁻⁶ M. The nutrient solution supplied to shoots was replaced by an ABA-supplemented one and gas exchange followed for at least 4 more hours.

**Effects of CO₂ enrichment**

A, E and g were measured successively on Q. pubescens under ambient (350) and enriched (1000 μmol-mol⁻¹) CO₂ mole fractions, both before and after ABA supply. Each measurement was made after at least 1 h of equilibration.

![](image)

**Fig. 1.** Effects of cutting a twig and plunging its cut end into a nutrient solution on A (open symbols), g (closed symbols) and ψ₇ (Q. robur).
Results

Effects of cutting

Cutting caused an immediate and steep decrease in stomatal conductance \((g)\) and net \(\text{CO}_2\) assimilation \((A)\) (Fig. 1), and a rapid increase of water potential \((\psi_w)\), the latter being a direct consequence of both a reduction in transpiration \((E)\) and the removal of all the resistances to water flux from root to shoots. These effects were immediate (appearing after less than 1 min) and only transient, vanishing in about 1 h. A new steady state was reached thereafter, with significantly lower \(A\) and \(g\), and was maintained for at least 3–4 h.

In as much as it displays a new steady state gas exchange rate, a cut twig is a valuable tool for studying effects of exogenous ABA in the absence of any water stress.

Effects of ABA application

At 10\(^{-4}\) M, the effects were very similar to those described above with two main differences: 1) there was a significant time lag before leaf reaction, which may be attributed to ABA diffusion into leaves; from the original records, we may estimate the delay to be 10.8 ± 1.9 min for \(A\) and 9.1 ± 1.2 min for \(g\) (Fig. 2a); 2) no recovery appeared during the 1st hours after application, even if \(A\) and \(g\) increased slightly after the first breakdown. Plotting these results on \(A\) vs \(c_i\) curves (Fig. 2b) reveals a strong reduction of mesophyll photosynthesis.

ABA reactions under increasing external \(\text{CO}_2\) molar fractions \((c_a)\)

\(c_a\) was temporarily increased to 1000 \(\mu\text{mol-mol}^{-1}\) just before and 1 h after ABA application. Results are shown as \(A\) vs \(g\) relations \((Q. \text{pubescens},\ Fig. 3)\). Decreasing reactions with concentrations below 10\(^{-4}\) M were observed. Increasing \(c_a\) caused additional stomatal closure even in the presence of ABA but did not promote the expected increase in \(A\). Furthermore, the application of ABA did not change the relationship between \(A\) and \(g\) for each \(c_a\); under constant humidity, this suggests that ABA affects both stomatal conductance and mesophyll assimilation.

Discussion

Cutting promoted quite immediate reactions by leafy shoots. These kinds of effects had been attributed to a hydropassive stomatal closure; but, like Myers et al. (1987) on Eucalyptus sp., we noticed that stomatal closure was accompanied by quasi constant \(c_i\) values, which reveals a reduction in mesophyll photosynthetic activity. These effects were reversible and the appearance of a new steady state enabled the use of cut twigs as an experimental tool for ABA studies.

At high concentrations of about 10\(^{-4}\) M, ABA had an important effect on stomata and photosynthesis on all tested oak species, although lower concentrations (10\(^{-6}\) M) had no effect.

Direct effects on mesophyll photosynthesis may be inferred from \(A\) vs \(c_i\) curves which show \(A\) reductions at constant \(c_i\) values, and from the constant \(A/g\) ratios at high \(c_a\). These results are in agreement with those of Raschke and Hedrich (1985). The \(c_i\) gradients across hypostomatous leaves (Parkhurst et al., 1985) are not large enough to modify these conclusions. Existence of ‘patchy behavior’ of stomata in response to ABA (Downton et al., 1988) could contradict these conclusions, but there is still not enough evidence to demonstrate the reality of this behavior.
Fig. 2. a. Time course of ABA effects at $10^{-4}$ M on $A$ (open symbols), $g$ (closed symbols) and $\psi_w$ of a cut twig supplied with a nutrient solution. b. $A$ vs $c_i$ curves for same data. Only steady state values have been sketched to avoid artifacts.
Fig. 3. Relations between $A$ and $g$ under low and high $c_A$. The ABA concentrations were $\Box$: $10^{-4}$ M; $\bigotimes$: $10^{-5}$ M; $\Delta$: $10^{-6}$ M (open symbols: values before ABA application; closed symbols: after ABA application).

References


