

Photosynthesis Related Characteristics of Upper and Lower Canopy Leaves of *Kandelia obovata*, a Mangrove Species in Taiwan

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ABSTRACT: To understand the light use properties of canopy leaves in *K. obovata*, the dominant mangrove species in the west coast of northern Taiwan, I compared photosynthesis related characters, such as leaf angles, leaf area, stomatal density, specific leaf area, and chlorophyll fluorescence, of upper and lower canopy leaves of the species. Leaves that had developed in exposed positions (upper canopy) exhibited significantly steeper leaf angles, smaller leaf area, higher stomatal density and lower specific leaf area than those in shaded positions (lower canopy). No significant difference was found in chlorophyll (Chl) content per unit leaf area and Chl *a/b* ratio between the two types of leaves. In contrast, compared to lower canopy leaves, upper canopy leaves had higher nitrogen content per unit leaf area indicating a higher light saturated CO₂ assimilation. Chlorophyll fluorescence measurement revealed that photosaturated electron transport rate (ETR) and the photon flux required to saturate ETR were significantly higher in upper canopy leaves than in lower canopy leaves. The plasticity in morphological and physiological response to variation in light regimes may represent an important mechanism for *K. obovata* to adapt to the saline and nitrogen limiting environment.

KEY WORDS: Mangroves, *Kandelia obovata*, Photosynthesis, Chlorophyll fluorescence.

INTRODUCTION

The level of irradiance is an important ecological factor affecting all photoautotrophic plants. Low light intensity poses stresses on plants because irradiance limits photosynthesis and thus net carbon gain and plant growth. In contrast, high light intensities may also be a stress for plants, particularly if other factors are not optimal (Björkman and Powles, 1984; Kao and Forseth, 1992). The ability of adjustment to variation in light regimes in trees with dense canopy structure, hence with leaves exposed to different light regimes, appears to be an important mechanism for maximum carbon gain (Küppers, 1989).

Kandelia obovata is the dominant mangrove species in the west coast of northern Taiwan (Liu, 1982; Sheue, et al., 2003). Factors such as high salinity and nutrient limitation have been shown to limit the growth of this plant in tidal waters (Huang and Chen, 1995; Kao and Chang, 1998; Kao et al., 2001). In a field trip to the mangrove swamp, I observed that inclination angles of leaves on the upper canopy differ from those of leaves on the

lower canopy of the forest. The inclination of leaf angle has direct effect on light interception. The display of leaves may be an adaptation to the saline environment in term of optimization of carbon gain and/or avoidance of light and temperature stress. The phenomenon prompted me to further examine the photosynthesis related characters of the canopy leaves.

The amount and kinetics of chlorophyll (Chl) *a* fluorescence emitted from leaves in response to actinic irradiation can be used to assess the primary photochemistry of photosynthesis (Krause and Weis, 1991). In particular, measurements of Chl *a* fluorescence can determine the potential quantum yield of photosystem II (PSII) (F_v/F_m), the efficiency of excitation capture by open PS II reaction centers (F_v'/F_m'), effective quantum yield of PSII (Φ_{II}), and the electron transport rate through PSII (ETR). These variables are tightly associated with the corresponding light responses and light-saturated rates of whole-leaf photosynthetic carbon dioxide uptake (Krall and Edwards, 1992; Björkman and Demming-Adams, 1994). In addition, the photochemical (qP, representing the fraction of PSII centers that are actually open at a given PFD) and non-photochemical quenching (qN) coefficients provide important information of photosynthetic activity. Measurements of various fluorescence

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parameters in response to light reveal how plants acclimate to variation in light intensity in their environment (Ishida et al., 1999; Ishida et al., 2001). In this study, the Chl fluorescence technique was used to assess the efficiency of photosynthetic energy conversion by the upper and lower canopy leaves of *K. obovata*.

In addition to leaf angle, leaf size and chlorophyll content also affect light interception, and nitrogen content affect photosynthetic activity. Hence, in this study I also measured these photosynthetic related characters. The objective of this study was to understand better the relationship between light-use properties and leaf display of *K. obovata* in the saline environment. I tested the hypothesis that morphological and physiological differences occur between upper and lower canopy leaves as a result of light environment.

MATERIALS AND METHODS

Twenty-four branches, 12 from upper (about 3 m height) and 12 from lower canopy, of *K. obovata* were collected from the mangrove forest in Waltz-wei Natural Reserve (25°10' N, 121°24' E), Taipei County in Oct., 1999. As soon as the branches were detached, they were placed in water and brought back to the laboratory. The most recent fully expanded leaves (the third or fourth pair of leaves) from each branch were used for following analyses.

Leaf angles from the horizontal were measured by aligning a hand-held clinometer (Suunto Co. PM-5/360 PC) with the plane of the lamina.

Twelve leaves, 6 from upper and 6 from lower canopy, were chosen for fluorescence measurements. To measure the fluorescence-PFD (photosynthetic photon flux density) response, leaves were held horizontally in a leaf-clip holder (2030-B, Walz, Effeltrich, Germany) (Biger et al., 1995). The PFD on leaves, provided by a halogen lamp (2050-H, Walz, Effeltrich, Germany), was adjusted from darkness to 1600 $\mu\text{mol m}^{-2} \text{s}^{-1}$ in steps of 50-200 $\mu\text{mol m}^{-2} \text{s}^{-1}$. The halogen lamp was equipped with a heat-reflectance filter to reduce heat generated by the lamp. PFD on the leaf was monitored with a microquantum sensor installed on the leaf-clip holder next to the spot where fluorescence is measured. After the leaf was exposed to the desired PFD for 10 min, the Chl *a* fluorescence of PSII was measured using a portable, pulsed amplitude modulated fluorometer (PAM 2000, Walz, Effeltrich, Germany). The values of F_v/F_m , F_v'/F_m' , the effective quantum yield of PSII

($\Phi_{II} = qP * F_v'/F_m'$), and the coefficient of photochemical and non-photochemical quenching [$qP = (F_m' - F_t)/(F_m' - F_o')$, $qN = (F_m - F_m')/(F_m - F_o')$], were computed (Schreiber, Schlina and Bilger, 1986), where F_v and F_m are the variable and maximal fluorescence, respectively, in dark adapted state, F_m' is the maximal fluorescence, F_o' is minimal fluorescence, F_v' is the difference between F_m' and F_o' , and F_t is the steady-state fluorescence in the light adapted state. During the fluorescence measurement, room temperature was maintained at 26°C while the leaf temperature was not controlled.

Following fluorescence measurement, a 2 * 2 cm² area from each experimental leaf was collected and total leaf Chl concentration was measured by extracting Chl from the leaf segment with 96% ethyl alcohol, and subsequent spectrophotometric (Model V-560, Jasco, Tokyo, Japan) analysis of the extract at wave-lengths of 649 and 665 nm (Wintermans and Motts, 1965).

A thin layer of nail polish was applied to the lower surface of a segment of the experimental leaves ($n = 6$) for estimates of stomatal density. After drying, the nail polish was peeled and scanned at 100 x magnification with a light microscope equipped with a calibrated ocular micrometer.

The area of leaves ($n = 12$, for each canopy layer) was determined with an area meter (LI-3100, Li-Cor, Nebraska), and each leaf was dried at 60°C for 48 h and then weighed. The specific leaf area (SLA) was calculated as leaf area per unit dry mass. These samples were then ground to a fine powder with a mortar and pestle. Total nitrogen (N) and carbon contents of samples were determined with an elemental analyzer (NA 1500, Fisons, Italy).

All statistical tests were performed with the computer software SYSTAT (statistical Solution, Cork, Ireland). Significant differences are reported as $P < 0.05$.

RESULTS

Leaves that had developed in exposed positions (upper canopy) exhibited significantly steeper leaf angles, smaller leaf area, less SLA and higher stomatal density than those in shaded positions (lower canopy) (Table 1). Lower canopy leaves had a 86 % and a 61 % increase in leaf area and SLA, respectively.

No significant difference in Chl *a/b* ratio and nitrogen (N) content per unit leaf dry weight was found between leaves on upper and lower canopy (Table 1). However, upper canopy leaves had significantly higher N content per unit leaf area. In

Table 1. A comparison of photosynthesis related characters of upper and lower canopy leaves of *K. obovata* (mean \pm s. e.).

Character	Upper Canopy Leaf	Lower Canopy Leaf
Leaf angle	65 \pm 2 ^a	36 \pm 2 ^b
Leaf area (cm ²)	14.6 \pm 0.9 ^a	27.2 \pm 1.8 ^b
Stomatal density (mm ⁻²)	142 \pm 6 ^a	124 \pm 5 ^b
SLA (cm ² g ⁻¹)	59.6 \pm 1.9 ^a	96.3 \pm 3.8 ^a
Chl <i>a/b</i>	3.1 \pm 0.1 ^a	2.9 \pm 0.1 ^a
Chl content (mg g ⁻¹)	4.2 \pm 0.2 ^a	7.1 \pm 0.4 ^b
Chl content (mg cm ⁻²)	0.052 \pm 0.003 ^a	0.058 \pm 0.004 ^a
N content (mg g ⁻¹)	19.4 \pm 1.2 ^a	18.8 \pm 0.9 ^a
N content (g m ⁻²)	3.28 \pm 0.21 ^a	1.96 \pm 0.08 ^b
C content (mg g ⁻¹)	495.1 \pm 3.6 ^a	439.6 \pm 5.5 ^b

* Means within a row followed by different superscripts are different at P = 0.05 (t-test)

contrast, lower canopy leaves had significantly higher Chl content per unit leaf dry weight than upper canopy leaves, however, no significant difference was found in Chl content per unit leaf area between the two layers of leaves (Table 1). Thus, variation in leaf Chl and N contents was mainly caused by variation in SLA.

Upper and lower canopy leaves had similar potential quantum yield of 0.81 \pm 0.02 and 0.82 \pm 0.02 (mean \pm s. e., n = 6), respectively, in dark-adapted condition. Under illumination, Fv'/Fm' and qP decreased with increasing PFD (Fig. 1A, B). The reduction rates in Fv'/Fm' and qP in response to increasing PDF were faster in lower canopy leaves than in higher canopy leaves. Consequently, at PFD > 200 $\mu\text{mol m}^{-2} \text{s}^{-1}$, upper canopy leaves had significantly higher qP and Φ_{II} (= (Fv'/Fm') * qP) than lower canopy leaves compared at the same PFD (Figs. 1B and C).

qN increased with increasing PFD and a more rapid rise in qN was measured in lower canopy leaves. Consequently, at PFDs of 200, 400, 600 and 800 $\mu\text{mol m}^{-2} \text{s}^{-1}$, lower canopy leaves had significantly higher qN than upper canopy leaves. qN saturated at PFDs of 800 and 1600 $\mu\text{mol m}^{-2} \text{s}^{-1}$ for lower and upper canopy leaves, respectively (Fig. 1B). No significant difference in light saturated qN was found between upper and lower canopy leaves, indicating similar capacity for non-photochemical quenching.

Both types of leaves had similar area based chlorophyll content and Chl *a/b* ratio (Table 1) indicating similar leaf absorptance. Hence, a standard leaf absorptance value of 0.84 was used and the electron transport rate through PSII was estimated (ETR = Φ_{II} * PFD * 0.5 * 0.84) (Genty et al., 1989). Figure 2 shows that the estimate ETR of upper canopy leaves increased with increasing PFD, reached saturation at PFD between 800 and 1200 $\mu\text{mol m}^{-2} \text{s}^{-1}$, then declined at PFD of 1600 $\mu\text{mol m}^{-2} \text{s}^{-1}$. The estimate ETR of lower canopy leaves also increased with increasing PFD, reached

saturation at PFD of 600 $\mu\text{mol m}^{-2} \text{s}^{-1}$, then remained constant as PFD increased further to 1600 $\mu\text{mol m}^{-2} \text{s}^{-1}$. The light-saturated ETR was significantly higher in upper canopy leaves than in lower canopy leaves.

DISCUSSION

Plants exhibit a wide range of adaptation, including biochemical, physiological and morphological adjustment that enables them to live successfully in different habitats. Results of this study show that morphological and physiological differences occur between upper and lower canopy leaves of *K. obovata*. The plastic response may confer this plant to grow in the tidal waters.

Leaves that had developed in upper canopy exhibited significantly steeper leaf angles, smaller leaf area and higher stomatal density than those in lower canopy (Table 1). Excessive irradiance and high temperature can adversely affect photosynthesis resulting in photoinhibition, particularly if other factors are not optimal. Upper canopy leaves of *K. obovata* may avoid conditions that predispose them to photoinhibition by reducing leaf size and developing leaves with steeper leaf angle. A steeper leaf angle would decrease the interception of excess light hence reduce leaf temperature, water loss and photoinhibition. In addition, steep leaf angles in the upper canopy leaves enhance light penetration in the canopy (Ehleringer and Werk, 1986; Forseth and Teramura, 1986; Gamon and Peracy, 1990). A reduction in leaf size lowers boundary layer resistance and provides more effective convective heat loss to the surrounding air (Givnish, 1987). In addition, increasing stomatal density would potentially increase transpirational heat dissipation and CO₂ uptake. In contrast to upper canopy leaves, leaves on the lower canopy their photosynthetic activity are more limited by light availability. The morphological characters of these leaves, more horizontally positioned, larger leaf area and higher SLA, would enhance their light interception.

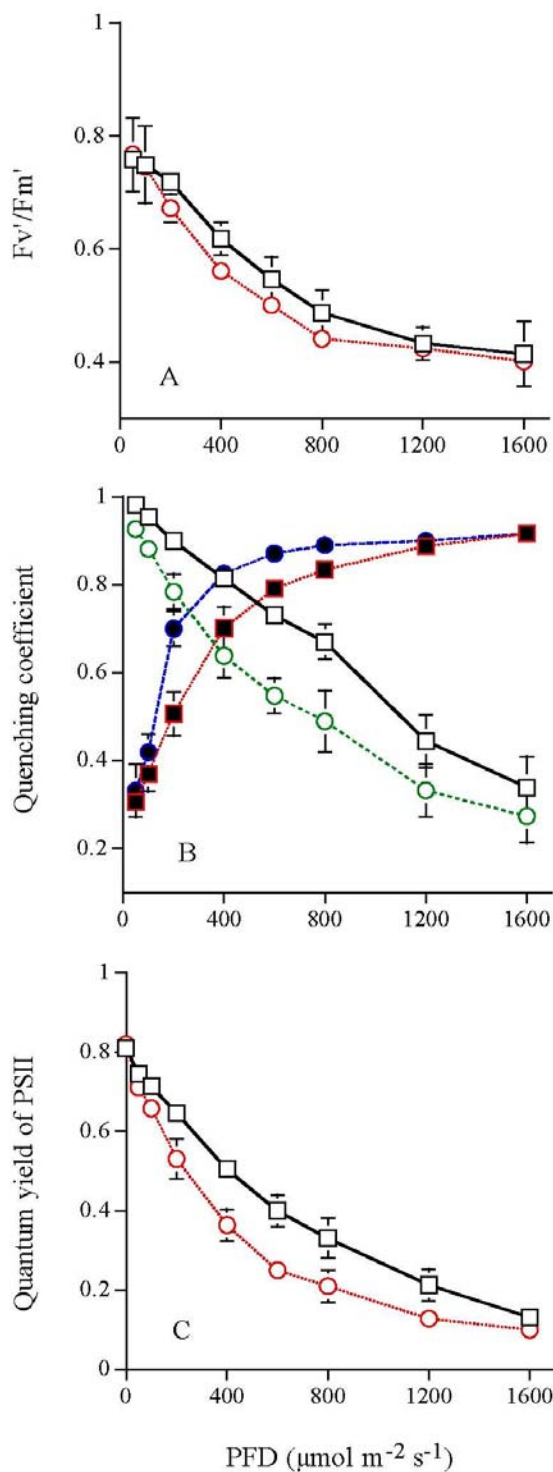


Fig. 1. The efficiency of excitation capture by open PS II reaction centers (F_v'/F_m') (A), photochemical (open symbols) and non-photochemical (closed symbols) quenching coefficients (B), and effective quantum yield of PSII (Φ_{II}) (C) of upper canopy (square) and lower canopy (circle) leaves of *K. obovata* in response to photosynthetic photon flux density (PFD) at the leaf surface. Bars represent s. e. (n = 6).

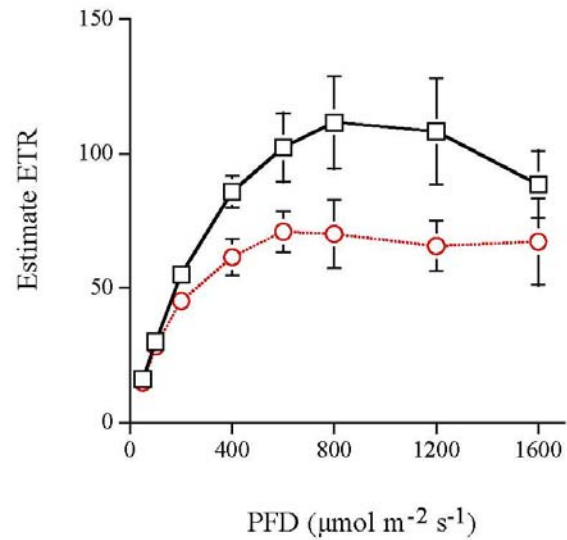


Fig. 2. Response of electron transport rate through PSII, estimated from chlorophyll fluorescence measurement, of upper canopy (square) and lower canopy (circle) leaves of *K. obovata* to photosynthetic photon flux density (PFD) at the leaf surface. Bars represent s. e. (n = 6).

No significant difference was found in Chl *a/b* ratio and Chl content per unit leaf area (Table 1) between upper and lower canopy leaves, which would reflect a similar light absorbance. In contrast, Chl fluorescence measurement revealed that upper canopy leaves had significantly higher q_P and Φ_{II} than lower canopy leaves compared at the same PFD (Figs. 1B and C), which indicates that upper canopy leaves adjust their photochemical capacity to maintain more reaction centers in an open (unreduced) state. Consequently, a higher photosaturated ETR was found in upper canopy leaves. The ETR values may be underestimated, since the absorbance value of *K. obovata* could be higher than the standard value (0.84) used to estimate ETR. Nevertheless, the estimates of ETR are valid for comparison purpose.

Positive correlation between photosaturated ETR and light-saturated rates of whole-leaf photosynthetic CO_2 uptake (A_{max}) was found (Krall and Edwards, 1992). Accordingly, fluorescence measurement suggests that upper canopy may have higher A_{max} than lower canopy leaves. Leaf N content analysis supports this suggestion. A positive correlation between leaf N content and photosynthetic capacity has been reported in a wide variety of plant species (Field and Mooney, 1986). The result that more nitrogen was allocated to upper canopy leaves than to lower canopy leaves (per unit area) (Table 1) suggest that upper canopy leaves had higher photosynthetic capacity. Mooney and Ehleringer (1978) have shown that leaves with high photosynthetic capacities can

have a high daily carbon in high-light environment. The higher carbon contents measured in upper canopy leaves than in lower canopy leaves (Table 1) may reflect this correlation. Thus, the differential distribution of N between upper and lower canopy leaves may increase the whole canopy carbon gain. In addition, the growth of *K. obovata* in tidal waters has been shown to be limited by nitrogen (Kao and Chang, 1998; Kao et al., 2001). Accordingly, it is expected that the plant would try to optimize its nitrogen use efficiency in this environment. Variation in foliar N content associate with different light environments results in a higher carbon gain for the whole canopy than for a canopy in which N content is uniformly distributed among leaves (Hirose and Werger, 1987; Elsworth and Reich, 1993; Evens, 1993; Hollinger, 1996). Thus, by allocating different N content to canopy leaves receiving different light regime, *K. obovata* may also increase photosynthetic N use efficiency for the whole canopy.

The result that lower canopy leaves had higher qN than upper canopy leaves at intermediate PFDs suggests that more energy is dissipated through non-photochemical quenching in lower canopy leaves than in higher canopy leaves at intermediate PFDs, due to the lower photochemistry ability of lower canopy leaves (indicating by lower qP and lower Fv'/Fm'). However, both type of leaves have similar light saturated qN indicating similar capacity of non-photochemical quenching ability. The ability of non-photochemistry quenching is related to the pool of xanthophylls (Demmig-Adams and Adams, 1992). Accordingly, the upper and lower canopy leaves of *K. obovata* may have similar size of xanthophylls pool. Increasing light stress was found to increase the pool size of the xanthophylls cycle (Demmig-Adams and Adams, 1996). Consequently, upper canopy leaves were found to have higher pool size of the xanthophylls cycle than lower canopy leaves in *Rhizophora apiculata* mangrove forest (Lovelock and Clough, 1992) and in a tropical tree species, *Acacia crassiparva* (Liu, et al., 2003). However, for *K. obovata* a reduction of the amount of solar radiation through the leaf inclination and the higher capacity of photochemical and assimilation activity in upper canopy leaves may reduce the need for increasing the size of xanthophyll pool in its upper canopy leaves. This could potentially save resource allocation to protect photosynthetic apparatus from light stress.

In conclusion, the plasticity in morphological and physiological response to variation in light regimes may represent an important mechanism for *K. obovata* to adapt to the saline and nitrogen

limiting environment. The hypothesis remains to be tested.

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水筆仔樹林上、下層葉光合作用相關特徵比較

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摘 要

水筆仔是北台灣西岸主要紅樹林樹種。水筆仔樹林的上層葉和下層葉可能因為接受到的光量不同，導致其在形態和生理反應上有所調整。本文比較該樹林上、下層葉的葉面角度、葉面積大小、氣孔密度、單位重葉面積、葉綠素螢光等與光合作用相關特徵，以了解水筆仔樹林對光使用特性。結果發現上、下層葉在形態上有顯著差異：上層葉葉面角度較大、面積較小、氣孔密度較高以及單位重葉面積較小。雖然兩者單位面積葉綠素含量和葉綠素 *a/b* 比沒有顯著差異，但上層葉單位面積有顯著較高的氮含量。葉綠素螢光測量顯示：相較於下層葉，上層葉其電子傳遞鏈速率在較高光量下才會達到飽和且有顯著較高的光飽和值。這些特徵差異顯示水筆仔為因應不同光環境，在葉形態和生理上有所調整。

關鍵詞：紅樹林、水筆仔、光合作用、葉綠素螢光。

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