Exogenous melatonin affects photosynthesis in characeae *Chara australis*

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Melatonin was found in the fresh water characeae *Chara australis*. The concentrations (~4 μg/g of tissue) were similar in photosynthesizing cells, independent of their position on the plant and rhizoids (roots) without chloroplasts. Exogenous melatonin, added at 10 μM to the artificial pond water, increased quantum yield of photochemistry of photosystem II by 34%. The increased efficiency appears to be due to the amount of open reaction centers of photosystem II, rather than increased efficiency of each reaction center. More open reaction centers reflect better functionality of all photosynthetic transport chain constituents. We suggest that melatonin protection against reactive oxygen species covers not only chlorophyll, but also photosynthetic proteins in general.

Introduction

Melatonin was first discovered in plants in two surveys of common fruits and vegetables from the market.1,2 In 1997, melatonin was reported in growing plants3 and a putative pathway for melatonin synthesis in plants was described in 2000.4 It was hypothesized that melatonin levels in plants may be timing mechanisms for circadian rhythm or seasonality with the highest level in dark periods and the lowest level during daylight.5 A diurnal rhythm in melatonin with a maximum in the dark phase and low levels during the day has been reported in the short-day plant *Chenopodium rubrum*, but changing the duration of the dark phase did not change the melatonin fluctuation.6 There was a 15–30-fold higher level of melatonin in etiolated (dark grown) seedlings as compared with light adapted tissues of *Hypericum perforatum* and *Arabidopsis thaliana*.7 Melatonin levels were higher in *Glycyrhiza uralensis* grown under red light than those plants grown under green or blue light.8 In water hyacinths a peak in melatonin was found late in the light phase, near sunset,9 while in field grown *Vitis vinifera* L. cv Malbec the peak occurred at dawn.10 If the grapes were shielded from sunlight throughout the day, the concentration of melatonin remained high.10 More recently, Byeon et al. found that increased melatonin and activity of the genes associated with melatonin biosynthesis were detected in detached rice leaves under constant high light with lower concentrations observed in constant darkness.11 These researchers hypothesized that melatonin biosynthesis is dependent on a light signal.11 However, none of the systems studied to date have provided definitive evidence of the role of melatonin in the circadian rhythms, plant light/dark responses or the timing mechanisms of plants.12-15

The current research was designed to investigate the potential interactions between melatonin and photosynthetic reactions in plant cells. For these studies, a model system was required that: (1) contains melatonin; (2) is responsive to melatonin signaling; (3) has active, independent and complete photosynthetic apparatus; and (4) allowed for separation of individual cells to reduce cell-to-cell signaling. We developed a model system using fresh water characeae *Chara australis*.

The plants consist of large cells up to 1 mm in diameter and several cm in length. Single cells are easily excised from the plants and contents of cell compartments can be extracted and analyzed. Thus the effects of melatonin can be studied on single cell level. The endogenous melatonin content of cells and cell compartments can be measured and accurate doses of exogenous melatonin can be easily administered. These studies have wide applicability to other plant studies since the *Characeae* are the sister group to the ancestors of all land plants16 and their use as experimental system has already established the fundamentals of plant cell electrophysiology.17

Based on the recently reported findings, we hypothesized that melatonin antioxidant properties facilitate greater photosynthetic efficiency. Our objectives were: (1) to establish that *Chara* sp contain melatonin and respond to melatonin exposure; and (2) to determine the rate and mechanism of melatonin effects on photosynthesis in *Chara*. The results of our study show that melatonin is highly conserved across ancient and modern plants with quantifiable concentrations of melatonin in *Chara*. 

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This is a consequence of the fact that the sum of all the quantum yields $\Phi_{L(PSII)}$, $\Phi_{L(F,D)}$, $\Phi_{L(NPQ)}$ is unity. The measurements were also performed for different intensities of excitation actinic light and the results always followed the same trends (control vs. melatonin-treated cells).

Before the measurement of quenching parameters, the fast O-J-I-P fluorescence transients were also measured for the dark-adapted cells. These measurements revealed that the maximal quantum yield of photochemistry of PSII for dark-adapted cells, $\Phi_{D(PSII)}$, was almost unaffected by melatonin treatment (0.75 for the control and 0.77 for melatonin-treated cells), but the total performance index, $P_{Itot}$, was by about 75% higher in the melatonin-treated cells (5.49) than in the control cells (3.13). The higher $P_{Itot}$ in melatonin-treated cells is in agreement with the higher amount of open PSII centers in these cells as mentioned above. Interestingly (but see discussion), we obtained a lower value of $ABS/RC$ parameter for melatonin-treated cells (1.01) than for the control cells (1.20) indicating that apparent antenna size of PSII was lower in melatonin-treated cells than for the control cells.

### Discussion

**Endogenous melatonin in Characeae.** Melatonin was found in many land plant species and in algae divisions of Chlorophyta, Rhodophyta and Dinoflagellata. We have now measured it in the...
important division of Charophyta in the characeae family, which is thought to have given rise to land plants. Concentrations of melatonin were similar to previously reported for leaf and shoot portions of higher plants such as St. John’s wort,4 Datura25 and Scutellaria28 but lower than has been observed in fruit such as grapes29 or cranberries.30 In contrast, concentrations in Chara were much greater than has been reported for lupin with the highest concentrations of melatonin found in root tissues.30 Tan et al. suggests that melatonin production dates back to purple nonsulfur bacteria and cyanobacteria, the precursors of mitochondria and chloroplasts, respectively.31 Thus finding melatonin in characeae is not a great surprise.

The role of exogenous melatonin in photosynthesis. To our knowledge, there are only two papers in which an effect of melatonin was explored by means of measurements of chlorophyll fluorescence.32,33 Wang et al. subjected detached apple leaves to dark-induced senescence. They found that ΦD(PSII) decreased more gradually in leaves supplied with 10 mM melatonin compared with controls, potentially indicating a role for melatonin in protection of the photosynthetic apparatus.32 Zhang et al. applied melatonin (at concentration of 50, 100, 300 or 500 μM) to cucumber seeds, which were also water-stressed by application of polyethylene glycol (PEG), i.e., melatonin was not applied to unstressed samples.33 Nevertheless the authors obtained very similar results as in our case: very small changes of ΦD(PSII) and of quantum yield of photochemistry of open reactions centers of PSII for light-adapted state, but an increase of qE, and hence also of ΦF(PSII) for the PEG-melatonin-treated samples as compared with the PEG-treated samples.33 The authors also observed a decrease of non-photochemical quenching of excitation energy, expressed by the NPQ parameter, for the PEG-melatonin treated samples.33

In agreement with other studies (reviewed in ref. 34), Zhang et al. also observed an increase of chlorophyll content (per fresh weight) when samples were treated by melatonin33 and Wang et al. and Arnao et al. reported that exogenous application of melatonin slowed the degradation of chlorophyll during senescence of apple52 and barley53 leaves, respectively. These findings are explained in literature as a protective role of melatonin against reactive oxygen species (ROS), which damages (oxidizes) the chlorophylls. However, we have measured a decrease of apparent antenna size of PSII (expressed by the ABS/RC parameter) in melatonin-treated cells, which might indicate a lower amount of chlorophylls in these cells. This seemingly contradictory result can be explained by the fact that ROS does not damage (oxidize) only chlorophylls, but also other components (proteins in general, including PSII and other photosynthetic proteins) of photosynthetic electron transport chain. Thus the application of exogenous melatonin lowers ROS damage of many photosynthetic components. If melatonin offers more protection to PSII proteins than to chlorophyll molecules, the antenna size might appear lower after melatonin exposure. The decreased amount of damaged photosynthetic proteins in melatonin-treated cells is also in agreement with an increased value of the qE parameter in these cells. The qE parameter is a measure of how many reaction centers of PSII are open during the illumination. The amount of open reaction centers reflects the functionality of PSII, as well as functionality of all other photosynthetic proteins. Hence if fewer proteins are damaged (oxidized) by ROS due to protective role of melatonin, more PSII are functional (open) in the light. This explanation is also in agreement with increased value of PItotal parameter in the melatonin-treated cells. To sum up, melatonin offers protection against ROS for chlorophylls as well as for the photosynthetic proteins in general.

Materials and Methods

Measurement of photosynthetic parameters. The excited chlorophyll molecules dispose of the absorbed light energy by: (1) transfer to a reaction center for photosynthetic primary charge separation, (2) fluorescence with longer wavelength; and (3) heat dissipation. The first process and the following photochemical electron transport reactions store energy into chemical bonds. Process (3) can be further divided into ever-present non-regulated heat dissipation (3-a) and regulated additional heat dissipation (3-b) as protection against excess light energy.

Chlorophyll a fluorescence signal was measured by modular version of Dual-PAM-100 (Walz) and by AquaPen (Photon Systems Instruments) fluorometers. Dual-PAM-100 was used for measurements of the quenching analysis by means of the saturation pulse method (reviewed in ref. 18) and the AquaPen was used for measurements of the O-J-I-P fluorescence rise curves (reviewed in ref. 19). To characterize the process: (1) we measured maximal quantum yield of photochemistry of photosystem II (PSII) for dark-adapted state ΦD(PSII)20 and the actual quantum yield of photochemistry of PSII for light-adapted state ΦF(PSII).21 The sum of quantum yield of non-regulated (constitutive) non-photochemical energy quenching by thermal dissipation and of energy quenching by fluorescence emission for light-adapted state Φ(F,D)22 measures processes (2) and (3-a)

The quantum yield of regulated (also called protective) non-photochemical energy quenching by thermal dissipation for light-adapted state Φ∗(NPQ) isolated process (3-b).22 The ABS/RC and PItotal parameters of the so-called JIP test reflect apparent antenna size (amount of functional chlorophyll) of PSII and potential for energy conservation of photons absorbed by PSII to reduction of photosystem I electron acceptors, respectively.23 AquaPen was used for measurement of ΦP(PSII), ABS/RC and PItotal and Dual-PAM-100 for ΦF(PSII), ΦF(F,D) and Φ∗(NPQ).

Melatonin was first diluted in less than 1% methanol and then in artificial pond water (APW in mM: 1.0 NaCl, 0.1 KCl, 0.1 CaCl2) to obtain the final melatonin concentration of 10 μM. Chara cells were treated with melatonin solutions for 20 h in darkness in a glass Petri dish. The cells were exposed to daylight for 1 h before start of the measurements. Both control and melatonin-treated cells were dark-adapted for 5 min. The measurement by AquaPen was performed, followed by the dark-adaptation again and by measurement by Dual-PAM-100. Each measurement was performed on 5–7 internodal Chara australis cells to increase the strength of the signal. Thus each point represents an average value. To evaluate the quenching analysis parameters their time course was followed, with the final steady-state value being

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the most important. In our measurements we assumed that the parameters reached their steady-state values after 3 min illumination. The results were obtained for intensities of excitation actinic and saturation light, respectively, at 70 μmol photons (of red light, λ = 635 nm) m⁻² s⁻¹ (Dual-PAM-100 measurements) and 1,500 μmol photons (of blue light, λ = 450 nm) m⁻² s⁻¹ (AquaPen measurements).

**Measurement of endogenous melatonin.** *Chara* cells were collected from a fresh water pond at Little Bay golf course in April 2011. The plants were more than meter high and the internodal cells about 1 mm in diameter. Sample cells were taken from top, middle and bottom part of the plants and placed into separate labeled containers. Some plants were pulled out carefully from the mud to isolate rhizoids (roots). Samples were brought to the laboratory. The cytoplasm enriched fragments were made by spinning long internodal cells at ~1 g for 30 min. The cytoplasm was moved to one end of the cell and the cells were quickly wilted and the cytoplasm-rich end was tied off with thread. The plants had a lot of male fruiting bodies attached to top branches. Samples included one plant top with fruiting bodies, one without and some fruiting bodies separately. The cells and fragments were taken from their medium and plunged into liquid nitrogen, then transferred into eppendorf tubes and stored in -80°C freezer until harvest and preparation was done over 8 h of daylight without recording the time of day.

The melatonin concentration was measured using previously published methods. In brief, melatonin and serotonin were extracted from frozen samples in complete darkness using only a single red light for safety. Individual samples were accurately weighed in 1.5 ml microfuge tubes, homogenized for 3 min in 100 μl methanol:water:formic acid (80:20:1 v/v) using a cordless motor Pellet Pestle grinder (Kontes) and disposable pestles (Kontes) and centrifuged for 3 min at 16,000 g (VWR, Galaxy 16DH Microcentrifuge) to settle particulate matter. The resulting supernatant was filtered (0.2 μm, Ultrafree-MC filtered centrifuge tubes; Millipore). Compounds were separated using a reverse phase C18 column (Waters BEH C18 column (2.1, 150 mm, 1.7 μm) and elution with 1% aqueous formic acid:acetonitrile (0.0–4.0 min, 95:5–5:95 v/v, 4.0–4.5 min, 95:5–95:5 v/v, 4.5–5.0 min, 95:5 v/v) at 0.25 ml/min using an Acquity ultra performance liquid chromatography (UPLC) at 30°C over a 5 min gradient with a 2 min cleaning and re-equilibration period. Eluted indoleamines were detected by time-of-flight mass spectrometry (ToF-MS; LCT Premier, Micromass, Waters Ltd.) using previously published optimized parameters and quantified by comparison to authenticated standards. The limits of detection (LOD) were 51 pg/μl and 430 pg/μl for melatonin and serotonin, respectively. The lower limits of quantification (LOQ) for melatonin was 543 pg/μl and serotonin had an LOQ of 978 pg/μl. Recovery of spiked melatonin in the sample matrix through the preparation and separation was 98% and recovery of serotonin was 86% as determined at 80% of the highest point in the linear quantification range.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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