The circadian night depression of photosynthesis analyzed in a herb, Pulmonaria vallarsae. Day/night quantitative relationships

Paolo Pupillo1 · Francesca Sparla1 · Bruno A. Melandri1 · Paolo Trost1

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Abstract
Although many photosynthesis related processes are known to be controlled by the circadian system, consequent changes in photosynthetic activities are poorly understood. Photosynthesis was investigated during the daily cycle by chlorophyll fluorescence using a PAM fluorometer in Pulmonaria vallarsae subsp. apennina, an understory herb. A standard test consists of a light induction pretreatment followed by light response curve (LRC). Comparison of the major diagnostic parameters collected during day and night showed a nocturnal drop of photosynthetic responses, more evident in water-limited plants and consisting of: (i) strong reduction of flash-induced fluorescence peaks (FIP), maximum linear electron transport rate ($J_{\text{max}}$, $\text{ETR}_{\text{EM}}$) and effective PSII quantum yield ($\Phi_{\text{PSII}}$); (ii) strong enhancement of nonphotochemical quenching ($\text{NPQ}$) and (iii) little or no change in photochemical quenching ($q_P$), maximum quantum yield of linear electron transport ($\Phi$), and shape of LRC ($\theta$). A remarkable feature of day/night LRCs at moderate to high irradiance was their linear-parallel course in double-reciprocal plots. Photosynthesis was also monitored in plants subjected to 2–3 days of continuous darkness (“long night”). In such conditions, plants exhibited high but declining peaks of photosynthetic activity during subjective days and a low, constant value with elevated $\text{NPQ}$ during subjective night tests. The photosynthetic parameters recorded in subjective days in artificial darkness resembled those under natural day conditions. On the basis of the evidence, we suggest a circadian component and a biochemical feedback inhibition to explain the night depression of photosynthesis in P. vallarsae.

Keywords Circadian rhythm · Chlorophyll fluorescence · Light response curve · Photosynthesis · Pulmonaria · Shade plants · Water stress

Abbreviations
Abs  Absorptivity
Chl  Chlorophyll
ETR  Photosynthetic electron transport rate
$\text{ETR}_{\text{EM}}$  Maximum extrapolated value of ETR
FIP  Flash-induced fluorescence peak
$F^m$  Maximum Chl fluorescence in dark-adapted plants
$F^m$  Maximum relative fluorescence under illumination
$F^o$  Minimum Chl fluorescence in dark-adapted plants
Ft  Current Chl fluorescence
Fv  Variable fluorescence
FvCB  Equation of Farquhar et al. 1980
$\Phi$  Quantum yield of photosynthesis (Farquhar et al. 1980)
$\Phi_{\text{PSII}}$  Effective quantum yield of PSII photochemistry
$J_{\text{max}}$  Maximum extrapolated rate of LRC (Farquhar et al. 1980)
$K_{\text{PAR}}$  Half-saturation PAR for an LRC
LRC  Light response curve
$\text{NPQ}$  Nonphotochemical quenching
PAM  Pulse amplitude modulated (fluorometer)
PAR  Photosynthetically active radiation
PPDF  Photosynthetic photon density flux
PSII  Photosystem II
$q_P$  Photochemical quenching
ROS  Reactive oxygen species
SD  Standard deviation
$\theta$  LRC smoothing factor (Farquhar et al. 1980)

* Francesca Sparla
francesca.sparla@unibo.it

1 Department of Pharmacy and Biotechnology, University of Bologna Alma Mater, Via Irnerio 42, 40126 Bologna, Italy
Introduction

A decrease of photosynthesis potential during night hours, including a drop of photosynthetic activity under artificial illumination that reproduces daily conditions, is long known. This effect, here defined as the circadian night depression of photosynthesis, was assigned to the expanding domain of circadian events following early reports on rhythmic photosynthetic activity in plants (Pallas et al. 1974; Kerr et al. 1985; Kloppstech 1985). However, most studies were focussed on circadian stomatal movements (Martin and Meidner 1971; Holmes and Klein 1986) and it was generally agreed that the daily changes in photosynthetic activity depended on control of leaf intercellular pCO₂ by stomata (Raschke 1975; Cornic and Briantais 1991). Therefore, the nature and very existence of biochemical oscillations of photosynthetic potential during a daily cycle and independent of light was in doubt, and to some extent remains so.

Plants are subject to a daily cycle of light and dark which has its own effects on metabolism regardless of the biological clock (Zaffagnini et al. 2019). These effects are largely explained in terms of reduction of protein disulfides during light-to-dark transitions and vice-versa, resulting in fine tuning of chloroplast enzyme activities and regulation of photosynthesis on multiple levels (Michelet et al. 2013; Yoshida et al. 2019). Electrons for disulfide reduction are provided either by the photosynthetic linear electron transport chain via thioredoxins f/m, or by the oxidative pentose phosphate pathway via NADPH thioredoxin reductase C (Cejudo et al. 2019; Zaffagnini et al. 2019). Thioredoxins are also involved in the ROS-dependent, dark oxidation of targets (Yochochi et al. 2021). The light/dark and circadian regulations of photosynthesis are distinct but strictly cooperating and overlapped under natural daily cycles, and influence each other with mechanisms that are not yet fully understood (Dodd et al. 2014).

Evidence has accumulated for a circadian control of many aspects of the photosynthesis process (Hennessey et al. 1993; Nakahira et al. 1998; Dodd et al. 2014). The use of transgenic luciferase as reporter enzyme (Millar et al. 1992; Matsuo et al. 2006; Philippou et al. 2019) revealed the night turnover of Cab polypeptides and other components of the photosystems under circadian control, with nuclear-encoded sigma factors as important mediators (Belbin et al. 2017). Delayed Chl fluorescence has been shown to oscillate according to a circadian rhythm in several plants (Gould et al. 2009). Moreover, the circadian clock underpins the synthesis of Rubisco small subunit, Rubisco activase and other photosynthetic enzymes (Martino-Catt and Ort 1992; Liu et al. 1996; Farré and Weise 2012) as well as components of the photosynthetic pathway (McClung et al. 2000), and drives post-translational modifications of chloroplast proteins (Booij-James et al. 2002; Söe and Mas 2014). It appears, therefore, that most stages of the photosynthesis process are under circadian control (Millar 2016). There are good reasons, therefore, to suspect that the night depression of photosynthesis investigated here is not merely a matter of stomata and may include a basic biochemical component (Hennessey et al. 1993), the more so as the circadian assimilatory response is independent of the stomatal cycle in arabidopsis mutants (Dodd et al. 2004).

Little information exists about circadian effects on photosynthesis of plants in nature (Resco de Dios et al. 2012; Millar 2016). Field research is made difficult by intricate relationships of the circadian clock with stomatal responses and the plant and soil water status (Flexas and Medrano 2002; Daszkowska-Golec and Szarejko 2013), as well as light, temperature and other environmental cues (Martino-Catt and Ort 1992; Jones et al. 1998). Moreover, plant sugars and photosynthetic products are known to modulate the circadian oscillator (Haydon et al. 2013; Frank et al. 2018); in particular, sucrose supports the robustness of circadian responses in light and darkness (Philippou et al. 2019). Metabolite effects can probably explain the extreme transiency of photosynthetic circadian responses in dark-kept plants (Hennessey et al. 1993; Nakahira et al. 1998; Booij-James et al. 2002).

In a previous paper (Recchia et al. 2017) we applied slow Chl fluorescence analysis using the sensitive Imaging PAM 2000 instrument to investigate inverse relationships between intensity of photosynthesis and length of the lifecycle in perennial understory herbs. By the same technique, we here present a fluorescence kinetic study of the night depression of photosynthesis and of prolonged darkness (“long night”) in one of these species, the lungwort Pulmonaria vallarsae apennina, a relatively shade tolerant herb of Italian uplands. This plant lends itself well to this type of experiments thanks to its long vegetation period and the large, enduring leaves that can undergo repeated PAM tests unharmed. The results are consistent with control of photosynthesis in this species by the circadian system under mild water stress, and suggest a biochemical limitation as the main factor responsible for the circadian night depression of photosynthesis.

Materials and methods

Plant growth

The experiments were carried out on two dozens of plants of Pulmonaria vallarsae Kern subsp. apennina Cecchi et Selvi (Italian lungwort, family Boraginaceae), a perennial understory herb, relatively shade tolerant and widely distributed on the Italian peninsula along Apennine reliefs (Cecchi 2015). The plants were collected in several hilly sites of
the Bologna district in summer 2018 and grown in 25 cm diameter, 27 cm high pots on clay compost soil with watering every 3 days in a naturally lit greenhouse under partial shading at the Orto Botanico of the University of Bologna.

**Chlorophyll fluorescence analysis and related experimental procedures**

Chl a slow fluorescence was assayed with the Imaging PAM 2000 fluorometer (Walz Gmbh, Effeltrich, Germany) and associated PIMR software via a dedicated computer as described (Recchia et al. 2017). One area of interest (3 mm diameter) was chosen on the lower (abaxial) surface of the leaf under study, which was held in place by a horizontal clip of 26 × 34 mm. PAM fluorometer tests were performed in laboratory or in greenhouse during late spring and summer months of four years (July 2018 to September 2021). Nighttime tests were carried out in darkness. For long experiments, plants were either transferred to a small laboratory greenhouse or to a nearby dark closet for the time required. Water was usually withheld for 2–3 days in advance of each trial and for its whole duration, since lavishly irrigated plants exhibit reduced night depression of photosynthesis (and reduced stomatal responses, Raschke 1975; Lawson et al. 2010).

Prior to each test, consisting of an activating light pretreatment followed by a light response curve (LRC) plants were held in the dark for 15 min. Minimum fluorescence in dark-adapted samples ($F_\text{o}$) was then obtained by applying measuring light of low frequency (1 Hz) and maximum fluorescence in dark-adapted samples ($F_\text{m}$) was obtained through an 800 ms blue saturation pulse of 2.4 mE m$^{-2}$ s$^{-1}$. The maximum quantum efficiency of PSII ($F_{\text{v}}/F_\text{m}$) was constant in all samples. Absorptivity was determined by the instrument ($\text{Abs} = 1 - \text{R/NIR}$) before each test.

The light pretreatment was conducted at actinic light of 100 µE m$^{-2}$ s$^{-1}$ with saturation pulses of 2.4 mE m$^{-2}$ s$^{-1}$ (duration 800 ms each) applied at 20 s intervals. This triggered 16 fluorescence peaks ($\text{FIP} = F'_m - F_t$) of variable height, i.e. ~4% to 42% of $F_\text{m}$ depending on the state of the plant (see Fig. 1). Accordingly, FIPs are used in this work as a probe of PSII activity. Other fluorescence parameters

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**Fig. 1** Activating light pretreatments (A, B) and subsequent LRCs (C, D) in a day/night experiment on a single *P. vallarsae* plant. A, morning test. Time-courses of $F_t$ (with FIPs) continuous trace; $\Phi_{\text{PSII}}$ yellow dots; $qP$ green dots; $\text{NPQ}$ red dots. B, the same leaf tested at evening. C, LRCs: kinetics of parameters $qP$ (cyan and blue dots) and $\Phi_{\text{PSII}}$ (green and grey); D, concomitant changes of $F'_m$ (yellow and orange) and $\text{NPQ}$ (violet and pink). All values relative to $F_\text{m} = 1.0$, except that $\text{NPQ}$ is divided by 4.

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were derived as follows: the effective quantum yield of PSII photochemistry is \( \Phi_{PSII} = (F'_{m} - F_{t})/F'_{m} \), nonphotochemical quenching \( NPQ = (F'_{m} - F_{m})/F'_{m} \), photochemical quenching \( qP = (F'_{m} - F_{t})/(F'_{m} - F'_{o}) \); photosynthetic electron transport rate \( ETR = \Phi_{PSII} \times PPFD \times 0.5 \times Abs \) (Björkman and Demmg 1987).

LRCs of electron transport rates (ETRs) were determined right after light pretreatments. LRCs were conducted at increasing steps of actinic light (from 1 to 460 \( \mu E \ m^{-2} \ s^{-1} \)). The duration of each step was 20 s and ended with a saturation pulse of 2.4 mE m\(^{-2} \) s\(^{-1} \). Light fluorescence related parameters \( \Phi_{PSII} \), \( qP \), \( NPQ \) and ETR were calculated as above by PAM fluorometer software.

In day/night experiments plants were usually assayed around 08.30 am and 11.20 pm, and in “long night” experiments plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h. The total number of experiments performed was 252 with about 1300 single plants were assayed every 12 h.

Data processing

LRCs were analyzed by nonlinear regression (CoHort software) using the function of the Farquhar-von Caemmerer-Berry (FvCB) model (Farquhar et al. 1980; Farquhar and Wong 1984) yielding parameters \( J_{\text{max}} \), \( \Phi \) and \( \theta \). In the FvCB model, \( J_{\text{max}} \) is the calculated rate of linear electron transport under saturating irradiance, \( \Phi \) is the maximum quantum yield of the linear electron transport and \( \theta \) is a curvature factor. LRCs were also represented in double-reciprocal plots with linear responses at moderate to high irradiances (PPFD > 100 \( \mu E \ m^{-2} \) s\(^{-1} \)), corresponding to rectangular hyperbolas in primary plots. When fitted to straight lines, LRC data give an estimate of \( ETR_{EM} \) (extrapolated maximum ETR for the high irradiance region of an LRC) and \( K_{PAR} \) (the PPFD at which \( ETR = ETR_{EM}/2 \)). \( ETR_{EM} \) approximated \( J_{\text{max}} \) with deviations within +15% of \( J_{\text{max}} \) values. Other fluorescence and photosynthetic parameters are provided by fluorometer-associated PIM software.

Results

Photosynthetic activity by day and night was investigated in leaves of potted, water-limited plants of \textit{Pulmonaria vallarsae} on the basis of slow Chl fluorescence responses, recorded and elaborated by the Imaging PAM fluorometer. A standard test consisted of an activating light pretreatment followed by an LRC, derived from the fluorescence induced by a series of light steps of increasing energy.

Two standard tests conducted on the same \textit{P. vallarsae} leaf by day and night are shown in Fig. 1 as a representative example. The same type of experiments was repeated on many plants, and the average values of fluorescence related parameters recorded at the end of the activating light pretreatments are listed in Table 1. In the morning (Fig. 1A), the leaf illuminated by actinic light of 100 \( \mu E \ m^{-2} \) s\(^{-1} \), responded to pulses of saturating light with a series of FIPs which soon reached their maximum height at 34% of \( F'_{m} \) (average value 26.2%; see Table 1). Simultaneously, \( qP \) and \( \Phi_{PSII} \) reached steady state values, current fluorescence \( F_{t} \) rapidly dropped and partially recovered, and the dissipation parameter \( NPQ \) made a jump followed by stabilization at a low level (Fig. 1A).

A quite different pattern was observed when the light induction pretreatment was administered during the night (Fig. 1B and Table 1). The most conspicuous difference consisted in small FIPs (10.1% of \( F'_{m} \) on average; see Table 1) which remained constant or nearly so during the whole pretreatment (low-photosynthesis state). At the same time there was a rapid rise of \( NPQ \) to an elevated, stable plateau symmetrical to a large drop of \( F_{t} \), while \( \Phi_{PSII} \) increased to its maximum which was significantly lower than during day. Photochemical quenching \( qP \) rose with slow kinetics to the same final value as in the morning (Fig. 1B). In sum, at the end of the light activating pretreatment, the sample measured during the day was characterized by higher values of FIPs (2.6-fold) and \( \Phi_{PSII} \) (1.4-fold) with respect to the night, while the sample measured during the night was typified by a higher level of \( NPQ \) (3.0-fold). On the other hand, the \( F_{t}/F'_{m} \) parameter commonly used to characterize

| Day | Night | Day/Night ratio |
|-----|-------|-----------------|
| \( F_{t} \), | 0.33 ± 0.03 | 0.22 ± 0.03 | 1.50 |
| FIP, | 0.26 ± 0.04 | 0.10 ± 0.01 | 2.58 |
| \( NPQ \), | 0.69 ± 0.13 | 2.08 ± 0.31 | 0.33 |
| \( \Phi_{PSII} \), | 0.44 ± 0.03 | 0.32 ± 0.03 | 1.38 |
| \( qP \), | 0.72 ± 0.06 | 0.69 ± 0.08 | 1.04 |
| \( J_{\text{max}} \), | 23.1 ± 1.2 | 13.2 ± 1.2 | 1.75 |
| \( \Phi \), | 0.155 ± 0.011 | 0.144 ± 0.015 | 1.07 |
| \( \theta \), | 0.76 ± 0.13 | 0.70 ± 0.15 | 1.09 |
| \( ETR_{EM} \), | 26.8 ± 1.8 | 15.1 ± 1.2 | 1.77 |
| \( K_{PAR} \), | 106 ± 13 | 50 ± 10 | 2.11 |

Table 1 Means ± SD (n = 73) of parameters \( F_{t} \), FIP, \( NPQ \), \( qP \), \( \Phi_{PSII} \) (A) and parameters \( J_{\text{max}} \), \( \Phi \), \( \theta \), \( ETR_{EM} \), \( K_{PAR} \) (B) for day/night PAM tests performed with \textit{P. vallarsae} plants under natural light/dark illumination. Group A data are final values of the activating light pretreatment, group B data are calculated from the LRCs. Units as in Figs. 1–2, but \( \Phi \) and \( \theta \) are calculated by FvCB equation.
stress subjected plants did not vary at all between day and night (not shown).

The pretreatments were followed by LRC tests at increasing light intensities. Trends of Chl fluorescence related parameters as a function of PPFD are reported in Fig. 1C-D. During the day, the slow descent of $F'_m$ with increasing irradiance was symmetrical to slowly rising $NPQ$, while $\Phi_{PSII}$ and $qP$ declined linearly. In the night test, most parameters were significantly lowered in sharp contrast to a quick, strong rise of $NPQ$. $F'_m$ underwent an abrupt biphasic fall, and the declining $qP$ trajectory showed a distinct night hump around 80–100 $\mu$E m$^{-2}$ s$^{-1}$ PPFD.

For both day and night tests, ETRs approximated a linear dependence on PPFD at low light intensities and an apparent hyperbolic response at higher light intensities (Fig. 2A). The linear responses in the low light range were very similar in day and night, while at high light intensities the hyperbolic responses diverged, with the day curve reaching much higher saturation levels. ETR data of each test were interpolated with the FvCB equation (Fig. 2A) allowing calculation of the relative parameters (Table 1). Mean LRCs for 73 day/night twin experiments are graphically summarized in Fig. 2A, while Fig. S1 reports single plant responses, with small differences between individual plants. The maximal ETR estimated as $J_{max}$ was on average 1.75-fold higher in day tests with respect to night tests, whereas the variations of the maximum quantum yield $\Phi$ and the curvature factor $\theta$ were very small (1.07 and 1.09, respectively). ETR data at moderate to high light intensities (PPFD > 100 $\mu$E m$^{-2}$ s$^{-1}$) could be interpolated by straight lines in double-reciprocal plots (Fig. 2B) with extrapolated maximum values defined as ETR$_{EM}$ having an overall day/night ratio of 1.77, similar to $J_{max}$. Like $J_{max}$ and ETR$_{EM}$, the half-saturation parameter of the LRC ($K_{PAR}$) was also higher during the day (Table 1). The near parallelism of double-reciprocal plots implies that ETR$_{EM}$ and $K_{PAR}$ tend to keep a similar ratio of the respective day/night values.

The daily trends of photosynthetic parameters were examined on the basis of 141 tests carried out at different times of the day, normalized for 05.30 am sunrise and represented at 2 h intervals in Fig. 3. Maximum rates of electron transport ($J_{max}$) showed a broad diurnal, almost symmetrical peak at 12–14 pm (Fig. 3A). The nocturnal value of $J_{max}$ was low and nearly constant between 9 pm and about 4.30 am and was followed by a sudden rise around dawn, indicating a very early onset of recovery of photosynthetic potential. This was confirmed by tests conducted before and after dawn on the same leaf in the absence of light (not shown). The FIP profile featured a sudden increase around sunrise with a sluggish ramp around 12–15 pm, and a quick drop in late afternoon and night (Fig. 3B), when $NPQ$ prevails (Fig. 1D).

The pre-dawn enhancement of activity observed in several experiments suggested that changes of photosynthetic potential can occur in the absence of light in *P. vallarsae*, so the effects of prolonged darkness were probed (“long night” trials). Plants grown under natural light–dark cycles were tested at day 0 at morning (0-d) and evening (0-n), then transferred to constant darkness and assayed in the dark every 12 h at each subjective morning and night, and finally brought back to the light once the morning $J_{max}$ value had dropped to nocturnal levels. Pooled results of 36 experiments, each lasting 48–72 h, are summarized in Fig. 4, which documents the oscillatory trajectory of most parameters during a “long night” and the transiency of the effect.

In detail, FvCB interpolation of the ETR data shows that the maximum electron transport rate ($J_{max}$) oscillates with circadian rhythmicity, with statistically supported day/night differences (Fig. 4 and Fig. S2). In the first subjective day
and \( \Phi_{\text{PSII}} \) to a lesser extent, oscillated in phase with \( J_{\text{max}} \). Pooled points over 2 h showed low values during the nights and attenuation of circadian oscillation towards the end of the experiments. By the same leaf, \( \Phi_{\text{max}} \) was very low in the daytime (at \( 0-d \)) and high and oscillating in subsequent darkness.

**Fig. 3** Daily courses of parameters \( J_{\text{max}} \) A and FIP B under natural light–dark cycle. Data of tests conducted in summer months of years 2018–2020, normalized to 05:20 am sunrise corresponding to July 20th in Bologna (\( n = 141 \)). Each bar represents mean \( \pm \) SD of 6 to 32 pooled points over 2 h.

in darkness (\( I-d \)), the \( J_{\text{max}} \) value (like \( \text{ETR}_{EM} \)) was fully recovered to the level of the previous morning in the light (\( 0-d \)). The effect was attenuated during the second subjective day (\( 2-d \)) but still visible if compared to night values, that were relatively constant over the whole experiment. No circadian rhythmicity affected parameters \( \theta \) and \( \Phi \), even though the latter was significantly lower during the whole “long night” with respect to the usual light/dark conditions (Fig. 4C) suggesting a light-dependent component of \( \theta \). The fluorescence peaks FIPs, too, while exhibiting a robust rhythm were 24% lower (at \( I-d \)) during the “long night” than under normal light regime (\( 0-d \), Fig. 4D). In fact FIP, and \( \Phi_{\text{PSII}} \) to a lesser extent, oscillated in phase with \( J_{\text{max}} \) showing low values during the nights and attenuation of circadian oscillation towards the end of the experiments. By contrast, \( qP \) showed only small, insignificant oscillations (not shown). \( NPQ \) behaved exactly symmetrical to \( J_{\text{max}} \), FIP and \( \Phi_{\text{PSII}} \) with opposite phasing; it was very low in the light (at \( 0-d \)) and high and oscillating in subsequent darkness. Despite rapid damping, these results indicate that most photosynthesis and fluorescence parameters of darkened \( P. \) \textit{vallarsae} leaves undergo oscillations of different strength between subjective days and subjective nights. Data within and between experiments also evidence a remarkable stability of \( J_{\text{max}} \), FIP and \( \Phi_{\text{PSII}} \) in the night and, conversely, of \( NPQ \) in the day (Fig. 4). Interestingly, the general features of the photosynthetic responses recorded for \( J_{\text{max}} \), FIP, \( \Phi_{\text{PSII}} \) and \( NPQ \) in subjective days and subjective nights were similar to responses in a normal light/dark cycle (compare Table 1 and Fig. 4).

Finally, an illustrative “long night” experiment is reported in Fig. 5A to highlight the oscillatory trends of photosynthetic electron transport, showing three selected LRCs in constant darkness (stages \( 1-d, 1-n, 2-n \)) and the final LRC after a brief recovery of the plant in the light (\( 3-d \)), all borne by the same leaf. Double-reciprocal plots (Fig. 5B) display straight/parallel patterns of the LRCs arranged in two distinct bundles for subjective day (high-photosynthesis state) and night (low-photosynthesis state). In fact, the ratios of subjective morning \( \text{ETR}_{EM} \) and \( K_{PAR} \) values to the respective nighttime values were almost identical in this and similar experiments, both parameters giving a day/night ratio of 2.2 for the first day in darkness (\( I-d/I-n \)). This also confirms the autonomy of most circadian photosynthesis responses from the presence of light during the first 1–2 days of darkness.

**Discussion**

Daily changes of photosynthetic activity were investigated by Chl fluorescence analysis using the Imaging PAM 2000 fluorometer in \textit{Pulmonaria vallarsae} subsp. \textit{apennina}, a perennial understory herb (lungwort) of Italian mountains. Thanks to the noninvasive PAM technique, changes of photosynthesis and fluorescence linked parameters of a single leaf could be followed for quite a time with repeated tests. Under natural light/dark cycles, LRCs taken by night in slightly water-stressed plants showed a strong depression of photosynthesis with respect to daytime values, somewhat recalling the response of heat exposed leaves (Sukhova et al. 2018). The night effect was smaller or even absent in fully watered plants, consequently only slightly water-stressed plants were used for the present experiments. In particular, night LRCs showed reduced electron transport rates at medium to high irradiances and little change at low irradiances. In terms of photosynthetic parameters this effect resulted in low nocturnal values of electron transport (\( J_{\text{max}} \) or \( \text{ETR}_{EM} \)), but constant or very small changes in maximum quantum yield (\( \phi \)), shape of the curve (\( \theta \)) and photochemical quenching \( qP \). Nonphotochemical quenching \( NPQ \) was strongly enhanced in night tests, thus probably contributing to a strong decrease of fluorescence (FIP, \( F_{I} \)) and a moderate decrease of effective PSII quantum yield (\( \Phi_{\text{PSII}} \)).
There is a plethora of biochemical impediments with potential damping effects on photosynthetic responses during night. The night blockade of Rubisco by active site decarbamylation and tight binding of inhibitory sugar phosphates (Taylor et al. 2022), and of Calvin-Benson cycle enzymes through redox post-translational modifications and formation of inactive complexes (Michelet et al. 2013; Gurrieri et al. 2021; Zimmer et al. 2021) might curtail the supply of substrates and acceptors for the linear electron transport and ATP synthase, the latter being itself dark-inactivated in an oxidizing cellular milieu (Tezara et al. 2002; Hisabori et al. 2013; Vaseghi et al. 2018; Hashida et al. 2018). This, in turn, might cause a nighttime feedback inhibition of PSII activity and photosynthetic potential, with an enhanced high energy state of thylakoids when illuminated, and consequent induction of energy dissipation mechanisms at the level of PSII antenna complexes (Laisk et al. 1992; Bassi and Dall’Osto 2021). This sequence of events is consistent with the high NPQ, low fluorescence responses and decreased ETR and ΦPSII. It is also consistent with the invariancy of the maximum quantum yield (Φ) of linear electron transport at low light intensities, with minimized NPQ and no metabolic control. It seems likely, therefore, that the night drop of most photosynthetic responses results from a persistent decrease of some key metabolite(s) and/or inactivation of ATP synthase and Calvin-Benson cycle enzymes, including Rubisco. It is also possible that a CO₂ limitation of Rubisco activity.

Fig. 4 Average results of “long night” experiments for LRC parameters $J_{\text{max}}$ A, $\Phi$ B, $\theta$ C of the FvCB function, and fluorescence related parameters FIP D, NPQ E and $\Phi_{\text{PSII}}$ F, means ± SD ($n=36$). Plants under natural light–dark regime were tested at ~11 am (0-d) and ~11 pm (0-n) of day 0, then transferred to constant darkness for 2 days (top black bar) and tested each day at 11 am (1-d, 2-d) and 11 pm (1-n, 2-n) in the dark. Two-tailed t-test was used for statistics: *** $p<0.0001$; ** $p<0.001$; * $p<0.05$.
due to stomata closure in the nighttime would reinforce this cascade of events.

Morning photosynthetic recovery in *P. vallarsae* does not depend on light in the short run and can be reiterated in constant darkness using an artificial “long night” of 2(-3) days, only interrupted by short tests every 12 h. Under these conditions, the photosynthetic activity during subjective days (i.e. in artificially darkened mornings) had the same features as the photosynthetic activity observed in naturally illuminated mornings: high levels of *J* max, *Φ* PSII and FIP, and low *NPQ*. Likewise, photosynthesis during subjective nights was depressed as under a normal light/dark cycle. The oscillatory changes of photosynthetic potential during a 52–72 h “long night” may be described as a daytime boost of photosynthetic activity (damping from the second day) over a low and relatively stable nighttime level of activity. The latter is presumably of metabolic origin since sugars and photosynthates are known to sustain the circadian rhythm (Haydon et al. 2013; Frank et al. 2018; Philippou et al. 2019).

A circadian oscillation of photosynthetic potential was sporadically described in different plants including green algae with different measuring techniques (Pallas 1974; Longergan 1981; Fredeen et al. 1991; Hennessey and Field 1991; Dodd 2014; Noordally and Millar 2015). Stomatal movements too are regulated by the circadian system (Martin and Meidner 1971; Holmes and Klein 1986) and there is little doubt that the present biochemical events are accompanied, to some extent, by stomatal responses. Circadian rhythms of stomatal movement, hydraulic conductance and growth are enhanced by drought (Caldeira et al. 2014), and we also observed that water abundant lungworts underwent a smaller drop of night photosynthetic potential, or none, compared to mildly water-stressed plants that had been grown under sparing irrigation. It thus appears that water stress enhances a number of plant circadian functions more or less related to photosynthesis. Early observations of circadian regulation of photosynthetic potential under constant levels of internal CO₂ (Hennessey and Field 1991; Fredeen et al. 1991) also supported the view that variable stomata conductance could not be the only determinant of the night depression of photosynthesis (Farré and Weise 2012). Nocturnal stomatal opening with high gas conductance is correlated with elevated photosynthesis performances and growth rates in several species, and has likely a genetic background (Caird et al. 2007; Resco de Dios et al. 2016).

The circadian nature of the night depression of photosynthesis is confirmed by the recovery process initiating at dawn with or without light, reminiscent of reports on early synthesis of antenna (Kloppstech 1985; Millar et al. 1992) and other plastid proteins (Dodd et al. 2014). A circadian rhythm of delayed Chl fluorescence under constant light has been recorded in arabidopsis and other species (Gould et al. 2009), also suggesting that the night depression effect involves photosynthetic functions besides stomata (see Dodd et al. 2004; Lawson et al. 2010). The expression of thioredoxins *f* and *m* mediating the light-dependent activation of Calvin-Benson cycle enzymes and ATP synthase is under circadian control (de Dios Barajas-López et al. 2011). Reactive oxygen species (Lai et al. 2012) and the redox state of 2-cys peroxiredoxins (Edgar et al. 2012), both potentially implied in the dark oxidation of redox-regulated proteins (Vaseghi et al. 2018), do also oscillate in arabidopsis leaves with circadian rhythm.

Circadian regulation of photosynthesis is mainly suggested in this paper by the recovery of a high photosynthesis state in subjective days in the absence of light. In fact, the high photosynthesis state established in either subjective or real days are similar and might in principle be achieved through analogous mechanisms. The light activation of  

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**Fig. 5** LRCs from a “long night” trial. **A**, a plant growing under natural light–dark regime was transferred at 08.00 pm of day 0 to constant darkness for 63 h and assayed every 12 h in darkness (see Fig. 4), then readmitted to light and watered, with a final test after 4 h light exposure (3-d). Four LRCs (1-d, 1-n, 2-n, 3-d) out of seven are shown for clarity. **B**, same data treated as double-reciprocal plots in the high light range (100–460 µE m⁻² s⁻¹ PPFD).
Photosynthetic metabolism is largely explained in terms of thioredoxin-mediated reduction of Calvin-Benson cycle enzymes (and related regulatory proteins Rubisco activase, CA1P phosphatease, CP12 and ATP synthase (Portis et al. 2008; Michelet et al. 2013; Hisabori et al. 2013; Gurrieri et al. 2021). In chloroplasts, thioredoxins and thioredoxin-regulated proteins are photosynthetically reduced by PSI through ferredoxin: thioredoxin reductase, or nonphotosynthetically by NADPH thioredoxin reductase C using NADPH derived from the oxidative pentose phosphate pathway (Zaffagnini et al. 2019). Light independent activation of redox-regulated chloroplast proteins is therefore possible, as also suggested by the high sensitivity of NTRC knockout mutants to prolonged darkness (Cejudo et al. 2019). The light-dependent and clock dependent regulation of photosynthesis do clearly interact, but whether they share a common mechanism of action still needs to be thoroughly investigated.

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Author contributions PP designed the study and performed the experiments; PP, PT, BAM and FS analysed the data and wrote the paper. All authors read, corrected and accepted the final manuscript.

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Declarations

Conflict of interest The authors have no conflict of interest to declare that are relevant to the content of this article.

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References

Bassi R, Dall’Osto L (2021) Dissipation of light energy absorbed in excess: the molecular mechanisms. Annu Rev Plant Biol 72:47–76. https://doi.org/10.1146/annurev-arplant-071720-015522
Belbin FE, Noonally ZB, Wetherill SJ, Atkins KA, Franklin KA, Dodd AN (2017) Integration of light and circadian signals that regulate chloroplast transcription by a nuclear-encoded sigma factor. New Phytol 213:727–738. https://doi.org/10.1111/nph.14176
Björkman O, Demmig B (1987) Photon yield of O2 evolution and chlorophyll fluorescence at 77°K among vascular plants of diverse origins. Planta 170:489–504. https://doi.org/10.1007/BF00402983
Booij-James IS, Swegle WM, Edelman M, Mattoo AK (2002) Phosphorylation of the D1 photosystem II reaction center protein is controlled by an endogenous circadian rhythm. Plant Physiol 130:2069–2075. https://doi.org/10.1104/pp.013441
Caird MA, Richards JH, Donovan LA (2007) Nighttime stomatal conductance and transpiration in C3 and C4 plants. Plant Physiol 143:4–10. https://doi.org/10.1104/pp.106.092940
Caldeira C, Jeanguenin L, Chaumont F, Tardieu F (2014) Circadian rhythms of hydraulic conductance and growth are enhanced by drought and improve plant performance. Nat Commun 5:5365. https://doi.org/10.1038/ncomms6365
Cecchi L, Selvi F (2015) Synopsis of Boraginaceae subfam. Boragineae tribe Boragineae in Italy. Plant Biosystems 149:630–677. https://doi.org/10.1080/11263504.2015.1057261
Cejudo FJ, Ojeda V, Delgado-Requerey V, González M, Pérez-Ruiz JM (2019) Chloroplast redox regulatory mechanisms in plant adaptation to light and darkness. Front Plant Sci 10:380. https://doi.org/10.3389/fpls.2019.00380
Cornic G, Briantais J-M (1991) Partitioning of photosynthetic electron flow between CO2 and O2 reduction in a C3 leaf (Phaseolus vulgaris L.) at different CO2 concentrations and during water stress. Planta 183:178–184. https://doi.org/10.1007/BF00197786
Daszkowska-Golec A, Szarejko I (2013) Open or close the gate – stomata action under the control of phytohormones in drought stress conditions. Front Plant Sci 4:1–16. https://doi.org/10.3389/fpls.2013.00138
de Dios B-López J, Serrato AJ, Cazalis R, Meyer Y, Chueca A, Reichheld JP, Sahrawy M (2011) Circadian regulation of chloroplastic f and m thioredoxins through control of the CA1 transcription factor. J Exp Bot 62:2039–2051. https://doi.org/10.1093/jxb/erq394
Dodd AN, Parkinson K, Webb AAR (2004) Independent circadian regulation of assimilation and stomatal conductance in the ztl-1 mutant of Arabidopsis. New Phytol 162:63–70. https://doi.org/10.1111/j.1469-8137.2004.01005.x
Dodd AN, Kusakina J, Hall A, Gould PD, Hanaoka M (2014) The circadian regulation of nitrogen assimilation and full-day stomatal conductance. J Exp Bot 65:1071–1080. https://doi.org/10.1093/jxb/eru450
Dodd AN, Kusakina J, Hall A, Gould PD, Hanaoka M (2014) The circadian regulation of assimilation and stomatal conductance in the ztl-1 mutant of Arabidopsis. New Phytol 162:63–70. https://doi.org/10.1111/j.1469-8137.2004.01005.x
Dodd AN, Kusakina J, Hall A, Gould PD, Hanaoka M (2014) The circadian regulation of nitrogen assimilation and full-day stomatal conductance. J Exp Bot 65:1071–1080. https://doi.org/10.1093/jxb/eru450
Edgar R, Green E, Zhao Y et al (2012) Peroxiredoxins are conserved markers of circadian rhythms. Nature 485:459–464. https://doi.org/10.1038/nature11088
Farquhar GD, Wong SC (1984) An empirical model of stomatal conductance. Austral J Plant Physiol 11:191–210. https://doi.org/10.1071/PP9840191
Farquhar GD, von Caemmerer S, Berry JA (1980) A biochemical model of photosynthetic CO2 assimilation in leaves of C3 species. Planta 149:78–90. https://doi.org/10.1007/BF00386231
Farré EM, Weise SE (2012) The interactions between the circadian clock and primary metabolism. Curr Opin Plant Biol 15:293–300. https://doi.org/10.1016/j.copb.2012.01.013
Flechas J, Medrano H (2002) Drought-inhibition of photosynthesis in C3 plants, stomatal and non-stomatal limitations revisited. Ann Bot 89:183–189. https://doi.org/10.1093/aob/mcf027
Frank A, Matioli CC, Viana AJC, Hearns TJ, Kusakina J, Belbin FE, Wells Newman D, Yochikawa A, Cano-Ramirez DL, Chembath A, Cragg-Barber K, Haydon MJ, Hotta CT, Ventinzi M, Webb AAR, Dodd AN (2018) Circadian entrainment in Arabidopsis by the sugar-responsive transcription factor bZIP63. Curr Biol 28:2597–2606. https://doi.org/10.1016/j.cub.2018.05.092
Fredeen AL, Hennessey TL, Field CB (1991) Biochemical correlates of the circadian rhythm in photosynthesis in Phaseolus vulgaris. Plant Physiol 97:415–419. https://doi.org/10.1104/pp.97.1.415
Gould PD, Diaz P, Hogben C, Kusakina J, Salem R, Hartwell J, Hall A (2009) Delayed fluorescence as a universal tool for the
measurement of circadian rhythms in higher plants. Plant J 58:893–901. https://doi.org/10.1111/j.1365-313X.2009.03819.x

Guerrero L, Fermani S, Zaffagnini M, Sparla F, Trost P (2021) Calvin-Benson cycle regulation is getting complex. Trends Plant Sci 26:898–912. https://doi.org/10.1016/j.tplants.2021.03.008

Hashida SN, Miyagi A, Nishiyama M, Yoshida K, Hisabori T, Kawai-Yamada M (2018) Ferredoxin/thioredoxin system plays an important role in the chloroplast NADP status of Arabidopsis. Plant J 95:947–960. https://doi.org/10.1111/tpj.14000

Haydon MJ, Mielczarek O, Robertson FC, Hubbard KE, Webb AAR (2013) Photosynthetic entrainment of the Arabidopsis thaliana circadian clock. Nature 502:689–692. https://doi.org/10.1038/nature12603

Hennessey TL, Field CB (1991) Circadian rhythms in photosynthesis: oscillations in carbon assimilation and stomatal conductance under constant conditions. Plant Physiol 96:831–836. https://doi.org/10.1104/pp.96.3.831

Hennessey TL, Freedan AL, Field CB (1993) Environmental effects on circadian rhythms in photosynthesis and stomatal opening. Planta 189:369–376. https://doi.org/10.1007/BF00194433

Hisabori T, Sunamura E, Kim Y, Konno H (2013) The chloroplast ATP synthase features the characteristic redox regulation machinery. Antiox Redox Sign 19:1846–1854. https://doi.org/10.1089/ars.2012.5044

Holmes MG, Klein WH (1986) Photocontrol of dark circadian rhythms in stomata of Phaseolus vulgaris L. Plant Physiol 82:28–33. https://doi.org/10.1104/pp.82.1.28

Jones TL, Tucker DE, Ort DR (1998) Chilling delays circadian pattern of sucrose phosphate synthase and nitrate reductase activity in tomato. Plant Physiol 118:149–158. https://doi.org/10.1104/pp.118.1.149

Kerr PS, Rufty TW, Huber SC (1985) Endogenous rhythms in photosynthesis of light-induced nuclear messenger RNAs. Plant Physiol 77:275–280. https://doi.org/10.1104/pp.77.2.275

Kloppstech K (1985) Diurnal and circadian rhythmicity in the expression of light-induced nuclear messenger RNAs. Planta 165:502–506. https://doi.org/10.1007/BF00398095

Lai AG, Doherty CJ, Mueller-Roeber B, Kay SA, Schippers JH, Dijkstra P (2012) Circadian clock-associated 1 regulates ROS homeostasis and oxidative stress responses. Proc Natl Acad Sci USA 109:17129–17134. https://doi.org/10.1073/pnas.1209481109

Laisk A, Klirats O, Oja V, Gerst U, Weis E, Heber U (1992) Analysis of oxygen evolution during photosynthetic induction and in multiple-turnover flashes in sunflower leaves. Planta 166:434–444. https://doi.org/10.1007/BF00195325

Lawson T, von Caemmerer S, Baroli I (2010) Photosynthesis and stomatal behaviour. In: Lüttge U, Beyschlag W, Büdel B, Francis D (eds) Prog Bot. Springer, Heidelberg

Liu ZR, Taub CC, McClung CR (1996) Identification of an Arabidopsis thaliana ribulose-1,5-bisphosphate carboxylase oxygenase activase (RCA) minimal promoter regulated by light and the circadian clock. Plant Physiol 112:43–51. https://doi.org/10.1104/pp.112.1.43

Lonergan TA (1981) A circadian rhythm in the rate of light-induced electron flow in three leguminous species. Plant Physiol 68:1041–1046

Martín ES, Meidner H (1971) Endogenous stomatal movements in Tradescantia virginiana. New Phytol 70:923–928

Martino-Catt S, Ort DR (1992) Low temperature interrupts circadian regulation of transcriptional activity in chilling-sensitive plants. Proc Natl Acad Sci USA 89:3731–3735. https://doi.org/10.1073/pnas.89.9.3731

Matsu S, Onai K, Okamoto K, Minagawa J, Ishiura M (2006) Real-time monitoring of chloroplast gene expression by a luciferase reporter: evidence for nuclear regulation of chloroplast circadian period. Mol Cell Biol 26:863–870. https://doi.org/10.1128/MCB.26.3.863-870.2006

Maxwell K, Johnson GN (2000) Chlorophyll fluorescence - a practical guide. J Exp Bot 51:659–666. https://doi.org/10.1093/jxb/51.3.345.659

McClung CR, Hsu M, Painter JE, Gagne JM, Karlsberg SD, Salomé PA (2000) Integrated temporal regulation of the photosynthetic pathway: circadian regulation of two Arabidopsis genes encoding serine hydroxymethyltransferase. Plant Physiol 123:381–392. https://doi.org/10.1104/pp.123.1.381

Michelet L, Zaffagnini M, Morisse S, Sparla F, Pérez-Pérez ME, Francia F, Danon A, Marchand CH, Fermani S, Trost P, Lemaire SD (2013) Redox regulation of the Calvin-Benson cycle: something old, something new. Front Plant Sci 4:470. https://doi.org/10.3389/fpls.2013.00470

Millar AJ (2016) The intracellular dynamics of circadian clocks reach for the light of ecology and evolution. Annu Rev Plant Biol 67:595–618. https://doi.org/10.1146/annurev-arplant-043014-115619

Millar AJ, Short SR, Chua NH, Kay SA (1992) A novel circadian phenotype based on firefly luciferase expression in transgenic plants. Plant Cell 4:1075–1087. https://doi.org/10.1105/tpc.4.9.1075

Nakahira Y, Baba K, Yoneda A, Shinma T, Toyoshima Y (1998) Circadian-regulated transcription of the psbD light-responsive promoter in wheat chloroplasts. Plant Physiol 118:1079–1088. https://doi.org/10.1104/pp.118.3.1079

Noordally ZB, Millar AJ (2015) Clocks in algae. Biochemistry 54:171–183. https://doi.org/10.1021/bi501088r

Pallas JE, Samish YB, Willmer CM (1974) Endogenous rhythm activity of photosynthesis, transpiration, dark respiration, and carbon dioxide compensation point of peanut leaves. Plant Physiol 53:907–911. https://doi.org/10.1104/pp.53.6.907

Philipou K, Ronald J, Sánchez-Villarreal A, Davis AM, Davis SJ (2019) Physiological and genetic dissection of sucrose inputs to the Arabidopsis thaliana circadian system. Genes 10:334. https://doi.org/10.3390/genes10050334

Portis AR Jr, Li C, Wang D, Salvucci ME (2008) Regulation of Rubisco activase and its interaction with Rubisco. J Exp Bot 59:1597–1604. https://doi.org/10.1093/jxb/erm240

Raschke K (1975) Stomatal action. Annu Rev Plant Physiol 26:309–340. https://doi.org/10.1146/annurev.pp.26.060175.001521

Recchia I, Sparla F, Pupillo P (2017) Photosynthetic properties of spring geophytes assessed by chlorophyll fluorescence analysis. Plant Physiol Biochem 118:510–518. https://doi.org/10.1016/j.plaphy.2017.07.020

Resco de Dios V, Goulden ML, Ogle K et al (2012) Endogenous circadian regulation of carbon dioxide exchange in terrestrial ecosystems. Global Change Biol 18:1956–1970. https://doi.org/10.1111/j.1365-2486.2012.02664.x

Resco de Dios V, Loik ME, Smith R, Aspinwall MJ, Tissue DT (2016) Genetic variation in circadian regulation of nocturnal stomatal conductance enhances carbon assimilation and growth. Plant Cell Environ 39:3–11. https://doi.org/10.1111/pce.12598

Seo PJ, Mas P (2014) Multiple layers of posttranslational regulation refine circadian clock activity in Arabidopsis. Plant Cell 26:79–87. https://doi.org/10.1105/tpc.113.119842

Sukhova E, Mudrilov M, Vodeneev V, Sukhov V (2018) Influence of the variation potential on photosynthetic flows of light energy and electrons in pea. Photosynth Res 136:215–228. https://doi.org/10.1007/s11110-017-0460-1

Taylor SH, Gonzalez-Escobar E, Page R, Parry MAJ, Long SP, Carmo-Silva E (2022) Faster than expected Rubisco deactivation in shade reduces cowpea photosynthetic potential in variable
light conditions. Nat Plants 8(118):124. https://doi.org/10.1038/s41477-021-01068-9

Tezara W, Mitchell VJ, Driscoll SP, Lawlor DW (2002) Effects of water deficits and its interaction with CO₂ supply in the biochemistry and physiology of photosynthesis in sunflower. J Exp Bot 53:1781–1791. https://doi.org/10.1093/jxb/erf021

Vaseghi M-J, Chibani K, Telman W, Liebthal MF, Gerken M, Schnitzer H, Mueller SM, Dietz K-J (2018) The chloroplast 2-cysteine per-oxiredoxin functions as thioredoxin oxidase in redox regulation of chloroplast metabolism. Elife 7:e38194. https://doi.org/10.7554/eLife.38194.001

Yokochi Y, Fukushi Y, Wakabayashi KI, Yoshida K, Hisabori T (2021) Oxidative regulation of chloroplast enzymes by thioredoxin and thioredoxin-like proteins in Arabidopsis thaliana. Proc Natl Acad Sci U S A 118:e2114952118. https://doi.org/10.1073/pnas.2114952118

Yoshida K, Yokochi Y, Hisabori T (2019) New light on chloroplast redox regulation: molecular mechanism of protein thiol oxidation. Front Plant Sci 10:1534. https://doi.org/10.3389/fpls.2019.01534

Zaffagnini M, Fermani S, Marchand CH, Costa A, Sparla F, Rouhier N, Geigenberger P, Lemaire SD, Trost P (2019) Redox homeostasis in photosynthetic organisms: novel and established thiol-based molecular mechanisms. Antioxid Redox Signal 31:155–210. https://doi.org/10.1089/ars.2018.7617

Zimmer D, Swart C, Graf A, Arrivault S, Tillich M, Proost S, Nikoloski Z, Stitt M, Bock R, Mühlhaus T, Boulouis A (2021) Topology of the redox network during induction of photosynthesis as revealed by time-resolved proteomics in tobacco. Sci Adv 7:eabj8307. https://doi.org/10.1126/sciadv.abj8307

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