DIVING-PAM-II

Underwater Chlorophyll Fluorometer Manual

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1 Safety Instructions

1.1 General Safety Instructions

- Read and follow safety and operating instructions prior to operation of the device. Pay attention to all safety warnings.
- Connect the device only to the power source indicated in operating instructions or on the device. If the device is not in use, remove the mains plug from the socket.
- Do not put the device near sources of heat.
- Expose the device to dust, sand, and dirt as little as possible.
- Ensure that neither liquids nor foreign bodies get inside the device.
- The device may only be repaired by the manufacturer.

1.2 Special Safety Instructions

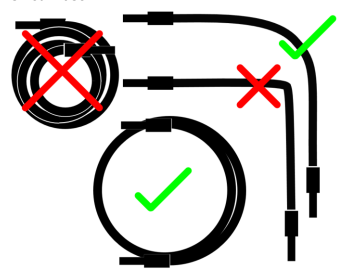
- The DIVING-PAM-II is a highly sensitive instrument which should be only used for research purposes, as specified in this manual. Follow the instructions of this manual to avoid potential harm to the user and damage to the instrument.
- The DIVING-PAM-II can emit very strong light! To avoid harm to your eyes, never look directly into the light port of the DIVING-PAM-II or its fiberoptics.
- Keep all ports and plugs clean (see Section 9.1, page 151).
 Do not expose open cable ends to water or high moisture.

- Do not force a plug into the wrong socket (cf. Fig. 3, page 12). Make sure that the connection is watertight (Fig. 4, page 14).
- Do not attempt to open pressure and temperature sensor (Fig. 3, page 12).
- Run all cables so that stepping on or stumbling over them is excluded.
- Do not open housing of the DIVING-PAM-II for repair.
 There are no serviceable parts inside the DIVING-PAM-II.
 Exception: battery change (Section 9.3, page 152).
- Before immersing the DIVING-PAM, make sure that all cable plugs are properly connected (the rippled cylinder of the cable end must be completely screwed in so that it presses against the rubber ring of the cable port: Fig. 4, page 14). Make sure that each unused cable port is properly sealed by a red riffled cap (screw in completely).
- Do not exceed the maximum diving depth of 50 m.
- The PC Interface Box DIVING-PAM-II/I is not watertight;
 keep it away from water or high moisture areas.
- At the interface box, orientate charge plug so that the red dot on the plug coincides with the red dot of the socket. Do not try to disconnect the plug by pulling at the cable. Disconnect plug by pulling at the rippled bushing of the plug.

1.3 Correct Handling of Fiber Optics DIVING-F

This fiber optics DIVING-F is made of delicate glass fibers. Each glass fiber extents over the entire length of the fiber optics. Do not sharply bend, stretch, or crush the fiber optics as this can break the glass fibers. Each broken fiber reduces light transmission by the fiber optics. If many fibers are broken, the fiber optics must be

replaced. Place protection caps on fiber ends when the fiber optics is not in use.



1.4 Instructions for Battery Handling

Only for DIVING-PAM-II with Lithium-Ion Battery

- The DIVING-PAM-II Info window indicates if a lithium battery is built in (Fig. 63, page 89).
- For long-term storage of the DIVING-PAM, discharge battery to about 30% of its capacity. The charge status is indicated in the top right corner of the Basic Data window (Fig. 24, page 40).

Only for DIVING-PAM-II with Lithium-Ion Battery

- For shipping of the DIVING-PAM-II:
 - Discharge battery to 30% (see above).
 - Use original DIVING-PAM-II/T transport case.
 - Download from Walz website the testing protocol for the DIVING-PAM-II battery.

https://www.walz.com/products/chl p700/diving-pam-II/downloads.html

Go to: Shipping the DIVING-PAM-II

Select: "Testing protocol"

- Print entire protocol (6 pages) and enclose this protocol in the box.
- Visibly label package with lithium battery label (downloadable at https://www.walz.com/prod-ucts/chl_p700/diving-pam-II/downloads.html).



Fig. 1: Lithium Battery Label

2 Introduction

2.1 Features DIVING-PAM-II

- The "Underwater Chlorophyll Fluorometer DIVING-PAM-II" has been designed for saturation pulse analysis of photosystem II in aquatic organisms. Typically, these organisms include sea grass species, macroalgae, zooxanthellae in corals, and cyanobacterial layers.
- Compared to its predecessor ("first-generation" DIVING-PAM), the chief technical advancements are the consistent use of energy-efficient LEDs, a far-red light source for selective excitation of photosystem I, an internal PAR sensor, a touchscreen for outdoor application, and WiFi communication.
- The fluorometer measures the efficiency of photosystem II under dark conditions (F_V/F_M) and in the light (Y(II)). Further parameters measured characterize photochemical fluorescence quenching (q_L, q_P), non-photochemical fluorescence quenching (q_N, NPQ, Y(NPQ), Y(NO)).
- The DIVING-PAM-II is equipped with a spectroradiometer which is calibrated to measure spectra of photon flux density. The software automatically integrates photon flux density in the range from 400 to 700 nm to obtain PAR, which, together with Y(II), is used to calculate relative rates of photosynthetic electron transport (ETR).
- The spectrometer can also record reflectance and fluorescence spectra of samples.
- An underwater oxygen sensor using optical detection of O₂ completes the range of accessories.

- The capacity of the internal memory corresponds to data of more than 27,000 data sets of saturation pulse analyses.
- A high-capacity lead acid battery lasts for up to 1300 saturation pulse analyses.
- In the lab, the DIVING-PAM-II can be combined with the same oxygen measuring system developed for the MINI-PAM-II (https://www.walz.com/products/chl_p700/mini-pam-II/accessories O2fluo configuration.html).

2.2 Structure of Manual

The DIVING-PAM-II fluorometer provides a vast range of settings and protocols. To make full use of these opportunities, become acquainted with terminology and principles of saturation pulse analysis (see Chapter 10, page 159). Section 10.4 (page 167) provides a list of review papers on PAM chlorophyll fluorescence and saturation pulse analysis.

Prior to working with the Diving-PAM-II, note the Safety Instruction (Chapter 1, page 1). Chapter 3 (page 9) deals with the setup of the basic system and the accessories available.

In the field, the DIVING-PAM-II is normally operated in the standalone mode. Chapter 4 (page 35) provides detailed instructions on how to use the screen interface including advises on fluorescence induction and light curve programs.

The DIVING-PAM-II can also be operated by the software Win-Control-3 running on a computer with Windows operating system. Installation of WinControl-3 is introduced in Chapter 5, page 91, and the features of WinControl-3 are dealt with in Chapter 6, page 97. Chapter 7 (page 145) provides hints for beginners, Chapter 8 (page 149) contains instructions for trouble shooting, and Chapter 9 (page 151) gives guidelines for maintenance of the device.

Further, technical information (Chapter 10, page 159) and guaranty conditions (Chapter 12, page 181) are provided.

This manual ends with lists of keywords (Chapter 13, page 185) figures (Chapter 14, page 191) and tables (Chapter 15, page 195).

3 Components and Setup

3.1 Basic System

3.1.1 Extent of Delivery (Basic System)

Item	Order Code	
Optoelectronic Unit	DIVING-PAM-II/B or -/R	
Fiberoptics	DIVING-F	
PC Interface Box	DIVING-PAM-II/I	
Power Supply	MINI-PAM-II/N	
Underwater Cable	DIVING-PAM-II/K5	
Miniature Spectrometer	MINI-SPEC	
PAR Calibration Block	160101439	
Distance Clip 60°	2010-A	
Dark Leaf Clip (3 pcs)	DIVING-LC	
Surface Holder	DIVING-SH	
Software	WinControl-3	
Transport Case	DIVING-PAM-II/T	
Standard USB-A to USB-B cable		
Silicone grease		
Fuses		
Manual		

3.1.2 System Overview

An overview to the principal components of the DIVING-PAM-II system and connections between them is given in Fig. 2.

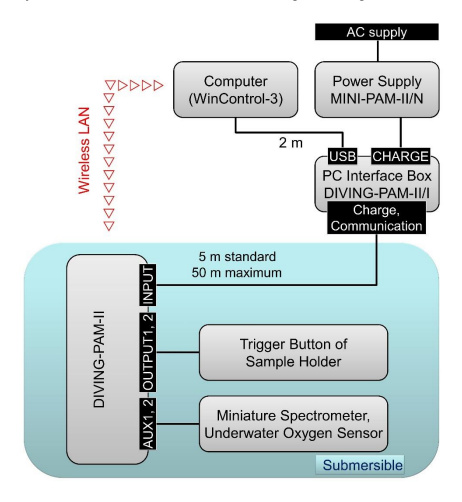


Fig. 2: DIVING-PAM-II: System Components

In Fig. 2, all submersible components of the system are drawn on aquamarine background.

Submersible components, which are part of the basic system, are the central unit of the DIVING-PAM-II, the Fiberoptics DIVING-F, which is inserted into the central optical port of the right end plate (not shown in Fig. 2), and the Miniature Spectrometer MINI-SPEC. Submersible accessories, which are not part of the basic system, are the Underwater Oxygen Sensor DIVING-PAM-II/O2, and the Universal Sample Holder DIVING-II-USH.

To measure light spectra and total photon flux in the visible range (PAR), the MINI-SPEC is plugged into the AUX 1 or AUX 2 port of the fluorometer's right end plate. The DIVING-PAM-II/O2 uses the same ports.

The Universal Sample Holder DIVING-II-USH includes a trigger cable which can be plugged into port OUT 1 or OUT 2 on the right end plate. When plugged in, the trigger button of the DIVING-II-USH has the same function of the START button on the right end plate of the fluorometer (Fig. 3).

The function of both the trigger button of the DIVING-II-USH and the START button depends on the window selected on the DIVING-PAM-II screen, e.g., when the window "Basic Data" is visible, a saturation pulse analysis will be triggered, or when the window "Induction Curve" is visible an induction curve will be performed.

Fig. 2 also outlines that the DIVING-PAM-II can be operated in the stand-alone mode or by the software WinControl-3 running on a computer with Microsoft Windows operating system. Operation by WinControl-3 requires connection between computer and fluorometer via the system's interface box (DIVING-PAM-II/I). Computer and interface box are linked by a standard USB cable; interface box and fluorometer are connected by a special underwater cable.

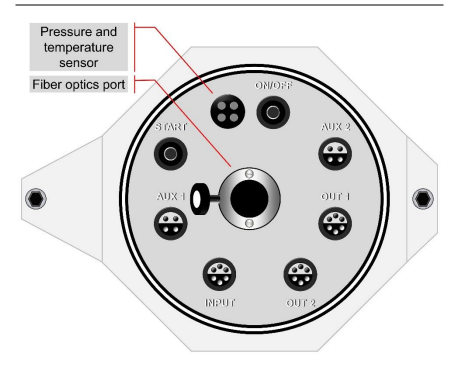


Fig. 3: Right End Plate of DIVING-PAM-II

Instructions on how to properly connect underwater cables are given by Fig. 4 (page 14). Note that the interface box is not submersible.

For communication between computer and fluorometer, the interface box does not need line power. When the interface box is connected to the mains, the DIVING-PAM-II battery is charged via the underwater cable. Advantages of WinControl-3 operation are the option to remotely control the DIVING-PAM-II, and to continuously record steady state chlorophyll fluorescence. (Note that continuous fluorescence is not recorded in the stand-alone mode.)

Table 1: Side panel of DIVING-PAM-II: Summary of Elements

Key or port (see Fig. 3)	Function	Comment
ON/OFF	On/off switch and Keyboard lock switch	Press briefly to switch on, press 2 seconds to switch off DIVING-PAM-II. When the DIVING-PAM-II is switched on, briefly pressing locks or unlocks keyboard (the red LEY LED indicates a lock state of the keyboard).
START	Trigger button	The button triggers the command displayed in the top right corner of the current display.
AUX 1/AUX 2	Connectors for auxiliary devices	Connect here MINI-SPEC, Underwater Oxygen Sensor DIVING-PAM-II/O2, or Fiber-Optic Oxygen Meter FireStingO2 (special adapter required).
OUT 1/OUT 2	Connectors for external light	Sockets for synchronized external light source (light is blanked out when PAM fluorescence is measured). The sockets work also as input for the trigger signal from Sample Holder DIVING-II-USH.
INPUT	Interface con- nector	To charge battery and to operate DIVING-PAM-II by WinControl-3.
Pressure and temperature sensor	Housing for pressure and temperature sensor. The digital sensor chip is second order temperature-compensated. Do not attempt open.	
Port for fiber- optics with locking screw	Optical window f	or internal light sources and fluorescence

As new feature, the DIVING-PAM-II has a Wireless LAN Interface built-in, which permits convenient data download at the experimental site. See Table 2, page 15.

Note that all peripherals may only be connected in dry and clean environment.

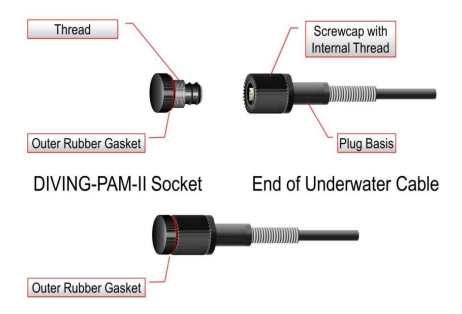


Fig. 4: Watertight Connection of Cable

- (1) Hold cable end at "plug basis".
- (2) Fully (!) insert plug in socket.
- (3) Tighten "screwcap".
- (4) Confirm that screwcap presses against "outer rubber gasket".

The waterproof fluorometer housing consists of a Plexiglas tube with Plexiglas end plates. The right end plate accommodates the fiberoptics port, various plug connectors, two push buttons (START and ON/OFF), and a combined pressure and temperature sensor (Fig. 3, page 12). All ports and buttons are explained in Table 1 (page 13).

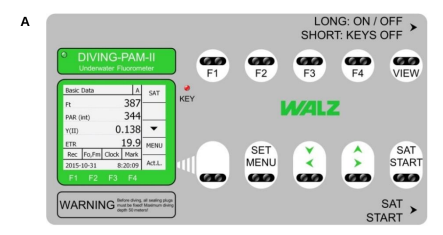
On the front of the DIVING-PAM-II, a compact, low-energy black/white screen displays measured data and provides field elements to control the fluorometer. The waterproof design of the fluorometer requires that these field elements are selected by optical switches which are arranged in two rows of 5 keys on the right

side of the screen (Fig. 5A, page 16). Just touching a key area activates the command of the associated element on the screen. The LED located right of the screen shines green when an optical switch is activated.

Table 2: WLAN Connection

Establish WLAN	Connection Computer to DIVING-PAM-II	
DIVING-PAM-II	Switch on WLAN When WLAN is set to "Auto" in window "Wireless", the WLAN is active 5 minutes after powering on the fluorometer; WLAN is always active when "On" is selected "Wireless" (Fig. 60, page 84). The "W" in the first line of Windows "Basic Data" indicates that WLAN is active.	
Computer	Select DIVING-PAM-II as access point Enter security key "password" Ignore error message "The connection is limited".	
Connect WinCon	trol-3 to DIVING-PAM-II	
Computer	Start Network Mode of WinControl-3. - Windows 7: Click "Start"→Select "All Programs", "WinControl-3". - Windows 10 and newer: Click tile "WinControl-3-Network Mode". - General alternative option: Search "WinControl-3 – Network Mode". The network mode opens the window below.	
Click "DIVING-PA Click OK Wait for DIVING-F by WinControl-3	Using auto-detection you can search for an IB4 interfac	

3.1.3 Fluorometer



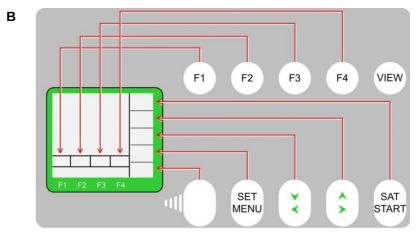


Fig. 5: Front View of DIVING-PAM-II

 ${f A},$ keys and screen. ${f B},$ relation of keys and fields elements.

The relation of optical switches and field elements are outlined in Fig. 5B (page 16). The function keys of the upper row of keys (F1 through F4) are associated with the 4 control elements located on the lower edge of the DIVING-PAM-II screen. The rightmost key in the upper row, the VIEW button, is not connected to with a field

element but changes appearance and properties in graphical windows (e.g., induction curve or spectral data) so that more numerical values are displayed and selection of individual points in a graph becomes possible.

The keys of the lower row from left to right correspond to the field elements on the right side of the screen from bottom to top. The function of the rightmost optical key of the lower row is identical to the START button on the right fluorometer end plate (Fig. 3, page 12) or the trigger button of the DIVING-II-USH (Fig. 17, page 28).

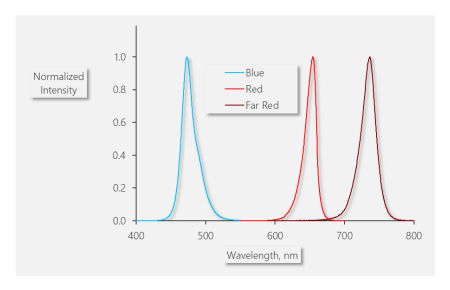


Fig. 6: Normalized Emission Spectra of DIVING-PAM-II LEDs

Typical LED emission spectra normalized to their maxima. The blue curve corresponds to the spectrum of the blue LED of the DIVING-PAM-II/B, the red curve represents the red LED in the DIVING-PAM-II/R. Both DIVING-PAM-II versions possess a far-red LED which emits maximally above 700 nm (right-most curve). Peak wavelength and full width at half maximum (in brackets) are indicated.

In the DIVING-PAM-II/R, a red LED acts as measuring light and as actinic light source. In the blue version (DIVING-PAM-II/R), a blue LED is used. Both versions of the DIVING-PAM-II are equipped with a far-red LED. Normalized emission spectra of blue, red, and far-red LEDs are shown in Fig. 6.

Another difference between the two versions is the spectral window for fluorescence detection. The BLUE version detects fluorescence at wavelengths > 630 nm but the RED version detects fluorescence at wavelengths > 700 nm (Fig. 7).

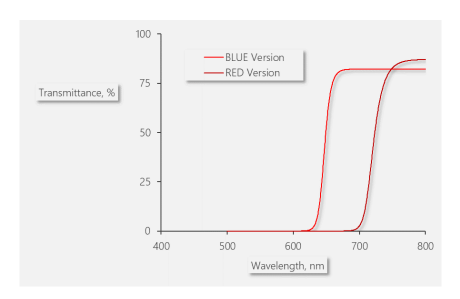


Fig. 7: Transmittance Spectra of DIVING-PAM-II Versions

Transmittance spectra of detection filters in the DIVING-PAM-II/B (BLUE Version, red line) and DIVING-PAM-II/R (RED Version, dark red line) are shown.

3.1.4 Miniature Spectrometer MINI-SPEC

The Miniature Spectrometer MINI-SPEC belongs to the basic system of the DIVING-PAM-II. The MINI-SPEC is calibrated to measure spectra of quantum fluxes. Integration of these spectra over the visible range results in PAR data like those recorded with Walz quantum sensors. Compared to standard quantum sensors, the spectrometer provides additional information on the spectral composition of photosynthetically active radiation which changes with water depth and can be influenced by photosynthetic organisms close by. During transport, the spectrometer is mounted on the fluorometer as show in Fig. 8, page 20.

The spectrometer is used to calibrate the internal PAR sensor of the DIVING-PAM-II (see Section 4.2.4.1, page 70). To this aim, the DIVING-PAM-II light guide and the entrance optics of the spectrometer are inserted in the PAR calibration block (Fig. 10, page 22). The light guide can be inserted either in the 60° or the 90° port according to the two possible orientations of the light guide in the Universal Sample Holder DIVING-II-USH. With both pieces fully inserted, the distance between fiber optics end and diffusing disk of the spectrometer matches the corresponding standard distances between fiber optics end and sample level in the DIVING-II-USH (7 mm, compare Fig. 16, page 26).

Replacing the entrance optics used for evaluation of light by the cap for fluorescence and reflection (Fig. 9, page 21) extents considerably the range of spectral information attainable by the miniature spectrometer.

In the fluorescence mode, a light guide in the cap leads light from a blue or a green LED located inside the spectrometer body to the sample. The excited fluorescence reaches the spectrometer body through a central hole in the cap. Green light penetrates deeper into photosynthetic tissue than blue or red (Terashima et al (2009)

Plant Cell Physiol 50:684-697). Therefore, the two excitation colors permit probing the effect of varying depths of penetration of excitation light on the shape of fluorescence spectra. The spectrometer automatically compensates for the spectral sensitivity of the detector.

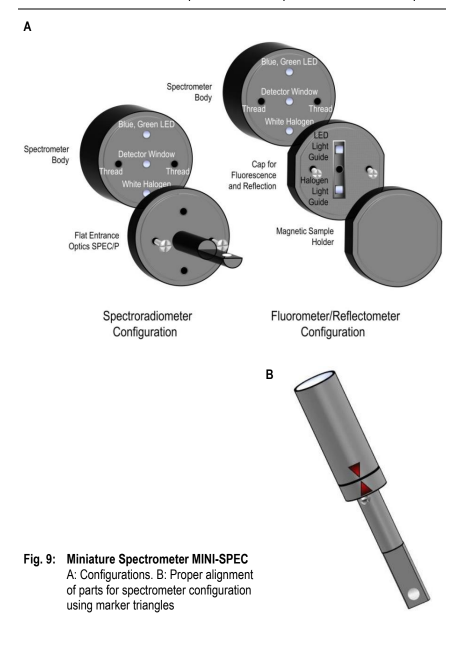
For reflection measurements, a halogen lamp is employed whose emission is transferred through another light guide to the sample (Fig. 9, page 21). From reflection measurements, reflectance (R) is derived by dividing a sample spectrum (r_{sample}) by the spectrum of a white light-diffusing reflector consisting of a fluoropolymer with very high diffuse reflectance in the entire visible range (r_{reference}):

$$R(\lambda) = \frac{r_{sample}(\lambda)}{r_{reference}(\lambda)}$$

The reference material is part of delivery; to maintain its optical properties, do not touch the white surface, keep away dirt, dust, and humidity, and store the reflection standard in a closed container when not used.



Fig. 8: DIVING-PAM-II with MINI-SPEC in Transport Position



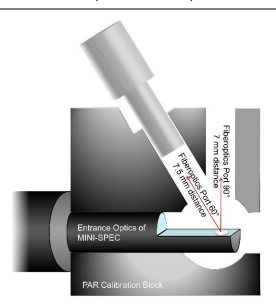


Fig. 10: PAR Calibration Block

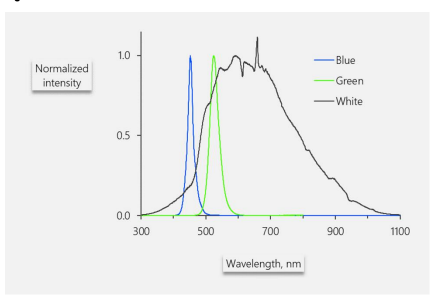


Fig. 11: Normalized Emission Spectra of MINI SPEC Light Sources

3.1.5 Distance Clip 60° 2010-A

The 2010-A clip positions the fiberoptics end-piece relative to the sample. The axis of the end-piece is positioned at a 60° angle relative to the sample plane. Two different spacer rings may be used to increase the distance between fiberoptics and sample. The distance between fiberoptics exit plane and sample has considerable influence on signal amplitude and effective light intensities (Fig. 14, page 25).

Normally, a specimen is examined above the viewing hole of the clip and it is hold between the jaws of the clip. In case of thick or bulky specimens, the sample is placed below the hole of the 2010-A clip.

From the 60° angle between sample and fiberoptics results slightly heterogeneous light intensities at leaf surface because the distance between fiber optics tip and leaf surface varies. The measured signal will be dominated by that part of the leaf, which receives maximal intensity, as this is most strongly excited and emits most of the fluorescence.

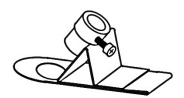


Fig. 12: Distance Clip 60° 2010-A

3.1.6 Dark Leaf Clip DIVING-LC

The Dark Leaf Clip DIVING-LC weighs 6.5 g and can be attached to most flat samples without detrimental effects. It is equipped with a miniature sliding shutter which prevents light access to the leaf during dark acclimation. The shutter is opened for the measurement after the fiberoptics has been inserted to prevent exposure to external light. Proper dark adaptation is essential for determination of the maximal quantum yield $F_{\text{V}}/F_{\text{M}}$.

The dark leaf clip plus adapter (Fig. 13, page 24) positions the fiberoptics tip at the relatively short distance of 3 mm above leaf surface. As a consequence, measuring light intensity and, thus, signal amplitude are higher than with the standard distance for vertical illumination of 7 mm (see Fig. 16, page 26). If signal saturation occurs at 3 mm distance, reduce measuring light intensity and gain. The background signal (F-Offset) will be automatically adjusted (cf. Section 4.2.1, page 57, and Section 6.10.4, page 136).



Fig. 13: DIVING-LC and Adapter

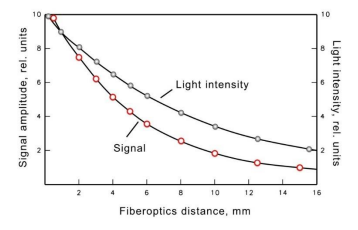


Fig. 14: Signal/Distance Relationship

Relationship between signal amplitude/light intensity and distance between fiberoptics tip and sample

3.1.7 Surface Holder DIVING-SH

To study corals or epilithic plants, the surface holder can be attached by three hooks and rubber bands to uneven, creviced surfaces. The three screws holding the rubber bands can be moved up and down to adjust the surface holder relative to the sample the desired distance.

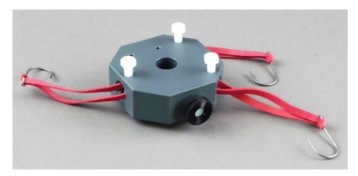


Fig. 15: .Surface Holder DIVING-SH

Achieving 7 mm Standard Distance Between Fiber and Sample



Fig. 16: The Standard Distance of 7 mm

Combinations of adapters and leaf clip to achieve the standard distance of 7 mm between fiberoptics end and sample level.

3.2 Accessories

3.2.1 Universal Sample Holder DIVING-II-USH

The Universal Sample Holder DIVING-II-USH was designed for underwater investigations of on samples like sea grass, macroalgae, corals, algal mats and periphyton.

The trigger button of the sample holder permits one-hand operation. The trigger signal is transmitted by a cable to ports OUT1 or OUT2 of the DIVING-PAM-II (see Table 1, page 13). The fiberoptics can be positioned perpendicularly to the sample or in the 60° arrangement. In both orientations, the distance between fiber end and sample surface corresponds to the standard conditions as shown in Fig. 10 (page 22) that is 7.5 mm for 60° and 7 mm for 90° angle between fiberoptics and sample surface.

Trigger cable, spectrometer cable and the fiber optics are loosely held together by a nylon-mesh-cover featuring a zipper. When not in use, the whole device can be hooked to the diver's jacket.

The general setup of the DIVING-II-USH is depicted Fig. 17, (page 28), connection to the DIVING-PAM-II is illustrated by Fig. 18 (page 29). The various modes of application are addressed in the following section.

DIVING-USH Applications

Leaves and macroalgae

With flat samples, the sample holder with clip attached can be used. The clip is connected to the holder with two screws (Fig. 17, page 28). The clip can be opened by a handle with a single finger of the hand holding the device.

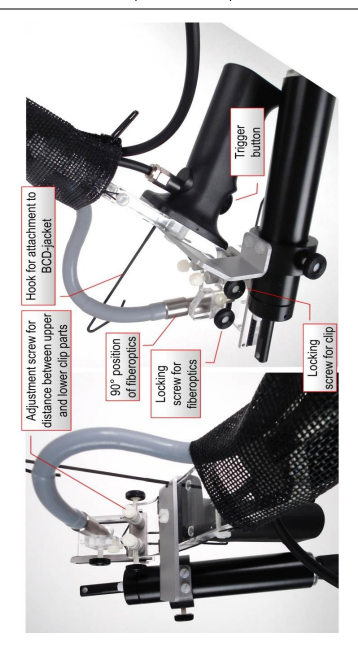


Fig. 17: Universal Sample Holder DIVING-II-USH

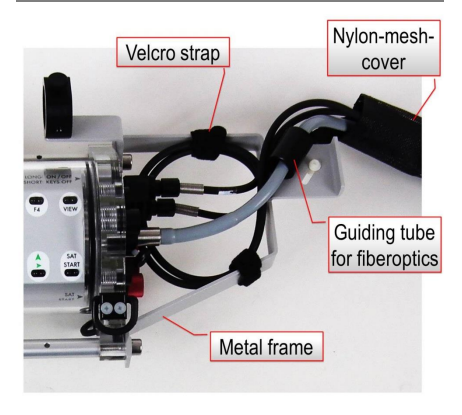


Fig. 18: Arrangement of Fiberoptics and Cables

Two rubber O-rings constitute the elastic part of the clip, holding the upper and lower parts together. The O-rings are held by nylon screws positioned sideward and at the lower side of the clip. O-rings can be hooked into to either of the two lower screws resulting in different stretching of the O-ring. Depending on stretching, the pressure of the clip put on a sample varies. Different pressures can be also obtained with O-rings of various thicknesses.

Furthermore, it is possible to adjust the distance between upper and lower clip part by two vertical nylon screws (Fig. 17, page 28). In this way, the clip can be adapted to the sample's pressure sensitivity and thickness. The stripe-like contact areas on the edges of the clip edge are covered with neoprene rubber foam (bottom jaw) and a plastic "pin-cushion" (top jaw), which assures a good grip even with slippery objects like kelp and other seaweeds.

The Fiberoptics can be mounted at angles of either 60° or 90° with respect to the sample plane. Changing the angle just takes a couple of seconds. At an angle of 60° ambient light can freely reach the site at which fluorescence is measured. The quantum flux density of this light can be measured the Miniature Spectrometer MINI-SPEC. Knowledge of quantum flux density is important for estimation of relative electron transport rate (ETR) and for evaluation of measured effective quantum yields.

The 90° arrangement is employed for illumination by the actinic light of the DIVING-PAM-II fluorometer. Well-defined light response and induction curves can be measured when the external light is fully excluded. For this purpose, an adapter is provided which slips over the fiberoptics metal endpiece (see Fig. 16, page 26, "Adapter"). One side of this adapter is covered with neoprene rubber, making provision for a light-tight, gentle contact with the sample. In order to prevent access of light from the bottom side, a "darkening plate" can be fixed in the lower part of the clip (part of delivery, not shown).

Bulky samples

For measurements of bulky samples (e.g., corals, sea anemones, periphyton and microphytobenthos), the bottom part of the clip can be removed. For this purpose, two rubber O-rings are pulled off. Then, the lower part is pulled out of its bearings. It is also possible to remove the entire clip and to use the adapter for dark acclimation only.

Sample holders for dark acclimation

Dark acclimation of a sample for at least several minutes is required for assessment of the maximal quantum yield (F_V/F_M) . For

this purpose, two different types of dark sample holders are available, the Magnet Sample Holder DIVING-MLC and the Dark Leaf Clip DIVING-LC. The former is suited for the study of relatively large and robust samples, whereas the latter is better suited for more fragile leaf-like samples.

3.2.2 Underwater Oxygen Sensor DIVING-PAM-II/O2

The DIVING-PAM-II/O2 sensor measured oxygen optically. A sensor spot carrying a fluorescent dye changes its fluorescence properties dependent on oxygen concentration. For Details see Instruction Manual for Oxygen Measurements:

https://www.walz.com/products/chl p700/diving-pam-II/downloads.html

Typical applications for the DIVING-PAM-II/O2 are measurements of bathymetric oxygen profiles and monitoring of long-term declines in oceanic oxygen concentrations associated with climate change



Fig. 19: Underwater Oxygen Sensor DIVING-PAM-II/O2

3.2.3 Magnet Sample Holder DIVING-MLC

The Magnet Sample Holder DIVING-MLC consists of two circular halves each containing magnets (Fig. 20, page 32). A flat sample sandwiched between the two halves is hold by magnetic pull. Typical samples are seagrasses, kelp and other large macroalgae.

The upper halve of the DIVING-MLC has a central hole which can be closed by a sliding shutter. The upper halve also provides a seat for the adapter DIVING-DA. With the shutter closed, the sample part below the central hole is shielded from light. Using an adapter DIVING-DA, the DIVING-PAM-II fiberoptics can be connected to the DIVING-MLC. Measurements with the still dark-acclimated sample can start after opening of the shutter.

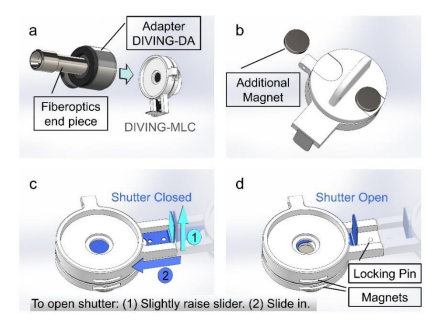


Fig. 20: DIVING-MLC

a, putting the fiberoptics with DIVING-DA adapter on a DIVING-MLC. **b**, positioning an additional magnet. **c**, DIVING-MLC with closed shutter. **d**, DIVING-MLC & open shutter.

The lower halve provides seats for additional magnets. The magnetic pull can be adapted to the sample by varying the number of additional magnets.

3.2.4 Miniature Fiberoptics DIVING-F1

The Miniature Fiberoptics DIVING-F1 has been designed to investigate small surfaces. The light guide consists of a single coated plastic fiber, which has an active diameter of 2 mm and a length of 1.50 m.



Fig. 21: Miniature Fiberoptics DIVING-F1

Fiber optics with adapters and Polishing Set (order number 000160103450, supplied with DIVING-F1).

3.2.4.1 Calibration of the internal light sensor and establishing the internal PAR list

Table 3: Miniature Fiberoptics DIVING-F1				
What is measured?	PAR at fiberoptics tip	PAR at sample level (Standard geometry, distance to sample level 7.5 mm, angle 60°. See Fig. 10, page 22.)		
Setup	Fiber touches center of the diffusing disk of the MINI-SPEC.	Fiber flush with adapter which is fully inserted in 60° port of the calibration block.		
Calibration of internal light sensor	Spec. Cal. Factor = 3.01 Spec. Cal. Factor = 1.22 See Section 4.2.4.1, page 70. The Spec. Cal. Factor can only be entered on the DIVING-PAM screen.			
Calibration of actinic light list	Go to "Actinic Light List" on MINI-PAM-II display (Section 4.1.10, page 52) or Settings Window of WinControl-3 (Section 6.10.3, page 136), execute "Calibrate".			

3.2.5 Underwater Cables DIVING-II/K25 and DIVING-II/K50

Underwater cables of two different lengths are provided for remote control and log-term measurements.

4 Stand-alone Operation

Stand-alone operation of the DIVING-PAM-II allows saturation pulse analysis of photosynthesis independent of a computer. Using water-tight reflection switches, single F_V/F_M or Y(II) measurements can be triggered, but more complex experiments like induction and light curves are also feasible. Continuous recording of steady state fluorescence, however, requires operation of the DIVING-PAM-II by WinControl-3 running on an external computer.

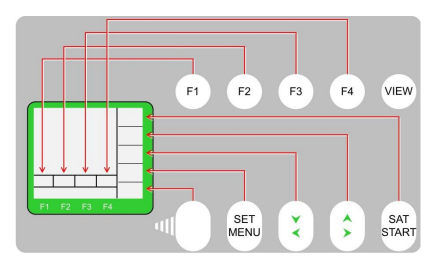


Fig. 22: Optical Switches and Display Fields

Note that "VIEW" is not linked to a display field but changes the appearance and opens navigation options of windows displaying graphical information (induction curve, light curve, spectra).

4.1 Top-Level Windows

The DIVING-PAM-II provides various windows for control and data display. Frequently used commands, fluorescence data and the actinic light list (PAR list) are presented in ten top-level windows (Table 4). Sections 4.1.1 to 4.1.10 will introduce all ten top-level windows in detail.

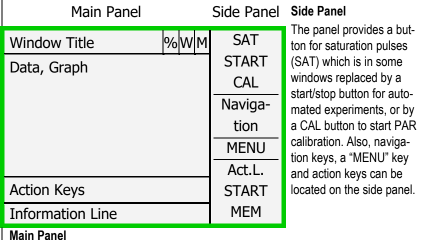
Accessible from all top-level windows is the Main Menu and its submenus which allow adjusting settings of the DIVING-PAM-II and its peripherals (Fig. 34, page 55).

Table 4: Overview of Top-Level Windows

Window		Content
1.	Basic Data	Minimum data set and basic action keys for filed operation.
2.	Primary Data	Data of last saturation pulse analysis and current levels of fluorescence and additional measured parameters.
3.	Quenching Analysis	Fluorescence levels of last saturation pulse analysis and corresponding fluorescence ratios (Y(II), F_V/F_M ,).
4.	Ft-Chart	Trace of fluorescence, total x axis interval: 25 or 125 s (see Fig. 60, page 84).
5.	Spectrometer	Measurement and display of light, fluorescence, or reflectance spectra.
6.	Actinic + Yield	Short illumination program with saturation pulse analysis.
7.	Induction Curve	Graphics of saturation pulse analysis of current curve.
8.	Light Curve	Graphics of saturation pulse analysis of current light response curve.
9.	Recovery Curve	Recovery graphics after induction curve and light response curve.
10	Actinic Light List	Set values of photosynthetically active radiation PAR in $\mu mol\ m^{\text{-}2}\ s^{\text{-}1}.$

All top-level windows consist of a "main panel" and a "side panel" (Fig. 23, page 37). The top line of the main panel displays window title and three small fields which from left to right display:

- Charge status of internal battery in %. When connected to the mains, "ext" is displayed.
- The WLAN status, where "W" indicates that the WLAN interface of the DIVING-PAM is active. WLAN can be activated via the item "Wireless" in menu "DIVING-PAM-II Settings" (Fig. 60, page 84).
- Mark. A capital letter, which is added as a mark to each saturation pulse analysis in the Report file.



Upper field: Window title, battery charge (%), WLAN status (W), and currently selected character to mark measurements. Central field: Numerical and/or graphical data. Lower fields: action keys and information line. Action keys are absent or differently arranged in some windows. The information line displays date + time or PAM activity or informs on critical fluorometer states.

Fig. 23: **Principal Screen Layout**

The bottom of the main panel provides various action keys. Depending on window, these action keys trigger saturation pulse analysis, control light conditions or affect graphic display. The "Information Line" at the bottom edge of the main panel shows current date and time.

The side panel provides arrow keys to switch windows, control keys for fluorometer functions, and a MENU key to access the main menu. Control keys are different between windows. For instance, the uppermost button in the sidebar triggers saturation pulse analyses in case of the first five top-level windows (cf. Table 4, page 36) but the same button starts automated experiments in windows Actinic + Yield, Induction Curve, Light Curve and Recovery Curve.

4.1.1 Basic Data

The Basic Data window (Fig. 24, page 40) displays a reduced set of data which is sufficient to monitor fast sampling under field conditions. Two data, Ft and PAR, represent current measurements but the Y(II), or F_V/F_M , and the ETR are derived from the last saturation pulse analysis.

The bottom of the Basic Data window provides keys for frequently used commands: the command Rec starts a new data set (Record), Fo,Fm determines maximum PS II photochemical yield, Clock starts repetitive triggering of saturation pulses or an automated experimental routine, and Mark opens the window "New Marker" in which the letter saved with each saturation pulse analysis ("mark") can be defined. To select the marker letter, use up and down keys in the window "New Marker". The currently selected letter is shown on the top edge of the window. Triggering

the SET key confirms current selection and returns to the Basic Data window.

The side panel of the Basic Data window offers four commands: the SAT key determines effective PS II photochemical yield, and the Act.L. represents a switch for actinic light. The MENU key leads to the Main Menu window as in all other top-level windows. Finally, a downward arrow key switches to the next window. Two arrow keys (up and down) are present in all other top-level windows except the last (Actinic Light List; Table 4, page 36) which requires only the upward key.

Comment on Rec (New Record)

Starting a new Record disconnects subsequent saturation pulse analyses from that of the previous Record. If the Rec is not followed by the command $\boxed{\text{Fo,Fm}}$, a $\boxed{\text{SAT}}$ command will only yield data of Y(II). In addition, q_P , and q_L will be calculated provided that the F_0 ' Mode is active. All parameters requiring F_0 and F_M (compare Table 27, page 168) cannot be calculated.

If in a new Record $\boxed{\text{Fo,Fm}}$ has been executed, and the F_0 ' Mode is inactive, the F_0 ' is calculated from F_0 , F_M and F_M ' (Section 10.2, page 162). Hence, all parameters of Table 27 can be calculated even when the F_0 ' Mode is inactive.

If a Record contains more than one F_0 , F_M determination, the last one will be used to calculate fluorescence coefficients. Such calculations, however, are valid only when measurements of F_0 , F_M and F_M are carried out with the same sample under the same conditions and settings.

In some cases, Fo,Fm and SAT measurements with different samples even result in fluorescence ratio parameters that exceed their valid range (compare Table 27, page 168). These invalid data will be displayed in grey on the screen.

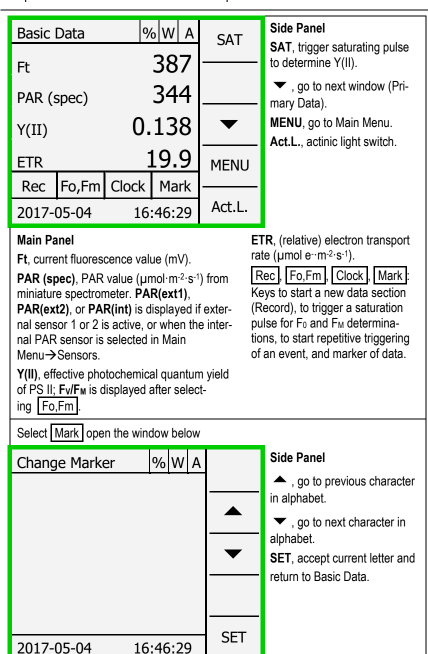
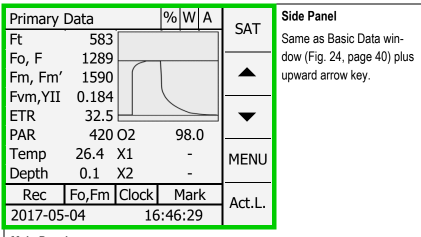


Fig. 24: Basic Data/Change Marker

4.1.2 Primary Data

The Primary Data window (Fig. 25) includes a monitoring graph displaying the fluorescence transient induced by the last saturation pulse. The window shows the "Basic Data" Ft, F_V/F_M or Y(II), ETR, and PAR. Additionally displayed are the F_0 and F_M signal levels (after F_0 , F_M) or the F and F_M ' signal levels (after F_0). Also, temperature, °C, and Depth are shown. When an optode oxygen sensor is connected, oxygen data are displayed (O2).



Main Panel

Ft, current fluorescence level

Displayed after Fo,Fm: Fo, Fm, and Fv/Fm; minimum and maximum fluorescence levels of dark-acclimated sample, maximum photochemical PS II yield. Displayed after SAT: F, Fm', and Y(II); fluorescence level just before saturation pulse and maximum fluorescence of light-exposed sample, effective photochemical quantum yield of PS II

Insert, fluorescence trace during last saturation pulse.

PAR, Temp, and **Depth**, photosynthetic active radiation in μ mol·m·2·s·1, temperature in °C, and diving depth in m, respectively.

O2, **X1**, **X2**, data measured by optional oxygen and auxiliary sensors, respectively.

Rec , Fo,Fm , Clock , Mark : see Fig. 24, page 40.

Fig. 25: Primary Data

4.1.3 Quenching Analysis

The window "Quenching Analysis" (Fig. 26, page 42) provides an overview on fluorescence levels and fluorescence ratio quotients calculated by the DIVING-PAM-II. Data line 2 to data line 4 (Fm'_Fm, Fo'_Fo, Y(II)_Fv/Fm), compare the data of the light exposed sample (left) with that of the dark-acclimated sample (right). All parameters of Fig. 26 are explained in Table 27 (page 168).

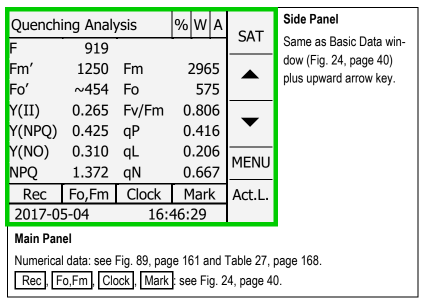


Fig. 26: Quenching Analysis

4.1.4 Ft-Chart

The Ft-Chart displays a 25 or 125 s interval of Ft where the right-most level of the graph corresponds to the current Ft value. The X axis interval can be adjusted in the menu "DIVING-PAM-II Settings (Fig. 60, page 84).

Continuous Ft values are not saved when the DIVING-PAM-II is operated in the stand-alone mode. Recording of Ft requires operation of the DIVING-PAM-II by WinControl-3.

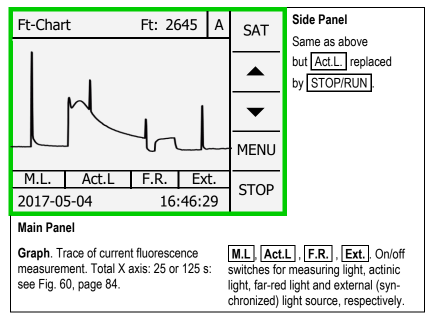


Fig. 27: Ft Chart

4.1.5 Spectrometer

The SPEC command of the Spectrometer window triggers reading data from the spectrometer. In the standard mode, the spectrometer is equipped with an optical entrance for external radiation (Fig. 9, page 21). Spectra of photon flux densities are measured in units of nmol photons m⁻² s⁻¹ nm⁻¹. PAR corresponds to the integral of the spectrum from 400 to 700 nm. Spectrometer-derived PAR will be used for ETR calculations if "Use Ext. PAR" in Menu Sensor Settings is checked (available by selecting "Sensors" in the Main Manu). Also, make sure that the spectrometer is selected as external PAR sensor (Main Menu → Sensors→Sensor Settings → Leaf Clip/Ext PAR Sett → External PAR sensor).

Note that the spectrometer measures the PAR of the DIVING-PAM-II internal light correctly only in the CAL mode (Fig. 33, page 54). Only in the CAL mode the LEDs are operated continuously. In the measuring mode, actinic light is pulse-width modulated and fluctuating PAR is displayed.

To record fluorescence or reflectance spectra, change spectrometer configuration as described in (Fig. 9, page 21). Then select operation mode: Main Menu → Sensor → Spectrometer → Operation Mode.

In menu Operation mode, the items "Fluores. (blue)", "Fluores. (green)" and "Reflectance" set the spectrometer mode to fluorescence spectra with blue excitation, fluorescence spectra with green excitation and reflectance spectra, respectively.

For reflectance measurements, first measure dark current in complete darkness, then the 100% reflection signal with the white reference material, and finally the sample.

In the Spectrometer window, touching the VIEW key displays a vertical cursor line as well as the x-y data of the intersection between cursor line and spectrum. Use the arrow keys to move through the spectra. Another VIEW command returns to the original function of arrow keys.

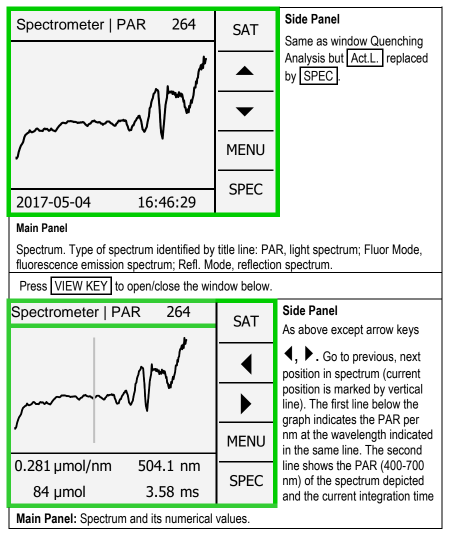
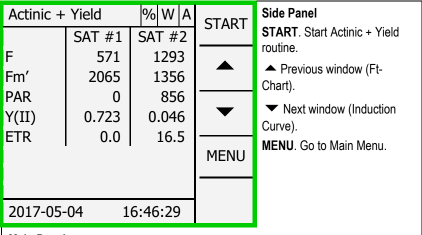


Fig. 28: Spectrometer

4.1.6 Actinic + Yield

The "Actinic + Yield" window is the first of four windows designed for starting and monitoring automated measuring routines. The parameters of the Actinic + Yield routine can be set in the menu "Program/Clock" (Fig. 40, page 64). The routine illuminates a sample with actinic light of a defined period. Depending on settings, saturation pulse analysis is carried out prior and after actinic light exposure, or only after actinic light exposure.

Because the Actinic + Yield experiment is rather short, it is frequently employed when light response of photosynthesis needs to be compared for many samples.



Main Panel

SAT #1, SAT #2. Column for data of first and second saturation pulse analysis, respectively.

F, **Fm**'. Fluorescence level before and at saturation pulse. For dark-acclimated samples: $F=F_0$ and $F_M'=F_M$.

Y(II). Effective photochemical quantum yield of PS II (F_V/F_M of dark-acclimated samples).

PAR, ETR. Photosynthetically active radiation (μmol·photons m⁻²·s⁻¹), electron transport rate (μmol·electrons m⁻²·s⁻¹).

Fig. 29: Actinic + Yield

4.1.7 Induction Curve

On this window, induction curves fluorescence induction experiments can be started and monitored (Fig. 30, page 48). The graphics panel provides a qualitative picture of induction curve properties. Because of the rather low screen resolution, it is difficult to quantitatively evaluate these graphical data.

The Induction Curve's numerical data can be accessed by the VIEW key. Navigation keys (◀, ▶.) permit passing from one saturation pulse analysis to another. The position in the induction curve of the current set of numerical data is indicated by a grey vertical line.

The parameters of an induction curve experiment, the number of saturation pulse analyses and the interval between them, can be adjusted in the menu "Induction Curve Settings" (Fig. 43, page 67), which is available over the Program/Clock line of the Main Menu (Fig. 34, page 55). In the "Induction Curve Settings" menu, a fluorescence recovery experiment (actinic illumination switched off) can be placed right after an induction curve.

The MEM of the Induction Curve window opens the "Induction Curve Memory" window. In this window, the navigation keys permit scrolling through stored induction curves.

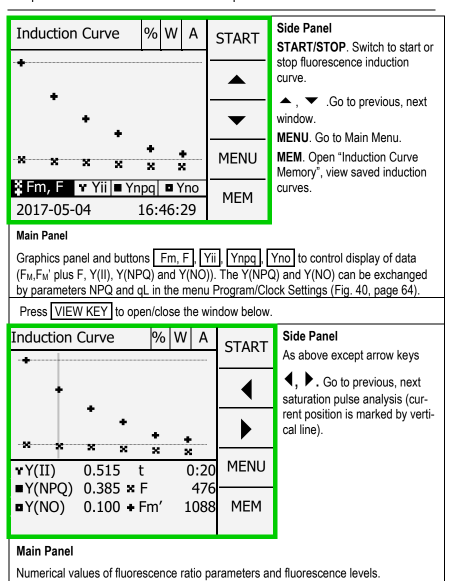


Fig. 30: Induction Curve

4.1.8 Light Curve

In analogy to the previous window (Induction Curve, 4.1.7), the "Light Curve" window provides buttons to start and stop a light curve, and it allows monitoring the light curve experiment (Fig. 31, page 50). Also, the Light Curve window provides numerical data of saturation pulse analysis which can be accessed by pressing the VIEW key. Navigation between different saturation pulses analyses and selection of displayed data works as described for the previous window (Induction Curve).

Light curve parameters (number and duration of light steps, initial PAR, and recovery curve) can be adjusted in the menu "Light Curve Settings" (Fig. 44, page 68). Selecting the MEM button and scrolling using the arrow keys on the side bar permits viewing stored light curves.

4.1.9 Recovery

Fluorescence recovery experiments can be automatically appended to an induction or a light curve (see item Program/Clock of the Main Menu; Fig. 34, page 55). When selected, the recovery curve is started automatically. Otherwise, recovery curves can be initiated (and cancelled) manually using the START/STOP button on the side panel of the Recovery window (see Fig. 32, page 51).

As in previous Induction and Light Curve windows, recovery data are represented graphically and numerically. The time course of recovery curves is determined: each curve last 39 min during which 7 saturation analysis are carried out. In case of a preceding induction or light curve, the last saturation pulse analysis of the induction or light curve corresponds to the first one of recovery.

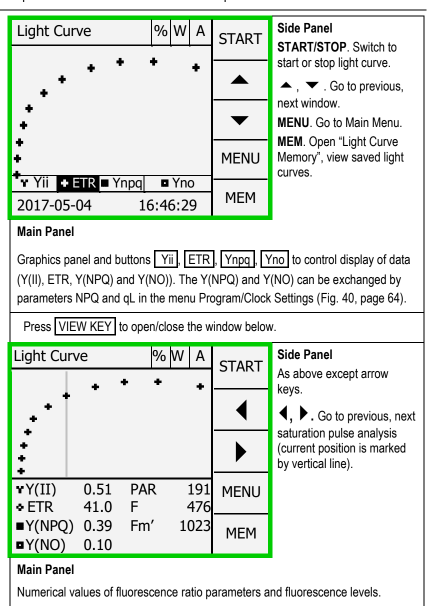


Fig. 31: Light Curve

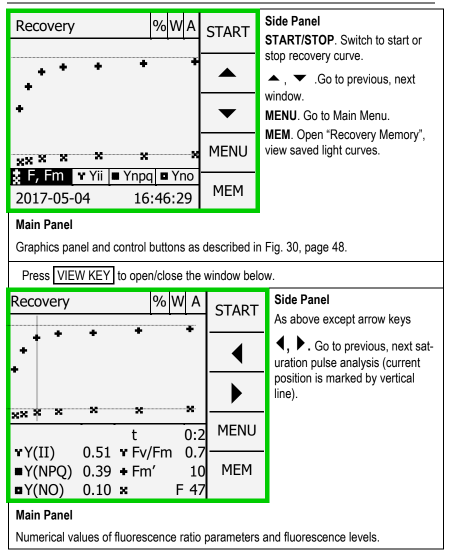


Fig. 32: Recovery

In recovery experiments, the interval between neighboring saturation pulse analysis roughly doubles with time (Table 5).

In the Recovery window, the MEM button opens the window "Recovery Memory" in which all recovery kinetics can be viewed independent if they are connected to an induction or light curve or if they represent separate experiments.

SAT number	Time in darkness, min
1	0.00
2	0:30
3	1:30
4	4:00
5	9:00
6	19:00
7	39:00

Table 5: Time Points for Recovery Curves

4.1.10 Actinic Light List

The window "Actinic Light List" contains PAR values (µmol m⁻² s⁻¹) which increase roughly exponentially with intensity settings. In the factory, blue or red LED emission is adjusted so that these PAR values apply for the sample level of the Universal Sample Holder DIVING-II-USH with the DIVING-PAM-II fiber optics fully inserted (i.e. distance between fiber optics tip and sample level is around 7.5 mm; angle between end piece of fiber optics and sample level: 60°). (Because of similar geometrical arrangement, these factory settings also apply to the Leaf Clip Holder 2035-B, 2010-A Distance Clip 60°, and the 2060-B Arabidopsis Leaf Clip when distance rings 2 mm plus 4 mm are used.)

The CAL command

The DIVING-PAM-II has an internal PAR sensor which is exposed to a small and constant fraction of LED emission. In the factory, this internal sensor is adjusted so that it displays the same PAR values as a calibrated cosine-response PAR sensor at sample level of a Universal Sample Holder DIVING-II-USH.

The CAL command varies LED emission until for each actinic intensity stetting (1 to 12) the readout of the internal PAR sensor matches the corresponding value of the light list. Therefore, after carrying out CAL, the PAR values of the light list apply to the sample level of a Universal Sample Holder DIVING-II-USH, the 2035-B leaf clip, and devices with similar geometry. This way the CAL command can compensate reduced light transmission of older fiber optics or intensity losses of LEDs used for long periods. Note that the entire PAR list can be multiplied by the "Actinic Factor" (Fig. 38, page 62).

Various measuring configurations

The use of the CAL command for other than factory configuration requires new adjustment of the internal sensor. For this purpose, a calibrated PAR sensor is positioned at sample level of the selected measuring setup and the internal PAR sensor is adjusted as described in Section 4.2.4.1, page 70 (see also Fig. 10, page 22). In short:

Using a calibrated PAR sensor which is not connectable to the DIVING-PAM-II requires manual adjustment. This is done by switching actinic light on and varying in the window "Int. PAR Sensor Settings" (Fig. 47, page 72) the "Calibration Factor" until internal PAR readout matches that of the external sensor.

If the calibrated PAR sensor feeds its data in the DIVING-PAM-II (e.g. the MINI-SPEC spectrometer or the PAR sensor of 2035-B

leaf clip), the internal PAR sensor can be calibrated automatically. To do so, select "Calibrate" in window "Int. PAR Sensor Settings" (Main Manu \rightarrow Sensors \rightarrow Internal PAR) and start automatic adjustment of calibration factor.

To calibrate the internal sensor with the MINI-SPEC spectrometer, use the PAR calibration block (Fig. 10, page 22).

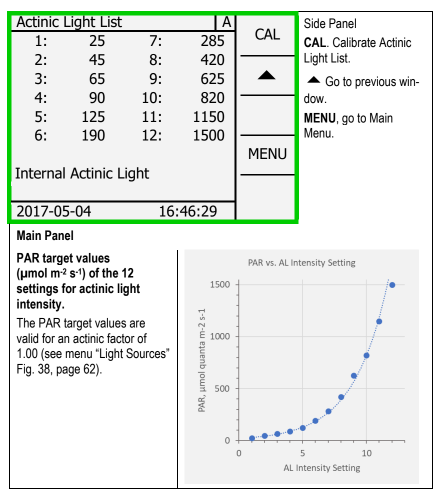


Fig. 33: Actinic Light List

4.2 Main Menu

The "Main Menu" (Fig. 34, page 55) forms the central hub to access settings, calibration data, hardware information and the memory of the DIVING-PAM-II. The Main Menu consists of seven items (Fig. 34, page 55). Selection of lines in the Main Menu is achieved by the arrow keys of the side panel. Most of these items lead to submenus that itself link to lower-level PAM Settings menus.

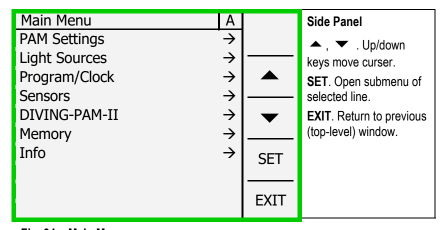


Fig. 34: Main Menu

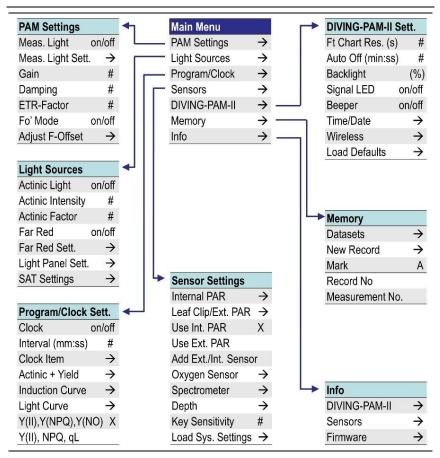


Fig. 35: Main Menu and Submenus

4.2.1 PAM Settings

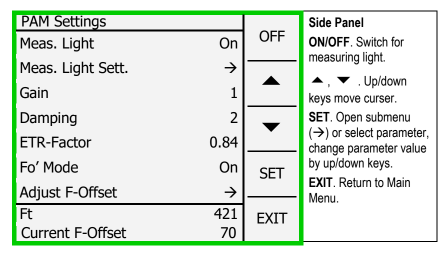


Fig. 36: PAM Settings

"PAM Settings" include adjustments of how the DIVING-PAM-II acquires PAM fluorescence. The menu contains seven items (Fig. 36, page 57), the window also displays the current Ft value and the currently active offset which is automatically subtracted from the raw signal.

Meas. Light

On/Off switch for measuring light (weak excitation light consisting of µs pulses). Measuring light can be switched on by selecting "Meas. Light" and selecting SET.

Meas. Light Sett.

This command opens the menu "Measuring Light Settings" in which measuring light intensity (Meas. Light Int.) can be selected by the SET key and then adjusted using the arrow keys. At constant frequency, measuring light intensity can be considered as being proportional to intensity settings 1 to 12.

The two remaining items of the menu "Measuring Light Settings" affect measuring light frequency. Selecting the line "Meas. Light. Freq." by $\boxed{\text{SET}}$ allows choosing between 5 frequency levels using arrow keys (see Table 6, page 59). The highest measuring light frequency of 100 Hz can be activated via the line "ML-F High". The ML-F High command overrules settings made under "Meas. Light. Freq." High measuring light frequency improves signal quality but bears the risk that its higher intensity drives photosynthesis, that is, the measuring light becomes actinic. In this case, F_0 measurements may be wrong.

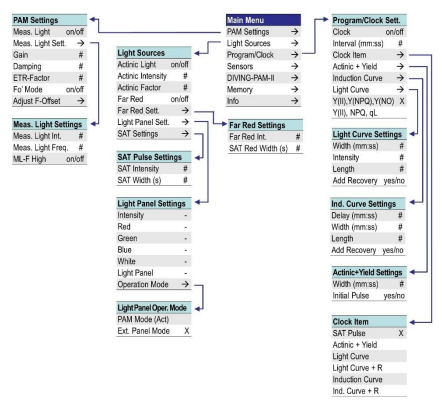


Fig. 37: Submenus PAM Settings
Light Sources, and Program/Clock

Setting	Frequency, Hz
1	5
2	10
3	15
4	20
5	25
high	100

Table 6: Measuring Light Frequencies

Measuring light frequency is automatically switched to high during a saturation pulse, that is, for F_M and F_M ' determinations. Measuring light frequency does not affect the frequency of Ft data acquisition by the WinControl-3 software.

The average PAR of measuring light at highest frequency and highest intensity setting was 1.5 µmol m⁻² s⁻¹ as measured by the PAR sensor of a 2035-B leaf clip, where the DIVING-PAM-II fiber was fully inserted in the fiber port. For the same geometrical arrangement, average measuring light intensities can be estimated using the subsequent equation:

$$I_{ML}(\frac{\mu mol}{m^2 \cdot s}) = 1.5 \ (\frac{\mu mol}{m^2 \cdot s}) \cdot \frac{f(Hz)}{100(Hz)} \cdot \frac{Int. \ Sett.}{12}$$

where I_{ML} , f and Int. Sett. is the current measuring light intensity in μ mol m⁻² s⁻¹, the current measuring light frequency, and the current intensity setting for measuring light, respectively.

Gain

Selecting "Gain" by the SET gives access to four electronic amplification factors (1 to 4) which can be adjusted by the arrow keys.

Damping

Damping is a software-based filter that specifically suppresses high frequency noise and, thus, can improve signal quality. Changing damping settings uses the same principle as described for "Gain". Default setting for damping is 2 (two). Changing damping to higher values can make the DIVING-PAM-II response slow.

ETR-Factor

This factor is used for ETR calculations and corresponds to the fraction of incident PAR absorbed by a leaf; its default value is 0.84 (cf. Section 10.3, page 166).

Fo' Mode

The "Fo' Mode" replaces after saturation pulses actinic by far-red light to quickly open PS II reaction centers. The measured F_0 ' is the minimum Ft during far-red illumination. Interval and intensity of far-red illumination can be adjusted in menu "Light Sources" (Fig. 38, page 62 or Section 6.10.2, page 135).

Adjust F-Offset

The "Adjust F-Offset" command determines the background signal for subtraction from the total signal. Background signals must possess the modulation characteristics of measuring light to be recognized by a PAM fluorometer. These signals can arise from:

- Fluorescence from suspension media or detector filter excited by measuring light.
- Traces of modulated excitation light transmitted by the detector filter.
- Non-optical modulated "electronic noise".

Usually, the background signal increases with measuring light intensity and signal amplification (gain). Therefore, the Adjust F-Offset command determines the background signal for all measuring

light intensities and all gain settings. The currently active offset is displayed in the bottom line of the PAM Settings window (Fig. 36, page 57).

Procedure

- Choose dim environment.
- Switch off any flickering light sources like fluorescent lamps or computer screens.
- Point fiber tip away from any objects, keep fiber tip clear.
- Run "Adjust F-Offset"

4.2.2 Light Sources

See Fig. 38, page 62.

Actinic Light

On/off switch for actinic light.

Actinic Intensity

Intensity regulation for actinic light. Select menu item by SET and choose setting using arrow keys. Settings 1 to 12 are available. Information on PAR is given in window "Actinic Light List" (Fig. 33, line 54).

Actinic Factor

Factor multiplying target PAR values in window "Actinic Light List". Factor range is 0.5 to 2.0.

Far Red

On/off switch for far-red light.

Far Red Sett.

Opens menu to adjust intensity and duration of far-red illumination. These settings are active in F₀' determinations

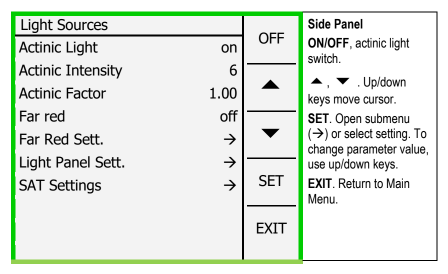


Fig. 38: Light Sources

Light Panel Sett.

Use SET to enter submenu "Light Panel Settings". The function of this window depends on the "operation mode". The operation mode is selected via the last line in the window (see Fig. 39, page 63).

Two operation modes are available: "PAM Mode (Act.)" and "Ext. Panel Mode". "PAM Mode (Act.)" means that actinic light is provided by an external source like the 2054-L External LED Source. (MINI-PAM-II accessory. Under dry lab conditions, the 2054-L lamp can be operated by the DIVING-PAM-II using a special adapter cable). "Ext. Panel Mode" means that the DIVING-PAM-II

internal light is used as actinic light to which an external light source may be added (see Table 21, page 131).

Only when "Ext. Panel Mode" is chosen, the item "Intensity" and the Light Panel On/Off switch are available in the previous window's (Light Panel Settings). In this case, "Intensity" determines the percentage of maximum intensity for all LED types (red, green, blue, and white). The numbers entered for an individual LED group correspond to percentages of the scale from zero to "Intensity". Typically, the maximum intensity of all four LED groups is greater than 5000 μ mol m⁻² s⁻¹. In "Ext. Panel Mode", the external actinic light can be operated in parallel.

Note: Operation of the external light source (2054-L External LED Source) requires line power.

Note: In absence of an external light source, "PAM Mode Act." is not available.

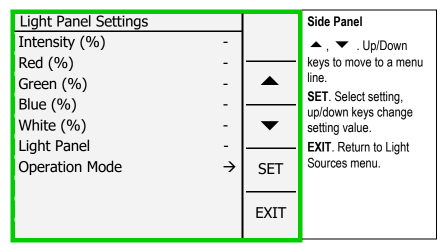


Fig. 39: Light Panel Settings

With "PAM Mode (Act.)", the total intensity of all four LED groups of the 2054-L is determined by the setting for actinic light intensity (see above). In the window "Light Panel Settings", the numbers for the four LED groups now represent ratio numbers determining the fraction with which each LED group contributes to total illumination.

SAT Settings

"SAT Settings" and SET opens the submenu "SAT Pulse Settings" in which relative intensity (1 to 12) and duration (width, 0.2 to 2.0 s) of saturation pulses can be set.

At sample level, intensity setting 12 corresponds to 6000 μ mol m⁻² s⁻¹ under standard geometrical conditions (distance fiber optics tip to sample level: 7.5 mm; angle between end piece of fiber optics and sample level: 60°). Saturation pulse intensity can be adjusted at increments of 500 μ mol m⁻² s⁻¹.

4.2.3 Program/Clock Settings

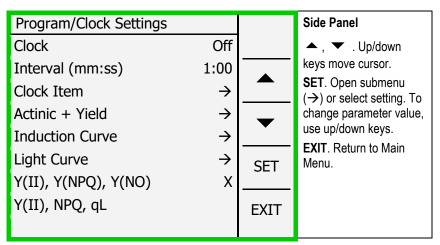


Fig. 40: Program/Clock Settings

The menu "Program/Clock Settings (Fig. 40, page 64) provides all options to configure automated experimental routines including Actinic + Yield, Induction Curve and Light Curve experiments as well as repetitive triggering of single measurements and experimental protocols.

Clock

On/off switch of clock. The "Clock" triggers repetitively an event at a defined interval. The interval is specified in "Clock Interval" and the event in "Clock Item".

Clock Interval

Adjust clock interval between 10 s and 60 min by selecting "Interval" (up/down keys and SET, respectively) and adjusting time interval (up/down keys).

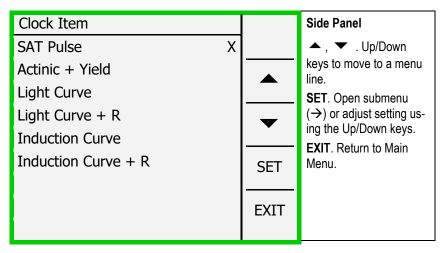


Fig. 41: Clock Item

Clock Item

Saturation pulse analyses and the programs Actinic + Yield, Induction Curve and Light Curve can be repetitively performed under clock control. Also, recovery experiments can be performed after induction and light curves (item Light Curve + R and Induction Curve + R, respectively). To select one of the six items in menu "Clock Item" (Fig. 41), move cursor to the item of interest and select SET. The selected item is then marked by an X ("SAT pulse is selected in Fig. 41).

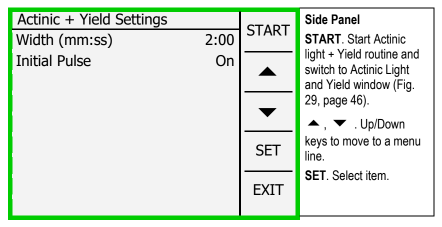


Fig. 42: Actinic + Yield Settings

Actinic + Yield

The behavior of the Actinic + Yield program is defined by two factors (Fig. 42, page 66): the duration (width) of actinic illumination (possible settings from 5 s to 5 min) and the option to start actinic illumination without preceding saturation pulse analysis (Initial pulse). Width of actinic illumination is adjusted as described above for clock interval and initial pulse is selected by the SET command. Further, actinic light intensity is adjusted in the window "Light Sources" (Fig. 38, page 62).

An Actinic + Yield routine can be started using the START button on the side panel of Fig. 42. In this case, the screen display will automatically switch to the Actinic + Yield experimental window (Fig. 29, page 46).

Induction Curve

Induction curve experiments are configured in the window "Induction Curve Settings" (Fig. 43, page 67).

<u>Delay</u> (range 5 s to 10 min) defines the dark interval between saturation pulse analysis with the dark-acclimated sample (F_0 , F_M determinations, Section 10.2, page 162) and beginning of actinic illumination.

<u>Width</u> (range 5 s to 10 min) is the time interval between two successive saturation pulse analyses during illumination.

<u>Length</u> is the number of saturation pulse analyses carried out during actinic illumination. Thus, the duration of actinic illumination is "Length - 1" times Width.

<u>Add recovery</u> appends a recovery curve to an induction curve (see Section 4.1.9, page 49 for information on recovery times).

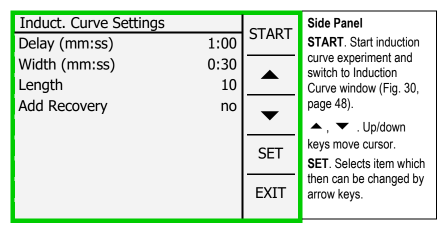


Fig. 43: Induction Curve Settings

Light Curve

Light Curves are defined in "Light Curve Settings" (Fig. 44, page 68). Properties of <u>Width</u> and <u>Add Recovery</u> are identical for light and induction curves.

<u>Intensity</u> specifies the actinic intensity setting for the first light step (range 1 to 5, for PAR values see Fig. 33, page 54).

<u>Length</u> is the number of light steps which can range from 2 to 12. If length = 5 and intensity = 2, 5 light steps with intensity settings 2, 3, 4, 5, and 6 will be performed. The time required for a light curve results from Length times Width.

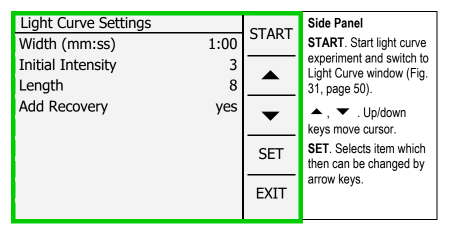


Fig. 44: Light Curve Settings

Select (Y(II), Y(NPQ), Y(NO)) or (Y(II), NPQ, qL)

The last two lines of the menu "Program/Clock Settings" affect graphical and numerical presentation of data in windows Induction Curve, Light Curve, and Recovery Curve. Selecting "Y(II), Y(NPQ) Y(NO)" displays three yield parameters that are used in analyzing energy partitioning. Choosing Y(II), NPQ, and qL displays the classical NPQ parameter and a parameter for indicating the reduction

state of PS II (q_L) . Selection between lines works as described for "Clock Item" (Fig. 41, page 65). See Table 27, page 168 for definitions of fluorescence parameters.

4.2.4 Sensors

Selection of "Sensors" in the Main Menu opens the window "Sensor Settings" (Fig. 45, page 69). This window includes six links leading to submenus (distinguished by →) and it allows to choose between the DIVING-PAM-II internal PAR sensor or the external sensor Miniature Spectrometer MINI-SPEC (Use Ext. PAR/Use Int. PAR). The 2035-B leaf clip (MINI-PAM-II accessory) can also be used as external sensor. Connect the 2035-B leaf clip using a special adapter cable (contact Walz) and operate the clip only under dry conditions). The currently active mode is marked by an X, to change selection, move cursor to the other sensor and select SET.

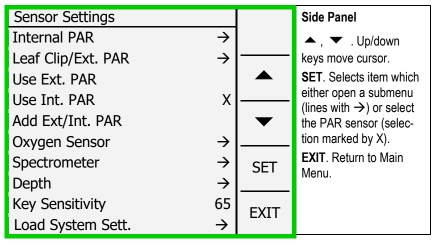


Fig. 45: Sensor Settings

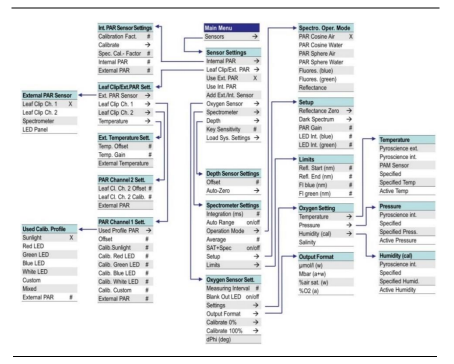


Fig. 46: Sensor Settings Menu and Submenus

4.2.4.1 Internal PAR

Selecting "Internal PAR" opens the window "Internal PAR Sensor Settings" (Fig. 47, page 72). This window is dedicated for calibration of the DIVING-PAM-II internal PAR sensor which receives continuously a small fraction of internal actinic light. The window displays the current "Calibration Factor", as well as the readouts (µmol m⁻² s⁻¹) of internal and external PAR.

The specific calibration factor (Spec. Cal-Factor) corrects for the effect of inhomogeneous illumination by actinic light of the diffusing disk of the miniature spectrometer in the PAR block arrangement (Fig. 10, page 22). The specific calibration factor is established at factory and automatically considered when the internal

sensor is calibrated against the miniature spectrometer, MINI-SPEC.

CALIBRATION

PAR Sensor compatible with DIVING-PAM-II: To calibrate the internal PAR sensor, connected to the DIVING-PAM-II a calibrated PAR sensor (e.g., MINI-SPEC), position diffusing disk of sensor (light entrance) at sample level and place fiber end as in the experimental situation, considering angle and distance (see comments in Section 3.1.4, page 19). The click Calibrate and establish calibration factor automatically.

Other PAR sensor: To calibrate the internal PAR sensor using an external PAR sensor which cannot be read by the DIVING-PAM-II, adjust calibration factor manually by selecting "Calibration Factor", pressing SET and changing the factor by arrow keys until internal PAR matches that of the external sensor.

Using the 2035-B leaf clip

The specific calibration factor is set to 1 when a 2035-B leaf clip is used for calibration of the internal light sensor. To use the 2035-B leaf clip for calibration, select "Leaf Clip/Ext. PAR" on window Sensor Settings (Fig. 45, page 69) to open window Leaf Clip/Ext. PAR Sett. (Fig. 48, page 73). On this window, select "Ext. PAR Sensor" and then "Leaf Clip Ch. 1" on window External PAR Sensor (Fig. 49, page 73).

When the blue light of a DIVING-PAM/B fluorometer is used for calibration of the internal sensor, the calibration factor for blue light (stored in the flash memory of the 2035-B leaf clip) will automatically be used. Similarly, when calibration is performed with a DIVING-PAM/R fluorometer, the "red" calibration factor will be used.

Proper calibration of internal sensor is apparent from same readouts of internal and external PAR (only 2035-B leaf clip; the MINI-SPEC shows a fluctuating signal when illuminated with the pulse-width modulated actinic light from the fluorometer). If calibration of the internal PAR sensor has previously been carried out but the two PAR values differ clearly, check if:

- (1) External PAR sensor is in the center of the actinic light beam.
- (2) Setup of previous calibration was different from the current one, for instance, in the previous calibration the external PAR sensor was not at standard distance from fiber tip.
- (3) External PAR sensor is connected, or incorrect PAR channel is selected in window "External PAR Sensor" (Fig. 49, page 73).

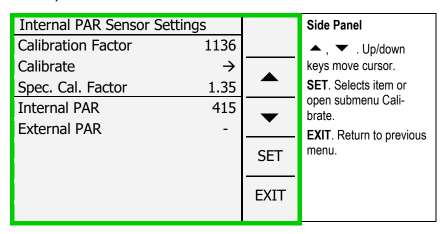


Fig. 47: Internal PAR Sensor Settings

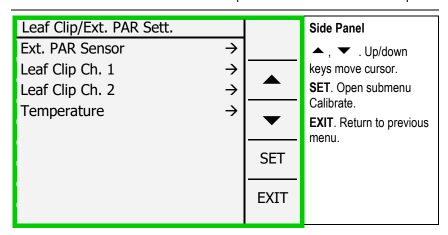


Fig. 48: Leaf Clip/External PAR Sensor

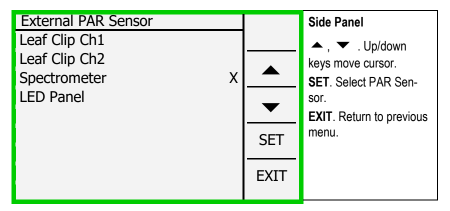


Fig. 49: External PAR Sensor

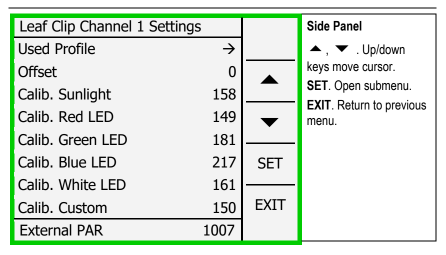


Fig. 50: Leaf Clip Channel 1 Settings

4.2.4.2 Leaf Clip/Ext. PAR

Selecting "Leaf Clip/Ext. PAR" opens the window "Leaf Clip/Ext. PAR Sett" (Fig. 48, page 73). The menu considers that various PAR sensors can be connected to the DIVING-PAM-II including the 2035-B leaf clip via a special adapter. The window provides four items:

<u>Ext PAR Sensor</u> opens a menu that selects the active external PAR channel (Fig. 49, page 73). Four possible PAR sensors are considered: PAR sensors 1 and 2 of 2035-B leaf clip, the miniature spectrometer, and a PAR sensor connected to the LED-Panel RGBW-L084. In the normal configuration of the DIVING-PAM-II, Spectrometer is selected.

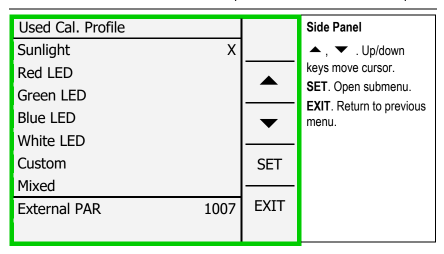


Fig. 51: Used Cal. Profile

Leaf Clip Ch. 1 opens "Leaf Clip Channel 1 Settings" which contains various calibration factors for PAR channel 1, that is, the mini quantum sensor of Leaf Clip Holder 2035-B. The different calibration factors ("Calib. Sunlight to Calib. White", see Fig. 50) were factory-established and are characteristic for each PAR sensor. By considering the spectral variations in sensitivity of the PAR sensor, these calibration values are optimized to measure various light qualities. Specifically, Calib. Sunlight is optimized to measure sunlight under clear skies. Calib. Red, Calib. Green, Blue and Calib. White LED applies to measurements of light from Red, Green, Blue, and White LEDs. Calib. Custom can be chosen by the user. The "Offset" in the window "Leaf Clip Channel 1 Settings" applies to all PAR measurements.

The item <u>Used Profile</u> in window "Leaf Clip/Ext. PAR" opens another menu (Used Cal. Profile, Fig. 51, page 75) in which a calibration factor or a calibration profile can be selected. In "Used Cal. Profile" the items "Sunlight" to "Custom" correspond to comparable items in the previous window., For example, selecting profile

"Sunlight" activates the factor "Calib Sunlight" for PAR measurements.

The profile "Mixed" adjusts the calibration factor dynamically depending on the contributions of the red, green blue and white LED of the External LED Light Source 2054-L (compatible with DIVING-PAM-II with special adapter for dry lab use).

<u>Leaf Clip Ch. 2</u> in window "Leaf Clip/Ext. PAR Sett" (Fig. 48, page 73) opens the calibration menu for a PAR sensor connect to the SMA socket on the side of the 2035-B leaf clip. This menu simply consists of slope (Calib,) and offset of the calibration line are given. In the lowest line, the PAR readout of the active PAR sensor is shown.

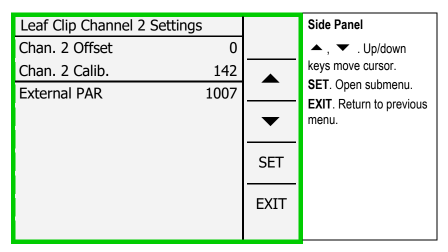


Fig. 52: Leaf Clip Channel 2 Settings

<u>Temperature</u> in window "Leaf Clip/Ext. PAR Sett" (Fig. 48, page 73) leads to the calibration data for the leaf temperature sensor of the 2035-B leaf clip (External Temp. Sensor Sett., Fig. 53, page 77). This window lists slope (Gain) and offset of the calibration line. Factory settings are Gain = 1.00, Offset = 0.

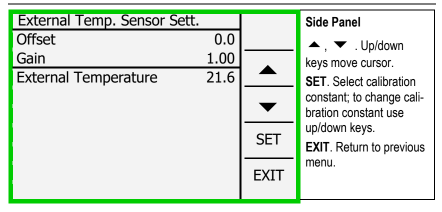


Fig. 53: External Temperature Sensor

4.2.4.3 Oxygen Sensor

The window "Oxygen Sensor Settings" controls the underwater oxygen sensor DIVING-PAM-II/O2. When the fiber-type oxygen sensor of the MINI-PAM-II oxygen package connected, the settings in the window apply to this device.

See separate manual for details:

https://www.walz.com/products/chl_p700/diving-pam-II/downloads.html

4.2.4.4 Spectrometer

The item "Spectrometer" in menu "Sensor Settings" leads to the window "Spectrometer Settings" (Fig. 54, page 79) which configures the miniature spectrometer.

<u>Integration time (ms)</u> determines the measuring time in ms for a single spectrum.

<u>Auto Range</u> in the active state ignores the integration time set above and determines the optimal integration time automatically.

<u>Operation Mode</u> opens another menu (Fig. 55, page 79) in which the type of spectrum (light, fluorescence, or reflectance). For light

spectra, the entrance optics (flat=cosine versus spherical) and the environment in which measurements are performed (air or water) can be specified.

The cosine configuration employs a diffusing disk as light entrance (Fig. 9). This configuration shows an approximate cosine response toward incoming radiation. The sphere configuration measures light from all directions with similar weight. At the time of writing of this manuscript, the spherical sensor is under development.

<u>Average</u> determines the number of measurements used to calculate the final spectrum.

<u>SAT+Spec</u> active links recording of a spectrum to a saturation pulse analysis.

<u>Setup</u> opens a menu for recording of the dark spectrum of the miniature spectrometer (Dark Spectrum \rightarrow). The dark current of the spectrometer at room temperature is measured in the factory and stored on the flash memory of the device. To newly establish the dark current, fully darken the entrance optics (cv. Fig. 9, page 21).

<u>Limits:</u> In the menu "Limits" short and long wavelength borders of reflectance spectra can be defined, and also the short wavelength limits of fluorescence emission spectra.

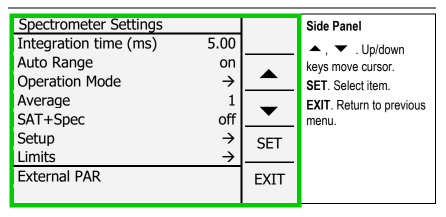


Fig. 54: Spectrometer Settings

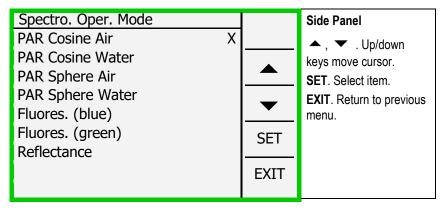


Fig. 55: Spectro. Oper. Mode

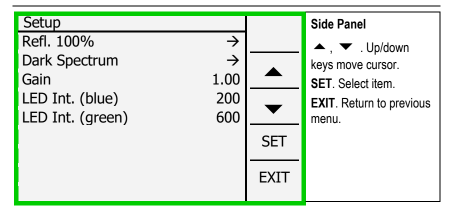


Fig. 56: Setup

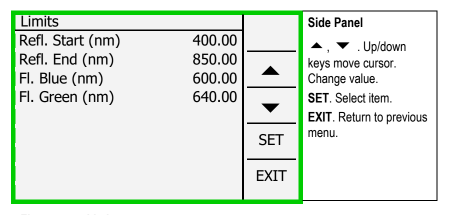


Fig. 57: Limits

4.2.4.5 Depth

Selecting Depth opens a submenu in which the offset for depth measurements is adjusted.

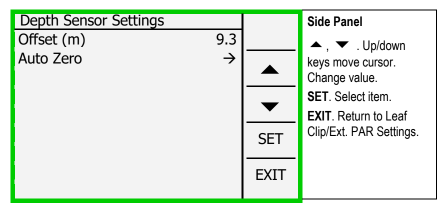


Fig. 58: Depth Sensor Settings

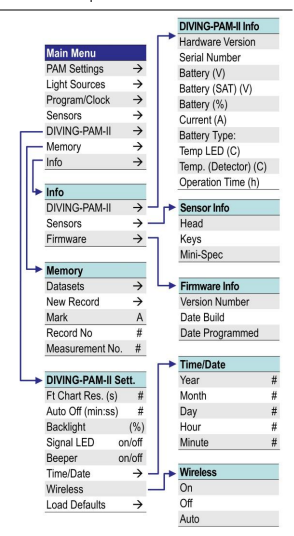
4.2.4.6 Key Sensitivity

Increasing the numerical value increases sensitivity of optical switches of the DIVING-PAM-II. Usually, a higher sensitivity is employed underwater compared to the use in air.

4.2.4.7 Load System Sett.

"Load System Settings" restores settings like calibration factor of internal PAR sensor (calibrated for geometry of 2035-B leaf clip), measuring light current and calibration factors of external devices like those stored on the 2035-B leaf clip. "Load System Settings" must not be confused with "Load Defaults" (Fig. 60, page 84).

Fig. 59: Memory and Info



4.2.5 DIVING-PAM-II

The menu "DIVING-PAM-II Settings" permits adjusting of various fluorometer settings.

Ft Chart Resolution (s)

Time resolution of Ft Chart (Fig. 27, page 43) can be either 0.2 or 1.0 s/dot corresponding to 25 or 125 s/total time axis.

Auto Off (min)

Time interval without saturation pulse analysis after which the DIVING-PAM-II powers off. The DIVING-PAM stays on when the clock is running or when communicating with an external computer either via cable or using WLAN.

Backlight (%)

Percentage of maximum intensity of the display's backlight LED array.

Signal LED

On/off switch for LED on top of the DIVING-PAM-II (green flash every 2 s, normal operation; green double flash every 2 s, clock-controlled operation; continuous light, saturation pulse analysis; green flash every 10 s, sleep mode).

Beeper

On/off switch for beeper which acoustically confirms keystrokes and saturation pulse analysis.

Time/Date

Menu for setting time and date.

Wireless

Select to open submenu "Wireless". Where:

On WLAN always on, which is battery-consuming on the long run

Off WLAN always off

Auto WLAN is activated with powering up the DIVING-PAM and deactivated after 5 minutes.

Load Defaults

The DIVING-PAM-II keeps on internal memory the default settings for the variables listed in Table 7, page 85. Virtually all parameters for which default settings are defined are also saved when the DIVING-PAM-II is shut off (Table 7). This way, the last settings are activated when powering on the DIVING-PAM-II. Defaults setting can be restored by "Load Defaults".

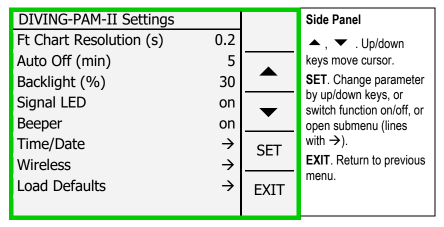


Fig. 60: DIVING-PAM Settings

Table 7: Default Settings

	Default Setting	Custom Setting (saved and restored)
Measuring Light Status Intensity Frequency Frequency high status	On 6, relative unit 3, see Table 6 Off	No Yes Yes No
Actinic Light Status Intensity Factor	Off 6, relative unit 1.00	No Yes Yes
PAM Signal Gain Damping	1, relative unit 2, relative unit	Yes Yes
Far Red Light Width 5 Intensity	5, s 8, relative unit	Yes Yes
Saturation Pulse Intensity	10, relative unit (≙ 5000 µmol m⁻² s⁻¹)	Yes
Width	0.6, s	Yes
Program Actinic Li Actinic light width Initial Pulse	ght and Yield 30, s Yes	Yes Yes
Program Induction Delay Width Length	Curve 40, s 20, s 12	Yes Yes Yes
Program Light Cur Width Intensity Length	ve 20, s 3, relative unit 8, light steps	Yes Yes Yes
Clock Item Interval	Saturation pulse 60, s	Yes Yes

Table 7: Default Settings

	Defects Cetting	Out to the Cattle of
	Default Setting	Custom Setting (saved and restored)
Hardware		(carros entarcoscos)
Signal LED status	On	Yes
Beeper status	On	Yes
Automatic power	15, min	Yes
down	10, 11111	100
Background light	60%	Yes
Graphics		
Ft chart time reso-	0.2, s/dot	Yes
lution	·, · · · · · · · · ·	
External Light		
Status	Off	No
Total intensity	1%	Yes
Red LED	10%	Yes
Green LED	10%	Yes
Blue LED	10%	Yes
White LED	10%	Yes
Stirrer		
Status	Off	No
Speed	10%	Yes
Pre-SAT off	10, s	Yes
Reverse	0, s	Yes
Interval mode	Off	Yes
Interval	2, min	Yes
Stirring interval	5, s	Yes
Stir in program	Off	Yes
F ₀ ' Mode		
Status	Off	Yes
PAR Sensor		
Status	Internal	No
Mark		
Character	Α	Yes

4.2.6 Memory

The "Memory" window provides an overview on stored saturation pulse analyses (Datasets). Importantly, here fluorescence kinetics induced by saturation pulses can be viewed. To see stored induction and light curves, use the MEM key of window "Induction Curve" (Fig. 30, page 48) and window "Light Curve" (Fig. 31, page 50), respectively. Also in this window, new records can be started, and the mark of saturation pulse data can be changed.

Deletion of data from the DIVING-PAM-II internal memory is possible using software WinControl-3.

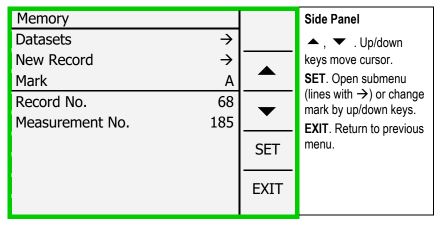


Fig. 61: Memory

4.2.7 Info

"Info" (Fig. 62, page 88) consists of three links to submenus of which three list hardware and software information:

- DIVING-PAM-II provides hardware information of the fluorometer including type of internal battery and operation hours.
- Sensors lists serial numbers of sensors connected to the DIVING-PAM-II.
- Firmware shows serial number and date of the firmware of the DIVING-PAM-II.

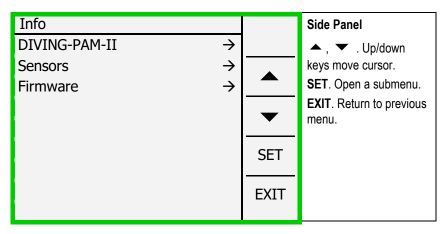


Fig. 62: Info

DIVING-PAM-II Info

Information available on this window is (1) hardware version of fluorometer, (2) serial number of fluorometer, (3) battery voltage at normal operation, (4) battery voltage during a saturation pulse, (5) charge status in percent, (6) present current consumption (7) type of built-in battery(Li-ion or lead acid), (8) temperature (in °C) of actinic LED, (9) temperature (in °C) of detector, and (10) operation time of device.

Note that the DIVING-PAM-II should be charged when the current battery voltage drops below 7.6 V.

DIVING-PAM-II Info		Side Panel
1 Hardware Version		EXIT . Return to previous
2 Serial Number		menu.
3 Battery (V)		
4 Battery (SAT) (V)		
5 Battery (%)		
6 Current (A)		
7 Battery Type		
8 Temp. LED (°C)		
9 Temp. Detector (°C)	FXIT	
10 Operation Time (h)	LVII	

Fig. 63: DIVING-PAM-II Info

5 WinControl-3 Installation

The WinControl-3 software is provided on a Walz USB flash drive. The WinControl-3 software are regularly optimized. The latest software version is available on the Walz website:

https://www.walz.com/products/chl_p700/diving-pam-II/downloads.html

WinControl-3 can be installed from the Walz USB flash drive or using the setup software downloaded from the Walz website.

5.1 Installation process

Installation of WinControl-3 is mostly automatic. Dialog boxes appearing during setup provide advice or allow configuration of WinControl-3. To install WinControl-3, proceed as follows:

- Close other programs as advised by the setup wizard.
- Execute setup file: double-click on file or right-click on file and choose "run" from context menu.
- A pop-up windows must appear which identifies the Heinz Walz GmbH as verified publisher.
- Accept default folder for program installation or choose different folder after clicking <u>Browse...</u>.
- Select "Standard" Installation. (The "JUNIOR-PAM Teaching Edition" runs only with JUNIOR-PAM fluorometers.)
- Install USB driver and select optional WinControl-3 links (icon or shortcut).
- Connect DIVING-PAM-II to computer and run PAM Firmware
 Update. If the current firmware* is outdated, PAM Firmware

Update will automatically replace it by the recent version. Running PAM Firmware Update after installation of WinControl-3 is important because new software properties may function only in the presence of the latest firmware.

*The term firmware denotes a piece of software residing on a flash memory of the DIVING-PAM-II. The firmware is integrated in the WinControl-3 software.

5.2 WinControl-3 Program Group

Setup of WinControl-3 creates the WinControl-3 program group (Table 8, p. 92) in the Windows Start menu. The WinControl-3 program group is comprised of 5 items. The items are introduced in sections 5.2.1 through 5.2.5.

Table 8: WinControl-3 in Windows Start Menu



WinControl-3



PAM Firmware Update



Uninstall WinControl-3



WinControl-3



WinControl-3 - Network Mode



WinControl-3 – Offline

5.2.1 PAM Firmware Update

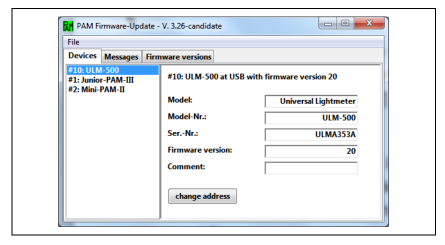


Fig. 64: PAM Firmware Update

Several devices connected to the same computer must have different addresses (channel numbers).

Initialization of PAM Firmware Update triggers a search for PAM devices connected to the computer. The result is displayed in the right panel of window "Devices" (Fig. 64, page 93). Each device name is preceded by its address number (between hash and colon).

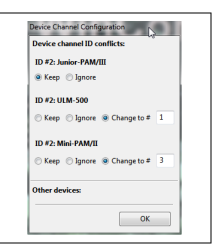
"PAM Firmware Update" compares the firmware in the device with the firmware included in the WinControl-3 software. If WinControl-3 includes newer firmware, the device is automatically updated.

"PAM Firmware Update" cannot update firmware of first-generation devices (DIVING-PAM, MICROFIBER-PAM, MICROSCOPY-PAM, MINI-PAM, WATER-PAM). In these devices, firmware resides on an EPROM chip and firmware update requires exchange of this chip.

A device can be selected by mouse click. The currently selected device is highlighted (white letters on blue background). The main panel of the window shows information on the device selected. The first four information lines define the hardware and software state of the device. The last line displays a comment associated with the device and typed in using the WinControl-3 software.

Devices with identical addresses cannot be operated simultaneously. If WinControl-3 detects identical addresses, the window "Device Channel Configuration" (Fig. 65, page 94) pops up offering a working address configuration and the option to change addresses manually. Note that address number is synonymous to channel number in the software WinControl-3.

Fig. 65: Device Channel Configuration



Address numbers can be changed manually via the button change address. Then, determine new address by picking a number from the drop-down list "New address:". If several devices are connected, the drop-down list offers only unused address numbers. The window "Messages" displays the protocol of activities including firmware update of devices. The window "Firmware Versions" compiles all software version provided by PAM Firmware Update.

5.2.2 Uninstall WinControl-3

This program removes WinControl-3 and all its links. It does not remove the USB driver software.

5.2.3 WinControl-3

This command starts WinControl-3 in the default mode. When devices which are compatible with the WinControl-3-type software are detected, WinControl-3 enters the measure mode. Clicking Offline-Mode interrupts the search process and WinControl-3 is started in the offline mode. When the search process fails to find compatible devices, a pop-up window provides three options: Yes triggers another search for devices, No launches the offline mode of WinControl-3, and Cancel ends the whole process. Measuring mode and offline mode, and several instances of WinControl-3 in the offline mode, can run in parallel.

After detection of the DIVING-PAM-II, measuring of fluorescence is automatically started. With a green leaf placed in the 2035-B leaf clip, fluorescence values of 400 to 600 are observed (Ft value, bottom of window). Check "Rec. Online" or click Start Online to display continuously the Ft on the Chart window. If the Ft is much lower than 400, make sure that the optical fiber is properly inserted. Click Autoscale if data are not visible. Trigger saturation pulse analyses by pressing Fo, Fm or SAT A healthy leaf, which was kept dark before, should show a value for F_V/F_M of 0.8 or higher.

5.2.4 WinControl-3 Network Mode

The Network mode requires MONITORING-PAM measuring heads and connection via a special interface. It is not available for the DIVING-PAM-II.

5.2.5 WinControl-3 Offline Mode

This command launches Wincontrol-3 without the initial search for available PAM devices.

6 WinControl-3 Operation

WinControl-3 offers the same functions as touchscreen operation of the DIVING-PAM-II, except that continuous data acquisition and light curve analysis is confined to WinControl-3.

WinControl-3 functions are arranged in 11 Windows (Table 9). The Chart window appears at software start. When more than one device is connected, the additional window "Moni-Bus" appears.

Table 9: Windows of WinControl-3				
	Window	Availability	Main panel	Content
1	Chart	Online and Offline	Graphics	Data versus time of all experiments
2	Induct. Curve	Online and Offline*	Graphics	Data versus time of fluorescence induction curves
3	Light Curve	Online and Offline*	Graphics	Data versus time of light response curves
4	SAT- Chart	Online and Offline*	Graphics	Saturation pulse kinetics
5	Spectrum	Online and Offline*	Graphics	Spectra
6	Report	Online and Offline	Alphanumerics	Saturation pulse data and data collected at the same time
7	Memory	Online only	Alphanumerics	Information on files stored on the DIVING-PAM-II flash memory
8	Batch	Online and Offline	Commands	Site for automatic execution of experiments
9	Control	Online only	Settings	Configuration of external light panel PAR sensor and stirrer control
10	Sensors	Online only	Settings	Control of PAR sensor, spectrometer, and oxygen sensor
11	Settings	Online only	Settings	Device settings
*Whe	*When data loaded.			

6.1 Chart Window

Fig. 66 divides the Chart window into six areas. Most of the side bars are also present when other windows are viewed.

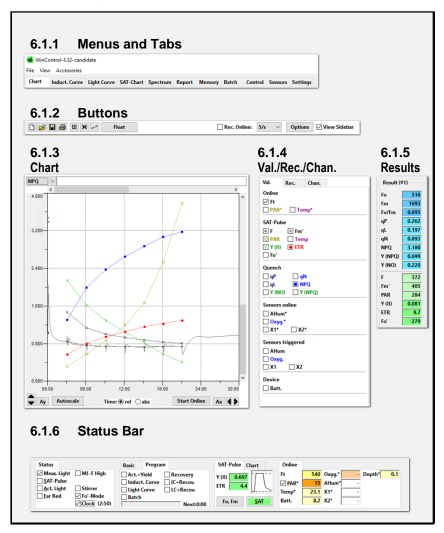


Fig. 66: Chart Window

6.1.1 Menus and Tabs

The top bar includes three menus (File, View, and Accessories) and all window tabs (compare Table 9, page 97). The three menus are explained in Table 10.

Table 10: Menu Overview		
Menu	Item	Comment
File	Load Data	Opens WinControl-3 files. Save current data first and switch off online recording of data.
	Save Data	Saves present data in WinControl-3 format.
	Save Settings	Saves all current instrument settings in a batch file. The settings can be restored by executing this batch file.
	Export Report	Exports the data displayed on Report window. For information on configuration of export data see Table 11 (page 100).
	Export Chart Record	Exports the data displayed on Chart.
	Quit	Exit WinControl-3.
View	Results Panel	Switches Results panel (Section 6.1.3) on or off.
	Status Panel	Switches Status panel (Section 6.1.6) on or off.
	Warnings	Prompts the display of the 'program starting time' and non-critical errors.
	Batch Win- dow	Switches Batch File window on or off.
Acces- sories	Temperature Units	Toggles between °Celsius and °Fahrenheit. The temperature unit affects only the numerical display on the Status bar (6.1.6, p.108).
	Plugins	Does not apply for the DIVING-PAM-II.
	Record File	Saves data continuously to reduce data loss in case of program failure. The command prompts for a folder in which the file should be saved. The file name is created automatically using data and time of record start: WinControl-Record-YYYY-MM-DD-hh_mm_ss.pam.

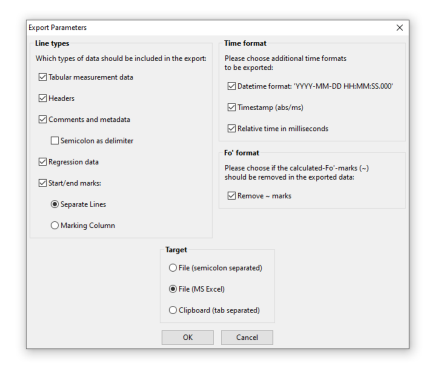


Fig. 67: Export Format for Reports

Table 11: Export Forma	at for Reports
Parameter	Action when checked
☑ Tabular measurement data	Exports data of saturation pulse analysis and all other data recorded at the same time.
☑ Headers	Exports column headers.
☑ Comments and metadata	Exports user and software comments (="metadata", e.g., device information).
☑ Semicolon as delimiter	Applies to comments and metadata. Information after a semicolon is written in a new cell.
☑ Regression data	Exports the cardinal numbers of light curves.

Table 11: Export Forma	t for Reports			
Parameter	Action when checked			
☑ Start/end marks				
Separate Lines	Writes start and end of an experiment in separate lines.			
O Marking Column	Writes start and end of an experiment in a separate column.			
☑ Datetime format	Exports date and time. Format: day/month/year hours:minutes:seconds.ms. In Excel, custom-formatting may be needed: dd/mm/yyyy hh:mm:ss.000.			
☑ Timestamp	Exports UTC (Coordinated Universal Time) time in ms. To convert into date/time use:			
	$=\frac{SourceCell + HoursTimeShift*3.6\cdot10^{6}}{86.4\cdot10^{6}} + 25569$			
	Format target cell, e.g.: dd/mm/yyyy hh:mm:ss.000			
☑ Relative time in ms	Export time of experiment in ms.			
☑ Remove "~" marks	Removes tilde (swung dash) signs. WinControl-3 marks calculated F´0 values by a tilde.			
File (semicolon separated)	Creates a file in which individual data are separated by semicolons.			
File (MS Excel)	Creates an Excel file. Note that columns "Date" and "Time" are differently formatted but contain the same information: dd/mm/yyyy hh:mm:ss.000			
O Clipboard (tab separated)	Copies the data to the clipboard. Data are separated by tab characters. Use the paste command to copy the data into a spread sheet program.			

6.1.2 Buttons

The elements of the buttons bar are introduced in Table 12 and Table 13.

Table 12: Button	IS	
lcon	Meaning	Comment
	Delete	Deletes all current data.
	Load	Same function as "Load Data" in the File menu (Section 6.1.1.).
	Save	Same function as "Save" in the File menu (Section 6.1.1.).
	Print	Prints current chart view.
	Grid on/off	Controls display of chart grid.
×	Lines	Connects data points with lines.
	Symbols	Controls the display of symbols.
Float	Add Chart	Creates an additional chart window with separate view settings.
☑ Rec. Online	Continuous recording	Controls continuous recording of fluorescence, PAR, and temperature.
5/s 🔻	Sampling frequency	Sets sampling frequency for online data. Open drop-down menu by the downward arrow. Available sampling frequencies are 5/s, 1/s, and 1/10s. 5/s is available only for fluorescence.
Options		See Table 13.
Or right-click on c	hart.	
☑ View Sidebar		Controls display of side bar (see 6.1.4, Val./Rec./Chan.)

Table 13: Options					
Menu Item	Present	Comment			
Export Record	Always	Exports the data displayed.			
Select cur- rent record	Always	Highlights data of a Record on the chart, and also on windows SAT-chart and Report.			
Split Selec- tion	After selection of interval.*	Put selection in separate record.			
Zoom to Se- lection	After selection of interval.*	Displays the interval enlarged.			
Export Se- lection	After selection of interval.*	Exports the data selected.			

^{*} How to select an interval on the chart: keep the left-hand mouse button pressed, move cursor across x-interval and release.

6.1.3 Chart

Fig. 68 outlines the tools to adjust the chart. Section 6.1.4 explains how to select data for display.

The scale of y- and x-axis can be manually changed: a vertical pair of arrows affects the y axis, and a horizontal pair of arrows adjusts the x-axis (see lower border of Fig. 68). Right next the two pairs of arrows are buttons for automatic axis scaling.

Manual and automatic y-axis scaling affects only the currently active y-axis unit. In contrast, the command Autoscale adjusts y-and x-axis so that all data fit in the chart area.

The y-axis unit can be picked from a drop-down menu located in the upper left corner of the chart. The active y-axis unit also determines the y-data of the cursor position on the chart.

A way to zoom in to a particular x-axis interval is to mark the x-interval of interest (see Fig. 68), open a context menu by right-click on the chart area, and select the command "Zoom to Selection".

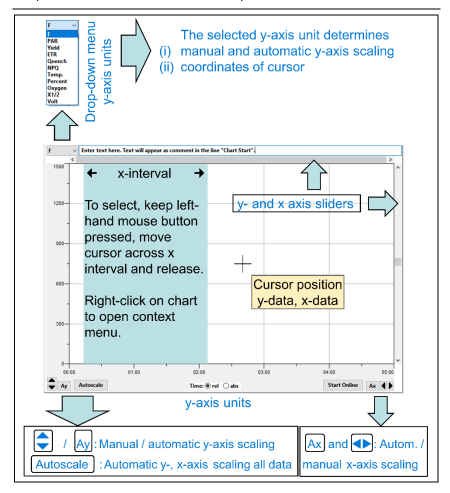


Fig. 68: Chart Tools

The selection of data on the chart results in highlighting the corresponding saturation pulse kinetics in the window SAT-Chart and the corresponding data lines in the window Report.

6.1.4 Val./Rec./Chan.

The field "Val. Rec. Chan." includes three different sidebars (Fig. 69). Data which should be displayed on the chart are selected on the Val. (= values) sidebar. The Val. sidebar distinguishes continuously recorded data (data groups: Online and Sensors Online) from saturation pulse data and data taken at the time of a saturation pulse (data groups SAT Pulse, Quench, and Sensors Triggered).

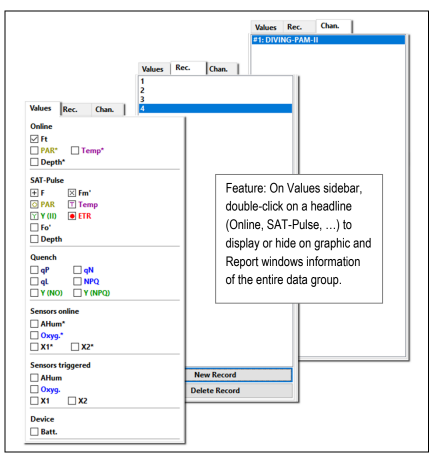


Fig. 69: Val. Sidebar

The Rec. (= Record) sidebar lists the number of individual charts of the present data file (= Report). Additional Records can be started by the button New Record and existing Records can be deleted by Delete Record. The latest Record, or the Record selected by mouse click, will be displayed on chart. The windows "Induction Curve" and "Light Curve" also possess the Rec. sidebar but in these cases the Induction Curve and Light Curve experiments, respectively, are listed.

The Chan. sidebar displays all devices detected by the WinControl-3 software. Only a DIVING-PAM-II was connected in Fig. 69.

On the Val. sidebar, the Ft represents a continuously recorded PAM fluorescence signal (online signal). The fluorescence level "F" is the corresponding saturation pulse signal (the F is the Ft measured right before the saturation pulse).

PAR* (photosynthetic active radiation, µmol photons m⁻²·s⁻¹) and Temp* (Temperature, °C) are two further online signal. They are distinguished by a superscript asterisk from their pendants which are measured together with saturation pulses. In the same way, data of additional sensors are marked (Fig. 69). The additional sensors available measure humidity (AHum) or oxygen concentration (Oxyg.). The sensor channels X1 and X2 are prepared for future use.

Besides F level fluorescence, the SAT-Pulse fluorescence levels are maximum fluorescence F_M ' (including F_M) and minimum fluorescence F_0 ' (F_0). Derived from these fluorescence levels is the photochemical quantum yield of photosystem II, Y(II) (including F_V/F_M). The electron transport rate (ETR, μ mol electrons/($m^2 \cdot s$) is calculated with Y(II) and PAR. The section "Quench" of the Val. sidebar includes six more saturation pulse parameters. All fluorescence parameters are explained in Chapter 10 (page 159).

6.1.5 Results

The Results sidebar (Table 14) shows numerical values of the current experiment. The upper part of the sidebar displays data of the Fo, Fm determination. These data are maintained throughout the current experiment. All other data are updated with each saturation pulse analysis. All fluorescence levels and ratios are explained in Chapter 10 (page 159).

Table 14:	Sidebar					
Parameter	Sample Data	Comment				
Fo	441					
Fm	1818	Fluorescence properties of the dark-acclimated sample.				
Fv/Fm	0.757					
qP	0.237					
qL	0.147					
qN	0.816	Fluorescence quotients describing the state of the light				
NPQ	2.212	acclimated sample.				
Y(NPQ)	0.616					
Y(NO)	0.278					
F	506	E				
Fm'	566	Fluorescence properties of the light-acclimated sample.				
PAR	285	Actinic light intensity, µmol m ⁻² s ⁻¹ .				
Y(II)	0.106	Fluorescence property of the light-acclimated sample.				
ETR	12.7	Relative electron transport rate, µmol m-2 s-1.				
Fo'	313	Fluorescence property of the light-acclimated sample.				

6.1.6 Status Bar

The "Status" field indicates the state of light sources and special functions (Table 15). The checkboxes function both as indicator and as on/off switches.

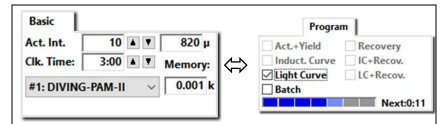
The "Basic" and the "Program" fields are stacked and can be selected by tabs. Actinic light and the clock frequency is adjusted on the Basic field, all automatic routines of the WinControl-3 software can be started in the Program field (Table 15).

The stacked "SAT-Pulse" and "Chart" field provide buttons to trigger F_V/F_M or Y(II) analyses, and graphic areas displaying fluorescence kinetics induced by saturation pulses. The "Online" field display live data numerically.

Table 15: Status Bar

Status ✓ Meas. Light	Meas. Light: Low frequency PAM measuring light.						
☐ SAT-Pulse ☐ Act. Light ☐ Stirrer ☐ Far Red ☐ Fo'-Mode ☐ Clock (0:00)	ML-F high: High frequency measuring light. Measuring light changes automatically to high frequency when actinic light is switched on.						
SAT-Pulse: Saturation pulse analysis to determine Y(II); equivalent to SAT button (see below).							
Act. Light: Actinic light to drive photos	ynthesis.						
Far-red: Light at wavelengths > 700 nr	n.						
Stirrer: Does not apply for the DIVING	-PAM-II.						
F_0 '-Mode: Automatically takes as F_0 ' fluorescence the minimum fluorescence in a period of far-red-light illumination following a saturation pulse.							
Clock: Repetitive trigger of the event s	pecified on						
Settings Window (Section 6.10, page	e 134). See below on how to adjust the inter-						

Table 15: Status Bar



Act. Int.: Setting and intensity of DIVING-PAM-II light source in μmol m⁻² s⁻¹.

Clk. Time: Time interval between automatically triggered events in minutes: seconds.

#1: DIVING-PAM-II: Device connected (channel).

Memory: Current file size in kilobytes (k).

Experimental Routines

Experimental routines are defined on the

Settings Window (Section 6.10, page 134). Most routines can be triggered by the clock.

Act.+Yield: Period of actinic illumination terminated by saturation pulse analysis.

Recovery: Dark phase with saturation pulse analyses performed at increasing intervals.

Induct. Curve: F₀, F_M determination followed by illumination by actinic light with repeated saturation pulse analysis.

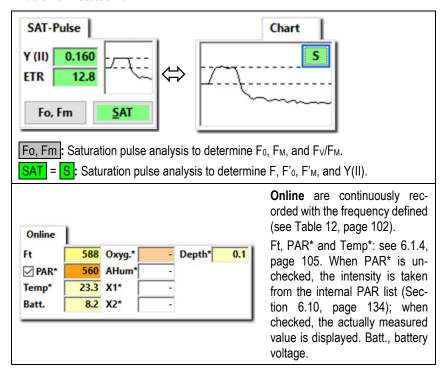
IC+Recov.: Induction curve plus dark phase with saturation pulse analyses performed at increasing intervals.

Light Curve: F_0 , F_M determination followed by illumination with stepwise increasing light intensities where each step is terminated by a saturatin pulse analysis.

LC+Recov.: Light Curve followed by a dark phase with saturation pulse analyses.

Batch: Execution of batch file program.

Table 15: Status Bar



6.2 Induct. Curve/Light Curve Windows

The icons and bars of the Chart window introduced above are also present in the Induction Curve and Light Curve windows. In contrast to the Chart window, which displays all data of a Record, the Induction Curve window displays individual induction curves, and the Light Curve window displays individual Light Curves. The same Record can include several Induction and Light Curves. Use up and down arrow keys to scroll through the list of curves.

Table 16 summarizes further differences between the three windows. The windows have different start buttons and only the Light

Curve windows possesses PAR as x-axis unit. The button Start IC starts an experiment in which a sample is exposed to a single defined light intensity, the button Start LC initiates a routine in which a sample is exposed to incrementally increasing light intensities. The x-axis radio button "Time" of the Light Curve window corresponds to the radio button "rel" (relative time) in the two other windows.

The Light Curve window allows fitting model functions to experimental data. The experimental data are the electron transport rates (ETR) plotted against the photon flux density, PAR. WinControl-3 provides two model functions called REG1 and REG2 which are introduced in Fig. 70 and Fig. 71, respectively. The function REG1 can decrease at high PAR values. Therefore, REG1 can consider photoinhibition of photosynthesis, where β is a photoinhibition parameter. In contrast, REG2 is a rectangular hyperbola which cannot describe photoinhibition.

Both models calculate the three cardinal parameters of a light curve:

- (i) α , electrons/photons: Initial slope of RLC which is related to the quantum efficiency of photosynthesis.
- (ii) ETR_m, μmol electrons m⁻²·s⁻¹: Maximum electron transport rate.
- (iii) I_K , μ mol photons $m^{-2} \cdot s^{-1}$: Idealized PAR value at which light-limited photosynthesis becomes light-limited.

The cardinal parameters are written into the Report. To export cardinal parameters separately, right click on the chart of the Light Curve window and select from the menu "Export Regression Data" (see Table 16). The item "Select current light curve" in the same menu highlights the data of the currently displayed light curve in the Chart and Report windows. A similar function is available for the window Induction Curve.

Table 16: Three Graphic Chart Window	Curve Win	dow			
Rec. Sidebar	Rec. Side	Rec. Sidebar			<u>r</u>
Val. Rec. Chan.	Val. Rec.	Chan.	Val.	Rec.	Chan.
1 2 3 Start Online	IC 1 IC 2 IC 3 	LC 1 LC 2 LC 3 			
<u>Lighting</u>	<u>Lightin</u>	<u>g</u>		Lighting	
PAR FREE Time	PAR	PAR			
Right click on chart Export Record Select current record	Right click o	Right click on chart Export Record Select current light curve Export Regression Data			
Val. Sidebar Quench qP qN qL NPQ Y (NO) Y (NPQ) Sensors online	Val. Side Quench qP qN qL NP Y (NO) Y (Sensors online	Val. Sidebar Quench qP qN qL NPQ Y (NO) Y (NPQ) Regression REG1 REG2 Sensors online			
x-axis units Time: ● rel ○ abs	x-axis uı Time: ⊚ rel	_	c-axis units me PA	ıR	

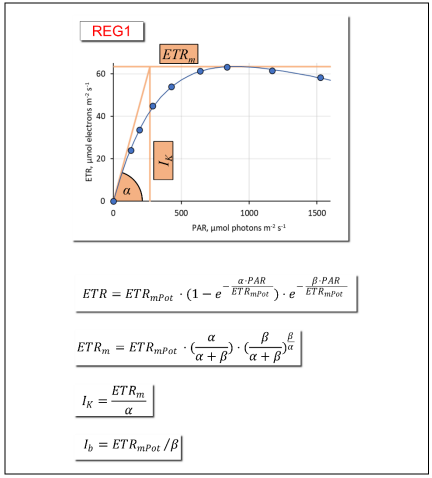


Fig. 70: Model Function REG1

The three cardinal points of the light curve are indicated (α , ETR_m , and I_K). The decrease at PAR>1000 µmol m⁻² s⁻¹ is frequently attributed to photoinhibition of photosystem II by strong light. I_b is the theoretical PAR at which the light curve reaches 1/e of ETR_{mPot}. ETR_{mPot} is the ETR_m in the absence of photoinhibition. According to: Platt T, Gallegos CL, Harrison WG (1980) Photoinhibition of photosynthesis in natural assemblages of marine phytoplankton. J Mar Res 38: 687-701

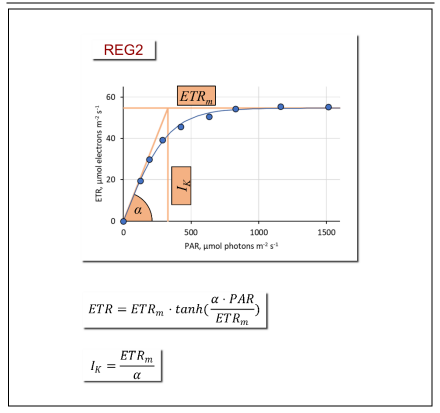


Fig. 71: Model Function REG2

The three cardinal points of the light curve are indicated (α , ETR_m , and I_K). According to Jassby AD, Platt T (1976) Mathematical formulation of the relationship between photosynthesis and light for phytoplankton. Limnol Oceanogr 21: 540-547.

6.3 SAT-Chart Window

The SAT-Chart window displays saturation pulse kinetics of fluorescence (Fig. 72). The automatically determined values of F and F_M ' are shown as dashed horizontal lines. Fluorescence traces are automatically scaled to fit into the coordinate system. The x-axis covers about 2 s, the saturation pulse starts at 200 ms, and the distance between data points is 50 ms. The mouse wheel scrolls through graphs.

Each graph is accompanied by a protocol panel. The protocol panel lists the pulse number (Nr) and the line number of the Report (Rep. Nr). Select a fluorescence trace by double-click in the protocol window. The selected graph automatically moves to the top of the window.

The Options menu (upper right corner of the SAT-Chart window) links the actual view of saturation pulse kinetics to the saturation pulse events selected in one of the three previous windows. The option "Follow Selection" automatically displays on top of the window the saturation pulse kinetics of marked events. When Follow Selection" is inactive, the option "Jump to Selection" brings the selected kinetics into view.

The Val. sidebar is similarly configured as described in Section 6.1.4 (page 105), except "Online Data" are omitted and the electron transport factor (ETR-F.) is added. The ETR-F. is the fraction of absorbed photons by the sample relative the PAR to which the sample is exposed. The ETR-F. is a factor of the equation for electron transport (Chapter 10, page 159). Each item selected on the Val. sidebar is numerically displayed below the corresponding saturation pulse kinetics.

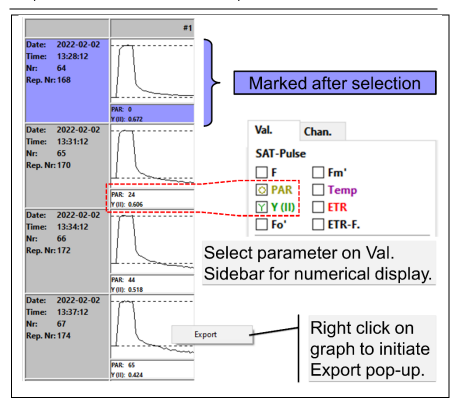


Fig. 72: SAT-Chart Window

Table 17: SAT Kinetics in Raw File

To extract a series of saturation pulse kinetics, open PAM file with spread sheet program, sort by SG and copy data right of SG.

	Α	В	С	D	 AD	AE	AF	AG	AH
1	Date	Datetime	Type	No.					
2	08/03/2022	08/03/2022 09:17:42.997	F	2	 SG	50	419	419	
3	08/03/2022	08/03/2022 09:17:44.389	F	3	 SG	50	383	383	
					SAT Graph Label	Time interval in ms	Satu	Data of ration pations	ulse

Single saturation pulse kinetics can be exported by right-click in its chart field. To export a series of saturation pulse kinetics, open the PAM file with a spread sheet program. In the text file, saturation pulse kinetics are preceded by an SG (Table 17). Sort the lines of the file by SG. Create a time scale by considering the neighboring points are spaced apart by 50 ms.

6.4 Spectrum Window

The window is active in the presence of spectral data. With the cursor placed inside the drawing area, or below the x-axis, the mouse wheel has a zoom function for the x-axis. When the cursor is placed left of the y-axis, the mouse wheel has a zoom function for the y-axis.

Right click on the drawing area to open a menu containing zoom and export commands (Fig. 73). The "Zoom to selection" is available after selection of an x-axis interval described before (Fig. 68).

An application menu is available in the top right corner of the window (Fig. 73). To change between the three principal applications (PAR, fluorescence, reflectance), the spectrometer MINI-SPEC/MP has to be reconfigured (see Fig. 9, page 21).

The dark current of the spectrometer at room temperature is measured in the factory and stored on the flash memory of the device. To newly establish the dark current, fully darken the entrance optics (cv. Fig. 9, page 21) and press Calibr. Dark.

The standard PAR configuration is "PAR cosine". This configuration employs a diffusing disk as light entrance (Fig. 9). The configuration shows an approximate cosine response toward incoming radiation. The spectrometer calibration differs between measurements in air and in water. Therefore, choose PAR cosine air or

PAR cosine water for proper measurements under the two conditions.

"PAR sphere" employs a spherical light entrance. The sphere configuration measures light from all directions with similar weight. At the time of writing of this manuscript, the spherical sensor is under development.

The setting "PAR Open sensor" is used by service engineers.

"Fluorescence (blue)" and "Fluorescence (red)" measures fluorescence emission spectra excited by blue or red light, respectively. Measuring "Reflectance" requires that the 100% reflection signal has been established with the highly reflective reference material provided with the spectrometer (Fig. 8, page 20).

The sidebar also display the Integration time (ms) which is the integrated measuring time for a single spectrum. Auto Range optimizes the integration time for a spectrum depending on incoming light. Auto Range ON is the default setting. Auto Range OFF is for special applications. When spectra look unrealistic, switch on Auto Range. Average determines the number of spectra averaged to yield the final spectrum. SAT+Spec measures a spectrum for saturation pulse analysis.

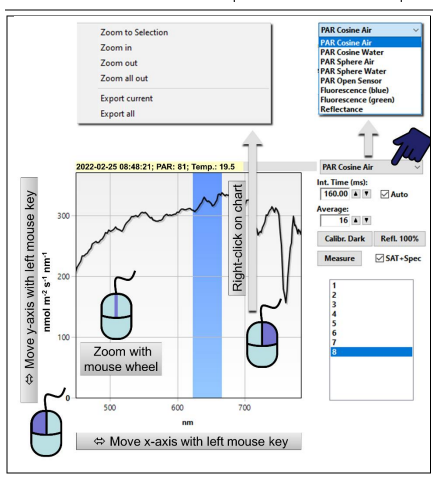


Fig. 73: Spectrum Window

6.5 Report Window

The Report window lists all data associated with saturation pulse analyses. Continuously recorded signals, saturation pulse kinetics and spectra are not listed. The fact that a spectrum was recorded is indicated by "SPEC" in the column "Type" (Table 18). The Report window also displays start and end of induction and light curves, as well as the cardinal points of light curves (Table 18).

The types of data displayed is controlled by the Val. side bar, which is configured as described for the SAT-CHART window (Section 6.3, page 115). Only data displayed on the Report table are exported.

Table 18: Report Table

Type: D, device. FO, F_0 and F_M determination. F, determination of F, F_0 ', and F_M '. SPEC, spectrum. SICWS and SICSE, induction curve start and end, respectively. SLCS and SLCE, light curve start and end, respectively. REG1 and REGS, cardinal parameters of regression analysis 1 and 2, respectively (see 6.2). **Mark**: Letters assigned to data (see 4.1.1, page 38). Hide Mark column via option menu. Here, the DIVING-PAM-II has the channel number #2. "#2" is preceding the cardinal values of light curve analysis, REG1 and REG2.

	Date	Time	Туре	No.	2:Mark	2:F	2:Fm	' 2	2:PAR	2:Y (II)	2:ETR
1			D			Device	Nr: #2,	DIVIN	IG-PAM-	II (UWFI	D0131A)
2	2022-03-08	09:07:11	SCHS								
3	2022-03-08	09:16:20	FO	1	Α	461	193	2	6	0.761	1.9
4	2022-03-08	09:16:50	F	2	Α	680	77	4	6	0.121	0.3
5	2022-03-08	09:17:24	SPEC		PAR: 12; Temp.: 21.2						
6	2022-03-08	09:17:42	SICS		Induction Curve start						
7	2022-03-08	09:17:48	SICE		Induction Curve end						
8	2022-03-08	09:17:59	SLCS		Light Curve start						
9	2022-03-08	09:18:07	REG1		#2: alpha: 0.362, ETRm: 13.99, lk: 38.634 (beta: 0.001, ETRmPot: 14.179)						
10	2022-03-08	09:18:07	REG2		#2: alpha: 0.272, ETRm: 13.665, lk: 50.280						
11	2022-03-08	09:18:05	SLCE		Light Cur	ve end					

The options menu of the Report window can be opened by clicking the Options button or by right-click in the Report field. The options menu contains various tools for handling the Report table. All menu items and their function are summarized in Table 19.

Command "Insert Settings" of the options menu writes the current device settings in the report. The abbreviations used for the various settings are compiled in Table 20.

Table 40. Ontions Many	
Table 19: Options Menu	
Options Menu Item	Comment
	General Report Management
Follow Selection	Automatically displays the data associated with events marked in other windows
Show Mark	Display marker letters (see "Mark" in Table 18)
Insert Settings	Writes settings of DIVING-PAM-II in Report table (see Table 20)
	Manipulation of All Data
Export All	For details see Section 6.1.1, page 99.
Delete All Measure Data	Self-explaining
Page Setup for Printing	Basic configuration of print layout. Checks if output fits on page
Preview Printing	Self-explaining
Print Report	Selects and configures printer, prints current Report
	Manipulation of Selected Data
	Requires that saturation pulse events on a chart or lines of the Report have been selected. To select, move the cursor with left mouse key pressed over events
Export Selected Lines	Confines export to lines of interest
Jump to Selection	Brings data into view, when "Follow Selection" is off

Table 19: Options Menu	
Options Menu Item	Comment
Delete Selected Data	As above. Affects only selected lines.
Preview Print Selection	As above. Affects only selected lines.
Print Selected Data	As above. Affects only selected lines.
Mark as Light Curve	Combines a series of saturation pulse analysis into a light curve and performs regression analysis REG1 and REG2. The series must not contain start or end marks of previous light or induction curves.

Table 20: Abbrevia	ations for Settings
MEA	Measuring Light Intensity
MI	Measuring Light Frequency
DI	Damping
GA	Gain
EF	ETR-Factor
FZ	F Offset
CW	Clock Interval
SI	Saturation Pulse Intensity
SW	Saturation Pulse Width
Al	Actinic Light Intensity
AF	Actinic factor
FRI	Far-red Intensity
FRW	Far-red Width
AW	Length of exposure of "Act. + Yield" routine
ICD	Delay of Induction Curve
ICW	Interval Between Saturation Pulses of Induction Curve
ICL	Length of Induction Curve
LCW	Step Length of Light Curve
LCI	Initial Intensity Setting of Light Curve
LCL	Length of Light Curve
LO	Offset of External PAR Sensor
LG	Calibration Factor of External PAR Sensor
LO2	Offset of External PAR Sensor 2
LG2	Calibration Factor of External PAR Sensor 2
TO	Offset of External Temperature Sensor
TG	Gain of External Temperature Sensor
ILO	Offset of Internal PAR Sensor

Table 20: Abbreviations for Settings					
ILG	Gain of Internal PAR Sensor				
PARGAIN_RED	Calibration Factor for Red LED of 2054-L				
PARGAIN_GREEN	Calibration Factor for Green LED of 2054-L				
PARGAIN_BLUE	Calibration Factor for Blue LED of 2054-L				
PARGAIN_WHITE	Calibration Factor for White LED of 2054-L				
PARGAIN_CUSTOM	Calibration Factor for Other light Sources				
ITO	Offset of Internal Temperature Sensor				
ITG	Gain of Internal Temperature Sensor				
MLC	Fm Correction Factor 1				
MLC2	Fm Correction Factor 2				
TRM	Trim Value for Measuring Light				
TRA	Trim Value for Actinic Light				
TRSA	Trim Value for Saturation Pulse				
TRFR	Trim Values for Far-red Light				

6.6 Memory Window

The Memory window accesses the data of the internal memory of the DIVING-PAM-II. The DIVING-PAM-II memory is a circular buffer which can store 512 files or 30 000 data entries maximally. The memory is full when one of the two limits is reached. For instance, if each file contains only two data entries, the full memory contains 512 files but only 1024 data entries, or, if each file contains 1000 data entries, then only 30 files fit into the memory.

When the memory is full, the oldest data are overwritten with the latest ones. Usually, data acquired in the offline mode are downloaded (6.6.2) and the memory is cleared (0) which avoids that all memory is used.

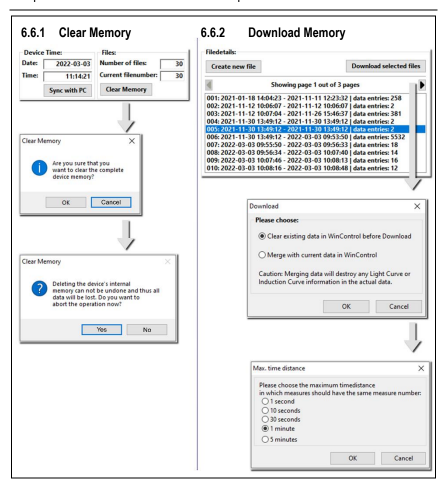


Fig. 74: Memory Window

6.6.1 Clear Memory

Clearing the memory is irreversible. To avoid inadvertent deletion of data, two warnings must be passed before the command Clear Memory is executed.

Above the Clear Memory button, memory information is given: "Number of Files" is the file count, "Current file number" is the serial number of the file to which data are currently written. The Current file number can be greater than the number of files when the memory is full and old files are overwritten by new ones.

The Clear Memory field display date and time of the internal computer of the DIVING-PAM-II. Click Sync with PC to apply the computer time settings to the DIVING-PAM-II.

6.6.2 Download Memory

A list of files is displayed in the Download field of the Memory window (Fig. 74). You can download individual or groups of files. To pick several files, hold down Ctrl key and click on files of interest. To pick consecutive files, hold down Shift key and click on first and last files of the row. Then click Download selected files.

Usually, data are written in an empty Report window. Selecting "Clear existing data ..." in the download dialogue box, empties the Report window.

When parallel measurements of two different fluorometers should be combined, choose "Merge with current data ...". The newly imported data will be sorted so that time points of import data matches time points of existing data. Two time points are considered as matching when their time difference is smaller as the interval selected from the box "Max. time distance" (cf. Fig. 74). The merge operation deletes the report entries for begin and of an induction or a light curve.

6.7 Batch Window

Batch files automatically execute experimental procedures. To activate the Batch window, click icon new batch or open existing batch file (Fig. 75, "Batch Start Buttons"). The click Edit and choose between "Add command" and "Record Macro". The command "Update indentation" is an automatic editing tool to improve readability.

6.7.1 Add Command

Add command leads to five groups of commands (Fig. 75). A command can be inserted into a batch file by left click.

Measure commands

Saturation pulse Saturation pulse analysis of light-ex-

posed sample

 F_0 , F_M determination Saturation pulse analysis of dark accli-

mated sample

Light Sources commands

Controls for actinic and measuring light. The light ON command asks for the light intensity: simple enter the light intensity setting in the grey-shaded field. New intensity settings can be chosen using the commend "Change intensity".

Settings commands

"Reset settings to default" installs the settings described in Table 7 (page 85).

Repetition commands

The group contains two commands. "Repeat specific number of times" and "Repeat indefinitely". Both commands write two lines on the batch file sheet. In case of the first command, these lines are:

Line 1:

for \$loopvar = 1 to count // Start repetition block with specified number of repetitions

Line 2:

next // End of repetition block

Write commands to be repeated between Line 1 and 2. Define how often the commands should be repeated by the number entered in field count.

The second command (Repeat indefinitely) repeats the commands placed between the two lines until the batch program is stopped manually.

Line 1:

while 1 // Start block repeating indefinitely

Line 2:

wend // End of repetition block

Timing

The command "Delay" inserts a time interval after the previous command has been terminated. The next command is executed when the time interval ends.

The command "Continue Delay" takes the end of the previous delay phase as starting point. Actions within this time interval are performed without affecting the interval defined for Continue Delay. A continuous time scale can be built by a series of Continue Delay commands.

"Wait until time of day" delays start of the batch program until the time specified.

6.7.2 Record Macro

The function "Record Macro" converts your manually entered commands into batch file lines. Simply click "Record Macro", perform experiment, and click "Record Macro" again.

6.7.3 Options

All items of the options menu of the Batch window are related to the MONI-DA data acquisition system.

Check MONI-DA Compatibility

Some batch file commands cannot be executed by the MONI-DA. This command searches for such incompatibilities.

Upload to Device

Transfers a batch file to the MONI-DA memory.

Export

Export the current batch file as "WinControl-3 Compiled Batch File" (*.wccb). This file format is for future use. Upcoming MONI-DA versions will be enabled to directly download wccb files, that is, without being connected to WinControl-3. The final goal is to install batch files by remote control.

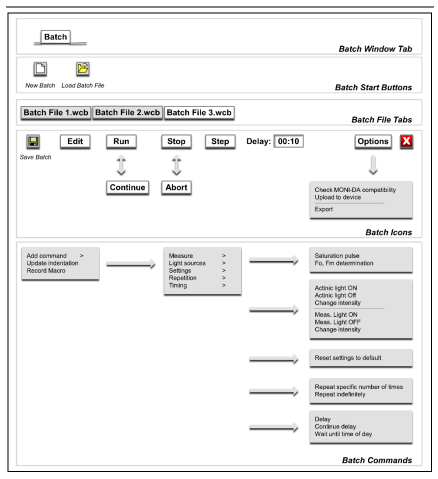


Fig. 75: Batch Window Overview

6.8 Control Window

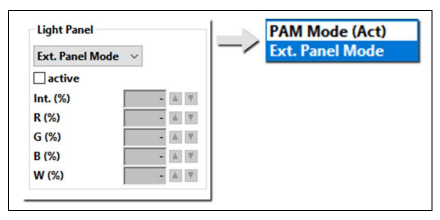


Fig. 76: Control Window

Fig. 76 shows the section of the Control Window which applies to the DIVING-PAM-II fluorometer. The drop-down menu on top of the Light Panel area consists of two points: "PAM Mode (Act.)" and "Ext. Panel Mode". Ext. Panel Mode is the standard mode in which the fluorometer's internal light source is used as actinic light to which an external light source may be added (see Table 21). Activating PAM Mode (Act.) selects the external light source as the sole actinic light.

With the External LED Light Source 2054-L is connected, the meaning of numerical inputs (Int. (%) to W (%) varies depending on the mode chosen (see Table 21). Typically, the maximum intensity of all four LED groups is 5000 µmol m⁻² s⁻¹ or higher. Operation of the 2054-L External LED Source requires line power.

Table 21: External Light Source		
	Ext. Panel Mode	PAM Mode (Act.)
Internal light source of DIVING-PAM-II		
Function	Main actinic source	Not available
Intensity setting	Actinic light intensity (Settings window)	Not available
External LED Light Source 2054-L		
Requirement	Not mandatory	Required
Function	Background illumination	Main actinic source
Intensity setting	Int %	Actinic light intensity (Settings window)
Control		
Int. (%)	Relative intensity (0-100%). The number multiplies the intensity of all light sources of the External LED Light Source 2054-L	Not active
R (%), G (%), B(%), W (%)	Absolute intensity in percent (is multiplied by Int. (%))	Factor defining the intensity with which a light source contributes to total intensity.

6.9 Sensors Window

External PAR Sensor

External PAR sensors are designed to measure the intensity of external light at sample level. External PAR sensors are employed to estimate electron transport rates driven by external artificial or natural light. In comparison, the internal PAR sensor measures the light intensity of internal light at sample level. Internal actinic light is used for short-term illumination (Act. + Yield), or when induction and light curve experiments are carried out.

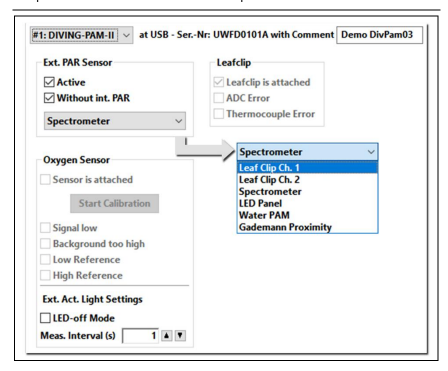


Fig. 77: Sensors Window

The checkbox "Active" selects sensing of external PAR. Checking "Without int. PAR" ignores measurements by the internal PAR sensor. The drop-down menu below these checkboxes lists external PAR sensors.

Drop-Down Menu

"Leaf Clip Channel 1" corresponds to the micro quantum sensor of the 2035-B clip (MINI-PAM-II accessory). Selectable when connected via special adapter cable (operation only under dry lab conditions). "Leaf Clip Channel 2" is the signal of a PAR sensor connected to the side port of the 2035-B clip. "Spectrometer" uses PAR spectra integrated over the visible range as PAR value. "LED panel" is the PAR sensor connected to the LED-Panel RGBW-L084. (Does not apply for External LED Light Source 2054-L).

"WATER-PAM" is available when working with this PAM fluorometer. "Gademann Proximity" is a sensor under development.

Leaf Clip

The checkbox of "Leafclip is attached" automatically indicates that a 2035-B clip is connected to the DIVING-PAM-II (see comment above). Erroneous conversion of the analogue measuring signal to digital information is signaled by "ADC Error". A defective thermocouple activates the checkbox of "Thermocouple error".

Oxygen sensor

This section, including "External Actinic Light Settings" is dedicated to the accessory "Underwater Oxygen Sensor DIVING-PAM-II/O2". For details, see separate manual:

https://www.walz.com/products/chl_p700/diving-pam-II/downloads.html

6.10 Settings Window

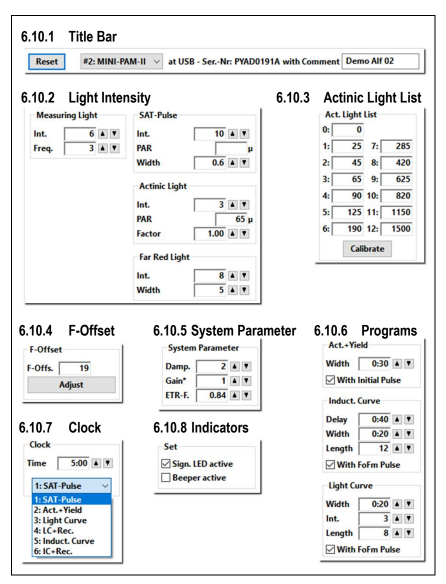


Fig. 78: Settings Window

6.10.1 Title Bar

The Reset button restores factory values for all settings of the current window. Located right of the Reset button is a drop-down menu listing all devices connected. The settings displayed on the current window belong to the device selected, and also the serial number which is displayed right of the drop-down menu. In the text box right of the serial number, up to 20 characters can be entered. Serial number and text box content are written in the first line of each record file.

6.10.2 Light Intensity

When actinic light is switched off, µs measuring pulses are delivered at 5 to 25 Hz depending on frequency setting "Freq." (Table 6, page 59). Switching on actinic light automatically increases the measuring light frequency to 100 Hz. The measuring light frequency can also be manually set to 100 Hz by checking "MF-F High" on the Status Bar (Table 15, page 108).

The intensity setting "Int." adjusts measuring light intensity. Twelve intensity levels are available. The numerical value of the setting is proportional to the measuring light intensity. How to estimate the integrated measuring light intensity for the frequency and intensity settings used is described in Section 4.2.1 (page 57).

Note that an increase in flash intensity increases the F_0 fluorescence level (the fluorescence intensity, in general is, proportional to the flash intensity). An increase in the number of flashes/pulse increases the F_0 level only if some PS II reaction centers become closed by the increased integrated measuring light intensity (the measuring light becomes actinic). If this is the case, the integrated

measuring light intensity must be reduced either by reducing intensity or by reducing frequency.

Like measuring light, the intensity of saturation pulses, actinic light and far-red light can be adjusted in 12 levels. The PAR of saturation pulses is indicated in newer versions of WinControl-3. The PAR of actinic light is taken from the light list (see below). Most of far-red emission is not absorbed by chlorophylls. Therefore, intensity data for far-red radiation (PAR or photon flux density) is not given as it would strongly overestimate its actual actinic effect.

6.10.3 Actinic Light List

Twelve actinic light levels are listed. These values are target PAR values. As delivered from the factory, the light emission of the LED is adjusted so that these target values are present at the sample level of a 2035-B Leaf Clip Holder. The target values can be varied by changing the "Factor" in the field "Actinic Light" (Section 6.10.2).

Calibrate newly adjusts the emission of the LED. Calibration must be performed when the sample level is closer or further away than the sample level (7 – 7.5 mm, see Section 3.1.4, page 19). Calibration is also necessary when the DIVING-F1 Miniature Fiberoptics is used. (See Section 3.2.4.1, page 34.)

6.10.4 F-Offset

Correction for background fluorescence. See Section 4.2.1 (page 57).

6.10.5 System Parameter

<u>Damping (Damp.)</u> Function for data smoothing. The smoothing effect increases with increasing numerical value, at the same time, but instrument response decreases See Section 4.2.1 (page 57).

<u>Gain Factor (Gain).</u> The Gain factor amplifies signal and noise. At low signal levels, increasing the gain improves digital resolution. See Section 4.2.1 (page 57).

<u>ETR-Factor (ETR-F.).</u> The ETR-Factor is the percentage of light absorbed by the sample and is a parameter for calculating the electron transport rate (Section 10.3, page 166).

6.10.6 Programs

Act. + Yield: Light exposure followed by saturation pulse analysis

"Width" is the duration of light exposure. The actinic light intensity adjusted in the field "Actinic Light" applies (Section 6.10.2, page 135). A saturation pulse analysis is always carried out at end of illumination. Checking "With Initial Pulse" places an additional saturation pulse analyses before illumination. The fluorescence levels of the initial saturation pulse analysis are defined as F_0 and F_M . All fluorescence ratio parameters requiring these two fluorescence levels (Table 27, page 168) are only available with initial saturation pulse.

Induct. Curve: Fluorescence curve following the onset of light

Three parameters determine the sequence of events. "Delay" indicates the dark time interval between F_0 and F_M determination and onset of light. The delay time can be adjusted between 5 seconds and 10 minutes. Default value is 40 seconds. "Width" is the

time interval between neighboring saturation pulses in the light period. Length is the number of saturation pulse analyses carried out. Time interval of actinic illumination is approximately "Width" times "Length". A dark period can be appended to an induction experiment (Table 15, page 108 and Table 5, page 52).

<u>Light Curve</u>: <u>Saturation pulse analyses after exposure to different</u> light intensities

The "Light Curve" program exposes a sample to consecutively increasing actinic light intensities. Three input values define the procedure. "Width" is the duration of exposure to a particular light intensity. "Int." is the first light level of the light curve. "Length" is the number of light steps. The duration of a light curve is determined by "Width" times "Length". A dark period can be appended to illumination (see above).

 \square With FoFm pulse: Both Induction and Light Curves can be performed without initial saturation pulse (F₀ and F_M measurement).

6.10.7 Clock

The clock utility repetitively triggers one out of 6 different measuring routines: Saturation pulse analysis, Act.+Yield, Induct. Curve, IC+Recov., Light Curve and LC+Recov (see Above). The interval time between two events (Clk. Time) can be adjusted between 10 s and 50 min. The time interval must be longer than the time required by the triggered event.

6.10.8 Indicators

The signal LED located above the touchscreen indicates the operational state of the DIVING-PAM-II as listed in Table 22. The

beeper acoustically marks begin and end of a saturation pulse analysis.

Table 22: LED Signal Code						
LED Color	Frequency	Process				
Measuring	m o d e					
Green	1 Hz	Normal operation				
Green double flash	1 Hz	Normal operation and clock running				
Green	continuous	Normal operation and saturation flash				
Additional codes during firmware update						
Green/red alternating	high	Waiting for software				
Red	continuous	Update running				

System information is displayed on the bottom of the Settings and the System Settings windows (see Table 23 below)

Table 23: System Information

System information displayed on bottom of Systems and Systems Settings window.

Headline	Model	Model Number	Serial Number
Display	DIVING-PAM-II (blue) or	DIVING-PAM-II/B or	UWFC####(A) or
	DIVING-PAM-II (red)	DIVING-PAM-II/R	UWFD####(A)
Information	Color version	Order code	S/N convention

6.11 System Settings Window

Different from all other windows of WinControl-3, the window System Settings is not represented in the row of tabs. To access system settings, open Settings window and click "System Settings" (Fig. 80). The functional elements of System Settings are shown in Fig. 79 together with their section numbers.

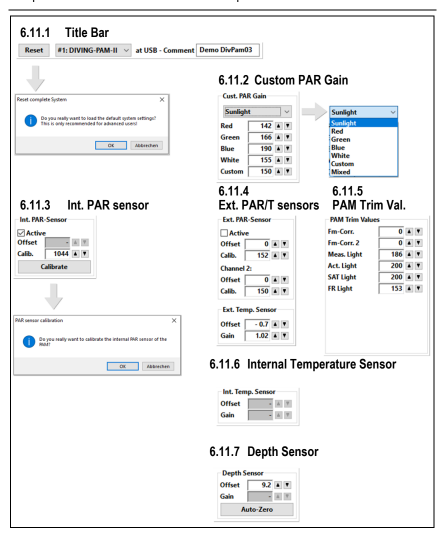


Fig. 79: System Settings

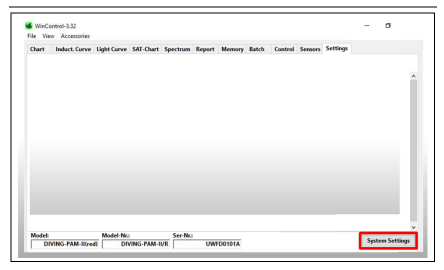


Fig. 80: System Settings Button

Click to open System Settings window.

6.11.1 Title Bar

The Reset button of the System Settings page recalls all calibration data established at the factory. This information is taken from the flash memories of the DIVING-PAM-II and of the external sensor (e.g., the 2035-B Leaf-Clip Holder).

The original calibration data cannot be changed by the user. The command Reset on the System Settings window is equivalent to the command "Load System Settings" of the DIVING-PAM-II menu (Section 4.2.4.7, page 81). The command Reset on the Settings windows (Section 6.10.1, page 135) is equivalent to the command "Load default" of the DIVING-PAM-II menu (Section (Fig. 60, page 84)).

The other elements of the Title Bar are described in Section 6.10.1, page 135.

Table 24: Custom PAR Gain

The table compiles the types of gain factors with their application, which are available for the PAR sensor of the 2035-B leaf clip or the PAR sensor of the 2065-M device.

Gain factor	Application		
Sunlight	Outdoor natural light. Factor displayed in "Ext. PAR Sen-		
	sor/Calib.".		
Red, green,	Red, green, blue, white LED light from MINI-PAM-II internal light		
blue, white	source or from 2054-L external light source.		
Mixed	2054-L external light source. The active factor is dynamically ad-		
	justed depending on the color mixture selected.		
Custom	User-defined factor.		

6.11.2 Custom PAR Gain

Different calibration factors for different light sources are provided. The calibration factor varies because of slight spectral variations in sensitivity of the PAR sensor. Table 24 summarizes the factors available.

6.11.3 Int. PAR sensor

The internal PAR sensor received a small fraction of the actinic light emitted by the internal LED of the MINI-PAM-II. The calibration factor of the internal PAR sensor is displayed in the "Calib." numerical field (Fig. 79).

The calibration of the internal PAR sensor is done by comparison with a calibrated external PAR sensor. The reading of the external PAR sensor depends on its position relative to the end of the optical fiber of the DIVING-PAM-II. Factory calibration was established with a 2035-B leaf clip in which the fiber was fully inserted.

For other configurations, the internal sensor must be newly calibrated. (See also Section 6.11.3, page 142.)

PAR sensor compatible with DIVING-PAM-II:

To calibrate the internal PAR sensor using the Miniature Spectrometer MINI-SPEC, proceed as described in Section 3.1.4, page 19, and click Calibrate.

To calibrate the internal PAR sensor using the PAR sensor of the 2035-B clip (MINI-PAM-II accessory, requires adapter cable, operation only under dry conditions), insert fiber into clip, position light sensor relative to fiber identically to the experimental situation (with the light sensor in the sample plane) and click Calibrate.

Other PAR sensor: To calibrate the internal PAR sensor using an external PAR sensor which cannot be read by the DIVING-PAM-II, proceed as described above but adjust calibration factor manually until the readout of the internal sensor matches that of the calibrated external sensor.

6.11.4 Ext. PAR/T sensors

The boxes "Ext. PAR Sensor" and "Ext. Temp. Sensor" (Fig. 79) display the factory-established calibration factors of the 2030-B clip or the 2065-M Mini Quantum/Temp.-Sensor, depending on which device is connected. In box "Ext. PAR Sensor", the "Channel 2" calibration applies to a second PAR sensor connected to the 2035-B leaf clip. Calibration factors must be manually entered when a PAR sensor does not support automatic installation of its calibration data.

6.11.5 PAM Trim Val.

Trim values have been set at the factory so that the DIVING-PAM-II meets its specifications.

Table 25: PAM Trim Values

The table compiles the types of gain factors with their application, which are available for the PAR sensor of the 2035-B leaf clip or the PAR sensor of the 2065-M device.

Туре	Application		
Fm-Corr.	Factor to compensate a signal decrease during a saturation pulse. The factor is determined with a Walz fluo-		
	rescence standard foil. Signal compensation is obso-		
	lete (value=0) for the latest generation of LEDs.		
Fm-Corr.2	Second factor to compensate a signal decrease during		
	a saturation pulse. Two factors are required when a bi-		
	phasic signal decrease occurs. Signal compensation is		
	obsolete (value=0) for the latest generation of LEDs.		
Meas. Light	Factors adjusting the intensities of measuring light/ac-		
Act. Light	tinic light/saturation pulse light/far-red light to meet the		
SAT Light	respective specification.		
FR Light			

6.11.6 Internal Temperature Sensor

Does not apply to the DIVING-PAM-II device.

6.11.7 Depth Sensor

Offset is adjusted in the factory under the prevailing conditions. Adjust offset using Auto-Zero if the readout above the waterline deviates from zero. Gain does not apply for the digital and temperature-compensated sensor of the DIVING-PAM-II.

7 Hints

7.1 Instrument Settings

Instrument settings are adjusted at the factory for optimum performance of the DIVING-PAM-II. For example, LED currents have been adjusted to meet target PAR values for the 60° measuring setup in the Universal Sample Holder DIVING-II-USH. For a different geometry, the internal PAR sensor requires recalibration to correctly measure internal actinic light (see Fig. 47, page 72). Estimation of PAR according to Fig. 14 (page 25) is rather inaccurate.

Also, the fluorescence offset (F-Offset) of your system has been measured and was saved on the DIVING-PAM-II memory. That means that the DIVING-PAM-II should show a fluorescence value close to zero in the absence of a sample. If this signal deviates clearly from zero, newly adjust F-Offset (see Fig. 36, page 57 ff).

7.2 Default settings

For fluorescence measurements with many macro algae and green leaves, default settings for measuring light and saturation pulses are well suited. Some samples require special settings. The following sections will provide some hints to adjust settings properly.

7.3 F₀ Fluorescence

Usually, measuring light intensity is adjusted to reach F_0 fluorescence levels around 500 mV (for a definition of F_0 see Section 10.2, page 162). Theoretically, the F_0 should stay below 640 mV.

Chapter 7 Hints

The latter upper value is derived from the assumption that the maximum F_V/F_M of any sample is 0.84 und from the fact that signal saturation occurs at 4000 mV (see equation below, Table 26). If required, measuring light can be adjusted (Fig. 36, page 57 or Section 6.10.2, page 135).

Table 26: Maximum F₀ of a Dark-acclimated Leaf

At low signal levels, signal height can be increased by increasing measuring light intensity. At too high intensities, the measuring light might drive photosynthesis to some degree. Therefore, test if switching on measuring light results in a stable signal or if any signal increase occurs. In the latter case, average measuring light intensity must be decreased either by reducing the amplitude of measuring light (measuring light intensity) or by reducing measuring light frequency or both.

7.4 F_M Fluorescence

The F_M and F_M ' levels are determined as the maximum of the fluorescence signal induced by a saturation pulse. Factory settings of saturation pulse width and intensity are adjusted to arrive at a plateau with many macro algae and green leaves (Fig. 81A). Some high light grown samples do not reach a plateau with standards settings (Fig. 81B). In this case, saturation pulse intensity or/and length should be increased. Also, fluorescence kinetics can

reach its maximum clearly before end of the saturation pulse (Fig. 81C). The latter does not result in erroneous F_M or F_M ' values because these values correspond to the maximum of fluorescence kinetics. In case of Fig. 81C, saturation pulse intensity or/and length might be decreased.

Some samples, particularly low light grown or senescing plants, exhibit with standard settings somewhat decreased F_V/F_M values but show normal fluorescence kinetics. These samples increase the F_V/F_M with decreasing saturation pulse intensity. Therefore, testing the F_V/F_M at saturation pulse intensities also below and above standard settings is important to optimize your saturation pulse settings.

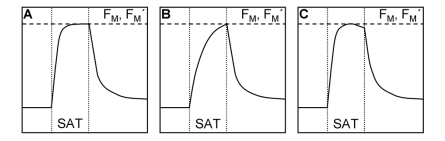


Fig. 81: Fluorescence Kinetics Induced by a Saturation Pulse

8 Trouble Shooting

8.1 Device does not switch off

Check if clock is running.

Check if device is connected to computer via USB or WLAN.

8.2 Device is not charged

Make sure that interface cable is connected to INPUT (there are several 6 pin connectors matching the interface cable, Fig. 3, page 12).

Make sure that the interface in connected to line power (Fig. 2, page 10).

8.3 "Overflow" message in window primary data

Reduce measuring light intensity and signal amplification (see "PAM Settings", Fig. 36, page 57)

8.4 Fluorescence Signal Noise

Check if fluctuating light sources (fluorescent tubes, computer screens) affect fluorescence.

Exclude that automatic scaling of Y axis after very low Y(II) has extremely amplified the fluorescence signal.

9 Maintenance

9.1 Cleaning

Depending on experimental site, the DIVING-PAM-II is exposed to various degrees of dirt. In marine environments, salt crystals may deposit on the fluorometer. To retain normal function, regular clearing is recommended. Suggested measures are:

- Shower off dirt and saltwater. Let air dry. Avoid scratches and do not wipe the surface unless all particles have been removed.
- Remove fiberoptics. Rinse fiberoptics and fiberoptics port with freshwater. Let air dry. Do not mechanically clean the fiberoptics port as this might scratch the fluorometer's optical window.
- Unscrew pushbuttons ON/OFF and START. Rinse port, plastic parts and spring. Let air dry. An aluminum tool to unscrew pushbuttons is part of delivery.
- Thoroughly rinse pressure sensor with freshwater. Let air dry. Only if particles remain after rinsing, remove protective cap. Use fingers to unscrew. When screwed very tightly, use small water pipe tongs (prevent scratches by wrapping the protective cap with a cloth). Rinse with freshwater. Never touch the sensor surface! Replace protective cap. Let air dry.

9.2 Battery Maintenance

Even when the fluorometer is not used, the status of the battery charge decays due to self-drain. Although self-drain currents are very low, the battery can become deeply discharged over long time periods. A deep discharge stresses the battery and can significantly reduce the battery capacity. Therefore, when the fluorometer is not used, charge battery every 6 months.

9.3 Battery Replacement

a Preparation

Disconnect all cables. Place DIVING-PAM-II on solid surface with display in front. Use paper towels or other padding to avoid scratches.



b Removing left end plate (01)

Use 6 mm Allen key (included in delivery) to remove the Allen screw connected to front handlebar.

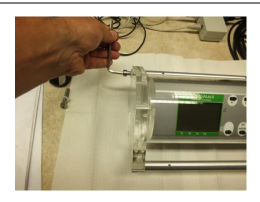


Fig. 82: Battery Replacement 01

c Removing left end plate (02)

Remove the Allen screw connected to rear handlebar.

NOTE FOR REASSEMBLY When reassembling the end plate, Tighten the two screws alternately and in small increments.



d Removing left end plate (03)

Take off end plate.



e Removing Plexiglas tube (01)

Pull out tube.

NOTE FOR REASSEMBLY Lightly grease sealing rings (lubricant included in delivery). Check correct position of sealing rings. Check if sealing rings are all flat and not folded or twisted.

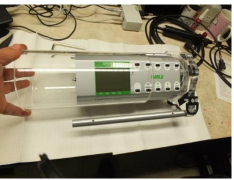


Fig. 83: Battery Replacement 02

f Removing Plexiglas tube (02)

Put tube aside.



g Disconnecting battery (01)

Turn fluorometer upside down.

Pull off battery plug connected to **RED** cable (+ sign, positive pole).



h Disconnecting battery (02)

Pull off battery plug connected to **BLACK** cable (sign, negative pole).

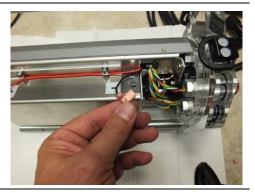
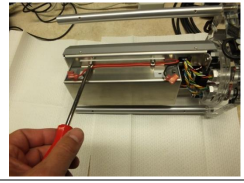


Fig. 84: Battery Replacement 03

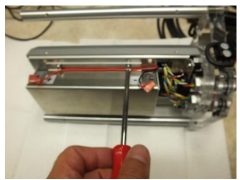
i Removing battery holder (01)

Use 6 mm Phillips screwdriver to undo the screw close to plus sign.



j Removing battery holder (02)

Undo screw close to minus sign.



k Removing battery holder (03)

Rotate fluorometer until the two remaining screws of the battery holder are easily accessible. Undo any of the two screws.

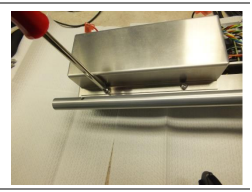
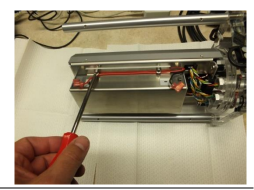


Fig. 85: Battery Replacement 04

I Removing battery holder (04)

Undo last screw of battery holder.



m Removing battery holder (05)

Lift off battery holder.



n Removing battery (01)

Put aside the four screws. Flip over battery holder.



Fig. 86: Battery Replacement 05

o Removing battery (02)

Release battery by gently striking the holder onto the hand.

NOTE FOR REASSEMBLY Match plus and minus signs of new battery and battery holder. Proceed in reverse order (n→a).

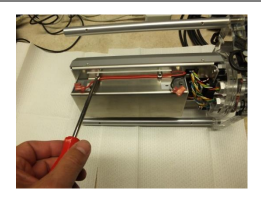


Fig. 87: Battery Replacement 06

10 Saturation Pulse Analysis

10.1 Pulse-amplitude Modulated (PAM) Fluorescence

The PAM principle is illustrated by Fig. 88. The top part shows the total fluorescence of a sample. µs-measuring flashes are given throughout the experiment starting with "Pulse on". These flashes cause the spikes in the fluorescence trace. From left to right, an external effect induces a "False Signal" of continuous fluorescence in the darkened sample. Then the sample is exposed to a period of actinic illumination ("Actinic on" and "Actinic off"), and, finally, the sample is kept in the dark again.

During actinic illumination, an effect of stray light on the fluorescence signal is additionally assumed. The fluorescence level at onset of stray light plus actinic light is denoted "Actinic F_0 ". The further increase of continuous fluorescence during illumination is denoted "Actinic F_V ", where the V stands for variable fluorescence. The "Actinic F_V " reflects changes of the fluorescence yield in the sample because stray light and actinic light are constant during the illumination period.

In Fig. 88, not only continuous fluorescence varies but also the amplitude of fluorescence spikes. PAM fluorometers ignore the changes of continuous fluorescence and measure only the amplitude of fluorescence spikes. This is achieved by subtracting the fluorescence level just before the μ s-measuring flash from the fluorescence level at the μ s-measuring flash. In Fig. 88, the PAM fluorescence amplitude during the initial dark phase is denoted "Pulsed F₀", and the maximum variable fluorescence at the end of actinic illumination is denoted "Pulsed F_V".

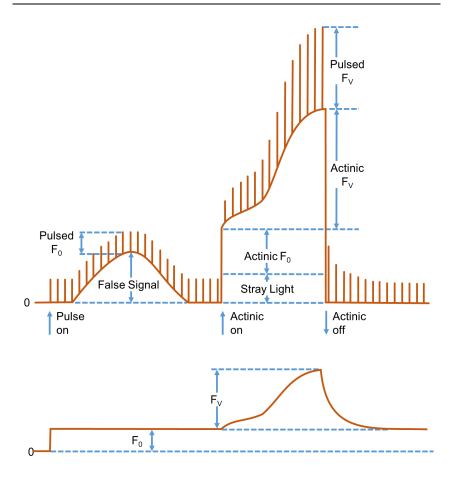


Fig. 88: Illustration of the PAM measurement principle
Figure redrawn from Dr. Ulrich Schreiber, Manual for PAM-101.

Because the µs-measuring flashes have constant amplitude, the varying amplitudes of fluorescence spikes is a measure of how efficient excitation light is converted into fluorescence. In other words, PAM fluorescence is proportional to the fluorescence yield.

The lower trace in Fig. 88 outlines the PAM fluorescence trace. Obviously, PAM fluorescence irons out the "False Signal" of total

fluorescence at the beginning of the experiment, and also the fluorescence jumps when actinic light is switched on and off. The course of continuous fluorescence within the range "Actinic F_V " resembles the corresponding trace of PAM fluorescence, because both measuring light and actinic illumination are constant.

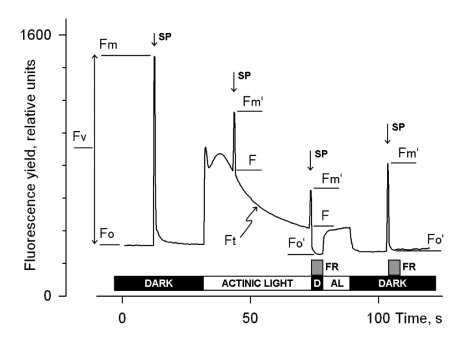


Fig. 89: Fluorescence Levels of Saturation Pulse Analysis

Y-axis (Fluorescence yield) corresponds to PAM fluorescence, see 10.1. AL, Actinic Light; D, dark; SP, Saturation Pulse; Ft, continuously recorded PAM fluorescence; FR, far-red illumination.

10.2 Saturation Pulse Analysis

The five principal levels of PAM fluorescence which are used for saturation pulse analysis are shown in Fig. 89. Two of these levels (F_0 and F_M) must be measured with the dark-acclimated sample. The three other levels (F_0 ', F, and F_M ') are measured with the actinic light-exposed sample or in a dark period following this light treatment. Some parameters of saturation pulse analysis require fluorescence measurement of the same sample in both the dark-acclimated and light-exposed state (Table 27, page 168).

Because PAM fluorescence is excited by μs pulses of <u>constant</u> amplitude, variations between fluorescence levels are usually interpreted as variation in chlorophyll fluorescence yield. This applies for variations between different types of fluorescence levels (e.g. between F_0 and F_M) and for variations of the same type of fluorescence level (e.g. the change of F_M ' during a fluorescence induction curve).

Measurements with Dark-Acclimated Samples

- **F**₀ Minimum fluorescence level excited by very low intensity of measuring light to keep photosystem II reaction centers open.
- **F_M** Maximum fluorescence level elicited by a pulse of saturating light (Saturation Pulse) which closes all photosystem II reaction centers.

Measurements with Illuminated Samples

 F_0 ' Minimum fluorescence level of illuminated sample. The F_0 ' is lowered relative to F_0 by non-photochemical quenching. The measuring routine for F_0 ' (see Fig. 89, page 161) determines the F_0 ' level during a dark interval following a Saturation Pulse. In this dark interval, far-red light is applied which selectively drives photosystem I. As

a consequence, electrons are removed from the intersystem electron transport chain and opening of photosystem II reaction centers is efficiently accelerated

If the F_0 ' Mode is switched off, the F_0 ' will be calculated according to Oxborough and Baker:

$$F_0' = \frac{1}{\frac{1}{F_0} - \frac{1}{F_M} + \frac{1}{F_M'}}$$

Oxborough K, Baker NR (1997) Resolving chlorophyll a fluorescence images of photosynthetic efficiency into photochemical and non-photochemical components - calculation of qP and Fv'/Fm' without measuring Fo'. Photosynth Res 54 135-142. https://doi.org/10.1023/A:1005936823310

In the Record table, the calculated value F_0 ' is preceded by a tilde sign (\sim).

- F_M ' Maximum fluorescence level of the illuminated sample. The F_M ' is induced by a Saturation Pulse which temporarily closes all photosystem II reactions centers. F_M ' is decreased relative to F_M by non-photochemical quenching.
- F The F corresponds to the momentary fluorescence level (Ft) of an illuminated sample shortly before application of a Saturation Pulse.

To quantify photochemical use and non-photochemical losses of absorbed light energy, fluorescence ratio expressions have been derived which use two or more of the five relative fluorescence yields introduced above. Table 27 (page 168) compiles the fluorescence ratio parameters available in the DualPAM software. Below, these parameters will be explained briefly.

F_V/**F**_M and **Y**(**II**) Maximum and effective photochemical quantum yields of photosystem II

The F_V/F_M and Y(II) estimate the fraction of absorbed quanta used for photosystem II photochemistry. F_V/F_M corresponds to the maximum photochemical yield of photosystem II, Y(II) is the effective photochemical yield of photosystem II. Measurements of F_V/F_M require that samples are acclimated to darkness or dim light so that all reactions centers are in the open state and non-photochemical dissipation of excitation energy is minimal.

In algae and cyanobacteria, however, the dark-acclimated state often is not showing maximal photosystem II quantum yield, as the photosystem II acceptor pool may be reduced in the dark by stromal reductants and, consequently, the so-called state 2 is formed exhibiting low photosystem II quantum yield. In this case, preillumination with moderate far-red light should precede determinations of F_0 and F_M .

The Y(II) value estimates the photochemical use of excitation energy in the light. It is lowered with respect to F_V/F_M by partial closure of photosystem II centers and various types of non-photochemical energy losses induced by illumination.

\mathbf{q}_{P} and \mathbf{q}_{L} $\,$ Coefficients of photochemical fluorescence quenching

Both parameters estimate the fraction of open photosystem II reaction centers. The q_P is based on the concept of separated photosystem II antenna units (puddle model), whereas the q_L assumes interconnected photosystem II antenna units (lake model) which was assumed to be present in leaves (*cf.* Kramer *et al.*, 2004). Determinations of q_P an q_L do not require fluorescence measurements with the dark-acclimated sample, except the F_0 ' mode is switched of and F_0 ' is calculated according to Oxborough and Baker (1997).

q_N and **NPQ** Parameters of non-photochemical quenching

Both parameters are associated with non-photochemical quenching of excitation energy, mainly involving a low thylakoid lumen pH- and a zeaxanthin-dependent quenching mechanism. The qN and the NPQ parameters require fluorescence measurements with the sample in the dark-acclimated and in the light-exposed states (cf. Table 27, page 168).

Calculation of NPQ (or SV_N; Gilmore and Yamamoto, 1991) corresponds to the Stern-Volmer equation for fluorescence quenching which predicts proportionality between fluorescence quenching (NPQ) and the concentration of fluorescence-quenching centers in the photosynthetic antennae (e.g. zeaxanthin).

Y(NO), Y(NPQ) and Y(II) Complementary photosystem II yields

Genty et al. (1996) and Kramer et al. 2004 have presented expressions describing the partitioning of absorbed excitation energy in photosystem II between three fundamental pathways the sum of which adds up to one:

- Y(NO) non-regulated losses of excitation energy including heat dissipation and fluorescence emission,
- Y(NPQ) regulated energy losses of excitation energy by heat dissipation involving ΔpH and zeaxanthin-dependent mechanisms, and
- Y(II) use of excitation energy for charge separation.

This concept of "complementary photosystem II quantum yields" is useful to analyze the partitioning of absorbed light energy in photosynthetic organisms. For instance, in the presence of strong light, a much higher Y(NPQ) than Y(NO) indicates that excess excitation energy is safely dissipated at the antenna level and that photosynthetic energy fluxes are well-regulated.

In variance, high values of Y(NO) would signify that excess excitation energy is reaching the reaction centers, resulting in strong reduction of photosystem II acceptors and photodamage, e.g. via formation of reactive oxygen species.

10.3 Relative Electron Transfer Rate (ETR)

Relative electron transfer rates for photosystem II are calculated according to:

$ETR(II) = PAR \cdot ETR-Factor \cdot P_{PS2}/P_{PS1+2} \cdot Y(II).$

The basic idea of the ETR equation is to multiply Y(II), the effective photochemical quantum yield of photosystem II, by an estimate for the photon flux density absorbed by all photosystem II in the sample. The latter estimate is derived from three numbers:

- (1) PAR Quantum flux density of photosynthetically active radiation (PAR) impinging on the sample.
- (2) ETR-Factor Sample absorptance (= 1 transmittance)

The ETR-Factor describes the fraction of incident photons absorbed by the sample. The most frequently used default value for green leaves is 0.84 meaning that 84% of incoming light is absorbed. The ETR-Factor can be lower in bleached leaves or leaves containing considerable amounts of non-photosynthetic pigments like anthocyanins.

(3) P_{PS2}/P_{PS1+2} Relative distribution of absorbed PAR to photosystem II

The default P_{PS2}/P_{PS1+2} is 0.5 which assumes the photosystem II contributes 50% to total sample absorptance. The P_{PS2}/P_{PS1+2} may deviate from the idealized factor of 0.5 depending on wavelength of light and acclimation status of the sample.

10.4 Reviews on Saturation Pulse Analysis of Photosystem II

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16 OH M : F (4004) OH - H II II

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Murchie EH, Lawson T (2013) Chlorophyll fluorescence analysis: a guide to good practice and understanding some new applications. J Exp Bot 64: 3983-3998. https://doi.org/10.1093/jxb/ert208

Schreiber U (2004) Pulse-amplitude-modulation (PAM) fluorometry and saturation pulse method: an overview. In: Papageorgiou GC, Govindjee (eds) Chlorophyll *a* Fluorescence: A Signature of Photosynthesis. Springer, Dordrecht, pp 279-319. https://doi.org/10.1007/978-1-4020-3218-9_11

Table 27: Fluorescence Ratio Parameters.

Source	Equation	Sample State	Range [Theory] [Experiment]
Maximum photochemical quantum yield of PS II (Kitajima and Butler, 1975)	$\frac{F_V}{F_M} = \frac{F_M - F_0}{F_M}$	Dark	[0, 1] [0, ~0.84]
Effective photochemical quantum yield of PS II (Genty et al., 1989)	$Y(II) = \frac{F_M' - F}{F_M'}$	Light	[0, 1] [0,~ 0.84]
Quantum yield of light-induced (ΔpH- and zeaxanthin-dependent) non-photochemical fluorescence quenching (Genty <i>et al.</i> 1996, Kramer <i>et al.</i> 2004)*	$Y(NPQ) = \frac{F}{F_M'} - \frac{F}{F_M}$	Dark and Light	[0, 1] [0, ~ 0.9]
Quantum yield of non-regulated heat dissipation and fluorescence emission: this type of energy loss does not involve the action of a transthylakoid ΔpH and zeaxanthin (Genty et al. 1996, Kramer et al. 2004)*	$Y(NO) = \frac{F}{F_M}$	Dark and Light	[0, 1] [0, ~ 0.9]
Stern-Volmer type non-photochemical fluorescence quenching (Bilger and Björkman, 1990; Gilmore and Yamamoto, 1991))	$NPQ = \frac{F_M}{F_M'} - 1$	Dark and Light	[0, ∞] [0, ~4]
Coefficient of photochemical fluorescence quenching (Schreiber et al. 1986 as formulated by van Kooten and Snel, 1990)	$q_P = \frac{F_M - F}{F_M' - F_0'}$	Light. If F ₀ ' calculated, Dark and Light	[0, 1] [0, 1]
Coefficient of photochemical fluorescence quenching assuming interconnected PS II antennae (Kramer et al. 2004)	$q_L = q_P \cdot \frac{F_0'}{F}$	As q _{P'.}	[0, 1] [0, 1]
Coefficient of non-photochemical flu- orescence quenching (Schreiber <i>et al.</i> 1986 as formulated by van Koo- ten and Snel, 1990)	$q_N = 1 - \frac{F_M' - F_0'}{F_M - F_0}$	Dark and Light	[0, 1] [0, ~0.95]

^{*} Kramer *et al.* (2004) have derived more complex equations for Y(NO) and Y(NPQ). Klughammer and Schreiber (2008) have transformed the equations by Kramer *et al.* (2004) into the simple equations of Genty *et al.* (1996).

Table 28: References Cited in Table 27

Bilger W, Björkman O (1990) Role of the xanthophyll cycle in photoprotection elucidated by measurements of light-induced absorbance changes, fluorescence and photosynthesis in leaves of *Hedera canariensis*. Photosynth Res 25:173-185. https://doi.org/10.1007/BF00033159

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https://doi.org/10.1007/BF00033156

11 Specifications

11.1 Basic System

11.1.1 Optoelectronic Unit

Design: Plexiglas tube with Plexiglas end plates, one with water-proof fiberoptics port

Mountings: Anodized aluminum rod (diameter 15 mm) mounted parallel to the fluorometer body below the screen at a distance of 30 mm, and another one mounted on the opposite side at a distance of 5 mm. Anodized aluminum holder for fiber optics and cables at one end of the fluorometer (maximum dimensions 20.5 cm x 18.5 cm x 5 cm; L x W x H). Two plastic loops to fasten a carrying belt, and a poly-oxymethylene (POM) ring with outer diameter of 46 mm to store the miniature spectrometer MINI-SPEC

Display: Backlit 160 x 104 dots (78 mm x 61 mm) transflective B/W screen

Control elements: 10 infrared reflection switches to operate control fields on the screen, pushbutton for saturation pulses, pushbutton to switch device on/off and to lock/unlock reflection switches

Pressure and temperature sensor: Module including a high linear pressure sensor and a temperature sensor. With gel protection and antimagnetic stainless steel cap

Ports equipped with watertight caps:

AUX1 and AUX2, 4-pole, for miniature spectrometer MINI-SPEC and, via special adapter, for MINI-PAM-II accessories Fiber-Optic Oxygen Meter FireStingO2 or Leaf Clip Holder 2035-B (connected via adapter cable)

OUT1 and OUT2, 6-pole, for operation of an external light source synchronized with PAM measuring light. Works also as input for or trigger signal from Sample Holder DIVING-II-USH

INPUT, 6-pole, for RS-485 communication and charging of internal battery. Works also as input for or trigger signal from Sample Holder DIVING-II-USH

Battery: Lead acid battery 8.0 V / 3.5 Ah (28 Wh) providing power for more than 1300 yield measurements

Maximum diving depth: 50 m

Operating temperature: -5 to +45 °C

Dimensions: Diameter 19 cm, length 39 cm

Weight: 3.9 kg

11.1.2 Light sources

DIVING-PAM-II/B (Blue Version)

Measuring light: Blue (470 nm) LED, standard modulation frequencies 5 to 25 Hz, adjustable in increments of 5 Hz, and 100 Hz, measuring light PAR at standard settings = $0.05 \, \mu$ mol m⁻² s⁻¹. Fluorescence at wavelengths greater than 630 nm is measured

Actinic light: Same blue LED as for measuring light, maximum actinic PAR = $3000 \, \mu \text{mol m}^{-2} \, \text{s}^{-1}$, maximum PAR of saturation pulses = $6000 \, \mu \text{mol m}^{-2} \, \text{s}^{-1}$ adjustable at increments of $500 \, \mu \text{mol m}^{-2} \, \text{s}^{-1}$.

DIVING-PAM-II/R (Red Version)

Measuring light: Red (655 nm) LED, modulation frequencies and PAR as described for DIVING-PAM-II/B. Fluorescence at wavelengths greater than 700 nm is measured

Actinic light: Same red LED as for measuring light, maximum PAR of actinic light and saturation pulses as described for DIVING-PAM-II/B

DIVING-PAM-II/B and DIVING-PAM-II/R

Far-red light: Peak emission at 735 nm

11.1.3 Data acquisition

Fluorescence: PIN photodiode protected by long-pass and a short-pass filters, 12 bit signal resolution

Other parameters: Piezo-resistive pressure sensor and temperature sensor. Pressure is converted in meters of diving depth, range 0 to -50 m, displayed at 0.1 m intervals. Temperature, range -10 °C to +60 °C, displayed at 0.1 °C intervals

Data storage: Flash memory, 8 MB, providing memory for more than 27,000 saturation pulse analyses

11.1.4 WLAN

Wireless LAN Interface, IEEE 802.11 b/g/n (2.4 GHz), Access Point Mode

11.1.5 Fiberoptics DIVING-F

Design: Randomized 70 μ m glass fibers forming single plastic shielded bundle with stainless steel adapter ends

Dimensions: Active diameter 5.5 mm, outer diameter 8 mm, length 150 cm

Weight: 340 g

11.1.6 PC Interface Box DIVING-PAM-II/I

Housing: Aluminum case with USB-B port, socket for power supply MINI-PAM-II/N, and waterproof 6-pole socket for RS-485 communication

Function: The interface box connects computer and DIVING-PAM-II. RS-485 serial data communication is used between box and DIVING-PAM-II, USB communication is employed between interface box and computer. Recommended maximum cable lengths are: 100 m RS-485 cable between DIVING-PAM-II and interface box, 2 m USB cable between interface and box computer.

Standard USB-A to USB-B cable included

Dimensions: 9.7 cm x 6.3 cm x 3.5 cm (L x W x H)

Weight: 270 g

Operating temperature: -5 °C to + 40 °C

11.1.7 Power Supply MINI-PAM-II/N

Input: 100 V to 240 V AC, 50 to 60 Hz

Output: 12 V DC, 5.5 A

Operating temperature: -5 to +45 °C, (non-condensing)

Dimensions: 13 cm x 5.5 cm x 3 cm (L x W x H)

Weight: 350 g including cables

11.1.8 Underwater Cable DIVING-PAM-II/K5

Underwater cable for RS-485 communication and charging of the DIVING-PAM internal battery

Length: 5 m

Weight: 500 g

11.1.9 Miniature Spectrometer MINI-SPEC

Design: POM tube, at one side, port for light detection, port for fluorescence excitation by blue (452 nm max) or green (525 nm max) LEDs, and port for white light from a tungsten lamp for reflection measurements; at the opposite side, 4-pole underwater socket.

Spectrometer: Hamamatsu micro-spectrometer, spectral range: 400 to 800 nm, spectral resolution: between 8 and 10 nm.

Maximum PAR: 4000 µmol m⁻² s⁻¹ for illumination having spectral characteristics similar to sunlight

Dimension: 3.25 cm diameter, 17.5 cm length max

Weight: 135 g

11.1.9.1 Flat Entrance Optics SPEC/P

Design: Hard-anodized aluminum rod of 10 mm diameter and 50 mm length, at one end with lateral light entrance through a 5 mm diameter diffusor and the opposite end inserted in a mounting plate (diameter 33 mm, height 5 mm). Aluminum rod with internal light guide

11.1.9.2 Fluorescence and Reflection Optics SPEC/R

Design: Spectrometer cap consisting of POM: maximum diameter 35 mm, height 13 mm, weight 16 g. With central 5 mm x 16 mm groove which accommodates at one end a Perspex light guide for fluorescence excitation by blue or green light, and at the other end a Perspex light guide for white light for reflection measurements. With 3 mm diameter central drilled hole as light channel to the detector window of the spectrometer. The sample is fixed between the cap part and another disk (maximum diameter 40 mm, height 10 mm, weight 8 g). The sample side of cap and disk is padded with foam rubber, both parts have magnets build-in to attract each

other and, thus, hold the sample. Including a 10 mm thick Zenith Polymer reflectance standard

11.1.9.3 PAR Calibration Block 000160101439

Design: POM block with drill hole for entrance optics of the Miniature Spectrometer. Oriented at an angle of 60° and 90° relative to the spectrometer port are drill holes for the Fiber Optics DIVING-F

Dimensions: 4.15 cm x 2 cm x 5 cm (L x W x H)

Weight: 40 g

11.1.10 Distance Clip 60° 2010-A

Design: Metal clip with fiber holder and 11 mm diameter sample

hole: 5.5 cm x 1.4 cm (L x W)

Fiber holder: 1.2 cm length, mounted 0.7 cm above base, with lateral screw to fix fiber optics. Angle between fiber optics axis and sample plane: 60°. Two spacer rings to vary the distance between fiber end and leaf surface

11.1.11 Dark Leaf Clip DIVING-LC

Design: Three clips made of white plastic with gasket contact areas and sliding shutter for light-tight closure.

Dimensions: Diameter 3.2 cm, length 8 cm

Weight: 6.5 g

11.1.12 Surface Holder DIVING-SH

Design: Holder made of grey PVC, equipped with 3 rubber bands and hooks to be attached to creviced surface (e.g. of coral); nylon screws for distance adjustment

Dimensions: $6 \text{ cm } \times 6 \text{ cm } \times 2.5 \text{ cm } (L \times W \times H)$

Weight: 95 g

11.1.13 Software WinControl-3

Program: WinControl-3 System Control and Data Acquisition Program (Windows 7, 8, 10) for operation of DIVING-PAM-II via PC, data acquisition and data analysis

Saturation Pulse Analysis: Measured: Ft, F_0 , F_M , F, F_0 ' (also calculated), F_M '. PAR, water temperature and depth (derived from pressure). Calculated: F_0 ' (also measured), F_V/F_M and Y(II) (maximum and effective photochemical yield of PS II, respectively), q_L , q_P , q_N , NPQ, Y(NPQ), Y(NO) and ETR (electron transport rate)

Fitting Routines: Two routines for determination of the cardinal points α , I_k and ETR_{max} of light curves

Programmed Features: Automatic determination of signal offset for all light intensities and all gain levels. Automatic calibration of internal PAR sensor against PAR from MINI-SPEC sensor

Communication Protocol: USB and IEEE 802.11 b/g/n

Computer Requirements: Processor, 1 GHz. RAM, 512 MB. Screen resolution, 1024 x 600 pixels. Interface, USB 2.0/3.0.

11.1.14 Transport Case DIVING-PAM-II/T

Design: Rugged, hard plastic outdoor case with wheels, pull-out handle and custom foam packing

Dimensions: 57 cm x 47 cm x 27 cm (L x W x H)

Weight: 7.7 kg

11.2 Accessories

11.2.1 Universal Sample Holder DIVING-II-USH

Design: Plexiglas bar (15 cm x 4.5 cm) with upward curved end possessing a port for positioning at 60° or 90° relative to the sample level the DIVING-PAM-II fiber optics. Mounted to the curved end is a 5.5 cm x 7.5 cm (W x H) sample clip consisting of a Plexiglas plate (lower part) and an aluminum frame open to the top (upper part). Featuring a 10 cm long plastic grip with button for triggering measurements via a 1.5 m trigger cable. Including a 1 m long tubular net with zipper to keep together trigger cable and fiber optics. With holder to for spectrometer MINI-SPEC

Dimensions: 25 cm x 4.5 cm x 21 cm (L x W x H)

Weight: 380 g

11.2.2 Underwater Oxygen Sensor DIVING-PAM-II/O2

Design: POM tube, at one end face, optical sensor spot fixed by a Perspex disk with central hole and PT100 resistance thermometer, at the other end face, 4-pole underwater socket. Temperature-compensated oxygen measurement by a high precision optical oxygen meter connected to the sensor spot by fiberoptics. Including a 2 m underwater cable 000130204945, two spare oxygen sensor spots OXSP5, and a holder (weight 75 g, maximum dimensions 6.5 cm x 6 cm x 12 cm, L x W x H) to attach both the DIVING-PAM-II/O2 and the spectrometer MINI-SPEC to the DIVING-PAM-II optoelectronic unit, consisting of the mounting brackets 000246001714 and 000246003914 and 1 ring holder 000244905514.

Dimension: 3.25 cm diameter, 17.5 cm length max

Weight: 135 g

11.2.3 Magnet Sample Holder DIVING-MLC

Design: Two plastic halves with magnets, one with sliding shutter made of spring steel and seat for the adapter DIVING-DA. The other halve with seats for additional magnets. Inner sides of both halves covered with black fabric. Including 4 additional magnets.

Dimensions: Diameter 30 mm, height 28 mm

Weight: 10 g

11.2.4 Surface Holder DIVING-SH

Design: Holder made of grey PVC, equipped with 3 rubber bands and hooks to be attached to creviced surface (e.g. of

coral); nylon screws for distance adjustment

Dimensions: $6 \text{ cm } \times 6 \text{ cm } \times 2.5 \text{ cm } (L \times W \times H)$

Weight: 95 g

11.2.5 Miniature Fiberoptics DIVING-F1

Design: Both ends with adapter which fits to the fiber port of the DIVING-PAM-II and the various accessories for fiber positioning of the DIVING-PAM-II system.

Dimensions: Active diameter 2 mm, length 1.5 m

11.2.6 Underwater Cable DIVING-II/K25

Dimensions: 25 m length, 6 mm diameter

Weight: 1.25 kg

11.2.7 Underwater Cable DIVING-II/K50

Dimensions: 50 m length, 6 mm diameter

Weight: 2.5 kg

Including charger **DIVING-II/L15**:

Input: 100 V to 240 V AC, 47 to 63 Hz

Output: 15 V DC, 4.65 A

Operating temperature: -5 to +45 °C, (non-condensing)

Dimensions: 13 cm x 5.5 cm x 3 cm (L x W x H)

Weight: 350 g including cables

Subject to change without prior notice

12 Guarantee

12.1 Manufacturer's Guarantee

Under this Manufacturer's Guarantee ("Guarantee"), subject to the Conditions and Instructions below, Heinz Walz GmbH, Germany ("Manufacturer"), guarantees (§443 BGB) to the end customer and user ("Customer") that all products supplied by it shall substantially conform in material respects to the Specifications for 24 months from the delivery date (date on invoice). In this Guarantee, "Specifications" means the product's features (as may be amended by Manufacturer from time to time), which are set out under the headings "specifications" and/or "technical specifications" within the product's respective brochure, data sheet, or respective tab on the Manufacturer's website for such product, and which may be included with the documents for the product when delivered. In case of an eligible guarantee claim, this Guarantee entitles the Customer to repair or replacement, at the Manufacturer's option, and this Guarantee does not include any other rights or remedies.

12.2 Conditions

This Guarantee shall not apply to:

- Any defects or damage directly or indirectly caused by or resulting from the use of unauthorized replacement parts and/or service performed by unauthorized personnel.
- Any product supplied by the Heinz Walz GmbH, Germany which has been subjected to misuse, abuse, abnormal use, negligence, alteration or accident.

- Damage caused from improper packaging during shipment or any acts of God.
- Batteries, cables, calibrations, fiberoptics, fuses, gas filters, lamps (halogen, LED), thermocouples, and underwater cables.
- Defects that could reasonably have been detected upon inspection of the product when received by the Customer and not promptly noticed within ten (10) days to Heinz Walz GmbH.
- Submersible parts of the DIVING-PAM or the underwater version of the MONITORING-PAM have been tested to be watertight down to the maximum operating depth indicated in the respective manual. Guarantee shall not apply for diving depths exceeding the maximum operating depth. Further, guarantee shall not apply for damage resulting from improper operation of devices, in particular, the failure to properly seal ports or sockets.

12.3 Instructions

- To obtain guarantee service, please follow the instructions below:
- The Walz Service Information Form available at https://www.walz.com/support/repair_service.html must be completed and returned to Heinz Walz GmbH, Germany.
- The product must be returned to Heinz Walz GmbH, Germany, within 30 days after Heinz Walz GmbH, Germany has received written notice of the defect. Postage, insurance, and/or shipping costs incurred in returning equipment for guarantee service are at customer expense. Duty and taxes are covered by Walz.

- All products being returned for guarantee service must be carefully packed and sent freight prepaid.
- Heinz Walz GmbH, Germany is not responsible or liable for missing components or damage to the unit caused by handling during shipping. All claims or damage should be directed to the shipping carrier.

12.4 Applicable law

This Guarantee is governed by German law. Place of jurisdiction is Bamberg, Germany.

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