

**Functional description** 

## photoLab<sup>®</sup> 6600 UV-VIS (spectroFlex 6600)



Spectrophotometer

ba75848e04 09/2011

### Accuracy when going to press

The use of advanced technology and the high quality standard of our instruments are the result of continuous development. This may result in differences between this operating manual and your instrument. Also,

we cannot guarantee that there are absolutely no errors in this manual. Therefore, we are sure you will understand that we cannot accept any legal claims resulting from the data, figures or descriptions.



#### Note

The latest version of the present operating manual is available on the Internet under http://www.WTW.com.

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#### 1 **Overview**

1.1 **Overview of the instrument** 

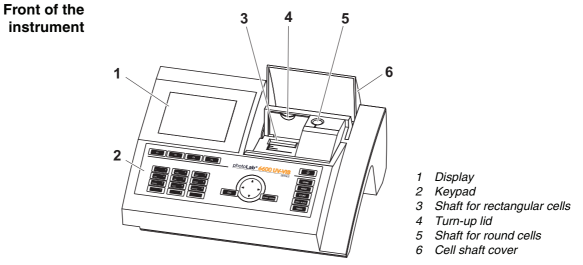
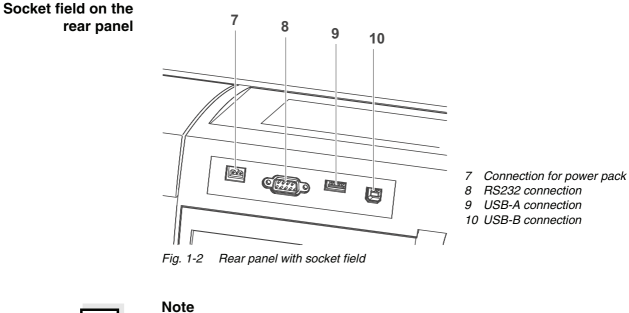


Fig. 1-1 Front of the instrument with operating elements

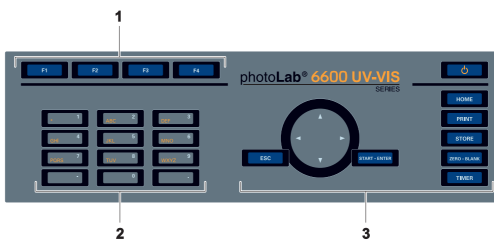




All connections comply with SELV.

#### Keypad 1.2

**Overview** 



- Function keys F1 to F4 (function menu-depending)
   Alphanumeric keypad
   Keys with dedicated function

Fig. 1-3 Keypad

**Key functions** The keys on the right side of the keypad have the following functions:

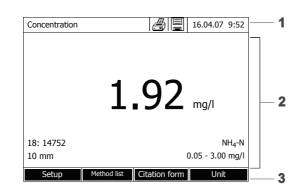
Кеу	Designation	Functions
Ģ	<on off=""></on>	<ul> <li>Switches on and off the photome- ter</li> </ul>
HOME	<home></home>	<ul> <li>Switches to the main menu from any operating situation. Actions that are not completed are can- celed.</li> </ul>
PRINT	<pre>PRINT&gt;</pre>	<ul> <li>Outputs the displayed measured value to an interface if the <i>Printer</i> symbol is displayed in the status line.</li> </ul>
STORE	<store></store>	<ul> <li>Saves a displayed measured value or spectrum if the Save symbol is displayed in the status line.</li> </ul>
ZERO • BLANK	<zero·blank></zero·blank>	<ul> <li>Starts one of the following measurements, depending on the operating situation:</li> <li>Zero adjustment</li> <li>Blank value measurement</li> <li>Baseline measurement</li> </ul>

Кеу	Designation	Functions
TIMER	<timer></timer>	- Opens the menu, <i>Timer</i> .
ESC	<esc></esc>	<ul> <li>Cancels the running action.</li> <li>Entries that have not yet been accepted are discarded.</li> </ul>
		<ul> <li>Switches to the next higher menu level.</li> </ul>
START • ENTER	<start.enter></start.enter>	<ul> <li>Starts an action (e.g. measure- ment)</li> </ul>
		<ul> <li>Opens a selected menu</li> </ul>
		<ul> <li>Confirms a selection or entry</li> </ul>
	< <b>▲</b> >or <▼>	
<	< <b>∢</b> >	<ul> <li>Deletes the character left of the cursor during character entries</li> </ul>
		<ul> <li>Moves the cursor to the left in a spectrum or kinetic diagram</li> </ul>
(Arrow keys)	<▶>	<ul> <li>Moves the cursor to the right in a spectrum or kinetic diagram</li> </ul>

**Function keys** The function keys F1 to F4 have different functions depending on the operating situation. The current functions are displayed in the function key menu at the bottom edge of the display (see section 4.2.1).

**Display elements** 

#### 1.3 Display



- 1 Status line (current state, date and time)
- 2 Display range for menus and measurement results
- 3 Function keys menu

Fig. 1-4 Display

Symbols in the	Symbol	Designation	Function
status line		Save	The <b><store></store></b> key is active. You can store the displayed data with <b><store></store></b> (see section 4.11).
		Printer	The <b><print></print></b> key is active. You can output to an interface the displayed data with <b><print></print></b> (see section 4.14).
		Progress bar	During the warm-up time (15 minutes) a progress bar appears on the display. The reproducibility of measured values is limited during the warm-up time (see section 4.14).

#### 2 Safety instructions

This operating manual contains basic instructions that you must follow during the commissioning, operation and maintenance of the photometer. Consequently, all responsible personnel must read this operating manual carefully before working with the meter. Keep this operating manual in the vicinity of the meter.

#### General safety instructions

Safety instructions in this operating manual are indicated by the warning symbol (triangle) in the left column. The signal word (such as "CAUTION") indicates the danger level:

#### WARNING

indicates instructions that must be followed precisely in order to prevent serious dangers to personnel.

# $\triangle$

#### CAUTION

indicates instructions that must be followed precisely in order to avoid slight injuries to personnel or damage to the instrument or the environment.



# **i**

#### Note

indicates notes that draw your attention to special features.

#### Note indica

indicates cross-references to other documents.

#### 2.1 Target group and user qualification

The photometer was developed for use in the laboratory. Carrying out photometric determinations with the aid of test sets frequently requires the handling of hazardous substances.

We assume that the operating personnel know how to handle hazardous substances due to their professional training and experience. The operating personnel must particularly be able to understand and correctly implement the safety labels and safety instructions on the packages and inserts of the test sets.

#### 2.2 Authorized use

The authorized use of the photometer consists exclusively of the carrying out of photometric measurements according to this operating manual. Follow the technical specifications of the cells in chapter 7 TECHNICAL DATA. Any other use is considered to be **unauthorized**.

#### 2.3 General safety instructions

The photometer is built and inspected according to the relevant guidelines and norms for electronic instruments (see chapter 7 TECHNICAL DATA). It left the factory in a safe and secure technical condition.



#### Note

The opening of the photometer or adjustment and repair work must only be performed by specialist personnel authorized by the manufacturer. Noncompliance invalidates any claim with regard to the warranty.

Function and operational safety

The smooth functioning and operational safety of the photometer can only be guaranteed if the generally applicable safety measures and the specific safety instructions in this operating manual are followed during operation.

The smooth functioning and operational safety of the photometer can only be guaranteed under the environmental conditions that are specified in chapter 7 TECHNICAL DATA.

If the photometer was transported from a cold environment to a warm environment, the formation of condensate can lead to the faulty functioning of the meter. In this event, wait until the temperature of the meter reaches room temperature before putting the meter back into operation.

**Safe operation** If safe operation is no longer possible, the photometer must be taken out of operation and secured against inadvertent operation.

Safe operation is no longer possible if the photometer:

- has been damaged in transport
- has been stored under adverse conditions for a lengthy period of time
- is visibly damaged
- no longer operates as described in this manual.

If you are in any doubt, contact the supplier of your photometer.

#### 2.4 Handling of hazardous substances

When developing test sets, WTW carefully sees that the tests can be carried out as safely as possible. Some hazards by dangerous substances, however, cannot always be avoided.



#### WARNING

Improper handling of certain reagents can cause damage to your health.

In any case follow the safety labels on the packing and the safety instructions of the package insert. Protective measures specified there have to be followed exactly.

Safety datasheets

The safety datasheets of the chemicals comprise all instructions on safe handling, occurring hazards, preventive actions and actions to take in hazardous situations. Follow these instructions in order to work safely.

#### 3 Commissioning

#### 3.1 Scope of delivery

- Spectrophotometer photoLab<sup>®</sup> 6600 UV-VIS
- Power pack connection cable
- Buffer batteries 4 x AA alkaline manganese (Mignon)
- Zero cell (16 mm, round)
- Short instructions
- CD-ROM with
  - Detailed operating manual
  - Analysis instructions
  - SpectralTransfer software
  - Language updates to install additional character sets (see section 4.20.3)

**Packing** This photometer is sent out in a protective transport packing.



#### CAUTION

Keep the original packing including the inner packing to protect the instrument against hard shocks if it has to be transported.

The original packing is also required for the proper return of the instrument if it has to be repaired.

Note that damage caused by improper transport voids all warranty claims.

#### 3.2 General notes on handling

The photoLab<sup>®</sup> 6600 UV-VIS photometer is an optical precision meter. Therefore, it should always be handled with care, especially in mobile use. Always protect the meter from conditions that could damage the mechanical, optical and electronic components. Heed the following points especially:

- The temperature and humidity during operation and storage must be within the limits specified in chapter 7 TECHNICAL DATA.
- The following influences always have to be avoided with the meter:
  - Extreme dust, moisture and wetness
  - Exposure to intensive light and heat
  - Fumes that are corrosive or contain high concentrations of solvents.
- For measuring, the meter must be placed on a flat surface.
- Spilled liquid or other material should be removed immediately (see section 5.2 CLEANING).
- If a cell has broken in the cell shaft, the cell shaft should be cleaned immediately (see section 6.1 ACTIONS IN THE CASE OF A BROKEN CELL).
- The cell shaft should always be closed when the photometer is not used.
- During transport of the photometer, the cell shaft has to be empty.
- For mobile use we recommend the transport case FC spectral 6000 (see section 8.1 ACCESSORIES).

#### 3.3 Initial commissioning

Perform the following activities:

- Insert the buffer batteries (see section 3.3.1)
- Connect the power supply (see section 3.3.2)
- Switch on the photometer (see section 3.3.3)
- Set the language (see section 3.3.4)
- Set the date and time (see section 3.3.5)
- Carry out a zero adjustment (see section 4.4)



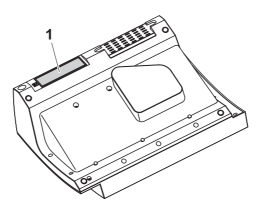
#### Note

When you set the language, date and time according to the mentioned sections of this operating manual you will quickly become familiar with the simple operation of the photoLab<sup>®</sup> 6600 UV-VIS. More detailed instructions on operation are given in section 4.2 GENERAL OPERATING PRINCIPLES.

#### 3.3.1 Inserting the buffer batteries

The buffer batteries supply the integrated clock while the photometer is switched off. Four alkaline manganese batteries (type AA or Mignon) separately included in the scope of delivery are used as the buffer batteries.

Insert the batteries as follows:



- 1 Turn the photometer upside down and place it on a soft surface.
- 2 Open the lid of the battery compartment (1).
- 3 Insert the four batteries in the battery compartment. Make sure that the poles of the batteries are in the correct position.

The  $\pm$  signs on the batteries must correspond to the  $\pm$  signs in the battery compartment.

4 Close the lid of the battery compartment.

#### Battery service life

The power consumption of the clock is very low. The lifetime of high quality batteries is at least 5 years.

#### 3.3.2 Connecting the power supply

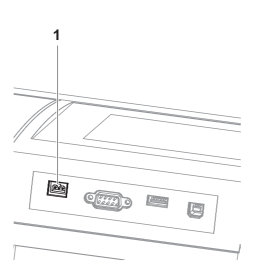
CAUTION

ly.

The power is supplied via the enclosed plug-in power pack. The power pack supplies the photometer with low voltage (12 VDC).

The line voltage of the usage location must fulfill the specifications stated on the power pack (the specifications are also given in chapter 7 TECHNICAL DATA). Always use the supplied 12 V original power pack on-

Connecting the plug-in power pack



- 1 Connect the miniplug of the power pack to the socket (1) of the photometer.
- 2 Connect the power pack to an easily accessible power socket.

The display illumination switches itself on and then off again.

# Operation with a mobile 12 V power source

You can also operate the photoLab<sup>®</sup> 6600 UV-VIS on the move and independent of the local power supply.

To do so, a 12 V power supply such as a commercial 12 V portable power source or a 12 V car battery and the ADA 12V available as an accessory is required (see section 8.1).

More detailed information on operation is available:

- in section 3.4.6 and
- in the operating manual of the ADA 12V .

#### 3.3.3 Switching on the photometer for the first time

During the initial commissioning, the photometer automatically guides you through the setting of the meter language, date and time after switching on (see following sections).

Language 16.04.07 9:52					
Englisch 🗸					
English					
Français					
Español					
Italiano					
Bulgarian/Български					
Česko					
Simplified Chinese/ 中文					
Traditional Chinese/ 繁體中文					
Greek/Ελληνικά					
Indonesian/Indonesia					

1 Press <ON/OFF>.

The photometer is switched on.

The display switches to the setting of the language (see section 3.3.4).

After the setting of the language the photometer carries out the self-test.

When the initial commissioning is completed, the photometer displays the *Home* menu each time after it is switched on and after the self-test (see section 4.1).

#### 3.3.4 Setting the language

During the initial commissioning the photometer automatically guides you to the setting of the meter language after switching on.

Language 16.04.07 9:52					
Englisch 🗸					
English					
Français					
Español					
Italiano					
Bulgarian/Български					
Česko					
Simplified Chinese/ 中文					
Traditional Chinese/ 繁體中文					
Greek/Ελληνικά					
Indonesian/Indone	sia				

- **1** Select a language with  $< \Delta > < \nabla >$ .
- 2 Confirm the selected language with **<START·ENTER>**.

The language has been set. The currently selected language is marked by a check.

The display switches to the setting of the *Date* and *Time* (see section 3.3.5).

After the initial commissioning, you can change the language in the *General* setup / Language menu at any time (see section 4.2.4).

#### 3.3.5 Setting the date and time

During the initial commissioning, the instrument automatically guides you to the setting of the time and date after the setting of the language.

Date/Time	Date/Time				
Date			16.04.200	)7	
Time			9:52:09		
			OK		

Date/Time	16.04.07 9:52
Date	16.04.2007
Time	9:52:09
Date	
23 .10.2006	
	OK

Date/Time

10 : 22 : 09

Date

Time

The Date/Time menu is open.

Using <▲><▼>, select a menu item and confirm or open it with <**START·ENTER**>.

- Select and confirm *Date*.
   The input field for the current date pops up.
- 2 Enter the current date with <0...9> and confirm.

The input field closes. The date is accepted.

- 3 Select and confirm *Time*.The input field for the current time pops up.
- 16.04.07
   9:52
   4
   Enter the current time with <0...9>

   16.04.2007
   and confirm.
   The input field closes.

The time is accepted.

After the initial commissioning, you can change the date and time in the *General setup / Date/Time* menu at any time (see section 4.2.4).

OK

#### 3.4 Connecting optional accessories

3.4.1 Communication interfaces

#### Connections

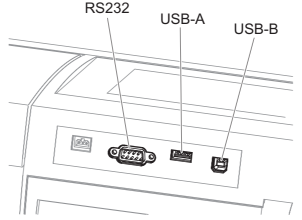


figure 3-1 Communication interfaces on the rear panel

You can connect the following accessories to the photometer:

- PC (see section 3.4.2)
- Printer (see section 3.4.2)
- USB storage media (see section 3.4.3)
- USB-PC keyboard (see section 3.4.4)
- Barcode reader (see section 3.4.5)
- ADA 12V(see section 3.4.6)



#### Note

If you want to connect several USB devices such as a USB-PC keyboard and a USB memory device to the meter, you can increase the number of USB-A sockets by a commercially available USB-2 hub with separate power supply.

#### 3.4.2 PC/printer

PC and printer can be connected to the photometer as follows:

Interface	PC	Printer	Functions
RS232	1	1	The data is sent to the interface with <b><print></print></b> .
			<ul> <li>If a printer is connected, the data is printed out.</li> </ul>
			<ul> <li>If a PC is connected, the data can be received with a terminal program (see section 4.14).</li> </ul>
USB-A		1	The data is printed out with <b><print></print></b> .
USB-B	1	-	Enables the direct connection of pho- tometer and PC. With this you can trans- mit measurement data to the PC (see section 4.12 and section 4.14) or update the photometer software (see section 4.20.1).
			The direct connection with the PC is established with the aid of the "Spectral- Transfer" program. The program is pro- vided on the supplied CD-ROM.
			More instructions on how to establish the connection are given in the operat- ing manual of the "SpectralTransfer" program (see CD-ROM).



#### Note

Suitable are all printers that can interpret the PCL-3 printer control language.

#### **Operation at RS232**

Connect the RS232 interface to the devices as follows:

- PC: with a commercially available zero modem cable
- Printer: with a commercially available RS232 printer cable

The cables are available in specialized computer shops.

Set up the following interface data at the PC/printer:

Baud rate	Selectable from 1200, 2400, 4800, 9600, 19200 The baud rate must agree with the baud rate set on the PC/printer.
Flow control ("handshake")	none
Parity	none
Data bits	8
Stop bits	1

#### 3.4.3 USB memory device

Using a USB memory device (such as a USB flash drive), you can

- update the meter software and method data (section 4.20)
- transmit data to the USB memory device (section 4.11 and section 4.12).

USB memory devices are connected to the USB-A interface.



#### Note

Please follow the instructions on using USB memory devices (see section 4.11.2).

#### 3.4.4 PC keyboard

With the PC keyboard it is possible to enter letters, e.g. to assign names for identification (ID).

In addition, the following keys of the PC keyboard are assigned with the following functions of the photometer:

PC keyboard	Photometer
Enter	<start.enter></start.enter>
Esc	<esc></esc>
F1 to F4	Function keys < <b>F1</b> > to < <b>F4</b> >

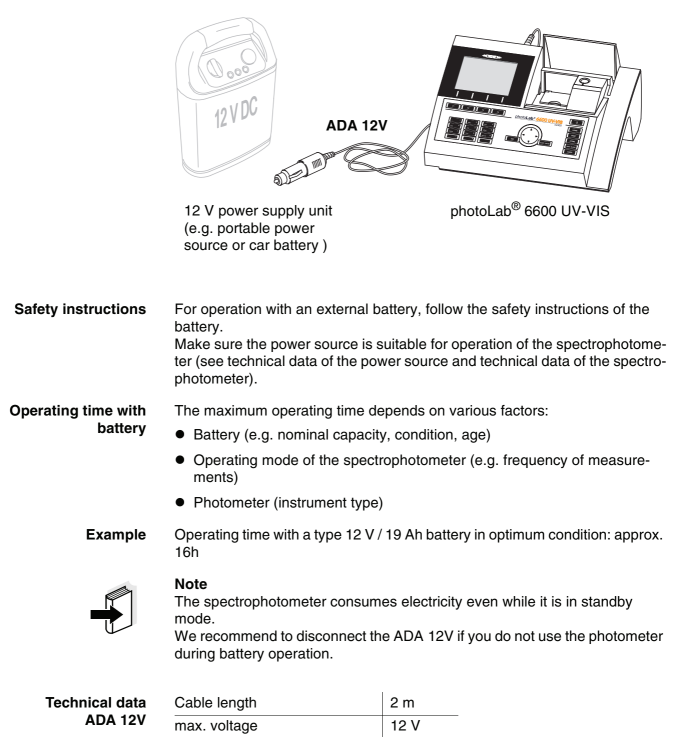
The USB-PC keyboard is connected to the USB-A interface.

#### 3.4.5 Barcode reader

The barcode reader enables the simplified entering of alphanumerical character strings and can be used in all operating situations that require the entry of text or numerals. The barcode reader is connected to the USB-A interface.

#### 3.4.6 Operation with the 12 V adapter ADA 12V

With the ADA 12V you can operate the photoLab<sup>®</sup> 6600 UV-VIS spectrophotometer on the move and independent of the local power supply. To do so, a 12 V power supply such as a commercial 12 V portable power source or a 12 V car battery is required.



8 A

max. current

#### 4 Operation

4.1 Switching on or off the photometer

#### Switching on

Starting the

Self test

Self test

Then press <START/ENTER>

Self test					16.04.07	9:5
Please make sure no	cell is i	nserted	and the	e o	over is clo	sed.
Then press <start e<="" td=""><td>ENTER:</td><td>&gt;</td><td></td><td></td><td></td><td></td></start>	ENTER:	>				
- <b>,</b> ,						
Setup					Info	
Login				I	16.04.07	9:5
Enter user nan	00					
Administrator						
/ annihistrator						

Please make sure no cell is inserted and the cover is closed.

1 Switch the photometer on with <**ON/OFF**>.

The display shows

- the *Self test* dialog (if the user management is not active).

or

 the Login dialog (if the user management is active).

With activated user management:

2 Login

Enter user name and password or register as a guest (see section 4.16.4).

Then the photometer displays the *Self test* dialog.

- **3** Remove all cells and close the cell shaft cover.
- 4 Start the self-test with **START-ENTER**>.

The photometer carries out the self-test.

Self test During the self-test, all cells must be removed and the cell shaft cover closed.

16.04.07 9:52

Self test	I	16.04.07	9:52
Keep cover closed	=1		
System test Filter test Lamp test Wavelength calibration			

The self-test includes:

 the test of the memory, processor, internal interfaces,

filter and lamp

 a calibration for each wavelength

After the self-test is completed, the main menu is displayed.



#### Note

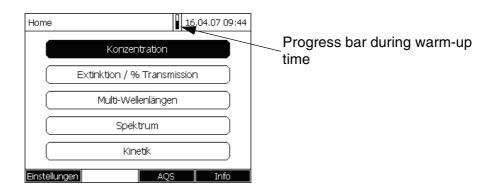
The result of the self-test can be viewed and printed with the *[Info]* function key (see section 4.18).

Warm-up time

After being switched on the photometer requires a warm-up time of 15 minutes. Reproducibility of measurement data is restricted during the warm-up time.

Therefore, do not measure during the warm-up time.

During the warm-up time, a progress bar appears on the display next to the date. The progress bar disappears as soon as the warm-up time is over.



AutoCheck With the AutoCheck function the photometer checks and calibrates the optical measuring unit. The AutoCheck is automatically carried out if measurement settings were changed since the last measurement, e.g.:

- if a different wavelength was selected or
- if a different method was selected.

If necessary, the photometer asks you to remove the cell from the cell shaft.

With unchanged measurement settings, the AutoCheck is carried out in the background at regular intervals of 5 minutes. The AutoCheck can only be carried out in the background if the cell shaft is empty. If a cell is in the cell shaft the AutoCheck is carried out only after the cell was removed.

#### Note

Remove the cell from the cell shaft after every measurement. Thus the photometer can carry out the regular AutoCheck.

Cells must be completely removed from the cell shaft.

Cells that are removed only half disturb the AutoCheck measurement and, as a consequence, falsify measured values until the next AutoCheck is carried out.

Plastic cells that are not recognized by the automatic cell recognition also disturb the AutoCheck.

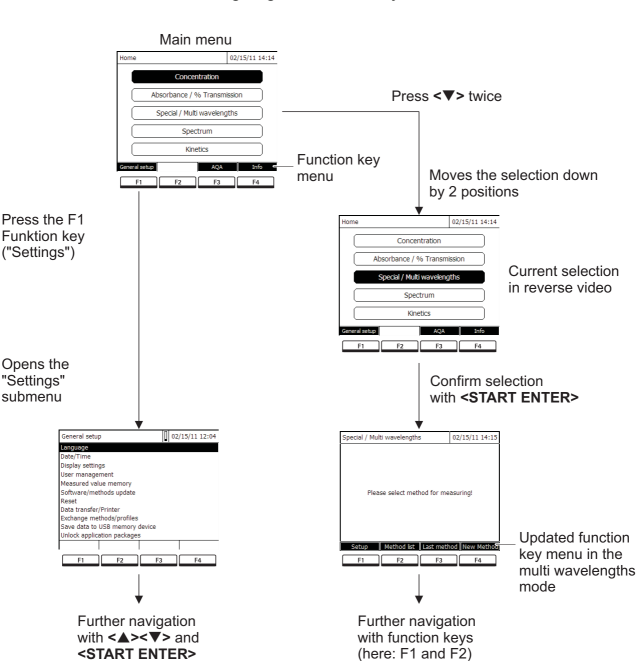


#### Note

During a running kinetic measurement the photometer cannot carry out any AutoCheck. That is why in this case a warm-up time of two hours is required. After this time the signal is stable enough so that the measurement accuracy is secured over a longer period of time.

**Display illumination** The photometer automatically switches off the display illumination if no key has been pressed for 5 minutes. The illumination is switched on again with the next keystroke. The function of the key becomes active only with the following keystroke.

**Switching off** To switch the photometer off, keep the **<ON/OFF>** key depressed until the photometer is switched off.



#### 4.2 General operating principles

#### 4.2.1 Navigating with function keys and menus

figure 4-1 Example of navigation with function keys (left) and "classical" menu navigation (right)

Use of the function keys The function keys F1 to F4 are below the display. Their functions change depending on the operating situation and mode. The current functions are displayed in the function key menu at the bottom edge of the display.

Apart from navigation, the function keys are also used for other operations:

- Opening a selection list or input field
- Executing a command (directly or with intermediate query)
- Switch over between two display options, such as absorbance ↔ transmission

Navigation with arrow keys (<▲><▼>) and <START·ENTER> These operating elements are used to select an item from a menu or list. The current selection is displayed in reverse video. Pressing of **<START·ENTER>** confirms the selection.

Apart from navigation, the **<START·ENTER>** key is also used for other operations:

- Opening a selection list or input field
- Confirming a selection
- Confirming entries of text and numerals
- Executing a command (directly or with intermediate query)
- Activating an item in a selection list ( $\checkmark$  = active)

#### 4.2.2 Display of navigation paths in short form

In this operating manual, the introductory navigation steps leading to individual menus or dialogs are clearly shown in a gray box. The box indicates a section of the menu tree.

Starting point of the description is always the main menu, which can be reached with the **<HOME>** key from any operating situation. From there navigation takes place downward.

Operating example: Navigation to the setting menu for the language The following example shows the elements of the menu tree with the relevant operating steps:

Jage	<home> [General setup]  - Language</home>	<ul> <li>Bold letters and angle brackets indicate a key on the photometer (except function keys).</li> <li>→ Press the "Home" key. The main menu is called up.</li> <li>Square brackets indicate a function key F1 to F4. The text between the brackets corresponds to the assignment according to the function key menu on the bottom edge of the display.</li> <li>→ Press the function key with the assignment "Settings"</li> <li>Text without brackets stands for a menu item indicated on the display (list item).</li> <li>→ Select the menu item with the arrow keys &lt;▲&gt;&lt;▼&gt;. The current selection is displayed in reverse video.</li> </ul>

Further navigation options:

- The **<ESC**> key moves you one level up in the menu tree.
- The **<HOME>** key directly calls up the main menu.



#### Note

If you are "lost" in a menu, press **<HOME>** and restart navigating from the main menu.



#### Note

The complete menu tree is given in the appendix of this operating manual.

#### 4.2.3 Entry of numerals, letters and characters

Numerals, letters, punctuation marks and special characters are entered with the alphanumeric keypad of the meter or using an external keyboard. Entries are required in operating situations such as the following:

- Entering the date and time
- Entering an ID e.g. when storing measurement data
- Selecting a method with the [Search] function
- Programming user-defined methods
- Entering user name and password
- Administrating users

#### **Character set** The following characters are available:

- Numerals 0 ... 9
- Letters A ... Z and a ... z
- Punctuation marks. -
- Special characters ° / + <sup>2</sup> <sup>3</sup> # %

#### **Operating principle** Entering characters is always possible if there is an input field on the display.



The numerals and characters (expect for the small letters) assigned to the keys of the alphanumeric keypad are printed on the keys. Example: With the **<7/PQRS>** key you can enter the following characters: 7, P, Q, R, S, p, q, r, s.

Select the required character by pressing the key several times (similar to a mobile phone). When pressing a key that is assigned to several characters once, the respective numeral appears first. To enter a numeral, one key-pressing is always sufficient.

When pressing the key for the first time a line pops up that displays all characters possible with this key. The currently selected character is highlighted.

A character is taken over in the input field if

- the character is highlighted for more than one second,
- the character is confirmed with <START.ENTER>,
- another alphanumeric key is pressed.



#### Note

During mere number entries (such as entering a wavelength), the keys of the alphanumeric keypad are assigned to the respective numeral only. Each keypressing directly enters the numeral (like a pocket calculator).

#### **Special characters**

**Operating example:** 

Entering the ID

Special characters are entered with the <1/\*> key.

The *Enter ID* input field appears if you press the **<STORE>** key while the storing symbol is visible. In the following example a measurement dataset with the ID "Test" is stored.

<b>E</b>	
Enter ID	
8	
8 TUVtuv	
-	
Enter ID	
Т	
8 T TUVtuv	
Enter ID	
Test_	

1 Press <**8/TUV**> several times until "T" appears in the input line.

Below the input field, a selection line pops up with all characters that are available for this key, e.g. 8 T U V t u v.

The currently selected character is highlighted.

After approx. one second the character is taken over and the selection line closed.

2 Complete the ID with <**A...9**> and confirm.

Correcting incorrect entries Using <<>>, erase all characters until you have reached the incorrect digit and repeat the entry from there.

Home	16.04.07 09:44				
	Konzentration				
	Extinktion / % Transmission				
	Multi-Wellenlängen				
	Spektrum				
	Kinetik				
Einstellun	jen AQS Info				

# General setup 16.04.07 9:52 Language Date/Time Display settings User managementg Measured value memory Software/methods update Reset Data transfer/Printer Exchange methods/profiles Save data to USB memory device Unlock application packages Index set

Language	16.04.07 9:52			
Deutsch ✓				
English				
Français				
Español				
Italiano				
Bulgarian/Български				
Česko				
Chinese/ 中文				
Traditional Chinese/ 繁體中文				
Greek/Ελληνικά				
Indonesian/Indonesia				

4.2.4 Detailed operating example: Changing the language

- 1 Call up the main menu with the **<HOME>** key.
- 2 Open the *General setup* menu with the F1 function key [Setup].
- 3 Using <▲><▼>, select the *Language* menu item and open with <**START**·ENTER>.

The *Language* menu shows a list with the available languages. The currently active language is marked by a check.

4 Select the required language from the list with <▲><▼> and confirm with <**START**.ENTER>.

The selected language is taken over immediately. The photometer moves up one menu level.

#### 4.3 Photometer settings and system administration

The general photometer settings are done in the **<HOME>** -> *General setup* menu. The general photometer settings comprise:

- Language (see section 4.3.1)
- Date/time (see section 4.3.2 and section 4.2.4)
- Display characteristics (see section 4.3.3)
- User management (see section 4.16)
- Administration of the measurement data memory (see section 4.11)
- Software and method update (see section 4.20)
- Reset of the settings to default values (see section 4.17)
- Settings for data transmission (see section 4.14.2)

#### 4.3.1 Language

The complete list of the available instrument languages is given in the *Language* chapter 7 TECHNISCHE DATEN menu of the photometer.



#### Note

If you want to set some special languages on your photometer (e.g. Chinese or Thai), a character set extension is required to display the characters (see section 4.20.3).



#### Note

How to set the language is described in detail in the operating example in section 4.2.4.

# 4.3.2 Date/Time

The date format is set automatically with the language setting. According to the locally usual version, the date format is displayed in the order, Day.Month.Year (*DD.MM.YY*) or Month/Day/Year (*MM/DD/YY* or *MM.DD.YY*).

	The <i>Date/Time</i> menu is open.
1	Select and confirm <i>Date</i> . The input field for the current date pops up.
2	Enter the current date with <b>&lt;09</b> > and confirm. The input field closes. The date is accepted.
3	Select and confirm <i>Time</i> . The input field for the current time pops up.
4	Enter the current time with <b>&lt;09</b> > and confirm.
	The input field closes. The time is accepted.
	3

# 4.3.3 Display settings

Here you can adjust the display contrast to the lighting conditions.

<HOME> [General setup] – Display settings

Display setting	s		16.04.07	9:52
Contrast			50 %	
1	r	7	1	
1			'	

- Select and confirm Contrast.
   A slide control for the display contrast appears.
- 2 Using <◀><►>, set the display contrast and confirm.

#### 4.4 Zero adjustment

A valid zero adjustment is required for the calculation of measured values in the modes, *Concentration, Absorbance / % Transmission, Special / Multi wavelengths* and *Kinetics*. With a zero adjustment, the absorbance of a cell filled with distilled water ("zero cell") is measured and stored.

Factory zero adjustment for concentration measurements For all measurements with WTW test sets (*Concentration* mode), a factory zero adjustment is available in the delivery condition. We recommend replacing it with a zero adjustment of your own. If a zero adjustment exists already for a method, the date and time of the last zero adjustment are displayed in the top right area of the display.

Concentration			16.04.07 9:52	
[ZERO 11.11.2010 11:11]				
or insert	elect method fo a barcoded ce AutoSelector.	5		
Setup	Method list	Last method	New Method	

### Zero adjustment for absorbance measurements

In the *Absorbance* mode, the zero adjustment has to be carried out separately for each cell type and each used wavelength. If a zero adjustment exists already for the inserted cell type at the selected wavelength, the date and time of the last zero adjustment are displayed in the top right area of the display.

Absorbance			16.04.07 9:52
		[ZERO 11	.11.2010 11:11]
	: measurement, ell or press <st< td=""><td></td><td></td></st<>		
525 nm	I	I	10 mm
Setup	Wavelength	Transmission	Reference

If no zero adjustment is available, the photometer will prompt you to carry out a zero adjustment.



#### Note

The cells must be absolutely clean and free of scratches.

Always use a cell of the same type for zero adjustment and measurement of the sample.

Notes on zero adjustment Zero adjustment with round cells:

- Only use clean, scratch-free round cells with distilled water. The minimum filling level is 20 mm. A ready zero cell is included in the scope of delivery of the photometer and PhotoCheck (see chapter 8 ZUBEHÖR, OPTIONEN).
- A ready zero cell can, in principle, be used any number of times. We recommend, however, to regularly check the zero cell for visible contamination and scratches and refill or exchange it if necessary (at least every 24 months).

Zero adjustment with rectangular cells:

- For rectangular cells, the zero adjustment must be carried out with the same cell type (manufacturer and glass type [e.g. optical glass, quartz glass]) that is used for measurement. This is important because cells of different manufacturers have a different absorption behavior. When changing the cell type repeat the zero adjustment with the new type.
- Prior to zero adjustment, clean the rectangular cell and fill it with distilled water. The minimum filling level is 20 mm.
- Rectangular cells always have to be inserted in the cell shaft with the same orientation for measurement and zero adjustment (e.g. cell printing on the left side ).

#### Note

Ordering information is given in chapter 8 ZUBEHÖR, OPTIONEN. The cells listed in the chapter 8 ZUBEHÖR, OPTIONEN are especially adapted to the WTW test set program. General requirements of the cells are given in chapter 7 TECHNISCHE DATEN. Note that the spectral transparency of the cell must be suitable for the intended application (example, quartz cell for UV range).

The zero adjustment takes place similarly in the *Concentration, Absorbance* / % *Transmission, Special / Multi wavelengths* and *Kinetics* modes.

1

ment.

Carrying out a zero adjustment

Concentration			16.04.07 9:52
Adjust	t		
Blank	value		
Zero a	adjustment		
3: A6/25			NH4-N
16 mm		(	0.20 - 8.00 mg/l
Setup	Method list	Citation form	Unit

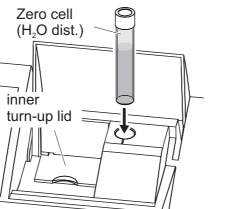
- Zero adjustment 16.04.07 9:52
  Please insert zero cell (distilled water)
  or press <START/ENTER>
- The zero adjustment window pops up.

In the respective mode, press the

Select and confirm Zero adjust-

<ZERO·BLANK> key.

2 In *Concentration* mode only:

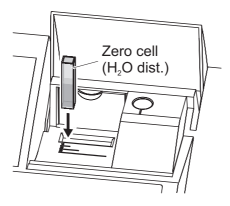


- **3** Close the inner turn-up lid.
- 4 Depending on the cell type, insert the zero cell as follows:

Round cell:

Insert the round cell in the round cell shaft so it touches the bottom.

If the inner turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.



Zero adjustmer	nt	16.04.07	9:52
	Zero adjustment successful		
		1	.0 mm
		OK	

Rectangular cell:

Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

The photometer automatically starts the zero adjustment and subsequently stores the value.

5 After a successful zero adjustment switch to measurement with [OK].

# Validity of the zero adjustment

The data of the zero adjustment is stored in the photometer separately for each cell type. As long as the data is valid, it is automatically used again after a temporary change to a different cell type. The validity depends on the respective mode:

Mode	Validity of the zero adjustment
<i>Concentration</i> (permanently programmed methods)	<ul> <li>Till the next zero adjustment</li> </ul>
Absorbance / % Transmission	<ul> <li>Till the next zero adjustment with the same wavelength *</li> </ul>
<i>Concentration</i> (user-defined methods) and	<ul> <li>Till the next zero adjustment for the same method *</li> </ul>
Special / Multi wavelengths	
Kinetics	Till another kinetic profile is loaded
	• Till the <i>Kinetics</i> mode is exited or the pho- tometer is switched off

\* After the wavelength or method respectively was temporarily exited the photometer displays that a zero adjustment is available and the time it was carried out. You can then decide whether to use this zero adjustment or carry out a new zero adjustment.

We recommend to repeat the zero adjustment in the following cases:

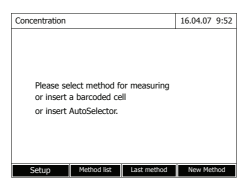
# When to repeat the W zero adjustment?

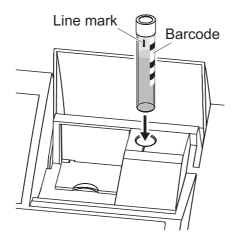
- If the photometer was subject to mechanical stress such as strong shock or transport
- If the ambient temperature changed by more than 5 °C since the last zero adjustment
- At least once per week
- If a new cell type (different manufacturer, different glass type is used)
- Basically each time you want to measure with the highest possible accuracy.

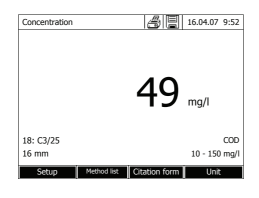
# 4.5 Measuring in *Concentration* mode

#### 4.5.1 Measuring cell tests with barcode

# <HOME> Concentration







Inserting a cell with barcode starts a measurement.

- **1** Open the cell shaft cover.
- 2 Close the inner turn-up lid.

If the inner turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.

3 Insert the barcoded round cell in the round cell shaft so it touches the bottom. When doing so, align the line mark with the notch at the front of the round cell shaft.

The photometer selects the method based on the bar code and automatically starts measurement.

- 4 Further options:
  - Select a different citation form with [Citation form], (e.g. NH<sub>4</sub> <-> NH<sub>4</sub>-N).
  - Select a different measuring unit with [Unit], (e.g. mg/l <-> mmol/l).
  - Make further settings such as dilution or blank value measurements with [Setup] (see section 4.5.6).

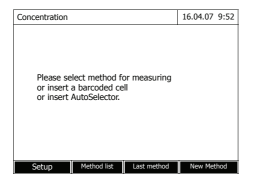
Display if the measured value is not within the measuring range (see section

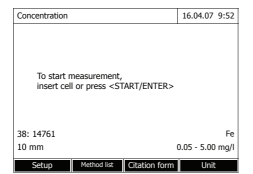
4.5.4).

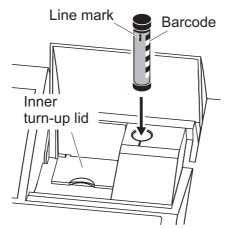
#### 4.5.2 Measuring reagent tests with AutoSelector

#### <HOME>

#### Concentration



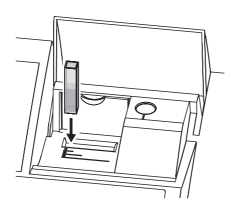




The method is selected by inserting the AutoSelector.

The photometer is ready to measure.

- **1** Open the cell shaft cover.
- 2 Insert the AutoSelector in the round cell shaft so it touches the bottom. When doing so, align the line mark with the notch at the front of the round cell shaft.
  - The photometer selects the correct method with the aid of the barcode.



- **3** Open the inner turn-up lid.
- 4 Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The correct measuring range is automatically selected when the rectangular cell (1, 2, 5 cm) is inserted.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

The photometer starts measuring automatically.

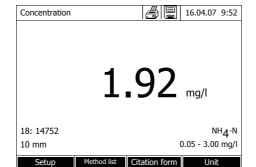
- **5** Further options:
  - Select a different citation form with [Citation form], (e.g. NH<sub>4</sub> <-> NH<sub>4</sub>-N).
  - Select a different measuring unit with [Unit], (e.g. mg/l <-> mmol/l).
  - Make further settings such as dilution or blank value measurements with [Setup] (see section 4.5.6).

Display if the measured value is not within the measuring range (see section 4.5.4).

# 4.5.3 Measuring reagent-free tests and user-defined methods

User-defined methods and reagent-free methods normally do not have a barcode and therefore, no automatic method recognition. In such a case, select the method manually:

<HOME> Concentration



Concentration	16.04.07	9:52
Please select method for measuring or insert a barcoded cell or insert AutoSelector.		
Setup Method list Last method Concentration	New Met	
To start measurement, insert cell or press <start enter=""></start>		
3: A6/25 16 mm	Ni 0.20 - 8.00	H <b>4</b> -N mg/I
Setup Method list Citation form		5,

inner turn-up lid 1 Select the method manually (see section 4.5.5).

The photometer is ready to measure.

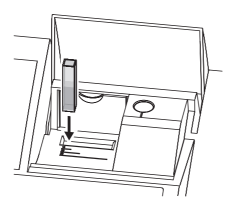
**2** Depending on the type, insert the cell as follows:

#### Round cell:

Close the inner turn-up lid.

Insert the round cell in the round cell shaft so it touches the bottom.

If the turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.



Concentration		<b>A</b>	16.04.07 9:52
	0.6	529	mg/l
1001: Nitrite 10 mm		0.0	NO2-N 150 - 1.000 mg/l
Setup	Method list	Citation form	Unit

Rectangular cell:

Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

- **3** Further options:
  - Select a different citation form with [Citation form], (e.g. NH<sub>4</sub> <-> NH<sub>4</sub>-N).
  - Select a different measuring unit with [Unit], (e.g. mg/l <-> mmol/l).
  - Make further settings such as dilution or blank value measurements with [Setup] (see section 4.5.6).

Display if the measured value is not within the measuring range (see section 4.5.4).

# 4.5.4 Exceeding the upper or lower limits of the measuring range

Measured value display if the measured value is outside the measuring range:

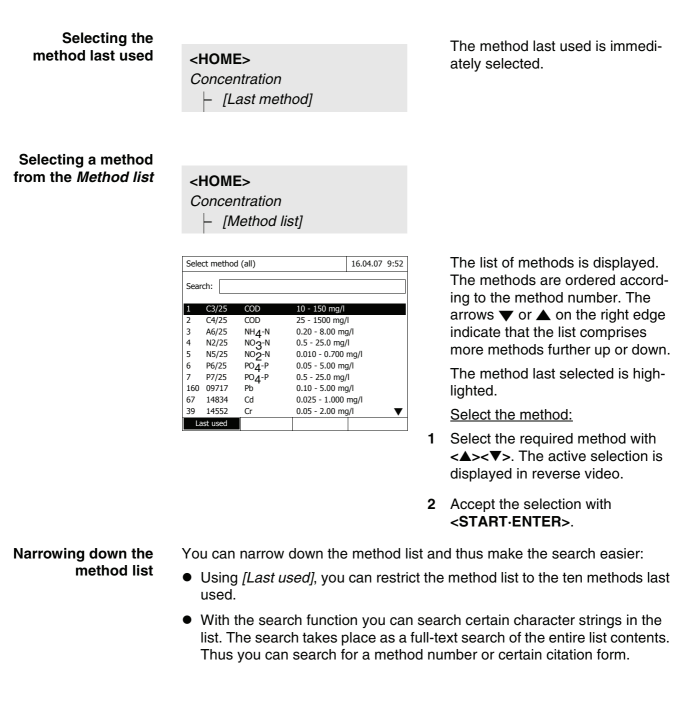
Ra	nge	Display	Example: MR: 10 - 150 mg/l
	LL < MV < UL	Measured value	128 mg/l
1	UL < <b>MV</b> < UL + 10%	Upper limit of measuring range exceeded by up to 10% and measured value	> 150 157 mg/l
	LL - 50% < <b>MV</b> < LL	Lower limit of measuring range undercut by up to 50% and measured value	< 10 7 mg/l
2	<b>MV</b> > UL + 10%	Upper limit of measuring range exceeded by more than 10%	> 150 mg/l
	<b>MV</b> < LL - 50%	Lower limit of measuring range undercut by more than 50%	< 10
3	Invalid measured value	Bars	
	e.g. <b>MV</b> < 0		mg/l

MR = Measuring range

UL = Upper limit value of the measuring range

LL = Lower limit value of the measuring range

MV = Measured value



# 4.5.5 Selecting a method manually

#### **Search function**

Select method (last used) 16.0			) 16.04.07 9:52
CO			
14	14540	COD	10 - 150 mg/l
23	14541	COD	25 - 1500 mg/l
All	methods		

Search for a character string:

Enter the character string to be searched for in the search window with **<A...9>**.

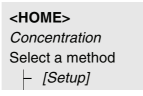
The list appearing below shows all hits containing the character string. The hit list is updated with each character that is entered.

#### Note

Note the case sensitivity when searching. It is not required or possible to enter inferior characters. When searching for chemical formulas, inferior characters are treated as normal characters. Example: The search for "NH4" shows all hits that contain "NH4" as well as "NH<sub>4</sub>".

#### 4.5.6 Settings for *Concentration* mode

Prior to measuring, check the settings for the selected method.



Concentration 16.04.07 9:52				
Dilution 🗸				
Sample blank value				
User-defined blank value				
Turbidity correction				
Display absorbance 🗸	Display absorbance 🗸			
AQA				
Edit method				
New method				
Measurement data memory				

The menu shows an overview of all settings.

Active settings are marked by a check.

Overview of the	Menu item	Explanation
settings	Dilution	Here you can set the dilution prior to measuring if you want to use a diluted sample.
		In the measured value display, the dilution is indi- cated in the form $[1 + x]$ (parts sample + parts dis- tilled water).
		For more detailed information on dilution, see section 4.5.7.
	Sample blank value	Here you can measure while taking a sample blank value into account.
		In the measured value display, measurements with sample blank value are marked by [SB] (Sample blank).
		For more detailed information on sample blank value, see section 4.5.8.
	User-defined blank value	If available, a user-defined reagent blank value is used.
		In the measured value display, measurements with a user-defined reagent blank value are marked by [BV/Lot number].
		For more detailed information on reagent blank value, see section 4.5.9.
	Turbidity correction	Activates/deactivates the automatic turbidity cor- rection.
		In the measured value display, measurements with automatic turbidity correction are marked by [TURB].
		For more detailed information on the automatic tur- bidity correction, see section 4.5.10.
	Display absorbance	Activates/deactivates the display of the absor- bance value in addition to the main measured value.
	AQA	Here you can view and change the AQA settings without discarding the current measurement.
	Edit method	Here you can edit user-defined methods.
	New method	Here you can create user-defined methods.
	Measurement data memory	Here you can view the measurement data memory.

#### 4.5.7 Measuring diluted samples

If the concentration of a sample exceeds the measuring range of a method, you can specifically dilute the sample so that the concentration of the diluted sample is in the measuring range of the method. Thus a valid measurement is possible.

After entering the factor for the dilution the meter converts the concentration to that of the undiluted sample.



#### Note

Optimum measurement results are achieved if the concentration of the diluted sample is in the middle of the measuring range of the method after diluting.

Setting the dilution

# <HOME>

Concentration

Concentration	16.04.07	9:52
Please select method for measuring		
or insert a barcoded cell		
or insert AutoSelector.		
Setup Method list Last method	New Met	hod
Concentration	16.04.07	9:52

Inserting a cell with barcode starts a measurement.

If a cell without barcode is used: Select the method manually (see section 4.5.5).

The photometer is ready to measure.

Concentration			16.04.07 9:52
	neasurement,		
insert cell or press <start enter=""></start>			
3: A6/25			NH4-N
16 mm			0.20 - 8.00 mg/l
Setup	Method list	Citation form	Unit

Sample + distilled water	
1 + _	
3: A6/25	<sup>NH</sup> 4 <sup>-N</sup>
16 mm	0.20 - 8.00 mg/l
Setup Method list Citation f	form Unit

- 1 Open the setting menu with [Setup].
- 2 Select and confirm *Dilution*. The input field for the dilution pops up.
- **3** Enter and confirm the dilution (**<0...9**>).

The entered dilution is taken into account with the next measurement.

The entered value for the dilution factor is valid for the selected method only. The dilution factor is erased if

- the photometer is switched off
- a different method is selected
- the factor 0 is entered in the *Dilution* menu.

If a dilution factor is active, it is indicated on the display during measurement in the form [1 + x].

#### 4.5.8 Sample blank value

By measuring and using a sample blank value, measurement errors due to coloring and turbidity of the sample matrix can be eliminated to a large extent.

The sample blank value is a characteristic of the sample (coloration) to be currently determined. It is determined by measuring the blank sample.

The blank sample is prepared from the sample by adding reagents to it and is especially adapted to measurement with the test to be measured. No chromophoric reagents, however, are added for the preparation of the blank sample.



#### Note

Due to the addition of reagents the sample is diluted. This can also change the pH value of the sample. For this reason the blank sample also has to be diluted and the pH value adjusted accordingly.

Validity The sample blank value applies to the next measurement only.

Single and multiple determination

The sample blank value can be determined by single or multiple determination. With multiple determination, the sample blank value is calculated as the median from the individual measured values.

Measuring the sample blank value

<home></home>
Concentration

Concentration			16.04.07	9:52
or insert	lect method fo a barcoded ce AutoSelector.	NI		
Setup	Method list	Last method	New Met	thod

Inserting a cell with barcode starts a measurement.

If a cell without barcode is used: Select the method manually (see section 4.5.5).

Concentration	16.04.07	9:52
To start measurement, insert cell or press <start enter=""></start>		
3: A6/25	N	H4-N
16 mm (	0.20 - 8.00	mg/l
Setup Method list Citation form	Unit	

Sample blank v	16.04.07	9:52		
	neasurement,   or press <st< th=""><td>ART/ENTER&gt;</td><td></td><td></td></st<>	ART/ENTER>		
3: A6/25 16 mm			N 0.20 - 8.00	H <b>4</b> -N ) mg/l

Sample blank	Sample blank value		16.04.07 9:52
	Last measure	d absorbance	
	0.115		
	Median		
	0.115 (1 M	1easuremen	t(s))
3: A6/25 16 mm			NH <b>4</b> -N 0.20 - 8.00 mg/l
Next meas.	Discard		Apply

Concentration			16.04.07	9:52		
				[SB]		
	To start measurement, insert cell or press <start enter=""></start>					
3: A6/25			N	H <b>4</b> -N		
16 mm		(	0.20 - 8.00	mg/l		
Setup	Method list	Citation form	Unit			

The photometer is ready to measure.

- 1 Open the setting menu with [Setup].
- 2 Select and confirm *Sample blank value*.
- **3** Insert the cell with a suitable blank sample.

The first single measurement for the sample blank value takes place.

The following data is displayed as the result:

- The measured absorbance from the (last) single measurement.
- The median from all single measurements carried out up to now.
- 4 If necessary, carry out further single measurements for the formation of the median with [Next meas.] or

discard the last single measurement with [Discard].

5 To accept the median value, press [Apply].

The photometer is ready to measure.

The use of the sample blank value is indicated by [SB] in the top right corner of the display.

#### 4.5.9 Reagent blank value

The evaluation of the photometric measurement always refers to the comparison value of a test sample without the substance to be determined (reagent blank value). Thus the influence of the basic absorbance of the reagents on photometric measurement is compensated for.

In practice, the reagent blank value is measured with the same amount of deionized water instead of sample.

Factory and userdefined reagent blank values With photometric concentration determination, the reagent blank value is a constant. The method data for all measurements with WTW test sets (*Concentration* mode) include an exactly determined reagent blank value. This value is overwritten if you measure the reagent blank value yourself (setting, *User-defined blank value*, see section 4.5.6).



Note

You can increase accuracy if you determine the reagent blank value with a test of a new lot and use the reagent blank value for all further measurements with this lot. This is especially recommended for measurements in the vicinity of the lower limit of the measuring range. To be able to attribute the reagent blank value in the measured value documentation later, you can enter the lot number of the reagent package (*Lot number*) during the blank value determination.

Validity The factory blank values always remain stored in the meter and can be activated at any time. The reagent blank values you measured yourself also remain stored in the meter until they are overwritten by a new blank value measurement.

#### **Single and multiple** determination The reagent blank value can be determined with single or multiple determination. With multiple determination, the reagent blank value is calculated as the median from the individual measured values.

User-defined For user-defined methods, you can activate the reagent blank value function as follows only:

Entry type	Function type	Reagent blank value possible?
Entry of a function	Linear	Yes
(with and without entering the ordi- nate intercept)	Nonlinear	No
Entry of value pairs or measure-	Linear	Yes
ment and storage of standard solu- tions	Parabola (second-order function)	Yes
(with entering/measuring and stor- ing E0)	Polygon line	No
Entry of value pairs or measure-	Linear	Yes
ment and storage of standard solu- tions (without entering/measuring and storing E0)	Parabola (second-order function) Polygon line Polygon line through zero	No

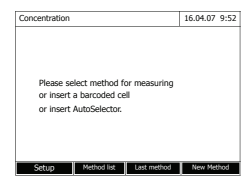


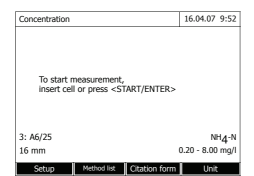
#### Note

If no value for E0 is stored during the entry of value pairs or the measurement and storing of standard solutions for a nonlinear function (parabola or polygon line), the message, *No blank value correction is intended for this method.* appears when the *User-defined blank value* function is activated. The blank value (E0) can be entered later by editing the method.

# Measuring the reagent blank value

<HOME> Concentration





Concer	Concentration			16.04.07 9:52
	Adjust			
Zero adjustment				
	Blank va			
3: A6/	25			NH4-N
16 mm	ı		1	0.20 - 8.00 mg/l
Se	tup	Method list	Citation form	Unit

Inserting a cell with barcode starts a measurement.

If a cell without barcode is used: Select the method manually (see section 4.5.5).

The photometer is ready to measure.

- 1 Using <**ZERO·BLANK**>, open the *Adjust* selection list.
- 2 Select and confirm *Blank value*.

The window for the measurement of the reagent blank value pops up.

The data of the last measurement appears in the measured value display.

Blank value	16.04.07 9:52
To start measurement, insert cell or press <sta< td=""><td>RT/ENTER&gt;</td></sta<>	RT/ENTER>
3: A6/25	NH4-N
16 mm	0.20 - 8.00 mg/l

Blank value			16.04.07 9:52
	Last measure	d absorbance	
	0.600		
	Median		
	0.600 (1 M	1easuremen	t(s))
3: A6/25			NH4-N
16 mm			0.20 - 8.00 mg/l
Next meas.	Discard		Apply

Blank value	16	6.04.07	9:52
	[BV/	'Lot num	nber]
To start measurement insert cell or press <s< td=""><td></td><th></th><th></th></s<>			
3: A6/25		NF	4-N
16 mm	0.20	0 - 8.00	mg/l
Setup Method list	Citation form	Unit	

**3** Insert the cell with the blank sample.

The first single measurement for the reagent blank value takes place.

The following data is displayed as the result:

- The measured absorbance from the (last) single measurement.
- The median from all single measurements carried out up to now.
- 4 If necessary, carry out further single measurements for the formation of the median with [Next meas.] or

discard the last single measurement with [Discard].

5 To accept the median value, press [Apply].

The *Lot number* entry field pops up.

6 Enter and confirm the *Lot number* (**<A...9>**). The blank value measurement is

completed.

The photometer is ready to measure.

The use of the reagent blank value is indicated by [BV/Lot number] in the top right corner of the display.

#### 4.5.10 Automatic Turbidity correction

The *Turbidity correction* function activates the automatic recognition and compensation of the light absorption caused by turbid substances.

After activating the function remains permanently switched on. Measured values that were measured with *Turbidity correction* are labeled with [TURB] (turbidity correction) on the display and in the documentation (printout and memory).

The *Turbidity correction* function is not active in the delivery condition.



#### Note

The setting for automatic turbidity correction is used with all methods where the automatic turbidity correction makes sense. The photometer automatically decides whether or not to use the function.

Switching on the turbidity correction

The automatic turbidity correction is activated and deactivated in the setting menu of the concentration measurement (see section 4.5.6 SETTINGS FOR Concentration MODE).

#### 4.5.11 Programming / modifying user-defined methods

**Overview** For *Concentration* mode, you can develop and store yourself user-defined methods under the method numbers 1001 to 1100. The photometer software supports you when creating the methods.

**Calibration data and calibration function** In photometry, the calibration function describes the dependency between the measured parameter (e.g. concentration) and the photometric measurement result (e.g. absorbance) of a sample. The knowledge of this dependency is a prerequisite for the development of a photometric method. The calibration function is usually determined by means of a series of measurements with standard solutions of known concentrations (nominal value), e.g. a 10-point calibration.



#### Note

In measuring operation, the reverse calibration function is used to output the measured absorbance as a concentration value.

Line types

The dependency between the nominal value and absorbance is often linear in a wide range as shown in the following example:

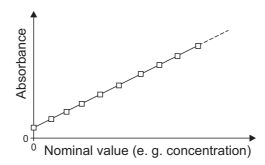
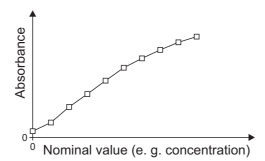


figure 4-2 Example of a linear calibration function after a 10-point calibration

In the case of a linear dependency, the calibration function is determined by means of linear regression. The slope and axis intercept (E0) are the characteristics of the calibration line.

In the case of a nonlinear dependency, the points of the measuring ranges can be connected to each other as a polygon line or approximated as a parabola:





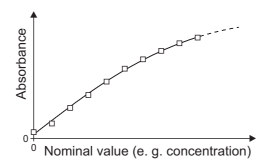


figure 4-4 Example of a parabola calibration function after a 10-point calibration

Determining the calibration function

You have the following options to create a method:

#### Measure and store:

Carry out a series of measurements with the following sample solutions while at the same the photometer takes over the values:

- Blank sample to determine the reagent blank value (with deionized water instead of sample, see section 4.5.9)
- at least one, up to ten standard solutions in different concentrations.

The photometer stores nominal value/absorbance value pairs of the individual measurements and determines the resultant characteristics of the calibration. When doing so, you can select the following line types: *Polygon line, Straight line* or *Parabola*.

#### • Enter as value pairs:

Entry of the value pairs, Nominal value (concentration) / Measured absorbance of an <u>already available</u> test series with the following sample solutions:

- Blank sample to determine the reagent blank value (with deionized water instead of sample, see section 4.5.9)
- at least one, up to ten standard solutions in different concentrations.

Based on the entered value pairs, the photometer determined the characteristics for the calibration. When doing so, you can select the following line types: *Polygon line, Straight line* or *Parabola*.

#### • Enter a function:

Entry of a function to calculate the concentration from the absorbance (reverse calibration function). You can enter on the photometer the coefficients of a polynomial equation of the following type:

$$c = a0 + a1 \cdot A + a2 \cdot A^2 + a3 \cdot A^3 + a4 \cdot A^4 + a5 \cdot A^5$$

with:

С	Measurement result, e.g. concentration
a0 to a5	Coefficients (input range 0.000 to 1000,000)
А	Absorbance



#### Note

Entering the formula is especially simple if you measure with a commercial test set for which the manufacturer has given the value for the coefficients a1. It is often called the "Factor" and corresponds to the reciprocal value of the slope of the straight line of the calibration function.

If a <u>linear</u> function (straight line) should be entered, it is necessary to enter the coefficients a0 and a1 to receive correct measured values. If the exact value for a0 is not known at the time the formula is entered, it is sufficient to enter the coefficient a1. In this case, the *User-defined blank value* function (in the *Concentration / Setup* menu) has to be activated to measure with this method.

Prior to measuring with this method, a blank value measurement has to be carried out. This procedure determines the value for a0, which then replaces the value from the programming of the method.

If the *User-defined blank value* function is not activated, the photometer uses the value zero for the coefficient a0.

More information on the entry of the formula (determination of coefficients)

Linear func- tion	If the value for a1 (slope of the reverse calibration function) is unknown, you can very simply program the method in the photometer by measuring/storing or entering the value pairs (see above).
	For entry as a formula, you can determine the coefficients of the reverse calibration function by linear regression. When doing so, the concentration has to be on the Y axis and the absorbance on the X axis.
	In the case of a linear function, the coefficients of the reverse calibration function can also be determined from the determined reagent blank value and the slope (m) of the calibration function (Y axis = absorbance, X axis = concentration). Proceed as described below.
	Explanation of the coefficients of the formula:
	<ul> <li>a0 = - E0*a1</li> <li>[E0 = reagent blank value (absorbance at concentration 0)]</li> </ul>
	<ul> <li>a1 = 1/m Reverse value of the slope of the calibration function (often referred to as "Factor") m = slope of the calibration function</li> </ul>
	<ul> <li>a2, a3, a4, a5 = further coefficients (when entering a linear function: zero)</li> </ul>
Nonlinear function	The coefficients of the reverse calibration function are determined by multiple regression. When doing so, the concentration has to be on the Y axis and the absorbance on the X axis.

# Further method data

Input field	Possible entries	
Number*	1001 1100	
Designation	Any name (max. 18 characters)	
Version	Any version designation (max. 18 characters)	
Wavelength*	Freely selectable (in nm)	
Cell*	16 (round), 10, 20 or 50 mm	
Citation form	e.g. PO4-P (max. 18 characters)	
Unit**	e.g. mg/l (max. 18 characters)	
Resolution*	0.001, 0.01, 0.1 or 1	
Lower and upper limit of the measuring range *	Any value between zero and the highest concen- tration of the used standard solutions	
Timer 0 to 3	Up to four analysis timers freely adjustable	
AQA2 target value	Any value within the measuring range	
AQA2 tolerance	Any	

\* necessary inputs \*\* default: mg/l

How to program user-defined methods

# <HOME> Concentration [Setup] [- New method

Edit method	16.04.07 9:52
Number	1001
Designation	Nitrite
Version	01
Wavelength	525
Cell	10 mm
Citation form	NO2-N
Unit	mg/l
Resolution	0.001
Calibration curve	Measure standard solutions
Method list Delete	Next

1 Enter the general method data here. The next available method number is already entered as the number.

You have the following options when filling out the input fields:

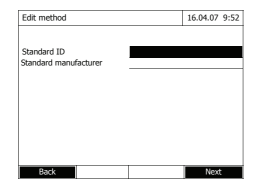
- Fill out all empty input fields one after the other
- Using *[Method list]*, select an already existing method as a model, give it a new method number and adjust the entries
- Using [Method list], select an existing method in order to change it (without changing the number).
- You can delete the method completely with [Delete].
- 2 Select the menu item, *Calibration curve*. Select the method for the determination of the calibration line. The following variants can be selected:
  - Measure standard solutions
  - Enter value pairs
  - Enter formula
- **3** Using *[Next]*, accept all entries on the page and switch to the next page.



# Note

During the following proceeding, you can return to the previous page at any time with *[Back]*, e. g. if you want to correct entries, add further value pairs or eliminate outliers.

#### Variant 1: Measure standard solutions



Edit method			16.04.07 9:52
	Target value		Absorbance
E0	0.000		
1			
Back	Add	Delete	Next

Edit meth	nod		16.04.07 9:52
	Target value		Absorbance
E0	0.000		
1	0.300		
2	0.600		
3	1.000		
Back	Add	Delete	Next

Edit meth	od	16.04.07 9:52
	Target value	Absorbance
E0	0.000	
1	0.300	
2	0.600	
3	1.000	
Back	Add Delet	e Next

- 1 Select and confirm *Measure standard solutions*.
- 2 Enter and confirm details of the standard solutions (optional).
- **3** Using *[Next]*, accept all entries on the page and switch to the next page.

The table for the measurement of standard solutions pops up.

In the first two lines of the table, the two value pairs (measuring points) that are at least required for a calibration are already prepared (reagent blank value E0 and any further nominal value).

4 Create further values pairs with [Add] as necessary.

You can delete a highlighted value pair with [Delete].

5 In the *Target value* column, enter the nominal values of the individual standard solutions.

Measuring the standard solutions:

6 Using the arrow keys <▲><▼> and <◀><▶>, navigate to the relevant input field in the Absorbance column and press <START.ENTER>.

Absorbance E0	16.04.07 9:52		The measurement display
			appears.
To start measurement,		7	Insert the cell with the respective standard.
insert cell or press <start enter=""></start>			The absorbance is measured. The result of the first single measurement is displayed.
525 nm	16 mm		
Absorbance E0	16.04.07 9:52	8	If necessary, carry out further sin-
			gle measurements for the forma- tion of the median with <i>[Next</i>
Last measured absorbance			meas.]
0.009 Median			or
			discard the last single measure-
0.009 (1 Measuremer	it(S))		ment with [Discard].
525 nm	16 mm	9	To accept the median value, press
Next meas. Discard	Apply		[Apply].



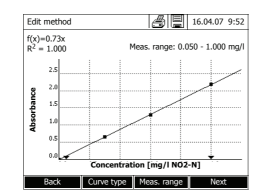
#### Note

If the zero standard concentration (reagent blank value E0) is not measured and stored, the photometer calculates the calibration line without this value. If the *User-defined blank value* function (in the *Concentration / Setup* menu) is activated for measuring with this method, the value for a0 is determined and replaces the calculated axis intercept from the programming of the method.

Edit metho	d		16.04.07 9:52
	Target value		Absorbance
E0	0.000		0.009
1	0.300		0.664
2	0.600		1.292
3	1.000		2.178
Back	Add	Delete	Next

- **10** Repeat the steps 6 to 9 until all input fields in the *Absorbance* column are filled out.
- **11** Using *[Next]*, accept all entries on the page and switch to the next page.

The value pairs are displayed in a diagram (standard: Polygon line).



Edit method

Timer 0

Timer 1

Timer 2

Timer 3

AQA2 target value

AQA2 tolerance

Back

The related formula f(x) and correlation coefficient  $R^2$  are displayed above the diagram.

- **12** If required, select a different line type for the line adjustment with *[Curve type]*.
  - Polygon line
  - Straight line
  - Parabola
- **13** If required, enter different measured value limits with *[Meas. range]*.
  - Lower limit
  - Upper limit
- **14** Using *[Next]*, complete the editing of the calibration line and proceed to the next page.

The timers and AQA2 data linked to the method are displayed.

- **15** If necessary, enter intervals for up to 4 timers.
- **16** If necessary, enter the AQA2 target value and AQA2 tolerance.
- **17** Complete the programming of the method with [Complete].

The method is programmed and selected for measuring.

# Variant 2: Enter value pairs

Unlike variant 1, the fields of the *Absorbance* column are filled out manually here. Accordingly, the steps 6 to 10 are not applicable here. Apart from that, the proceeding is identical to variant 1.

16.04.07 9:52

00:00:00

00:00:00

00:00:00

00:00:00

1.00 mg/l

0.10 mg/l

Complete

Variant 3: Enter formula

0.605
1,000 mg/
3.000 mg/

1 Select and confirm *Enter formula*.

Input fields for the coefficients (a0 ... a5) of the formula are displayed.

2 Enter and confirm the factors.

If no value is entered for a coefficient the photometer automatically uses the value 0.



#### Note

Entering the formula is especially simple if you measure with a commercial test set for which the manufacturer has given the value for the coefficients a1. It is often called the "Factor" and corresponds to the reciprocal value of the slope of the straight line of the calibration function.

If a linear function (straight line) should be entered, it is necessary to enter the coefficients a0 and a1 to receive correct measured values. If the exact value for a0 is not known at the time the formula is entered, it is sufficient to enter the coefficient a1. In this case, the *User-defined blank value* function (in the *Concentration / Setup* menu) has to be activated to measure with this method. Prior to measuring with this method, a blank value measurement has to be carried out. During this procedure the value for a0 is determined and replaces the previous value.

Edit method	16.04.07 9:52
Timer 0	00:00:00
Timer 1	00:00:00
Timer 2	00:00:00
Timer 3	00:00:00
AQA2 target value	2.000 mg/l
AQA2 tolerance	0.200 mg/l

- **3** Enter and confirm the measuring range limits.
- 4 Complete the entering of the formula with [Next].

The timers and AQA2 data linked to the method are displayed.

- 5 If necessary, enter intervals for up to 4 timers.
- 6 If necessary, enter the AQA2 target value and AQA2 tolerance.
- 7 Complete the programming of the method with [Complete].

The method is programmed and selected for measuring.

#### 4.5.12 The IQ-LabLink procedure

The IQ-LabLink procedure enables the secure data exchange between the

sensors of the IQ SENSOR NET online measuring system and a photometer (such as the

photoLab<sup>®</sup> 6000 series) with the aid of a commercial USB memory device.

During the matrix adjustment of the sensor, the measurement data of the IQ SENSOR NET sensor are adjusted to the photometrically determined reference data.

Previously, the reference data had to be entered manually during the matrix adjustment. With the aid of the IQ-LabLink procedure, the reference data can now be directly transferred to the IQ SENSOR NET sensors by means of a USB memory device, without manual input and with no risk of confusion.

System requirements for the IQ-LabLink procedure

- IQ SENSOR NET:
  - Terminal/controller with USB-A interface and software for the IQ-Lab-Link procedure (e.g. MIQ/TC 2020 XT)
  - Online sensor with software for the IQ-LabLink procedure (e.g. VARiON<sup>®Plus</sup> 700 IQ)
  - Photometer:
    - Photometer with software for the IQ-LabLink procedure (e.g. photoLab<sup>®</sup> 6000 series)

Course of the IQ-LabLink procedure

#### Step 1 on the IQ SENSOR NET terminal:

Automatic creation of a job file on the USB memory device with current sensor values, parameters, designation of the measuring location and automatic allocation of a job reference number for clear identification.

#### Step 2 on the photometer:

Automatic recognition of the job files, menu-guided measurement of all required parameters, storage of the determined data in the job file.

#### Step 3 on the IQ SENSOR NET terminal:

Automatic recognition of the job files, complete reading of all data required for the matrix adjustment on keypressing.



#### Note

To carry out the matrix adjustment of an online sensor with the IQ-LabLink procedure requires operating steps on both instruments: IQ SENSOR NET system and photometer of the photoLab<sup>®</sup> 6000 series.

The detailed description of the cross instrument operating steps for the matrix adjustment with the IQ-LabLink procedure on theIQ SENSOR NET and the photometer is given in an additional operating manual. This operating manual can be downloaded from the Internet under www.wtw.com.

# 4.6 Measuring the Absorbance / % Transmission

#### 4.6.1 General information

The absorbance or transmission respectively is measured without the use of any methods or profiles. All settings are configured during measurement.

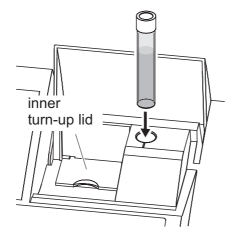
Measuring against the Reference absorbance The absorbance or transmission can alternatively be measured against the absorbance of the zero adjustment or against a *Reference absorbance* determined by yourself (see section 4.6.3 MEASURING AGAINST THE REFERENCE ABSORBANCE).

### 4.6.2 Measuring the absorbance or transmission

# <HOME>

Absorbance / % Transmission

Absorbance			16.04.07	9:52
To start measu insert ceil or p		ART/ENTER>		
300 nm				
Setup Wav	elength	Transmission	Referen	nce



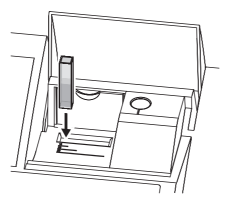
The settings of the last measurement are active.

- 1 Using *[Wavelength]*, change the wavelength as necessary.
- 2 Using [Absorbance] <-> [Transmission], you can switch over between absorbance and transmission measurement.
- **3** If necessary, use or measure a reference measurement with [*Reference*] (see section 4.6.3).
- 4 Depending on the type, insert the cell as follows:

#### Round cell:

Insert the round cell in the round cell shaft so it touches the bottom.

If the inner turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.



Absorbance	0.8	860	16.04.07 9:52
489 nm			10 mm
Setup	Wavelength	Transmission	Reference



Rectangular cell:

Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

The photometer starts measuring automatically.

5 Using [Absorbance] <-> [Transmission], switch over the display from Absorbance to Transmission or vice versa.

## 4.6.3 Measuring against the Reference absorbance

Each time the photometer is switched on, the absorbance or transmission is measured against the absorbance of the zero adjustment as a basis. You can, however, also determine a *Reference absorbance* and use it as the basis.

The *Reference absorbance* refers to the adjusted wavelength. The measured value remains stored until

- the photometer is switched off
- the cell type is changed
- the wavelength is changed
- a new reference value is measured
- it is deleted manually ([Reference] / Delete).
- the Absorbance / % Transmission measuring mode is exited

10 mm

# Single and multiple determination

The Reference absorbance can be determined with single or multiple determination. With multiple determination, the mean value is calculated as the median from the individual measured values.

Measuring the Reference absorbance

Absorbance / % Transm	ISSION	
Absorbance	16.04.07	9:5
To start measurement, insert cell or press <start enter=""></start>		

Wavelength Transmissio

<HOME>

Alexande

489 nm

The settings of the last measurement are active.

1 Start the reference measurement with [Reference].

If a value for the reference absorbance is already stored, it can be deleted or overwritten by a new reference measurement.

After the reference absorbance value has been deleted, the photometer measures against the absorbance of the zero adjustment.

Reference absorbance	16.04.07	9:52
To start measurement, insert cell or press <start enter=""></start>		
489 nm	1	0 mm

Reference a	osorbance		16.04.07	9:52
	Last measured absorb	ance		
	0.232			
	Median			
	0.232 (1 Measure	emer	it(s))	
489 nm			1	.0 mm
Nevt meas	Discard		Ann	b.

Absorbance			16.04.07	9:52
			Reference:	:
	neasurement, I or press <st< td=""><td>ART/ENTER&gt;</td><td></td><td></td></st<>	ART/ENTER>		
489 nm			1	0 mm
Setup	Wavelength	Transmission	Refere	nce

2 Insert the cell with the reference sample.

The first single measurement for the Reference absorbance is carried out.

The following data is displayed as the result:

- The measured absorbance from the (last) single measurement.
- The median from all single measurements carried out up to now.
- 3 If necessary, carry out further single measurements for the formation of the median with [Next meas.] or

discard the last single measurement with [Discard].

4 To accept the median value, press [Apply].

The photometer is ready to measure.

The reference absorbance is displayed in the top right corner during absorbance or transmission measurement.

# 4.7 Special / Multi wavelengths methods

#### 4.7.1 Basic information on Special / Multi wavelengths measurements

In the Special / Multi wavelengths mode of the photoLab<sup>®</sup> 6600 UV-VIS, you can carry out measurements with special methods and functions.

You can use the following functions for these methods:

- Measurements at different wavelengths
- Multiple measurements at one wavelength (e.g. before and after adding a reagent)
- Use of procedure variables. Procedure variables provide a value that has to be entered prior to each measurement on the photometer (e.g. volume, pH value or temperature)
- Check whether a value meets a condition.
   With a condition you can check a value for validity (e.g. absorbance value, procedure variable or the result of a formula).
- Formula editor for the convenient programming of any user-defined methods

The method list in the Special / Multi wavelengths mode comprises:

Special methods

- preprogrammed multi wavelengths methods
- preprogrammed special methods
- special methods programmed by the user



#### Note

If you program any special methods yourself, you can use all extended functions of the Special / Multi wavelengths mode.

# 4.7.2 Programming / modifying the Special / Multi wavelengths methods

#### Note

For multi wavelength methods, you can use the method numbers 2001 to 2050. All special methods can also be selected in the method list of the concentration mode.

The creation of a user-defined method is done in the following steps:

Enter the general method data • Method number, method name, unit etc. Enter the wavelengths for absorbance measurements ( $A_{x nm}$ ) Minimum 1, maximum 10 Define the procedure variables  $(K_X)$  (optional) Procedure variables are used to take into account any influence quantities that cannot be measured by the photometer. The values for these procedure variables have to be entered for all measurements with the method, e.g. the temperature or pH value. Enter the formula to calculate the measurement result Enter the formula with which you want to calculate the measurement result in the formula editor. Enter an additional condition (optional) • Conditions are used to check the measurement result for validity. The condition is entered with the formula editor.

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#### Example: Determination of chlorophyll a according to Nusch

The chlorophyll determination is based on two measurements (before and after adding an acid) of the optical density (= absorbance) of the extract of an aqueous sample at 665 nm.

Chlorophyll a ( $\mu g/l$ ) = 29.6 \* ( $A_{(before) 665 nm} - A_{(after) 665 nm}$ )\*( $V_{Extract}/V_{Sample}$ )

# <u>with:</u>

<u></u>	
A(before) 665 nm	1st absorbance measurement at 665 nm (before adding the acid)
A <sub>(after) 665 nm</sub>	2nd absorbance measurement at 665 nm (after adding the acid)
V <sub>Extract</sub>	Volume of the extract (in ml)
V <sub>Sample</sub>	Volume of the aqueous sample (in ml)

**Converted equation** For entry on the photometer, assign names that you can enter in the formula editor on the photometer to the variables of the equation.

 $R = 29.6 * (A_{665nm} - A_{665nm_2}) * (K_1/K_2)$ 

## with:

R (chlorophyll a (µg/l))	Result (concentration chlorophyll A in µg/l)
A <sub>x nm</sub> (= A <sub>(before) 665</sub> nm) A <sub>x nm_2</sub> (= A <sub>(after) 665 nm</sub> )	Variables for absorbance. These values are measured by the photometer. Here: Two measurements at the same wave- length, at different points of time.
K <sub>1</sub> (= V <sub>Extract</sub> )	Procedure variables
K <sub>2</sub> (= V <sub>Sample</sub> )	K1 = Volume of the extract (in ml)
	K2 = Volume of the aqueous sample (in I)
Numerals	Freely selectable numerical values

# <HOME> Special / Multi wavelengths – [Setup] – Edit method

Edit method		16.04.07 9:52
Number		2001
Name		Chlorophyll a
Version		1.0
Citation form		Chl a
Unit		µg/I
Resolution		0.1
Cell		10 mm
Lower limit of measuring		0 µg/l
Upper limit of measuring		1000 µg/l
Method list	Delete	Next

Wavelength			16.04.07 9:52
Wavelength 1			665 nm
Back	Add	Delete	Next

1 Enter the general method data here. The next available method number is already entered as the number.

You have the following options when filling out the input fields:

- Fill out all empty input fields one after the other
- Using *[Method list]*, select an already existing method as a model, give it a new method number and adjust the entries
- Using [Method list], select an existing method in order to change it (without changing the number).
- You can delete the method completely with [Delete].
- 2 Using [Next], accept all entries on the page and switch to the next page.

Enter the wavelengths for the absorbance measurements  $(A_x n_m)$ .

**3** Add another wavelength with *[Add]*.

Delete a highlighted wavelength with [Delete].

4 Using *[Next]*, accept all entries on the page and switch to the next page.

Procedure variables	16.04.07 9:52
Procedure variables are variables who values have to be entered during the measurement (e.g. temperature or pl	course of the
If a procedure variable is required to Create a procedure variable (K) with	
Back Add	Next
Procedure variables	16.04.07 9:52
K 1 K 2	V (extract) V (sample)
Back Add De	lete Next
Back Add De	lete Next
Back Add De	lete Next 16.04.07 9:52
	16.04.07 9:52 t an operation, function
Formula entry Use the <operators> softkey to select</operators>	t an operation, function Pi).
Formula entry Use the <operators> softkey to select or constant (e.g.: +, -, *, tan, log, e, Use the <variables> softkey to select</variables></operators>	t an operation, function Pi).
Formula entry Use the <operators> softkey to select or constant (e.g.: +, -, *, tan, log, e, Use the <variables> softkey to select certain wavelength or a procedure va</variables></operators>	16.04.07 9:52 t an operation, function Pi). : an absorbance at a riable.
Formula entry Use the <operators> softkey to select or constant (e.g.: +, -, *, tan, log, e, Use the <variables> softkey to select certain wavelength or a procedure va Enter numerals via the keyboard. You can erase the last entry with &lt;◀</variables></operators>	16.04.07 9:52 t an operation, function Pi). : an absorbance at a riable.

Create all required procedure variables.

5 Create a procedure variable required for the formula with [Add] and enter a designation, e.g. the measured parameter.

or

Using *[Next]*, accept all entries on the page and switch to the next page.

6 Add another procedure variable with *[Add]*.

or

Delete a highlighted procedure variable with [Delete].

7 Using *[Next]*, accept all entries on the page and switch to the next page.

Enter the formula.

8 Enter any numbers with <0...9>.

Use [Operators], <**▲**><**▼**> <**◀**><**▶**> and <**START**·ENTER> to enter an operator, a function or a constant.

Use [Variables], <**▲**><**▼**> <**◀**><**▶**> and <**START**·ENTER> to select a variable.

The formula is displayed after each step.

Using <◀> you can delete the last element of the formula.

Use [Back] to quit the formula editor.

Formula entry	16.04.07 9:52	9	Selec The c is dis
Variables A(665 nm) K1 (V extract (ml)) K2 (V sample			15 015
Back Operators Variables	Next Back		
Formula entry	16.04.07 9:52	10	Add a The c
R = 29.6 * (A665nm - Back Operators Variables	Next		is dis
Formula entry	16.04.07 9:52	11	Selec
Variables A(665 nm)			A <sub>665 m</sub> ment. The c is dis
K1 (V extract (ml)) K2 (V sample (	(ml))	12	To me same Selec The n
Back Operators Variables	Next Back		up. Enter ment, suren confir The c is dis
Formula entry	16.04.07 9:52	13	Comp The c

9 Select and confirm the variable. The current version of the formula is displayed.

**10** Add an operator. The current version of the formula is displayed.

11 Select and confirm the Variable A<sub>665 nm</sub> for the second measurement.

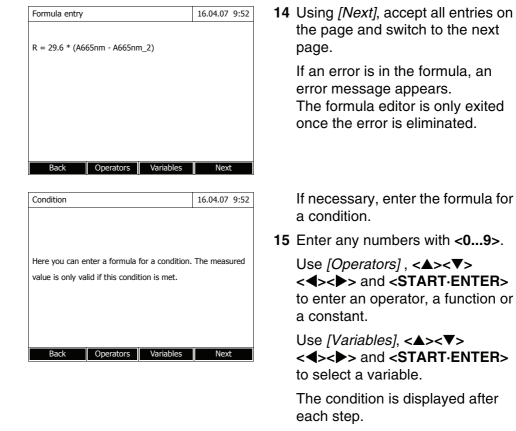
The current version of the formula is displayed.

12 To measure once again at the same wavelength:
 Select the underscore (\_).
 The measurement input field pops up.
 Enter the index for the measure-

ment, e.g. 2 for the second measurement at this wavelength, and confirm.

The current version of the formula is displayed.

**13** Complete the formula. The current version of the formula is displayed.



Using << > you can delete the last element of the condition.

Use [Back] to quit the formula editor.

- **16** Complete the condition.
- **17** Complete the programming of the method with *[Next]*.

Condition	16.04.07 9:52
<sup>A</sup> 665 nm <sup>&lt; 2</sup>	
b5	
Back	Next

Edit method			16.04.07 9:52
Sequence			Designation
Measurement Measurement			
Back			Next
Special / Multi	wavelengths		16.04.07 9:52
V extract (ml)			
Press <start enter=""> to enter the value</start>			
2001:Chl a 10 mm			Chlorophyll a

Method list Citation form

Setup

If the formula includes several measurements at the same wavelength (measurement sequence), you can assign names to the individual measurements of the sequence.

- **18** Enter the names for the individual measurements of a sequence.
- **19** Complete the programming of the method with [*Next*].

The method is programmed and selected.

The photometer is ready to measure.

#### Special / Multi wavelengths [Method list] The list of methods is displayed. 16.04.07 9:52 Select method (all) The methods are ordered according to the method number. 2001 Protein 2002 DNA purit Select the method: **1** Select the required method with < > < V >. The active selection is displayed in reverse video. **2** Accept the selection with <START.ENTER>. Last used The photometer is ready to measure. Narrowing down the If the list is very long, you can narrow down the method list and thus make method list the search easier as follows: • Using [Last used], you can restrict the method list to the ten methods last used. • With the search function you can search certain character strings in the list. The search takes place as a full-text search of the entire list contents. Thus you can search for a method number or certain citation form. Search function Select method (last used) 16.04.07 9:52 Search for a character string: Enter the character string to be Pro\_ searched for in the search window 2001 Chl a Chlorophyll a with **<A...9**>. The list appearing below shows all hits containing the character string. The hit list is updated with each character that is entered. All methods Note Note the case sensitivity when searching.

# 4.7.3 Selecting a Special / Multi wavelengths method

<HOME>

To select a method for Special / Multi wavelengths measurements, proceed as follows:

Special / Multi wavelengths	16.04.07 9:52
Please select method for measuring!	
Setup Method list Citation form	Unit

Special / Multi wavelengths

<HOME>

Special / Multi wavelengths

4.7.4 Carrying out Special / Multi wavelengths measurements

16.04.07 9:52

- **1** Select the required method with
  - [Method list] (see section 4.7.3).

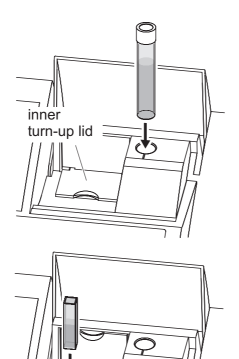
For methods with procedure variables: Enter the values of all procedure variables one after the other.

If necessary, carry out a zero measurement.

V extract (ml)		
Press <start enter=""> to enter the value</start>		
2001:Chl a 10 mm	Chlorophyll a 0.00 - 1000.00 µg/l	
Setup Method list Ci	itation form Unit	
Special / Multi wavelengths	16.04.07 9:52	
Measurement 1		
Zero measurement require	ed!	
2001:Chl a 10 mm	Chlorophyll a 0.00 - 1000.00 μg/l	

Setup Method list Citation form Unit

Special / Multi	wavelengths		16.04.07 9:52
Measurement 1			
To start measurement, insert cell or press <start enter=""></start>			
2001:Chl a 10 mm			Chlorophyll a
Setup	Method list	Citation form	Unit



The photometer is ready to measure.

2 Depending on the type, insert the cell as follows:

## Round cell:

Insert the round cell in the round cell shaft so it touches the bottom.

If the inner turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.

# Rectangular cell:

Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

Special / Multi wavelengths			16.04.07 9:52
V extract (ml) V sample (ml) Measurement 1	100 ml	:00	
Proceed with <start enter=""></start>			
2001:Chl a 10 mm			Chlorophyll a
Setup	Method list	Citation form	Unit

Special / Multi wavelengths			16.04.07 9:52
Measurer	ment 2		
To start measurement, insert cell or press <start enter=""></start>			
2001:Chl a 10 mm			Chlorophyll a
Setup	Method list	Citation form	Unit

Special / Multi	wavelengths		16.04.07 9:52
V extract (ml) V sample (ml) Measurement 1 Measurement 2	A(665 n) = 0.6		
	1.	78	mg/ml
Start new analysis with <start enter=""></start>			
Setup	Method list	Citation form	u Unit

An intermediary result is displayed if there are several measurements.

The photometer is ready for the next measurement.

3 Start the measurement.

The photometer is ready to measure.

The result is displayed.

If an entered condition is not met, no measured value is displayed.

4 If necessary, start a new measurement with the method.

# 4.8 Spectrum

#### 4.8.1 General information

With the Spectrum function, the absorbance or *Transmission* in dependency of the wavelength is measured and recorded. The wavelength range can be freely selected within the measuring range of the photometer. The increment is 1 nm.

A spectrum is recorded without using any methods or profiles. All settings are configured during measurement.

- **Baseline** A baseline has to be recorded before a spectrum is recorded. The baseline has to cover at least the wavelength range of the spectrum to be recorded. Once the baseline is measured, it remains stored in the photometer until
  - a new baseline is recorded
  - the Spectrum mode is exited or the photometer is switched off
- **Settings** You can record a spectrum with standard settings without opening the setting window.

The following settings are possible for a spectrum:

Input field	Possible entries
Wavelength start	190* 1100 nm
Wavelength stop	190 1100* nm
Mode	Absorbance* or Transmission
Smoothing	Yes* or No
Scaling	Auto* or Manual
Scaling: Auto*	During measurement, the instrument adjusts the axis scaling (minimum and maximum value of the axis) to the measured values. The entire curve is always visible.
Scaling:Manual Y-axis min Y-axis max	The axis scaling (minimum and maximum value of the axis) is set manually.

\* default setting



#### Note

You can store the current settings as a profile with *[Save]*. You can load a stored profile with *[Open]*. Profiles for spectra have the file extension, ".profil".

9000000000000000000000000000000000000	
and the second a baseline first (<2ERO>).         1       Adjustment of wavelength range under (<2ERO>).         4       Adjustment of wavelength range under (<2ERO>).         4       400       600       800         90       000       000       1000         Wavelength [nm]       Setup       Open         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbar         Smoothing       A         Scaling       A         Spectrum       16.04.07       9         4.0       Apply         Spectrum       16.04.07       9         4.0       Apply         Spectrum       16.04.07       9         4.0       Apply         Spectrum       16.04.07       9         4.0       Apply       Apply         Spectrum       16.04.07       9         4.0       Apply       Apply         You have to record a baseline first (<2ERO>).       Adjustment of wavelength range under (<2ERO>).         4.0       600       800       1000         Wavelength [nm]       Open         Setup       Open         S	
Spectrum         2       You have to record a baseline first ( <zero>).         1       Adjustment of wavelength range under (<zero>).         -1.0       600         400       600         Wavelength [nm]         Setup       Open         Spectrum       16.04.07         Wavelength start       1100         Mode       Absorbar         Smoothing       3         Scaling       A         Spectrum       16.04.07         9       4.0         4.0       Apply         9       4.0         4.0       Apply         9       4.0         4.0       Apply         9       4.0         4.0       Apply         9       4.0         9       4.0         9       0         9&lt;</zero></zero>	·····
2       You have to record a baseline first ( <zero>).       1         1       Adjustment of wavelength range under (<general setup="">.       0         -1.0       600       800       1000         Wavelength [nm]       0pen         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbal         Smoothing       0         Scaling       A         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbal         Smoothing       0       0         Spectrum       16.04.07       9         Vaulation       16.04.07       9         Spectrum       16.04.07       9         Spectrum       16.04.07       9         Spectrum       16.04.07       9         Spectrum       10       0       0         2       You have to record a baseline first (<zero>).       0       0         1       Adjustment of wavelength range under (<zero>).       0       0         -1.0       0       00       000       1000         Wavelength [nm]       0       0       0       0<!--</td--><td>·····</td></zero></zero></general></zero>	·····
2       You have to record a baseline first ( <zero>).       1         1       Adjustment of wavelength range under (<general setup="">.       0         -1.0       600       800       1000         Wavelength [nm]       0pen         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbal         Smoothing       0         Scaling       A         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbal         Smoothing       0       0         Spectrum       16.04.07       9         Vaulation       16.04.07       9         Spectrum       16.04.07       9         Spectrum       16.04.07       9         Spectrum       16.04.07       9         Spectrum       10       0       0         2       You have to record a baseline first (<zero>).       0       0         1       Adjustment of wavelength range under (<zero>).       0       0         -1.0       0       00       000       1000         Wavelength [nm]       0       0       0       0<!--</td--><td>·····</td></zero></zero></general></zero>	·····
Centeral setup2.         -1.0       600       800       1000         Wavelength [nm]       Open         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbai         Smoothing       A         Scaling       A         Spectrum       16.04.07       9         Wavelength start       Mode       Absorbai         Scaling       A       Apply         Spectrum       16.04.07       9         Image: A start (ZERO>)       Intervention of wavelength range under (ZERO>).       Adjustment of wavelength range under (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>)       Intervent (ZERO>)         Image: A start (ZERO>)       Intervent (ZERO>)       Intervent (ZERO>)         Image: A start (ZERO>)       Intervent (ZERO>)       Intervent (ZERO>)	
Centeral setup2.         -1.0       600       800       1000         Wavelength [nm]       Open         Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbai         Smoothing       A         Scaling       A         Spectrum       16.04.07       9         Wavelength start       Mode       Absorbai         Scaling       A       Apply         Spectrum       16.04.07       9         Image: A start (ZERO>)       Intervention of wavelength range under (ZERO>).       Adjustment of wavelength range under (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>).       Intervent of start (ZERO>).         Image: A start (ZERO>)       Intervent of start (ZERO>)       Intervent (ZERO>)         Image: A start (ZERO>)       Intervent (ZERO>)       Intervent (ZERO>)         Image: A start (ZERO>)       Intervent (ZERO>)       Intervent (ZERO>)	
400         600         800         1000           Wavelength [nm]         Open           Spectrum         16.04.07         9           Wavelength start         1100           Mode         Absorbar           Smoothing         3           Scaling         A           Spectrum         16.04.07           9         4.0           Spectrum         16.04.07           9         4.0           Spectrum         16.04.07           2         You have to record a baseline first ( <zero>).           1         Adjustment of wavelength range under -1.0           400         600         800           Wavelength [nm]         0pen           Spectrum         16.04.07         9</zero>	
400     600     800     1000       Wavelength [nm]     Open       Spectrum     16.04.07     9       Wavelength start     1100       Mode     Absorbar       Smoothing     A       Scaling     A       Spectrum     16.04.07       You have to record a baseline first     ( <zero>).       1     Adjustment of wavelength range under       -1.0     600     800       400     600     800       Spectrum     16.04.07     9</zero>	—
Setup     Open       Spectrum     16.04.07     9       Wavelength start     1100       Mode     Absorbai       Smoothing     A       Scaling     A       Spectrum     16.04.07       Spectrum     100       Spectrum     16.04.07       Spectrum     16.04.07       You have to record a baseline first     (       ( <zero>).     1       Adjustment of wavelength range under     (       General setup&gt;.     0       400     600       800     1000       Wavelength [nm]     0       Setup     0</zero>	
Spectrum       16.04.07       9         Wavelength start       1100         Mode       Absorbai         Smoothing       A         Scaling       A         Spectrum       16.04.07       9         Spectrum       ( <zero>).       1         Adjustment of wavelength range under       (<zero>).      </zero></zero>	
Wavelength start Wavelength stop 1100 Mode Absorbai Smoothing Scaling A	
Wavelength stop     1100       Mode     Absorbat       Smoothing     A       Scaling     A       Spectrum     16.04.07       Spectrum     16.04.07       You have to record a baseline first ( <zero>).     A       Adjustment of wavelength range under <general setup="">.     0       -1.0     600     800       400     600     800       Steup     Open       Spectrum     16.04.07</general></zero>	:52
Mode     Absorban       Smoothing     Smoothing       Scaling     A       Spectrum     16.04.07       Spectrum     16.04.07       You have to record a baseline first     ( <zero>).       1 Adjustment of wavelength range under     (<zero>).       1 Adjustment of wavelength range under     (<zero>).       1 Adjustment of wavelength range under     0       Setup     Open       Setup     Open</zero></zero></zero>	
Smoothing Scaling A Spectrum 16.04.07 9 Spectrum 16.04.07 9 2 You have to record a baseline first ( <zero>). 1 Adjustment of wavelength range under -1.0 400 600 800 1000 Wavelength [nm] Setup Open Spectrum 16.04.07 9</zero>	
Spectrum 16.04.07 9  4.0  Spectrum  2 You have to record a baseline first ( <zero>).  1 Adjustment of wavelength range under (<zero>).  1 Adjustment of wavelength range under -1.0  400  600  Wavelength [nm]  Setup  Open  Spectrum  16.04.07 9</zero></zero>	Yes
Spectrum       16.04.07 9         4.0	uto
Spectrum       16.04.07 9         4.0	
90 minute     4.0	
Spectrum       2 You have to record a baseline first ( <zero>).       1 Adjustment of wavelength range under <general setup="">.       -1.0     600       400     600       Wavelength [nm]       Setup     Open</general></zero>	:52
You have to record a baseline first ( <zero>). Adjustment of wavelength range under -1.0 400 500 Wavelength [nm] Setup Open Spectrum 16.04.07 9</zero>	
2 You have to record a baseline first ( <zero>). 1 Adjustment of wavelength range under <general setup="">. </general></zero>	
-1.0         400         600         800         1000           Wavelength [nm]         Setup         Open	
-1.0         400         600         800         1000           Wavelength [nm]         Setup         Open	
400         600         800         1000           Wavelength [nm]         Open           Setup         16.04.07         9	
Wavelength [nm]       Setup     Open       Spectrum     16.04.07 9	
Setup Open Spectrum 16.04.07 9	_
4.0	
	:52
3.0	:52
2.0	:52
to 50 1.0	:52
	:52
0.0	:52
-1.0 <u>: : : :</u> 300 400 500 <b>Wavelength [nm]</b>	:52
2	:5

## 4.8.2 Recording the Spectrum

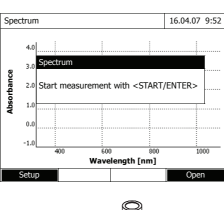
A message containing operating instructions is displayed.

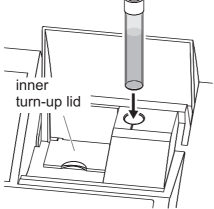
- 1 Open the setting menu with [Setup].
- 2 Select the start and end point of the spectrum to be recorded and the mode (*Absorbance* or *Transmission*).
- 3 Accept all entries with [Apply].

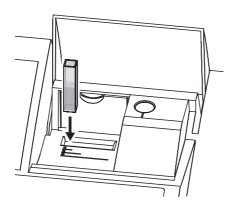
A message containing operating instructions is displayed.

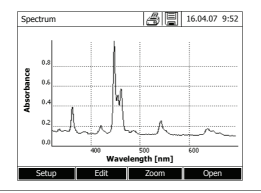
Recording the baseline:

- 4 Press the <ZERO·BLANK> key. The photometer records the baseline.
- 5 Wait until the baseline is completely recorded.









The photometer is ready to measure after the baseline has been recorded.

Recording the spectrum:

6 Depending on the type, insert the cell as follows:

# Round cell:

Insert the round cell in the round cell shaft so it touches the bottom.

- 7 Close the inner turn-up lid.
- 8 Start the measurement with <START·ENTER>.

After the spectrum has been recorded, the following message appears: *Recording of spectrum is completed.* 

# Rectangular cell:

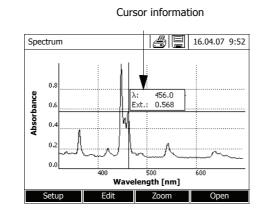
Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

- 9 Close the cell shaft cover.
- **10** Start the measurement with **<START·ENTER>**.
- **11** Wait until the spectrum is completely recorded.

At the end of the recording the following message appears: *Recording of spectrum is completed.* 

12 Confirm the message with <START.ENTER>.



The cursor appears at the absolute maximum of the spectrum.

- **13** You have the following options:
  - Immediately edit the spectrum (see section 4.8.3)
  - With **<PRINT>**, you can output the spectrum to a connected printer as a graphic.
  - You can save the spectrum as a \*.csv file with <STORE>. As the storage location, you can select the photometer (*Internal DataB folder*) or a USB memory device connected to the USB-A connection (*USB memory*). Stored spectra can be recalled and edited at any time (see section 4.8.3).

## 4.8.3 Loading/editing a spectrum

A spectrum can be edited immediately after measurement. Stored spectra can be loaded and edited as well.

The following tools are available for editing:

- Cursor function for incremental moving along the curve with indication of the x and y values
- Zoom function to scale up a section
- Mathematical functions for various evaluating and calculating operations. The functions are described from page 93.

Loading a stored spectrum

Open (Inter	nal DataB folder)	16.04.07 9:5
26.02.07	Holmium.csv	
23.02.07	K2Cr2O7_340nm.csv	

<HOME>

Spectrum

The list with the spectra stored in the exchange memory is displayed.

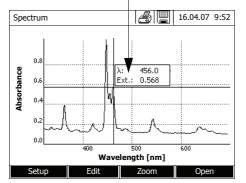
- 1 If necessary, you can select a different memory location for the spectrum with *[Location]* (USB memory device at the USB-A connection).
- Select the required spectrum.
   The original view of the curve is displayed.

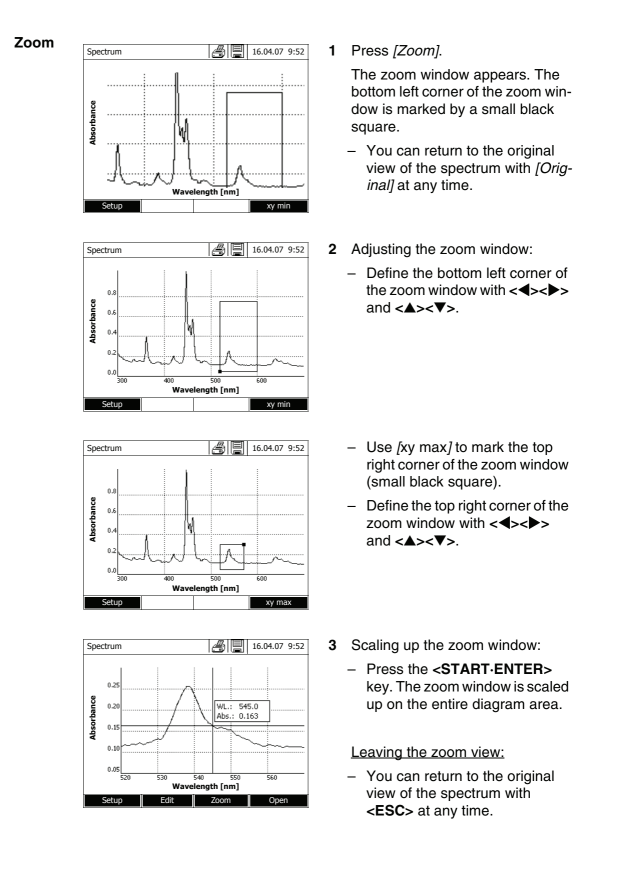
The cursor consists of a horizontal and vertical line that cross each other on a point of the curve. A box names the x and y values of the point of the curve.

Move the cursor along the x axis (wavelength) with  $< \P > < \triangleright >$ . You can scan and evaluate the curve point after point.

#### Cursor

Cursor information





Open the selection of mathematical functions with [Edit]:

Edit

#### 93

• Extreme values (zoomed area)

Highlights the extreme values (minimum and maximum values) of the displayed spectrum.

• Mark points

Opens an edit mode where you can highlight individual points of the spectrum.

With the *[Mark]* function key you can highlight individual points. The wavelength and measured valued are displayed at the highlighted point.

With the [Delete] function key you can remove individual points.

- *Delete all marks* Erases all highlighted points in the spectrum.
- Original

Displays the original, unedited spectrum.

• Integral

Calculates the area between the zero line and curve within a freely selectable wavelength interval [X1,X2].

Derivative

Calculates the derivative of the total spectrum. To calculate the second and third derivative, the function can be carried out several times.

- Compare spectrum Loads a second spectrum into the same diagram for direct comparison.
- Add spectrum
   Adds a stored spectrum to the current spectrum.
- Subtract spectrum Subtracts a stored spectrum from the current spectrum.
- Divide spectrum (ratio)
   Divides the absorbance or % transmission values of the current spectrum
   by the values of a stored spectrum
- Add fixed value Adds a constant absorbance or % transmission value to the current spectrum.
- *Multiply fixed value* Multiplies the absorbance or % transmission values of the current spectrum by a constant value.



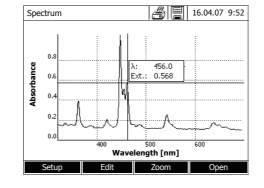
#### Note

The addition, subtraction and division of two spectra always applies to the common wavelength range of both spectra only.

## 4.8.4 Saving / exporting a spectrum

The saving of a spectrum saves both the edited and the original spectrum. Consequently, the original spectrum can be restored from each stored spectrum.

## Saving



- Record a spectrum (see section 4.8.2) or Load a stored spectrum (see section 4.8.3).
- 2 If necessary, connect a USB memory device to the USB-A interface.
- 3 Open the save dialog with **<STORE>**.
- If necessary, change the storage location with [Location]: Internal DataB folder.
   Exchange folder in the instrument or USB memory: USB memory device connected at the USB-A connection.
- **5** If necessary, change the file name.
- 6 Save the file with <START.ENTER>.

## **Export to a PC** Export a stored spectrum to a PC: see section 4.12.3

## 4.9 Kinetics

The Kinetics function enables the temporal tracing of the absorbance or transmission of a sample at a certain wavelength.

The photometer automatically calculates the slope between two adjacent measuring points from the available measurement data. The catalytic activity can also be determined and displayed if required.

To record the kinetics, the photometer carries out single measurements at regular intervals (measuring interval) and stores the measured values as a time function.

All settings for a recording are administrated as a profile. Profiles can be created, stored, edited and deleted. Each measurement requires a respective profile.

#### 4.9.1 Creating/editing profiles for Kinetics recordings

# Note

Profiles for Kinetics records are stored under the numbers 4001 to 4020. In the delivery condition, a profile is stored for demonstration purposes.

A profile for a Kinetics recording comprises the following data:

Input field	Possible entries
Number *	4001 4020
Name	Any name (max. 18 characters)
Mode*	Absorbance or Transmission
Wavelength*	Freely selectable (in nm)
Duration*	Total duration in the format hh:mm:ss (hours:minutes:seconds)
Interval*	Measuring interval = time interval between two successive single measurements in the format hh:mm:ss (hours:minutes:seconds)
	Exception: With the setting, <i>Measurements/interval</i> : <i>Max/</i> <i>interval</i> the interval is defined differently (see below).
Delay	Time between the start of the recording and the start of the first single measurement
Scaling	Auto or Manual

#### ba75848d04 09/2011

Input field	Possible entries
Scaling: Auto**	During measurement, the instrument adjusts the axis scaling (minimum and maximum value of the axis) to the measured values. The entire curve is always visible.
Scaling:Manual Y-axis min Y-axis max	The axis scaling (minimum and maximum value of the axis) is set manually.
Measurements/interval	1/interval or Max/interval
	Here you define how many measurements are carried out per interval.
	This setting has an impact on the calculation of the slope of the individual intervals (see section 4.9.6).
Catalytic activity	Yes or No
	Here you determine whether the catalytic activity should be calculated.
	The catalytic activity is a measure for the amount of substance that is converted per time unit. To accelerate the substance conversion, a cata- lyst or enzyme (biological catalyst) is used in most cases.
Catalytic activity: Yes	
Factor Unit Resolution	The catalytic activity or enzymatic activity is calcu- lated from the slope of the curve.
riesolution	Cat. A. = mean value Slope $[\Delta/min]$ * Factor
	Here you can enter the value for <i>Factor</i> .
	The calculated value for the catalytic activity is displayed in the menu, [Edit] / Slope & catalytic activ- ity, together with the unit and resolution selected here.
* necessary inputs	·

\*\* default: Auto

# Creating/editing a profile

<home> Kinetics – [Setup] – Edit profi</home>	le
Edit profile (1 of 2)	16.04.07 9:52
Number	4001
Name	NADH
Mode	Absorbance
Wavelength	340 nm
Duration	02:00:00
Interval	00:00:30
Delay	00:01:00
Scaling	Auto

Next

Profile list Delete

Edit profile (1 of 2)	16.04.07 9:52
Measurements/interval	1/interval
Catalytic activity	Yes
Factor	1.000
Unit	cat
Resolution	0.01
Back	Complete
Duck	compicte

1 Enter the data for the profile here. The next available profile number is already entered as the number.

You have the following options when filling out the input fields:

- Fill out all empty input fields one after the other
- Using [*Profile list*], select an already existing profile as a model, give it a new profile number and adjust the entries
- Using [*Profile list*], select an existing profile in order to change it (without changing the number).
- You can delete the profile completely with [Delete].
- 2 With [Next] you can switch to further settings.
- **3** Enter further data for the profile here.
- 4 Accept all entries with [Complete].
  - The profile is created and selected. The photometer is ready to measure.



## Note

The *Catalytic activity* function is only available if the Absorbance mode was selected.

4.9.2	Loading	а	profile	for	<b>Kinetics</b>	recording

To load a profile for Kinetics recording, proceed as follows:

<home></home>		
Kinetics		
[Profile list]		

Selec	t profile (	all)		16.04.07	9:52
4001	NADH		Absorbance		
4002	ADH		Absorbance		
		1	1	r	
Las	st used				

The list of profiles is displayed. The profiles are ordered according to the profile number.

Selecting a profile:

- Select the required profile with
   <▲><▼>. The active selection is displayed in reverse video.
- 2 Accept the selection with <**START·ENTER**>.

The photometer is ready to measure.

Narrowing down the list of profiles

If the list is very long, you can narrow down the profile list and thus make the search easier as follows:

- Using [Last used], you can restrict the profile list to the ten profiles last used.
- With the search function you can search certain character strings in the list. The search takes place as a full-text search of the entire list contents. Thus you can search for a profile number or name.

## **Search function**

Select profile (last used)		16.04.07 9:52
NA_		
4001 NADH	Absorbance	
		-1
All profiles		

#### Search for a character string:

Enter the character string to be searched for in the search window with **<A...9>**.

The list appearing below shows all hits containing the character string. The hit list is updated with each character that is entered.

# Note

Note the case sensitivity when searching.



## 4.9.3 Recording the Kinetics

#### Note

During the recording, the photometer cannot carry out any regular self-test or self-calibration (AutoCheck), because the recording would have to be interrupted for this. A warm-up time of at least two hours is required for the photometer to measure reliably during the recording.

<home> Kinetics</home>		
Kinetics	16.04.07	9:52
Please select a profile for mea	_	
Setup Profile list	Oper	1

 
 Kinetics
 16.04.07
 9:52

 To start measurement, insert cell or press <START/ENTER>

 4002
 Absorbance

 Setup
 Profile list
 Open

inner turn-up lid Note the warm-up time of at least 2 hours for kinetic recordings.

1 Select the required profile with [*Profile list*] (see section 4.9.2).

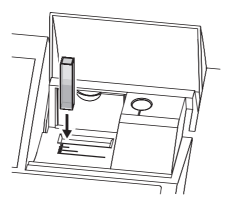
The photometer is ready to measure after the profile has been selected.

**2** Depending on the type, insert the cell as follows:

#### Round cell:

Insert the round cell in the round cell shaft so it touches the bottom.

If the inner turn-up lid is opened too wide, a message prompts you to close the inner turn-up lid.



Kine					6.04.07 9:52
4002 Duration: 00:00:24			Num		surements: 4 val:00:00:06
	1.0				
	0.9				
8	0.8				
Absorbance	0.7				
d d	0.6				
Abs	0.5				
	0.4				
	0	50	100	150	200
		-	Time [s	I	
					Stop

## Rectangular cell:

Open the inner turn-up lid.

Insert the rectangular cell vertically so it touches the bottom and left edge of the cell shaft. The opaque sides of the rectangular cell must point to the front and back.

The photometer has an external light recognition. If there is too much external light, a message prompts you to close the cell shaft cover.

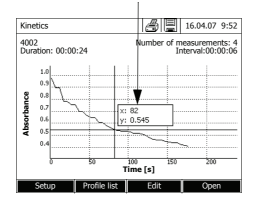
The photometer starts recording automatically.

**3** Wait until the recording is finished.

Stopping the recording:

- Use [Stop] to terminate the recording prematurely. The curve recorded up to this point can be stored and edited (see section 4.9.6).
- Use **<ESC>** to completely cancel measurement. The curve recorded up to this point is discarded.

## Cursor information

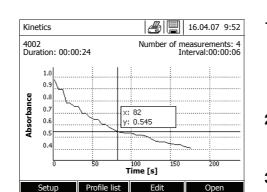


4 After the specified *Duration* has expired, the cursor appears.

You have the following options:

- You can move the cursor along the curve and have the measurement data for each point displayed (see section 4.9.6)
- With **<PRINT>**, you can output the kinetic curve to a connected printer as a graphic.
- You can store the kinetic curve with **<STORE>** (see section 4.9.4).
- Execute further functions to edit the kinetic record (see section 4.9.6)
- Close the kinetic record with **<ESC>**.

Saving



# 4.9.4 Saving / exporting a Kinetics record

1 Carry out the kinetic recording (see section 4.9.3) or

Load a stored kinetic record (see section 4.9.4).

- 2 If necessary, connect a USB memory device to the USB-A interface.
- 3 Open the save dialog with <STORE>.
- If necessary, change the storage location with [Location]: Internal DataB folder.
   Exchange folder in the instrument or USB memory: USB memory device connected at the USB-A connection.
- **5** If necessary, change the file name.
- 6 Save the file with <START.ENTER>.

**Export to a PC** Export a stored kinetic record to a PC: see section 4.12.3

Example of a kinetic recording (\*.csv file)

6 4001 1 1 525 1280913092 59 5 1 0.000 0.301 0 1.000 µkat 2 Device: Serial number:Software: User: photoLab 6600 UV-VIS 09130512 1.30-WTW-1.60 Administrator Start time Wavelength [nm] 04.08.2010 11:11 525 Time [s] Absorbance 0,092 0 0,077 5 10 0,073 0,069 15 . . . . . . .

Line 1 - explanations:

Column	Value	Explanation
1	6	Version of the file format for the CSV file
2	4001	Profile number
3	1	Measurement of absorbance (0) or transmission (1)
4	1	Measurement once per interval (0) or as often as possible (1)
5	525	Wavelength (in nm)
6	1280913092	Start time (internal data format)
7	59	Duration (in sec)
8	5	Interval time (in sec)
9	1	Scaling automatic (0) or manual (1)
10	0.000	Minimum for manual scaling
11	0.301	Maximum for manual scaling
12	0	Enzymatic activity Off (0) or On (1)
13	1.000	Factor for enzymatic activity
14	µkat	Unit of enzymatic activity
15	2	Decimal points for enzymatic activity

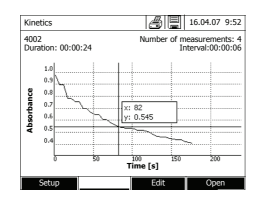
## 4.9.5 Loading a Kinetics record

You can load and view stored Kinetics records.

Loading a stored Kinetics record

<HOME> Kinetics – [Open]

		16.04.07	9:52
T			
26.02.07	Enzyme kinetics.csv		
24.02.07	ADH.csv		
24.02.07	kinetics_4002_070224_1410.csv	,	
Location	Delete		
Location	Delete		



The list with the stored Kinetics records is displayed (*Internal DataB folder*).

- 1 With [Location] select the memory location of the kinetic record (Internal DataB folder or USB memory for a USB memory device at the USB-A connection).
- 2 Select the required Kinetics record.

The curve is loaded.

You have the following options:

- You can move the cursor along the curve and have the measurement data for each point displayed (see section 4.9.6)
- With **<PRINT>**, you can output the kinetic curve to a connected printer as a graphic.
- You can store the kinetic curve with **<STORE>** (see section 4.9.4).
- Execute further functions to edit the kinetic record (see section 4.9.6)
- Close the kinetic record with **<ESC>**.

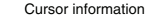
# 4.9.6 Editing a Kinetics record

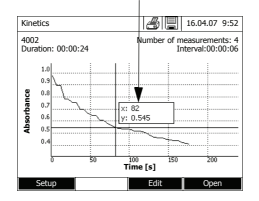
The following functions are available for kinetic records:

- Moving along the curve with the cursor
- Displaying a list with the slopes of the curve for each interval
- Scaling the Y-axis of the diagram

- Combined display of two kinetic records in one graphic
- Display of the difference of two kinetic records

Cursor





The cursor consists of a horizontal and vertical line that cross each other on a point of the curve. A box names the x and y values of the point of the curve.

Move the cursor along the x axis (time axis) with <◀><►>. You can scan and evaluate the curve point after point.

# Slope of the curve & catalytic activity

The function, Slope & catalytic activity indicates the slope of the kinetic curve in the individual intercepts (intervals) of the curve.

An intercept corresponds to the Interval entered in the profile.

- 16.04.07 9:52 Kinetics 0.63 cat Slope [ $\Delta$ /min] ( $\Delta$ / Time 0.000 5 s 0.000 10 s 0.000 15 s 0.000 20 s 0.000 25 s 0.000 30 s
- 1 Indicate the slope of the kinetic curve in the individual intercepts (intervals) of the curve with [Edit]/ Slope & catalytic activity.

If the calculation of the catalytic activity was selected when the profile was created it is displayed here together with the slope.



#### Note

Interval

1

2

3

4

5

6

Back

The Slope & catalytic activity function is only available if the kinetic recording was done in the Absorbance mode.

The displayed slope for an interval is determined as follows, depending on the slope:

Measurements/interval	Slope
1/interval	Slope, converted to the interval, "1 minute"
Max/interval	Slope of the straight line determined by linear regression in an interval, converted to the interval, "1 minute"

**Scaling of the Y-axis** You can manually determine the scaling of the Y-axis with [Setup]/Scaling/ Manual.

*Compare kinetics* For direct comparison, you can load a second kinetic record into the same diagram with *[Edit] / Compare kinetics.* 



## Note

The *Compare kinetics* function can only be carried out if both kinetic records were made in the Absorbance mode.

# Subtract kinetics

You can subtract a stored kinetic record from the current kinetic record with [*Edit*] / *Subtract kinetics*.



#### Note

The *Subtract kinetics* can only be carried out if both kinetic records were made with the following settings:

- Mode: Absorbance
- Measurements/interval:1/interval
- Equal interval

# 4.10 Timer

You can use the timers to remind you by an acoustic signal of a time interval that has expired.

The photometer has two types of timers:

- The User defined timer is a timer that can be freely assigned. The interval and name can be freely set. Only one freely assignable timer is available. It cannot be erased (see section 4.10.1).
- Analysis timer are timers permanently stored in the photometer. The names and intervals of the analysis timers are stored in the method data of a measuring method (*Concentration* mode). The number of available analysis timers corresponds to the number of reaction times prescribed in the analysis instructions of the programmed methods (see section 4.10.2).

The photometer administrates all timers in the timer overview.

The timer overview (the *Timer* menu) is opened with the **<TIMER>** key. The *Timer* menu can be opened in any operating situation. Operation of the timer does not disturb any other functions. The timer overview can be exited with the **<ESC**> key.

When the *Timer* menu is opened for the first time, only the user-defined timer is in the timer overview. You can include analysis timers into the list or remove them according to your requirements (see section 4.10.2).

The timer overview displays the status of each timer and, of a started timer, the remaining time of the specified time interval.

All timers are started manually.

As soon as one single timer has been started the timer symbol appears on the display in all operating modes.

When a timer has been started it is given the timer status, *Active*. When the specified time interval has expired the timer status changes from *Active* to *Expired* and an acoustic signal sounds.

In the timer status *Expired* the acoustic signal sounds until the timer is stopped manually.

After the stop, the timer status changes to *Inactive* and the acoustic signal is switched off.

#### 4.10.1 User defined timer

If you want to manually enter time intervals, use the User defined timer function.

# <TIMER>

Timer			16.04.07 9:52
Designation		Time	Status
User defined time	r	00:15:00	Inactiv
A6/25 - 1		00:15:00	Inactiv
Start	Stop	Edit	Add

The *Timer* menu is open.

- **1** Highlight the User defined timer.
- 2 If necessary, change the name and time of the timer with [Edit].
- **3** Start the highlighted timer with *[Start]*.

The status of the timer is *Active*. When the specified time interval has expired, an acoustic signal sounds and the status changes to *Expired*.

4 Stop the highlighted timer with [Stop].

The status of the timer changes to *Inactive*. The acoustic signal is switched off.

#### 4.10.2 Analysis timer

Between the individual steps of a measurement, reaction times often have to be observed. The length of the reaction time is defined in the relevant analysis instructions.

For all required reaction times, the analysis timers with the corresponding time intervals are stored in the instrument. The names of the analysis timers include the method name and a current number so several timers within a method can be distinguished from each other.

To be able to use an analysis timer for a method you have to load it first in the timer overview.

To do so, first select the required method and then add the available analysis timers to the timer overview so they can be started as necessary.

The timer overview always comprises the free timer and the selected analysis timers.

# <TIMER>

Timer		16.04.07 9:52
Designation	Time	Status
User defined timer	00:15:00	Inactiv
A6/25 - 1	00:15:00	Inactiv

1 Select the required method in the *Concentration* mode.

Manual selection of the method (see section 4.5.5).

2 Open the Timer menu.

The *Timer* menu is open.

3 If necessary, add a new timer to the list with [Add].

#### Note:

The [Add] function key is only displayed if a method is selected for which analysis timers were programmed but are not yet displayed in the list of timers.

- 4 Highlight an analysis timer.
- 5 If necessary, remove the analysis timer from the list with [*Remove*].
- 6 Start the highlighted timer with [Start].

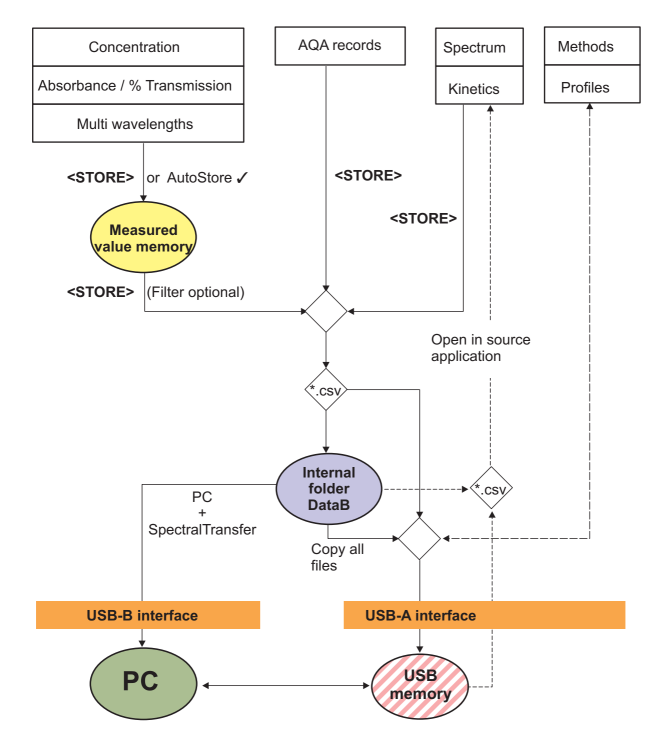
The status of the timer is *Active*. When the specified time interval has expired, an acoustic signal sounds and the status changes to *Expired*.

7 Stop the highlighted timer with [Stop].

The status of the timer changes to *Inactive*. The acoustic signal is switched off.

# 4.11 Memory





Measurement data	Save, back up, export		
Concentration, Absorbance / % Transmis- sion	Measurement datasets of these measuring modes are first stored in the measured value memory of the photometer (1000 memory locations) with <b><store></store></b> or <i>AutoStore</i> .		
Special / Multi wavelengths	The measured value memory is available from the <i>Measurement data memory</i> menu. Here you can view, filter and export into a PC- readable file (*.csv) the stored measurement datasets ( <b><store< b="">&gt;).</store<></b>		
	Csv files of these measuring modes cannot be reimported to the photometer.		
	Measurement datasets of these measuring modes can also be stored to a pdf file (see section 4.11.11).		
Spectrum Kinetics	You can store and export measurement data of these measuring modes directly as a PC-readable file (*.csv) with <b><store></store></b> .		
	Csv files of these measuring modes can be reimported and displayed on the photometer.		
	Measurement data of these measuring modes can also be stored to a pdf file (see section 4.11.11).		
AQA records	You can store and export measurement data of these measuring modes directly as a PC- readable file (*.csv) with <b><store></store></b> .		
	Csv files of records cannot be reimported to the photometer.		
	Measurement data of these measuring modes can also be stored to a pdf file (see section 4.11.11).		
User-defined methods / pro- files	Method data and profile data are stored and exported with the <i>Exchange methods/profiles</i> function in the <b><home></home></b> / <i>General setup</i> menu.		

For each export procedure you can select the location where the PC-readable files (\*.csv, \*.pdf) should be stored: either to the photometer (*Internal DataB folder*) or an external memory (*USB memory*). On an external memory, the data are stored in the pHotoLab\_6600 directory.

The files stored in the photometer (*Internal DataB folder*) can later be transferred to a connected PC or to an external memory (*USB memory*).

#### 4.11.2 Instructions on using USB memory devices

The safety of data stored on USB memory devices depends on the quality of the memory device and the data transmission. Data is stored partly or not at all if for example:

- The power supply of the external memory device is interrupted during the write process, or
- The external memory device is prematurely disconnected from the photometer during the data backup.

To prevent a data loss we recommend the following:

- Save all data internally in the photometer first.
- After performing a backup leave the USB memory device connected to the photometer for some time.
- Check whether the stored data is complete, e.g. on a PC.
- Use the USB memory device for data transport but not for permanent data storage.

	4.11.3 Measurement datasets					
Elements of a	A complete measurement dataset consists of:					
measurement dataset	<ul> <li>Consecutive number (is automatically assigned by the photometer)</li> </ul>					
ualasei	Date/time					
	<ul> <li>Identification (e.g. ID or "AutoStore")</li> </ul>					
	User name					
	• Measured parameter, e.g. method number, dilution, wavelength (depending on the measuring mode)					
	<ul> <li>Measured value with unit and, if necessary, citation form</li> </ul>					
<b>Operations</b> with	Measurement datasets can be					
measurement datasets	<ul> <li>stored (see section 4.11.4)</li> </ul>					
Ualaseis	<ul> <li>displayed and printed (see section 4.11.6)</li> </ul>					
	• filtered, i.e. selected or hidden based on certain criteria (see section 4.11.7 and section 4.11.8)					
	<ul> <li>deleted (see section 4.11.9).</li> </ul>					
the storage is full	You can erase measurement datasets (see section 4.11.9), or overwrite the oldest dataset with the next storing procedure. A security prompt appears before a dataset is overwritten. To backup the measurement data, you can transmit the measurement datasets from the measurement data memory to the internal DataB folder or a USB memory device connected to the USB-A connection and archive them further from there (see section 4.12.3).					
	4.11.4 Saving measurement datasets manually					
	After each measurement, you can store the measurement data manually with the <b><store></store></b> key. It is stored in the measurement data memory. The memory symbol in the header indicates that the measurement data displayed on the screen is ready to be stored. With the measuring modes, <i>Concentration, Absorbance / % Transmission</i> , and <i>Special / Multi wavelengths</i> you have the additional option to automatically store all new measured values at the time of the measurement ( <i>AutoStore</i> , see section 4.11.5).					
Storing with identification (ID)	When storing manually, an input field for the identification (ID) appears after pressing the <b>STORE</b> > key. Here you can enter an individual combination of alphanumeric characters for later easier identification of the measurement datasets. 30 digits are available for this.					

The following measurement data are stored in the measured value memory

lf

automatically (see section 4.11.5) or manually (with the **<STORE>** key, see section 4.11.4):

- Concentration
- Multi wavelengths
- Absorbance / % Transmission

The data stored in the measured value memory can be filtered with filter criteria and then exported to the PC-readable \*.csv format.

The photometer automatically offers a file name during the storage procedure.

Example: Saving data from the measured value memory

<home></home>
Concentration,
Absorbance / % Transmission, o
Special / Multi wavelengths
IO at an I

- [Setup]
   Measurement data
  - memorv

Save (Internal DataB folder)

- 1 If necessary, set the filter criteria with [Setup].
- 2 Open the save dialog with **<STORE>**.

The photometer automatically proposes the location *Internal DataB folder* and a file name.

- **3** If necessary, change the location with *[Location]* (*USB memory*).
- 4 If necessary, change the proposed file name.
- 5 Save the measurement data with **<START·ENTER>**.

The data are stored.

If the photometer (*Internal DataB folder*) is selected as the location, the data can then be copied to a USB memory device (see section 4.12.1).

MData_1.c			

16.04.07 9:52

#### 4.11.5 Saving measurement datasets automatically

For the measuring modes, *Concentration*, *Absorbance / % Transmission*, and *Special / Multi wavelengths* you can record every measured value automatically (*AutoStore*).

All automatically stored measurement datasets are given the ID "AutoStore". The "AutoStore" ID is overwritten if the same measured value is manually stored afterwards (**<STORE>**).

This ensures that every measurement dataset is stored in the data memory only once.

Activating Activate the *AutoStore* function as follows: *AutoStore* 

# <HOME>

Concentration, Absorbance / % Transmission, or Special / Multi wavelengths

– [Setup]

 Measurement data memory
 → Setup

The available functions are displayed.

Select and confirm AutoStore.
 The AutoStore function is active (✓).

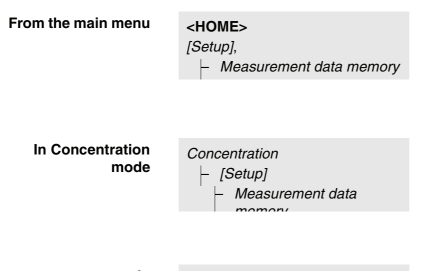


#### Note

The *AutoStore* setting is valid for all three measuring modes, *Concentration*, *Absorbance / % Transmission*, and *Special / Multi wavelengths*.

#### 4.11.6 Displaying measurement data memory

Depending on the operating situation, you can recall the measured value memory as follows:



In Absorbance / % Transmission mode Absorbance / % Transmission [ [Setup] [ Measurement data

In Special / Multi wavelengths mode Special / Multi wavelengths [Setup] Measurement data memory

Each of these options indicates the contents of the measurement data memory as a list as follows.

Measurement data memory		16.04.07 9:52
27.03.07 14:00 3.50 mg/l Ni	Administrator	AutoStore
27.03.07 14:05 3.64 mg/l Ni	Administrator	AutoStore
27.03.07 14:10 3.69 mg/l Ni	Administrator	AutoStore
27.03.07 14:15 3.72 mg/l Ni	Administrator	AutoStore
27.03.07 14:20 3.72 mg/l Ni	Administrator	AutoStore
27.03.07 14:25 3.75 mg/l Ni	Administrator	AutoStore
27.03.07 14:30 3.73 mg/l Ni	Administrator	AutoStore
27.03.07 14:35 3.80 mg/l Ni	Administrator	AutoStore
27.03.07 14:40 3.78 mg/l Ni	Administrator	AutoStore
Filter 🗸		
Memory space usage: 9/		
Setup Single value	Delete	

Measurement datasets can be

If there are more datasets available than can be displayed, the arrows  $\blacktriangle$  and  $\blacktriangledown$  are displayed additionally.

*Filter*✓ indicates that filter settings are active. In this case, only those datasets are displayed that correspond to the selected filter criteria (see section 4.11.7).

#### Options

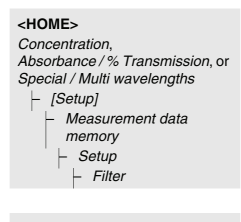
- displayed in short form as a list or in details as individual values ([List] <-> [Single value])
- filtered (see section 4.11.7 and section 4.11.8)
- deleted (see section 4.11.9).
- with <STORE>, you can store the entire displayed list as a \*.csv file in the internal DataB folder or on a USB memory device connected to the USB-A connection. The filter settings apply to the storing process. You can freely select the file name. Thus you can, e. g. store in a separate file and systematically archive measurement data of a certain period.
- with **<PRINT>**, the entire displayed list can be printed. The filter settings apply to the print process.

#### 4.11.7 Filtering measurement datasets

The functions to display, delete and download stored measurement datasets refer to all stored measurement datasets that correspond to the specified filter criteria.

#### Filter criteria The following filter criteria can be set:

- *Mode* (measured parameter)
- User
- *ID* (identification)
- *Date* (date *from* ... *to* ...)
- *Method* (for the measured parameters, *Concentration* and *Multi wavelength*)





- Mode (Concentration, etc.)
- User (**<A...9**>)
- *ID* (**<A...9**>)
- Date (from ... to ...)
- Method

Measurement data memory	o i	16.04.07 9:52
27.03.07 14:00 3.50 mg/l Ni	Administrator	AutoStore
27.03.07 14:05 3.64 mg/l Ni	Administrator	AutoStore
27.03.07 14:10 3.69 mg/l Ni	Administrator	AutoStore
27.03.07 14:15 3.72 mg/l Ni	Administrator	AutoStore
27.03.07 14:20 3.72 mg/l Ni	Administrator	AutoStore
27.03.07 14:25 3.75 mg/l Ni	Administrator	AutoStore
27.03.07 14:30 3.73 mg/l Ni	Administrator	AutoStore
27.03.07 14:35 3.80 mg/l Ni	Administrator	AutoStore
27.03.07 14:40 3.78 mg/l Ni	Administrator	AutoStore
Filter 🗸		
Memory space usage: 9/		
Setup Single value	Delete	

The filter setting menu is displayed.

- **1** Set the filter criteria.
- 2 If necessary, deactivate any selected filter criteria with [Reset entry].
- **3** Confirm the filter selection with *[Apply]*.

The *Measurement data memory* list is displayed.

The following information is displayed additionally:

- Current memory occupancy
- Active filter criteria (*Filter* ✓)



#### Note

Alternatively, you can hide measurement datasets that meet the specified filter criteria with the Selected values: invert selection function (see section 4.11.8).

#### 4.11.8 Inverting filters

With the Selected values: invert selection function you can hide all measurement datasets that correspond to the specified criteria of the filter (see section 4.11.7).



#### Note

You can use this function to select and delete measurement datasets no longer used.

#### <HOME> Concentration. Absorbance / % Transmission, or Special / Multi wavelengths [Setup] Measurement data memory Setup Selected values: invert selection 16.04.07 9:52 Measurement data memory AutoStore 27.03.07 14:00 3.50 mg/l Ni Administrator 27.03.07 14:05 3.64 mg/l Ni 27.03.07 14:10 3.69 mg/l Ni Administrator AutoStore AutoStore Administrator 27.03.07 14:15 3.72 mg/l Ni AutoStore Administrator AutoStore 27.03.07 14:20 3.72 mg/l Ni Administrator AutoStore 27.03.07 14:25 3.75 mg/l Ni Administrator

Administrator

Administrator

Administrator

Delete

AutoStore

AutoStore

AutoStore

27.03.07 14:30 3.73 mg/l Ni

27.03.07 14:35 3.80 mg/l Ni

27.03.07 14:40 3.78 mg/l Ni

Memory space usage: 9/ Setup

Single value

Filter 🗸

The Measurement data memory list is displayed. All measurement datasets corresponding to the filter criteria are hidden.

#### 4.11.9 Erasing stored measurement datasets

If you no longer need any stored measurement datasets, you can erase them individually or altogether.

<home> Concentration, Absorbance / % Transmission, or Special / Multi wavelengths [Setup] Measurement data memory</home>				
Measurement data memory	6 E	16.04.07 9:52		
27.03.07 14:00 3.50 mg/l Ni 27.03.07 14:05 3.64 mg/l Ni 27.03.07 14:15 3.69 mg/l Ni 27.03.07 14:15 3.72 mg/l Ni 27.03.07 14:20 3.72 mg/l Ni 27.03.07 14:25 3.75 mg/l Ni 27.03.07 14:35 3.80 mg/l Ni 27.03.07 14:35 3.80 mg/l Ni 27.03.07 14:40 3.78 mg/l Ni	Administrator Administrator Administrator Administrator Administrator Administrator Administrator Administrator	AutoStore AutoStore AutoStore AutoStore AutoStore AutoStore AutoStore AutoStore AutoStore AutoStore		
Filter ✓ Memory space usage: 9/ Setup Single value	Delete			

The *Measurement data memory* list is displayed.

The filter settings used last are active.

#### Erasure functions

The following erasure functions are available.

- Erasing an individual measurement dataset
- Erasing all measurement datasets **1** of the displayed list
- **1** Highlight a measurement dataset.
- 2 Remove the highlighted measurement dataset with [Delete].
  - Open the setting menu with [Setup].
  - 2 Select and confirm *Delete memory (selected values only)*.

All measurement datasets corresponding to the current filter criteria are erased.

or

• Erasing all measurement datasets

Select and confirm *Delete memory (all values)*.

All measurement datasets are erased.

#### 4.11.10 Saving kinetic recordings, spectra and AQA files

After the following measurements, the *Save* dialog opens and prompts you to save the data in a \*.csv file:

- Kinetics
- Spectrum
- AQA3/MatrixCheck

If the data are not saved in \*.csv format, they are lost when the measuring mode is terminated.

# Note

During a kinetic recording, the current measurement is always saved in the file, "KineticsBackup.csv" for safety reasons.

#### 4.11.11 Saving data as a pdf file

All data that can be printed (printer symbol on the display) can also be saved as a pdf file. The pdf file contains the data that are also output to a USB printer. Kinetic recordings and spectra are stored in the pdf file as a graphic.

To store data as a pdf file, use the **<PRINT>** key as for printing. When doing so, the pdf print has to be set as the printer in the menu, **<HOME>**/*General setup*/*Data transfer*/*Printer*/*Function of PRINT key*.

Subsequently, enter a file name and select the storage location (internally folder DataB or USB memory device).

# 4.12 Saving / exporting files

If you want to back up or process measurement data files outside the photometer, you can copy them to external media.



#### Note

Please follow the instructions on using USB memory devices (see section 4.11.2).

#### 4.12.1 Copying all measurement data files to a USB memory device

Even if no PC is directly connected to the photometer, you can very simply transfer all measurement data files from the photometer (*Internal DataB folder*) to a connected USB memory device.

<home></home>
---------------

[Setup]

 Save data to USB memory device

When the data saving procedure is finished, a message appears.

1 Confirm the message with **<STORE>**.

All measurement data files from the photometer (*Internal DataB folder*) have been transferred to the USB memory device.

The complete folder structure from the photometer is created on the USB memory device. The individual measurement data files are stored in subfolders sorted by measurement data types.

# 4.12.2 Copying user-defined methods / profiles to a USB memory device

#### <HOME>

[Setup]

Exchange methods/profiles
 /Store to USB memory
 device

A list is displayed that includes all user-defined methods and profiles available on the photometer. All methods and profiles are checked off with a checkmark. All methods and profiles checked off are saved.

 If necessary, select individual methods/profiles with <▲><▼> and remove the checkmark with<START.ENTER>.

These methods/profiles will not be saved.

2 Start the save process with [Store].

A message appears when the data have been saved.

3 Confirm the message with **<START**.**ENTER**>.

The save process is completed. The data are stored in the *Exchange\_Method\_Profile* folder on the USB memory device. The individual files with the methods/ profiles are in subfolders.

Already existing files with identical names are overwritten without confirmation prompt.

#### 4.12.3 Copying files to a PC

You can copy from the photometer to a PC the following data:

- Measurement data
- Spectra
- Kinetic recordings
- AQA records
- User-defined methods
- Profiles

After saving measurement data in \*.csv or \*.pdf format, you can copy them to a PC. Measurement data in csv format can be directly imported to and processed in spreadsheets such as Microsoft<sup>®</sup> Excel<sup>®</sup>.

#### Note

Depending on the country variant, some spreadsheet programs require a certain decimal separator for the correct import of numerical values (comma or point). The decimal separator can be selected in the following menu:

<HOME> -> General setup -> Data transfer/Printer -> Decimal separator for csv-Files.

Files containing measurement data can be copied to a PC in the following ways:

- By using a USB memory device as a temporary storage (see section and section 4.12.1). Subsequently, you can connect the USB memory device to a PC and read out the data.
- By means of the "SpectralTransfer" program (see operating manual of the "SpectralTransfer" program). The "SpectralTransfer" program and the corresponding operating manual is provided on the enclosed CD-ROM.
- Using the "pHotoLab Data spectral" program (see operating manual of the "pHotoLab Data spectral" program). The "pHotoLab Data spectral" program is available as an accessory.

# 4.13 Importing files

You can import to a pHotoLab<sup>®</sup> 6xxx spectrophotometer the data that were created with the same or another pHotoLab<sup>®</sup> 6xxx spectrophotometer, and the data that were saved to a USB memory device or a PC.

You can import the following data:

- Spectra
- Kinetic recordings
- User-defined methods
- Profiles

# 4.13.1 Importing spectra or kinetic recordings from a USB memory device

You can import to the photometer any spectrum or kinetic recording by opening an externally stored spectrum or kinetic recording with the Open function of the photometer.

#### 4.13.2 Importing methods / profiles from a USB memory device



#### Note

When importing methods make sure that your photometer supports the wavelengths of the imported methods.

#### <HOME>

[Setup]

 Exchange methods/profiles
 /Import from USB memory device

> A list is displayed including all user-defined methods and profiles stored in the corresponding subfolders of the Exchange directory on the USB memory device. All methods and profiles are checked off with a checkmark. All methods and profiles checked off are imported.

 If necessary, select individual methods/profiles with <▲><▼> and remove the checkmark with<START.ENTER>.

These methods / profiles are excluded from importing.

2 Start the import with [Import].

A confirmation prompt appears before any data on the photometer are overwritten.

A message appears when the data have been imported.

3 Confirm the message with **<START**.**ENTER**>.

The import is completed. The imported methods / profiles are available on the photometer.

# 4.13.3 Importing files from a PC

You can import files from the PC to the photometer in the following ways:

- By means of the "SpectralTransfer" program (user-defined methods only) (see operating manual of the "SpectralTransfer" program). The "SpectralTransfer" program and the corresponding operating manual is provided on the enclosed CD-ROM.
- Using the "pHotoLab Data spectral" program (see operating manual of the "pHotoLab Data spectral" program). The "pHotoLab Data spectral" program is available as an accessory.

# 4.14 Printing data (RS232, USB)

#### 4.14.1 Printer and terminal programs

**Usable printers** Data can be printed with the following printers:

- Matrix printer connected to the RS232 interface
- Standard printer (ink or laser) connected to the USB-A interface



Note

Suitable are all printers that can interpret the PCL-3 printer control language.

The printer symbol 
indicates that the display contents can be printed. To print, press <PRINT>.

**PC + terminal** program The data can also be received by a PC with terminal program instead of a printer. For this the PC is connected to the photometer via the RS232 interface. The output is identical to that of a matrix printer.

**pdf file** As an alternative, you can also output the print data to a pdf file.



#### Note

In den following paragraphs, "Print" means:

- output to a printer (RS232 interface)
- output to a PC + terminal program (RS232 interface)
- output to a USB printer
- output to a pdf file.

#### 4.14.2 Settings for data transmission

Settings are possible for the data transmission to a printer or PC.

Decimal separators<br/>for CSV filesFor the output of CSV files you can select either a comma or a point as the<br/>decimal separator. The setting is made in the following menu:

<HOME> -> General setup -> Data transfer/Printer -> Decimal separator for csv-Files -> Comma (12,34) or Point (12.34).

Short and long<br/>versionWhen printing measurement datasets, you can select a short or long version<br/>with different information contents. The setting is made in the following menu:

<HOME> -> General setup -> Data transfer/Printer -> Data format (print) -> Short or Extended.

**Baud rate for RS232 interface** The baud rate can be set for printers that are operated at the RS232 interface. Adjust the photoLab<sup>®</sup> 6600 UV-VIS to the baud rate of the printer. The setting is made in the following menu:

<HOME> -> General setup -> Data transfer/Printer -> Baudrate for RS232 printer -> 1200 ... 19200.

#### **Printer** Here you can set which function is assigned to the **<PRINT>** key:

- Output to a USB printer
- Output as pdf file

The setting is made in the following menu:

<HOME> -> General setup -> Data transfer/Printer -> Function of PRINT key -> USB printer or PDF file.

#### 4.14.3 Printing measurement datasets

This section describes how to print measurement datasets of the measuring modes, *Concentration*, *Absorbance / % Transmission*, and *Special / Multi wavelengths*.

By means of sample printouts, the printed information is described below:

Concentration

```
21 05.06.07 14:05:41 C4/25 844 mg/l COD Inlet
Administrator 0.005 02.06.07 11:02:13 2 AQA1: 9 AQA2: 14
```

and Special / Multi wavelengths mode

Structure of the lines from left to right:

1st line:

[Consecutive no.] [Date] [Time] [Method name] [Measured value] [Unit] [Citation form] [Dilution] [ID or "AutoStore"]

2nd line (long version only):

[User] [Reagent blank value] [Date of blank value measurement] [Time of blank value measurement] [Lot ID of blank value measurement] [AQA1: label] [AQA1: record no.] [AQA2: label] [AQA2: record no]



# Note

Optional elements (e.g. dilution or ID) are output only if they were really used for measurement or storage.



14 05.06.07 11:25:01 445 nm 0.609 Absorbance AutoStore Administrator 0.133 02.06.07 09:59:01 AQA1: 9

Structure of the lines from left to right:

mode 1st line:

[Consecutive no.] [Date] [Time] [Wavelength] [Measured value] ["Absorbance" or "Transmission" mode ] [ID or "AutoStore"]

2nd line (long version only):

[User] [Value of reference absorbance] [Date of reference absorbance] [Time of reference absorbance] [AQA1: label] [AQA1: record no.]



#### Note

Optional elements (e.g. ID or reference absorbance) are output only if they were really used for measurement or storage.

#### 4.14.4 Printing Kinetics records

# Sample printout

Г

photoLab 05.06.07 320 nm		09130512	1.30-WTW-1.60	Administrator
Time [s] 6 17 25 35  (etc.)	(	Absorbance ),092 ),077 ),073 ),077	2	

# Structure of the lines from left to right

1st line:

[Instrument type] [Series number] [Version of meter software and method data] [User]

2nd line:

[Start date] [Start time]

3rd line:

[Wavelength]

6th and following lines:

Passed time with related measured value



#### Note

If you output a kinetic recording to a USB printer or pdf file (not to the RS232 interface), the current graphic representation is shown on the display.

# 4.14.5 Printing spectra

#### Sample printout

photoLab 6600 UV-VIS 091305 07.06.07 09:47:00	12 1.30-WTW-1.60	Administrator
Wavelength [nm] 320 321 322 323 324 		Absorbance 0,238 0,240 0,241 0,240 0,239 
(etc.)		

#### Structure of the lines from left to right

# 1st line:

[Instrument type] [Series number] [Version of meter software and method data] [User]

# 2nd line:

[Start date] [Start time] [Wavelength]

5th and following lines:

Wavelength with related measured value



#### Note

If you output a spectrum to a USB printer or pdf file (not to the RS232 interface), the current graphic representation is shown on the display.

#### 4.15 Analytical quality assurance (AQA)

#### 4.15.1 **General information**

The target of the analytical quality assurance (AQA) is to secure correct and precise measurement results.

Settings for AQA checks are only available for users of the user group, administrator.

Every registered user can carry out the AQA check (see also section 4.16.1).

Analytical quality assurance (AQA) can be carried out in two steps independent of each other:

- AQA1: Monitoring of the photometer
- AQA2: Monitoring of the total system. It comprises the photometer, the used test, the accessories and the user's way of working.

The monitoring includes a check procedure that has to be successfully repeated by the user within a certain period (AQA interval).



#### Note

Note

The AQA monitoring is not active in the delivery condition.

AQA in measured value documentation

All values that are measured after a passed check and within the AQA interval are given the addition *Protocol ID* in the measured value documentation. This addition is used to identify the relevant AQA inspection record.

#### 4.15.2 Photometer monitoring (AQA1)

At least one set of test standards such as PhotoCheck or  $CertiPUR^{\mathbb{B}}$  is required for the photometer monitoring.

The administrator specifies which test standard has to be used as the minimum requirement for the AQA1 monitoring.

The extent of the monitoring can be enlarged with further test standards.



#### Note

Settings for AQA checks are only available for users of the user group, administrator. Event registered user can carry out the AQA check (see also section 4.16.1)

Every registered user can carry out the AQA check (see also section 4.16.1).

#### Spectroquant<sup>®</sup> PhotoCheck

The PhotoCheck consists of 12 test standards in duplicate, 2 zero cells and 2 cells to check the barcode reader. Each PhotoCheck package contains a lot dependent test certificate with all nominal values (absorbances) and tolerances of the test standards. These values are entered in the photometer during the configuration of the AQA1 check.

CertiPUR<sup>®</sup> test standards

Each CertiPUR<sup>®</sup> standard is provided with a lot dependent test certificate with all nominal values (absorbances) and tolerances of the test standards. The values were preset in the factory.



#### Note

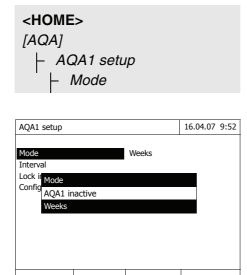
Observe the shelf life of the test standards. The values in the photometer always have to be checked when a new package of test standard is used. If necessary, adjust the values at the photometer.

Overview of the photometer monitoring Photometer monitoring (AQA1) consists of the following parts:

- Configuring settings in the AQA1 setup menu.
  - Activate AQA1
  - Specify AQA1 Interval
  - Activate/deactivate the meter lock for missing or expired AQA1 check
  - Define the extent of the AQA1 monitoring by activating or deactivating the individual test standards.
  - Enter the nominal values, tolerances and lot numbers for the individual test standards
- Carrying out the AQA1 check. The photometer compares the results with the nominal values while taking into account the tolerances.

The steps are described in detail below.

#### Activating AQA1 The AQA1 monitoring is activated in the *Mode* menu:



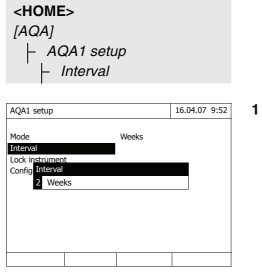
Select and confirm Weeks.

AQA1 is active. The *Interval* setting indicates *Weeks* as the interval unit.

# Defining the AQA1 Interval

The AQA1 Interval defines the interval between two AQA1 checks. When an interval has expired, the following consequences become effective:

- Warning and loss of the AQA1 labeling
- Locking of the photometer against all measurements (if activated).



 Enter a numeric value (2 to 52 weeks) (<0...9>) and confirm
 The *Interval* defined for the AQA1 check is active.

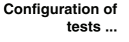
# Configuring the lock of the photometer

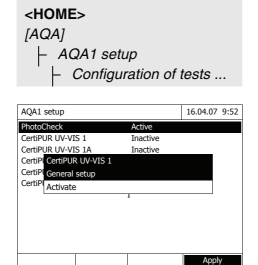
Here you configure whether or not the photometer will be locked against all measurements if there is no valid AQA1 check or the interval for the AQA1 check has expired.

<home></home>
[AQA]
– AQA1 setup
├ AQA1 setup ├ Lock instrument

AQA1	setup	16.04.07	9:52
Mode Interva	Lock instrument		
Config	Should the instrument be locked for measurements if AQA1 check is inva has expired?	further alid or	
	No		
	Yes		

1 Select and confirm *Yes*. The photometer is locked against all measurements if the AQA1 check is invalid or the AQA1 interval has expired.





All possible test standards or test standard sets are listed.

- 1 Select and confirm a test standard or test standard set.
- 2 Adjust and confirm the extent of the monitoring with *Activate* or *Deactivate*.
- **3** Confirm the test standard (set) once again.
- 4 Switch to the adjustment of the nominal values and tolerances with *Setup*.

PhotoCheck		16.04.07 9:52
Lot number:		10010115
		HC616115
Use by		16.04.2008
	Target value	Tolerance
445/1	0.196	± 0.020
445/2	0.500	± 0.030
445/3	0.998	± 0.040
445/4	1.508	± 0.050
525/1	0.197	± 0.020
525/2	0.495	± 0.030
525/3	0.992	± 0.040
525/4	1.496	± 0.050
		Apply

Example, PhotoCheck:

- 5 Using <▲><♥> and <◀><▶>, select the Lot number, Target value or Tolerance entries and open them for editing with <START.ENTER>.
- 6 Enter and confirm the required value (<0...9>)
- 7 Accept all values with [Apply].

# Carrying out the AQA1 check (example: PhotoCheck)

The AQA1 check comprises the check with all test standards activated in the menu, *AQA menu / AQA1 setup / Configuration of tests ...* for AQA1 (see page 136).

First, a zero adjustment for all wavelengths takes place. Subsequently, the first individual checks with the selected test standards take place (e.g. PhotoCheck).

<home></home>	
[AQA]	
– AQA1 check	

PhotoCheck	16.04.07	9:52
Reference measurement		
Please insert zero cell (distilled water).		

The photometer is ready for the zero adjustment.

 Insert the zero cell. The cell is automatically recognized and the zero adjustment is started for all wavelengths.

After the successful zero adjustment, the photometer is ready to measure the PhotoCheck test standard 445/1.

PhotoCheck 445/1	16.04.07	9:52
Please insert PhotoCheck 445/1		

2 Insert the cell. The cuvette is automatically recognized and the measurement started.

> After measuring, the measurement result, Target value, Tolerance and an evaluation (OK or failed) are displayed.

> The photometer offers to repeat the measurement if the check failed.

> If the check was successful, the measurement of the next Photo-Check test standard, e.g. 445/2, appears on the display.

- Measure all test standards in the same way.
   After all test standards are successfully measured, the check is passed.
- **Test record** A test record is displayed after the check. It can be printed and stored as a file (in the internal DataB folder or USB memory device at the USB-A connection, see section 4.11.1).

Sample printout:

photoLab 6600 AQA1 Protocol ID Executed by: Executed Valid until:	UV-VIS 09130512		Administrator OK 9 Administrator 22.05.2007 26.06.2007
PhotoCheck 445-1 445-2 445-3 445-4 525-1  (etc.)	OC479094 0.200 +- : 0.500 +- : 1.000 +- : 1.500 +- : 0.200 +- :	100 200 200 200	OK 0.192 0.511 1.006 1.526 0.247 



#### Note

Afterwards you can view the last AQA1 test record under AQA1 info.

#### 4.15.3 Total system monitoring (AQA2)

For the total system monitoring, standard solutions with a defined analyte content are required (preferably certified Spectroquant<sup>®</sup> CombiCheck standards).



#### Note

Settings for AQA checks are only available for users of the user group, administrator. The AQA check can be carried out by any registered user.

Spectroquant<sup>®</sup> CombiCheck Spectroquant<sup>®</sup> CombiCheck standards are multiparameter standards ready to use, i. e. they can be used for several test sets (methods).

In addition to the CombiCheck standards, one parameter standard solutions can also be used. They are prepared by dilution to the respective end concentration. The end concentration should be in the middle of the measuring range.



#### Note

The suitable CombiCheck standards and one parameter standards are listed in the WTW catalog or on the Internet.

Total system monitoring (AQA2) consists of the following parts:

- Configuring the general settings in the AQA2 setup menu.
  - Activate AQA2
  - Select the AQA2 interval unit (Weeks or Measurements)
  - Activate/deactivate the measurement lock for missing or expired AQA2 check. The measurement lock is effective for all methods that were activated for AQA2 monitoring
- Selecting the method to be activated for AQA2
- Configuring the method-specific settings in the AQA2 setup menu.
  - Activate AQA2
  - Specify AQA2 Interval
  - Enter the nominal value, tolerance and designation (standard ID) for the test standard
- Carrying out the AQA2 check. During the check the test is carried out with the standard solution as the sample while the other conditions are the same. The photometer compares the result with the nominal value while taking the tolerance into account.

The steps are described in detail below.

Overview of the total system monitoring

General AQA2 settings

<home> [AQA]  - AQA2 setu</home>	ıp		
AQA2 setup		16.04.07	9:52
Mode Lock methods Method	Weeks Yes		
Method list			

- 1 Select and confirm *Mode*. The *Mode* selection field pops up.
- 2 Select and confirm *Weeks* or *Measurements*.

AQA2 is active. For all methods, the AQA2 intervals are entered either in weeks or number of measurements.

**3** Accept the general settings with *[Apply]*.

I	•

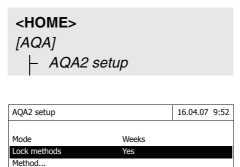
# Note

Method list

When the mode (*Weeks* or *Measurements*) is changed, all AQA2 intervals are reset to the preset values.

#### Locking the method

Here you configure whether or not a method will be locked against measurement if there is no valid AQA2 check or the interval for the AQA2 check has expired.



- 1 Select and confirm *Lock methods*.
- 2 Select and confirm *Yes*.

The method lock is enabled.

Each method will be locked if the AQA2 check is invalid or the AQA2 interval has expired.

Activating AQA2 monitoring for a method

<home></home>
[AQA]
– AQA2 setup
– Method

AQA2 setup	16.04.07 9:52
Method	3: A6/25
AQA2	AQA2 active
Interval	12 Weeks
Target value	4.00 mg/l NH <sub>4</sub> -N
Tolerance	0.50 mg/l NH <sub>4</sub> -N
Standard ID	•
Method list	

- 1 Select a method (see section 4.5.3).
- 2 Select and confirm AQA2.
- **3** Select and confirm *AQA2 active*. AQA2 is active for this method.

Defining the AQA2 Interval, nominal value and tolerance The AQA2 Interval defines the interval between two AQA2 checks. When an interval has expired, the following consequences become effective:

- Warning and loss of the AQA2 labeling
- Locking of the method against measurement (if activated).

Setting range:

1 to 12 weeks (default: 12 weeks) or

1 to 10000 measurements (default: 200 measurements)



#### Note

The unit of the AQA2 interval (Weeks or Measurements) is defined in the line, *Mode* (see page 140).

AQA2 setup	16.04.07 9:52
Method	3: A6/25
AQA2	AQA2 active
Interval	12 Weeks
Target value	4.00 mg/l NH <sub>4</sub> -N
Tolerance	0.50 mg/l NH⊿-N
Standard ID	- <b>-</b>
Method list	

- 4 Select the *Interval* and enter the AQA2 interval.
- 5 If necessary, adjust the values for *Target value* and *Tolerance*.

6 Optional: Select *Standard ID* and enter a designation. The designation is recorded in the AQA2 documentation.

Repeat the steps 1 to 8 if you want to configure further tests for AQA2.

Carrying out the AQA2 check for a method

<HOME> [AQA] - AQA2 check

AQA2 check			16.04.07	9:52
Target value	2.00			
	measurement, Il or press <st <="" th=""><td>ART/ENTER&gt;</td><td></td><td></td></st>	ART/ENTER>		
3: A6/25			N	H <b>4</b> -N
16 mm			0.20 - 8.00	mg/l

- 1 Carry out the check like a normal measurement (see section 4.5.1 to 4.5.3).
- 2 Insert the cell or start measurement with <START·ENTER>.

After the measurement is completed, the result and its evaluation are displayed.

If the check failed, it is possible to repeat the measurement.

If the check was successful, the *AQA2 check* function is finished.

**Test record** A test record is displayed after the check. It can be printed and stored as a file (in the internal DataB folder or USB memory device at the USB-A connection, see section 4.11.1).

Sample printout:

```
photoLab 6600 UV-VIS 09130512 1.30-WTW-1.60 Administrator
AQA2
                   OK
Protocol ID
                   32
Executed by:
                    Administrator
Executed
                    21.05.2007
Valid until:
                   13.08.2007
Method
                    6: P6/25 PO4-P
Standard ID
                   CC10 OC557775
Target value
                    0.80 +- 0.08 mg/l
Measured value
                    0.84
                           mg/l
```



#### Note

Later you can view the last AQA2 test records for all methods monitored with AQA2 under AQA2 info.

#### 4.15.4 AQA3/MatrixCheck

The *MatrixCheck* is used to check if the photometric determination is disturbed by other substances present in the sample (sample matrix). The MatrixCheck can be carried out by spiking or diluting:

The photometer enables a simplified MatrixCheck with the aid of the Spectroquant<sup>®</sup> CombiCheck R-2 addition solution. The MatrixCheck can be carried out immediately. The volumes required for the sample and standards are displayed on the screen. The MatrixCheck is then carried out with a single spiking.

For the MatrixCheck with a standard of your own, however, you can enter the number of spikings or dilutions yourself (max. 3).



#### Note

Settings for AQA checks are only available for users of the user group, administrator. The AQA check can be carried out by any user.

# MatrixCheck by<br/>spikingFor the MatrixCheck by spiking, the photometric determination is repeated<br/>after a defined amount of analyte has been added to the test sample in the<br/>form of standard solutions.

The nominal value for the determination is calculated from the added amount of analyte, provided that there is no disturbance due to the sample matrix. After the photometric determination the measured value is compared to the nominal value and the recovery rate is calculated. A matrix disturbance is likely if the recovery rate is less than 90 % or more than 110 %.

# MatrixCheck by<br/>dilutingFor the MatrixCheck by dilution, the photometric determination is repeated<br/>after the test sample has been diluted with distilled water.

The nominal value for the determination is calculated from the dilution, provided that there is no disturbance due to the sample matrix. After the photometric determination the measured value is compared to the nominal value and the recovery rate is calculated. A matrix disturbance is likely if the recovery rate is less than 90 % or more than 110 %.

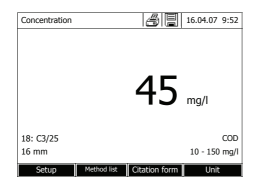
Practical instructions	<ul> <li>After evaluating the measured value of the sample the photometer suggests for the MatrixCheck to spike or dilute the sample and standard with suitable volumes.</li> <li>You can change the suggested values of the volumes for the sample and standard. The photometer checks your entries and informs you of errors (e.g. if a nominal value is outside the measuring range of the test). For each spiking or dilution, the relevant nominal concentration value is displayed.</li> </ul>			
	<ul> <li>To be able to reliably recognize matrix effects by <u>spiking</u>, the <u>volume</u> increase after spiking should be <u>small</u>.</li> </ul>			
	<ul> <li>To be able to reliably recognize matrix effects by <u>diluting</u>, the <u>dilution factor</u> should be <u>high</u>.</li> </ul>			
	<ul> <li>You can carry out the MatrixCheck as a series of measurements, consist- ing of up to three determinations with different spiking volumes or dilutions respectively.</li> </ul>			
	<ul> <li>Prepare all test sample solutions simultaneously at the beginning of the series of measurements.</li> </ul>			
Overview of the AQA3/MatrixCheck	The MatrixCheck consists of the following parts:			
	<ul> <li>Configuring settings in the AQA3/MatrixCheck setup menu.</li> </ul>			
	<ul> <li>Specify the maximum deviation from the nominal value after spiking or diluting (default setting: 10%)</li> </ul>			
	<ul> <li>Carrying out the AQA3 / MatrixCheck</li> </ul>			
Specifying the maximum deviation from the nominal value	The assessment of the recovery rate is determined with the maximum deviation from the nominal value. The assessment of the recovery rate is displayed next to the recovery rate after the check has been carried out.			
	<home> Concentration [- [Setup] [- AQA [- AQA3/MatrixCheck</home>			

setup

- Maximum difference

AQA3/MatrixCheck setup	16.04.07 9:52
Maximum difference	10%
Maximum difference	
10.0 %	
	<u> </u>
	1

Carrying out the MatrixCheck



Dilute		Delete	Next
10	0	45	
10	0	45	
10	0	45	
Sample [ml]	Standard [ml]	Target value [mg/l]	
Standard ID Standard co		0 0 mg/ICOD	
Method Sample con	centration	1: C3/25 45 mg/ICOD	
MatrixChecl	k (Spike)		16.04.07 9:5

- 1 Enter and confirm a numerical value.
  - The setting is active.
- 2 Exit the menu with **<ESC>**.

- **1** Measure the original sample without spiking or diluting it (see section 4.5.1 to 4.5.3).
- 2 The measured value is displayed.
- **3** Open the setting menu with *[Setup].*
- 4 Select and confirm AQA.
- 5 If necessary, check the settings in the menu, *AQA3/MatrixCheck setup*.
- 6 Select and confirm *AQA3/Matrix*-*Check*.

The display for the MatrixCheck opens up.

If the spiking with the standard values of the CombiCheck R-2 suggested by the photometer would cause the measuring range to be exceeded, the MatrixCheck by diluting is automatically suggested.



## Note

The following description shows the proceeding for the MatrixCheck by spiking. To switch to the MatrixCheck by dilution, use the *[Dilute]* function key. The proceeding is similar there, but the entry of the Standard ID and Standard concentration is not applicable.

MatrixCheck (Spike)			16.04.07 9:52
Method Sample cond	entration	1: C3/25 45 mg/ICOD	
Standard ID Standard co	ncentration	COD 1500 400 mg/ICOD	
Sample [ml]	Standard [ml]	Target value [mg/l]	
10 10 10	0.5 1 1.5	62 77 91	
Dilute		Delete	Next

7 In the *Standard ID* entry field, select the simplified MatrixCheck with the CombiCheck standard solution or enter a designation for another standard solution used.

If the CombiCheck is selected, no more entries are required (continue with step 10).

8 Enter the concentration of the used standard solution in the *Standard concentration* entry field.

#### Specifying the series of measurements:

- 9 Enter the volumes of sample and standard of the individual test sample solutions in the columns, *Sample [ml]* and *Standard [ml]*. The nominal value is calculated after each entry.
  - You can delete a measurement from the series of measurements with [Delete].

Note that all nominal values have to be within the measuring range of the test.

**10** Using *[Next]*, accept all entries on the page and switch to the next page. The entries are checked by the photometer.

The photometer is ready to carry out the series of measurements.

#### Carrying out the series of measurements:

According to the program, the samples are measured top down. You can, however, select the samples yourself and thus change the order with < > < V >.

**11** Use *[Measurement]* to proceed to the measurement of the (first) sample.

MatrixCheck (Spike)				16.04.07 9:52	
Method 1: C3/25 Sample concentration 45 mg/ICOD					
Sample [ml]	Stand [ml]		arget value mg/l]	nom [mg	ninal //]
10	0.5	6	52	58	
10	1	7	7		
10	1.5	9	1		
Back	K	Measurem	ne		Complete

MatrixCheck			16.04.07	9:52
Method Sample concent Sample Standard	ration	1: C3/25 45 mg/ICOD 10 ml 0.5 ml		
	easurement, or press <st< th=""><td>ART/ENTER&gt;</td><td></td><td></td></st<>	ART/ENTER>		
16 mm				-
Back				

MatrixCh	eck			Ð		16.04.07 9:52
Method Sample concentration				1: C3/25 45 mg/IC		
Sample [ml]	Stan [ml]	dard	Target [mg/l]		nom [mg	ninal //I]
10	0.5		62		58	94 % 🗸
10	1		77			
10	1.5		91			
Back	ζ	Measu	reme			Complete

The measurement display appears.

**12** Insert the cell with the respective sample.

The sample is measured.

After the measurement, the recovery rate is displayed in the right table column.

The assessment of the recovery rate is displayed next to the recovery rate ( $\checkmark$  or X ).

The criteria for the assessment are determined in the menu, *AQA3/MatrixCheck setup / Maximum difference*.

- **13** If necessary, repeat the steps 11 and 12 for the remaining samples.
- 14 Use [Complete] to complete the MatrixCheck.

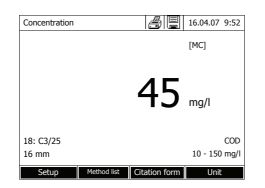
The Save dialog box pops up.

**15** If necessary, change the storage location with *[Location]*: *Internal DataB folder*. Exchange folder in the instrument or

USB memory:

USB memory device connected at the USB-A connection.

- **16** If necessary, change the file name.
- 17 Save the file with <START.ENTER>.



The display returns to the measured value display of the original sample without spiking / dilution.

The [MC] status indicator is displayed. A MatrixCheck was carried out for this measured value.

**Test record** The result of the MatrixCheck is displayed in a test record. You can print this record and save it as a file.

To save the file in the photometer, select the *Internal DataB folder* as the location. To save the file in an external USB memory device at the USB-A connection, select *USB memory* as the location (see section 4.11.1).

Sample printout:

photoLab 6600 UV-VI	S 09130512 1.30-WTW-1.60	Administrator
- MatrixCheck	OK	
Protocol ID	7	
Method	1: C3/25 COD	
Sample concentration	n45 mg/lCOD	
Standard ID	COD 1500	
Standard concentrat	ion400 mg/lCOD	
Sample	Standard	Target value
Actual value		
ml	ml	mg/lmg/l
10	0.5	625894% OK
10	1	777192% OK

## 4.16 User management

The functions of the user management are only available for users of the user group, *Administrator*.

An administrator can

- activate or deactivate the user management for the meter
- create, change or delete individual user accounts.

#### 4.16.1 User levels and user rights

The photoLab<sup>®</sup> 6600 UV-VIS allows the management of up to 100 users. Every user is member of a user group with defined user rights.

**User groups** There are three hierarchical user groups:

- Administrator (top level)
- User (user account registered by the administrator)
- Guest (user without user account)

Administrators and users log in to the photometer with their user name and password. Guests can optionally enter a name for their login. Thus, documented measured values can later be assigned to the user.

User rights in detail	Action	Administrator	User	Guest
	Select methods	✓	~	1
	Carry out measurements	✓	~	<ul> <li>✓</li> </ul>
	Store measurement data	✓	~	<b>√</b>
	Check photometer (AQA1)	✓	~	$\bigcirc$
	Check total system (AQA2)	✓	~	$\bigcirc$
	AQA1 measured value labeling	✓	~	1
	AQA2 measured value labeling	✓	~	$\Diamond$
	Edit user-defined methods	✓	~	$\Diamond$
	Exchanging methods / profiles	✓	0	$\Diamond$
	Change AQA settings	✓	0	$\Diamond$
	Clear the memory	✓	0	$\Diamond$
	Set the date and time	✓	0	$\Diamond$
	Administrate users	<ul> <li>✓</li> </ul>	Ø	$\Diamond$
	Reset photometer settings	✓	0	$\Diamond$
	Carry out software update	<ul> <li>✓</li> </ul>	0	Ø



#### Note

You can also switch off the user management and reactivate it as necessary.

To do so, you need administrator rights. If the user management is switched off, the user name and password do not have to be entered. Each user has full rights.

### 4.16.2 Activating or deactivating the User management function

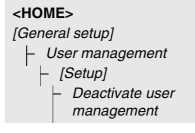
Each user can activate the user management function. If the function is deactivated, each user has administrator rights.

Only members of the user group, administrator can deactivate the user management function.

If the function is active, each user has to log in to the photometer. After the login, the user has certain rights depending on the user group.

# 

Deactivating the user management function



Select and confirm Yes.
 The user management function is active.

Activating the user management creates an administrator user account. The user name is "Administrator". The preset password is "admin". Change this password as soon as possible.

The user management function is inactive.

Each user has administrator rights.



#### Note

If the user management is deactivated by a user of the *Administrator* user group, all user accounts that were set up are lost. The password for the administrator is reset to "admin".

#### 4.16.3 Creating, changing or deleting a user account

When the user management function is active, a user with administrator rights can administrate user accounts.

Creating a user<br/>accountDuring the creation of a user account, the Name, whether or not the user<br/>belongs to a User group and the Password are defined.

<home></home>
[General setup]
– User management
└

User management		16.04.07 9:52	
Name	User group		
Administrator			
Admin2	Administrator		
Enter us	ser name		
A_			
		<b>.</b>	
Setup	Add	Delete	Change

The input field for the new user name pops up.

1 Enter the user name (**<A...9**>) and confirm.

The selection field for the user group (*Administrator / User*) pops up.

- 2 Select and confirm the user group. The input field for the password pops up.
- 3 Enter the password (**<A...9>**) and confirm.

The user account is created and appears in the list of user accounts.

Editing a userWhen a user account is changed, the User group and Password can be<br/>changed.

<HOME> [General setup] User management

User managen	User management		
Name	User group		
Administrator	Administrator		
Admin2	Administrator		
User gro	oup		
User			
Adminis	strator		
Cotup	Add	Delete	Change
Setup	Add	Delete	Change

- 1 Select a user account.
- 2 Press [Change] to edit the user account.

The selection field for the user group (*Administrator / User*) pops up.

**3** If necessary, select and confirm another user group.

The input field for the password pops up.

4 If necessary, enter (**<A...9>**) and confirm another password.

The user account is changed and appears in the list of user accounts.

## Deleting a user account



- 1 Select a user account.
- 2 Delete the user account with [Delete].

A security prompt appears: *Confirm deletion* ?

**3** Confirm the security prompt. The user account is deleted.

#### 4.16.4 Login with active user management

To be able to always assign measurement data to a user, the administrator can activate the user management function. After doing so, the photometer can only be operated after login with a user name. Depending on the authorization class (administrator, user, guest), important settings are released for changes or locked.



#### Note

The user management function is not active in the delivery condition of the photoLab<sup>®</sup> 6600 UV-VIS. Every user can carry out all functions.

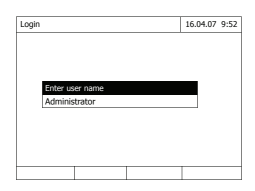
Activating the user management creates an administrator user account. The user name is "Administrator". The preset password is "admin". Change this password as soon as possible.

Make sure to use the correct spelling (upper and lower case) of user name and password for the login.

After logging in to the *Administrator* group with a user name, you can create further users or administrators or switch off the user management function.

The *Login* window with the *Enter user name* prompt appears after the meter has been switched on and after a user has logged off.

In the following example, a user will log in with the user name, "Administrator".



Login		16.04.07	9:52
Enter pa	issword		
admin			

Home (Admir	nistrator)		16	.04.07 13:56
Konzentration				
E	Extinktion / % Transmission			
Multi-Wellenlängen				
Spektrum				
Kinetik				
Einstellungen	Abmelden	AQS	;	Info

The photometer is switched on. The *Login* dialog is displayed.

1 Enter the user name (**<A...9>**) and confirm.

The input field for the password pops up.

If the user name is not known (or incorrectly spelled) it is possible to log in without a password as a guest with restricted rights (see section 4.16.1).

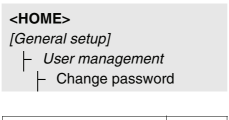
2 Enter the password (**<A...9>**) and confirm.

If the password is written correctly (note upper and lower case), the *Home* main menu opens up. The entered user name is displayed.

#### 4.16.5 Changing the password

The administrator sets up user accounts and assigns a password to each user account.

As soon as any user has successfully logged in with the password, they can change the password for their user accounts themselves.



User management	16.04.07 9:52
Old password	
	-

- 1 Enter and confirm the old password.
- 2 Enter and confirm the new password.

The password is changed.

## 4.17 Reset

You can reset (initialize) the measurement settings or all settings.



Note

The *Reset* function is available for users of the user group, Administrator only.

You have the following options of resetting the photometer settings:

<ul> <li>Reset configuration</li> </ul>	All settings except for the measure- ment data memory, user-defined methods and measured blank val- ues are deleted.
<ul> <li>Delivery condition</li> </ul>	All settings (including measurement data memory and user-defined methods) are deleted and the pho- tometer is reset to the delivery con- dition.

## <HOME>

[General setup] - Reset

The menu where to select the reset type (*Delivery condition* / *Reset configuration*) is displayed.

**1** Select and confirm the reset type.

The reset is carried out.

## 4.18 Photometer information ([Info])

The following photometer information is displayed:

- Photometer designation
- Version number of the meter software/method data
- Hardware version
- Series number of the meter
- Registered user
- Hardware status (for service purposes)
- Memory status

## <HOME> [Info]

Info	16.04.07 9:52
Model designation:	photoLab <sup>®</sup> 6600 UV-VIS
Serial number:	. 07440001
Software/methods version:	1.30-WTW-1.60
Build:	04/03/09 11:57
Hardware version:	0-
Hardware status:	FF 00000000
Lamp counter	12
System test	1
Filter test	1
Wavelength calibration	1
Free internal memory space	
Registered user	

The meter information and result of the self-test are displayed and can be printed.

## 4.19 Lamp counter

The photometer counts the operating hours of the lamp. The information on the operating hours of the lamp is given in the *Info* menu.

The number quoted there corresponds to the number of flashes.



#### Note

tometer.

4.20

Only members of the user group, *Administrator* may carry out any software and method updates.

The software and method update is used to continuously update your pho-

The update comprises

• the newest firmware (meter software)

Software and methods update

• new or changed method data



#### Note

User-defined data (such as settings, user-defined methods or measured data) are not changed by a software and methods update.

The current software version is available on the Internet under http://www.WTW.com.

The software can be transmitted to the photometer as follows:

- by means of a USB memory device as a temporary storage (section 4.20.1).
- by means of a USB connection between a PC and the photometer (section 4.20.2).

## 4.20.1 Update using a USB memory device

Store the new software required for the update on the USB memory and connect it to the photometer.

#### Execution

- 1 Connect the USB memory device to the PC.
- 2 Unpack the contents of the downloaded exe or zip file <u>with the</u> <u>entire folder structure in the main</u> directory (top level) of the USB memory.



#### Note

Make sure the folder structure of the files is retained during the unpacking process.

If you use a program such as WinZip to unpack the files, the option, "Nutze Ordnernamen" or "Use Folder Names" must be set. Details are given in the documentation of your unpack program.

The USB memory must have the folder "Update" on the top level. The Update folder comprises several subfolders.

The following steps are carried out at the photometer.

- <HOME>

   [General setup]

   Software/methods update

   Software/methods update

   Software/methods update

   Select source of update data:

   USB memory device

   PC

   Cancel

3 Connect the USB memory device

4 Switch on the photometer if nec-

to the photometer.

essary.

5 Using <s><t>, select USB memory device as the source and press <START.ENTER>.

The update process takes approx. five minutes.

Subsequently, the photometer switches itself off and then on again.



#### Note

If the update cannot be carried out, an error message appears on the display. Check whether the "Update" folder with its subfolders is stored on the USB memory device (top level).

## 4.20.2 Update using a PC

**Requirements** The following is required for the update:

- A free USB connection on the PC
- A USB cable (type A type B)
- The SpectralTransfer program. It is on the CD-ROM provided with the photometer.
- The ActiveSync<sup>®</sup> program.
   It is available free of charge in the download area of Microsoft<sup>®</sup> on the Internet.
- The current photometer update file. It is available on the Internet under http://www.WTW.com. The update file comprises:
  - the newest firmware (meter software)
  - new or changed method data.
- at least 3 MB free storage on the photometer.

#### Note

You can view the currently available free storage on the photometer in the Info menu.

If less free storage is available than required for the update, the update is not possible by means of a PC at the moment.

You can either erase measurement data from the photometer until enough free storage is available, or you can carry out the update by means of a USB memory device. The free storage is required during the update only. After a successful update, this storage is released again.

**Execution** The execution of the software and method update with a PC is described in detail in the operating manual of the SpectralTransfer program. The operating manual (pdf file) is on the CD-ROM provided with the photometer.

The update process takes approx. five minutes. Subsequently, the photometer switches itself off and then on again.

### 4.20.3 Language update

If you want to set some special languages on your photometer (e.g. Chinese or Thai), a character set extension is required to display the characters.

You can install the additional character sets with a corresponding language update. After the installation, the character sets will occupy some of the storage space of the photometer:

- Chinese: 2 MB
- Thai: 0.3 MB

The language updates are on the CD-ROM provided with the photometer.



#### Note

A language update cannot be undone. Therefore, we recommend to carry out the language updates only if they are really required.

**Before the update** Before carrying out the language update, make sure the current software version is installed on the photometer. It is available on the Internet as an update. Download this software update and install it before starting the installation of the language updates.

**Requirements** Free storage space on the photometer is required, depending on the character set to be installed and the installation procedure:

	Storage space required for installation via		
Character set	USB storage medium	PC	
Chinese	2 MB	4 MB	
Thai	0.3 MB	0.6 MB	



#### Note

You can view the currently available free storage space on the photometer in the

Info menu (F4 key). If less free storage is available than required for the update, the update is not possible. You can back up and erase from the photometer measurement data so that enough free storage is made available.

**Execution** The update is executed in the same way as a software and method update and takes approx. 2 minutes. All files required for the update are in a zip archive or a self-unpacking exe file ("FontXXXXXX.zip" or "FontXXXXX.exe") on the CD-ROM. It also includes a Readme file with detailed installation instructions for the language update.

# 5 Maintenance and cleaning

5.1 Exchanging the buffer batteries

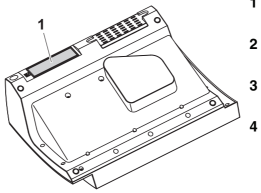
## CAUTION

There is a risk of explosion if unsuitable batteries are used. Only use leakproof alkaline manganese batteries.



#### Note

If you leave the photometer switched on during the exchange or insert the new batteries within a minute after taking out the old ones, the date and time are retained in the photometer.



- 1 Turn the photometer upside down and place it on a soft surface.
- **2** Open the lid of the battery compartment (1).
- **B** Remove the old batteries from the battery compartment.
- Insert the four new batteries in the battery compartment. Make sure that the poles of the batteries are in the correct position.
   The ± signs on the batteries must correspond to the ± signs in the battery compartment.
- **5** Close the lid of the battery compartment.

**Battery service life** The power consumption of the clock is very low. The lifetime of high quality batteries is at least 5 years.

**Disposal of batteries** Dispose of the batteries at a suitable facility according to local legal requirements. It is illegal to dispose of the batteries with household refuse.

Within the European Union, the batteries are removed at a specialized treatment center at the instrument's end of life. The instruments are taken to one of those specialized treatment centers via the recycling system set up for this purpose.

## 5.2 Cleaning

Especially after a cell has broken or after a reagents accident, the photometer should immediately be cleaned (see also section 6.1 ACTIONS IN THE CASE OF A BROKEN CELL).

## 5.2.1 Cleaning the enclosure

## CAUTION



The housing components are made out of synthetic materials (ABS, PMMA and PC). Thus, avoid contact with acetone, ethyl alcohol and similar detergents that contain solvents. Any splashes must be wiped off immediately.

Clean the photometer enclosure as follows:

- If the housing surface is dirty, wipe it with a soft cloth and mild soapy water.
- Remove any chemicals splashes as soon as possible.
- For disinfection, you can use isopropanol for cleaning for a short time.

## 5.2.2 Cleaning the cell shaft



## CAUTION

The surface areas of the cell shaft are made of synthetic material (PPO/ PS, PMMA). Thus, avoid contact with acetone, ethyl alcohol and similar detergents that contain solvents. Any splashes must be wiped off immediately.



#### Note

If a cell has broken, the cell shaft has to be cleaned immediately. To do so, proceed as described in section 6.1.

Normally, it is not required to clean the cell shaft routinely. Remove dust and slight contamination with a moist, lint free cloth. Use isopropanol <u>briefly</u> to remove persistent coatings (e.g. reagent remains). Especially clean the bottom parts of the lateral surfaces of the rectangular cell shaft where the light barriers for the automatic cell recognition are located.

#### 5.2.3 Cleaning the detector lens

Normally, it is not required to clean the detector lens routinely. Cleaning the detector lens can be necessary in the following cases:

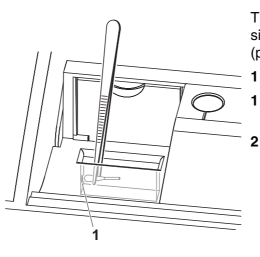
- If the lens is visibly smudged, e.g. after a cell has broken or after a reagent accident (see also section 6.1 ACTIONS IN THE CASE OF A BROKEN CELL).
- If, due to environmental influences or reagent contamination, the photometer displays the message, *Wavelength calibration* during the self-test after being switched on (see section 6.2)



#### Note

If the lens is often smudged (error, *Wavelength calibration* during the self-test), check whether the correct operating conditions are observed. Follow the details in section 3.2 for this purpose.

Proceed as follows to clean the detector lens:



The detector lens is on the front left side of the rectangular cell shaft (pos. 1).

- Switch off the photometer.
- 1 Cut off one end of a customary cotton swab (approx. 2 cm).
  - Grasp the cut-off end with the tip of a pair of tweezers or small pliers. Clean the lens with the dry head of the swab. To do so, move the head from the center of the lens outward in circles. If there are persistent coatings, moisten the swab with a little deionized water or isopropanol.



#### Note

After recommissioning, carry out the photometer monitoring for all measurements (see section 4.15.2).

# 6 What to do if ...

6.1 Actions in the case of a broken cell

## WARNING

Cells can contain dangerous substances. If the contents are released, follow the safety instructions of the package insert. If necessary, take corresponding protective measures (protective goggles, protective gloves etc.).



## CAUTION

Do not turn the photometer upside down to remove the liquid! When doing so, the liquid could come into contact with electronic components and damage the photometer.

The photometer has a drain device through which the contents of a broken cell can drain off without causing any damage.

Proceeding after a cell has broken

- **1** Switch off the photometer and disconnect it from the power supply.
- 2 Let the liquid drain off into a suitable container and dispose of it properly according to the instructions of the reagent package.
- **3** Carefully remove all broken glass, e.g. with tweezers.
- 4 Carefully clean the cell shaft using a moist, lint-free cloth. If there are persistent coatings, use isopropanol <u>for a short time</u>. Especially clean the bottom parts of the lateral surfaces of the rectangular cell shaft, where the light barriers for the automatic cell recognition are located.
- 5 Let the cell shaft dry.

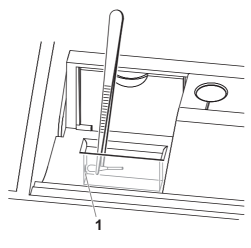


#### Note

After recommissioning, carry out the photometer monitoring for all measurements (see section 4.15.2).

If, after recommissioning, an error occurs during the wavelength calibration, the detector lens is probably smudged. In such a case, clean the lens as follows:

# Cleaning the detector lens



The detector lens is on the front left side of the rectangular cell shaft (pos. 1).

- **1** Switch off the photometer.
- 2 Cut off one end of a customary cotton swab (approx. 2 cm).
- 3 Grasp the cut-off end with the tip of a pair of tweezers or small pliers. Clean the lens with the dry head of the swab. To do so, move the head from the center of the lens outward in circles. If there are persistent coatings, moisten the swab with a little deionized water or isopropanol.

## 6.2 Error causes and remedies

Instrument does not react to keystroke	Cause	Remedy
	<ul> <li>Operating condition undefined or EMC load unallowed</li> </ul>	<ul> <li>Processor reset:</li> <li>Press the &lt;<b>ON/OFF</b>&gt; and</li> <li><b><esc< b="">&gt; key simultaneously.</esc<></b></li> </ul>

Acoustic signal on keystroke	Cause	Remedy
REYSTICKE	<ul> <li>The key does not have any func- tion in the current operating state</li> </ul>	<ul> <li>Press a different key</li> </ul>

Measuring range undercut or exceeded

Cause	Remedy
<ul> <li>Method not suitable</li> </ul>	<ul> <li>Select method with suitable mea- suring range</li> <li>Dilute the sample</li> </ul>



#### Note

In *Concentration* mode you can display the current absorbance value as an additional information (*[Setup]/Display absorbance*, see also section 4.5.6).

Self-test does not start.	Cause	Remedy
The instrument displays Please remove cell	<ul> <li>A cell is inserted in one of the cell shafts</li> </ul>	<ul> <li>Remove the cell</li> <li>Then press the</li> <li><b>START-ENTER</b>&gt; key</li> </ul>
	<ul> <li>A foreign object is inserted in one of the cell shafts</li> </ul>	<ul> <li>Remove foreign object</li> <li>Then press the</li> <li><b>START-ENTER</b>&gt; key</li> </ul>
	<ul> <li>The instrument has to carry out a new adjustment for the rectangu- lar cell recognition</li> </ul>	<ul> <li>Press the <b>START·ENTER</b></li> <li>key.</li> </ul>
	<ul> <li>The rectangular cell shaft is con- taminated</li> </ul>	<ul> <li>Clean the cell shaft (see section 5.2.2 and section 6.1)</li> <li>Restart the instrument</li> <li>If necessary, confirm the <i>Please remove cell</i> message with </li> <li><start-enter>.</start-enter></li> </ul>
	<ul> <li>Instrument defective</li> </ul>	<ul> <li>Return instrument to service department</li> </ul>

#### Obvious measu

bviously incorrect measured values	Cause	Remedy
	- Cell contaminated	- Clean the cell
	<ul> <li>Dilution set incorrectly</li> </ul>	<ul> <li>Set the dilution</li> </ul>
	<ul> <li>Selected method not suitable</li> </ul>	<ul> <li>Select different method</li> </ul>
	- Zero measurement incorrect	<ul> <li>Perform zero measurement</li> </ul>
	<ul> <li>Blank value incorrect</li> </ul>	<ul> <li>Remeasure the blank value</li> </ul>
	<ul> <li>Cells that are not recognized (e.g. some plastic cells) disturb the AutoCheck measurement and, as a consequence, falsify measured values until the next AutoCheck is carried out</li> </ul>	<ul> <li>Use suitable cells (see section 7.1 and section 8.1)</li> </ul>
Fluctuating measured values	Cause	Remedy
	<ul> <li>Cell shaft cover open</li> </ul>	<ul> <li>Close the cell shaft cover</li> </ul>

1

Self test failed.	Cause	Remedy
	- System test: Instrument defective	<ul> <li>Return instrument to service department</li> </ul>
	- Filter test: Instrument defective	<ul> <li>Return instrument to service department</li> </ul>
	- Wavelength calibration:	
	<ul> <li>Foreign particle in the cell shaft</li> </ul>	<ul> <li>Remove foreign object</li> </ul>
	<ul> <li>Lens smudged</li> </ul>	<ul> <li>Clean the lens (see section 5.2.3 or section 6.1). If this happens repeatedly, check the operating conditions (see section 3.2)</li> </ul>
	<ul> <li>Instrument defective</li> </ul>	<ul> <li>Return instrument to service department</li> </ul>
		1
Connected printer does not print	Cause	Remedy
	<ul> <li>Printer not suitable</li> </ul>	<ul> <li>Connect a printer that can inter- pret the printer control language PCL-3</li> </ul>

# 7 Technical data

## 7.1 Measurement characteristics

Measuring principle Single-beam spectrophotometer

Light source	Lamp type	Xenon flashlamp
	Average lifetime	5 x 10 <sup>8</sup> flashes, corresponding to at least 13000 h in permanent operation
Monochromator	Туре	Grating monochromator with step motor
	Wavelength range	190 - 1100 nm
	Max. scan speed	approx. 3300 nm/min
	Wavelengths calibration	Automatic
	Accuracy	± 1 nm
	Reproducibility	± 0.5 nm (checkable, e.g. with holmium oxide filter)
	Resolution	1 nm
	Spectral band width	4 nm
Photometric measurement	Light sensor	Photo diode
	Measuring range	A = -3.300 to A = +3.300
	Linearity	< 1 % for A ≤ 2.000 * in the range 340 nm 900 nm
	Accuracy *	- 0.003 A for A < 0.600
		$-$ 0.5 % of the reading for 0.600 $\leq\!\!A\!\!\leq$ 2.000
	Reproducibility *	± 0.002 at A = 1.000
	Resolution	$\Delta A = 0.001$
	Scattered light	< 0.1 % transmission at 340 and 408 nm
	. tin the verse 000mm 1000	

• \* in the range 200nm ... 1000nm

Usable cells

Round cells	<ul> <li>Outer diameter: 16 mm</li> <li>Inner diameter: 13.6 mm</li> <li>Flat cell bottom</li> </ul>
Rectangular cells *	<ul><li>Path length: 10 mm, 20 mm and 50 mm</li><li>Maximum width: 12.6 mm</li></ul>
Minimum filling level	20 mm
Minimum filling volume	Round cell 16 mm: 4 ml Rectangular cell, 10 mm: 2 ml Rectangular cell, 20 mm: 4 ml Rectangular cell, 50 mm: 10 ml
Cell recognition	Automatic for most types

• Suitable are rectangular glass cells with opaque lateral surfaces (see section 8.1). Plastic cells with clear or serrated lateral surfaces are not reliably recognized by the automatic cell recognition.

Cells that are not recognized disturb the AutoCheck measurement and, as a consequence, falsify measured values until the next AutoCheck is carried out.

Warm-up timeAt least 15 min for single measurements2 h for kinetic measurements with the highest possible precision

## FCC Class A Equipment Statement

<u>Note:</u> This equipment has been tested and found to comply with the limits for a Class A digital device, pursuant to Part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instruction manual, may cause harmful interference to radio communications. Operation of this equipment in a residential area is likely to cause harmful interference in which case the user will be required to correct the interference at his own expense.



#### Note

Changes or modifications not expressly approved by the manufacturer could void the user's authority to operate the equipment.

**Measuring modes** 

- Concentration
  - Measurement with permanently programmed methods, adjusted to the WTW-test set program
  - Automatic method selection if test sets with barcodes are used
  - Program support for the creation of additional user-defined methods (max. 100)
  - Citation forms and units method dependent
  - Display of the absorbance value can be added
  - Method data update possible via Internet
- Absorbance / % Transmission
  - Measurement against own reference absorbance value possible
- Multi wavelengths
  - Freely definable calculations from up to four individual absorbance values at different wavelengths
  - Calculations can be stored as methods (max. 50)
- Spectrum
  - Absorbance or % transmission mode
  - Limits freely selectable within the wavelength range
  - Increment: 1 nm
  - Recording duration for the complete wavelength range: < 7 min
  - Settings can be stored as profiles (max. 20)
  - Evaluation functions: Cursor scanning, zoom, min./max. recognition, peak area determination, derivation, smoothing, multiplication by constants, addition of constants, addition and subtraction of spectra, formation of the quotient of two spectra
- Kinetics
  - Absorbance or % transmission mode
  - Minimal adjustable scan interval: 1 s (if the absorbance of the test sample is high, the scan interval is extended due to the longer duration of the individual measurements)
  - Settings can be stored as profiles (max. 20)
  - Evaluation functions: Cursor scanning, zoom, min./max. determination, slope calculation (for an interval or total), enzymatic activity

Memory for measured values	Memory capacity	<ul> <li>1000 single measured values from the measuring modes, concentration, absorbance / % transmission and multi wavelengths</li> <li>4 MByte internal memory, sufficient for approx. 100 spectra and 400 kinetic curves (sample values based on the following assumptions: All spectra over a wavelength range of 600 nm and all kinetic curves with 150 single values each)</li> </ul>
	Output options	USB memory device, printer, PC
	File formats	ASCII, *.csv
Monitoring	AQA1	Check of the photometer
functions	AQA2	Check of the total system
	AQA3	Check of the sample matrix
User management	Can be switched off	yes
	User accounts	3 hierarchical levels (administrator, user, guest)
	Password protection	for administrators and users

# 7.2 Measured value documentation and quality assurance

## 7.3 General meter data

Dimensions	404 x 197 x 314 mm (width x height x	depth)	
Weight	approx. 4.5 kg (without plug-in power supply)		
Housing type of protection	IP 30		
Electrical protective class	III		
Test mark	CE, cETLus		
Allowed environmental conditions	Temperature	Operation: +10 °C to + 35 °C (41 °F to 95 °F) Storage: -25 °C to + 65 °C (-13 °F to 268 °F)	
	Humidity	Yearly mean: $\leq 75 \%$ 30 days/year:95 %Other days:85%	
	Climatic class	2	
Power supply	Power pack	Type: FRIWO FW 7362/12 Input: 100 - 240 V $\sim$ / 50 - 60 Hz / 0.70 A Output: 12 V = / 2.5 A Length of the connection cable to the photometer: 2.0 m	
Guidelines and norms used	EMC	EC directive 2004/108/EC EN 61326-1 EN 61000-3-2 EN 61000-3-3 FCC Class A	
	Instrument safety	EC directive 2006/95/EC EN 61010-1	
	Climatic class	VDI/VDE 3540	
	IP protection	EN 60529	
Communication	RS232	1 x 9-pin D-sub	
interfaces	USB	<ul> <li>1 x USB-A (for printer, USB memory devices, keyboard or bar code reader)</li> <li>1 x USB-B (for PC)</li> </ul>	

**Other features** • Drain

- Drain for spilled cell contents
- Photometer software update and method data update possible via Internet

Available languages

- German (Englisch)
- English
- Français
- Español
- Italiano
- Bulgarian/Български
- Česko
- Simplified Chinese/ 中文 \*
- Traditional Chinese/ 繁體中文 \*
- Greek/Ελληνικά
- Indonesian/Indonesia
- Japanese/ 日本語
- Magyar
- Malay/Melayu
- Norsk
- Polski
- Portuguése
- Russian/Русский
- Slovenščina
- Thai/ ภาษาไทย \*
- Turkish/Turkce
- Dansk
- Română
- Nederlands

\* These languages require additional character sets (for details, see section 4.20.3 LANGUAGE UPDATE)

# 8 Accessories and options

## 8.1 Accessories

Cells for the WTW	Description	Model	Order no.
test set program	25 empty round cells (16 mm)	RK 14/25	250 621
	Rectangular cell, 10 mm	REK 10	250 605
	Rectangular cell, 20 mm	REK 20	250 600
	Rectangular cell, 50 mm	REK 50	250 614
	Rectangular cell, quartz, 10 mm	REK 10 Quarz	250 606
	Rectangular cell, quartz, 20 mm	REK 20 Quarz	250 601
	Rectangular cell, quartz, 50 mm	REK 50 Quarz	250 615
	100 PMMA disposable rectangular cells, 10 mm	REC 10 PMMA	250 607
	Half micro cell, optical special-pur- pose glass, 50 mm	REC 50/2	250 616
	Support for 10 mm plastic cells	PL6-10 SIC	250 213

	Description	Model	Order no.
Case and cable for mobile use	Case for photoLab <sup>®</sup> 6000 / spectroFlex series	FC spectral 6000	250 212
	Adapter for 12 V (car) operation	ADA 12V	902 760

Application packages	Description	Model	Order no.
packages	Brewing application package (German/Englisch)	PL6-BREW	250 214

## 8.2 Test equipment

Test equipment	Description	Model	Order no.
	Test equipment for AQA1	PhotoCheck 14693	250 490
	Test equipment for pipette volume	PipeCheck 14692	250 498



## Note

Standard solutions for the WTW test set program are listed in the WTW catalog or on the Internet.

## 8.3 Optional equipment

The following optional extensions are available in specialist shops:

- USB barcode reader (hand scanner)
- USB PC keyboard

## 8.4 Connection cable:

**PC** You can connect a PC to the photoLab<sup>®</sup> 6600 UV-VIS in one of the following ways:

	Description	Order no.	
	<ul> <li>Cable with USB-B and USB-A plug</li> </ul>	Specialist shops	
	<ul> <li>Zero modem cable 9-pin (D-sub socket) - 9-pin (D-sub socket)</li> </ul>	820 070	
USB printer	You can connect a USB printer to the photoLab <sup>®</sup> 6600 UV-VIS:		
	Description	Order no.	
	<ul> <li>Cable with USB-B and USB-A plug</li> </ul>	Specialist shops	
Thermoprinter	You can connect the P3001 thermoprinter to the photoLab the following ways:		
	Description	Order no.	
	<ul> <li>Cable with GenderChanger</li> </ul>	250 745	
	or:		
	<ul> <li>Cable, 9-pin (socket) - 9-pin (plug)</li> </ul>	Specialist shops	
		•	
Needle printer	You can connect an LQ300 needle printer to the photoLab one of the following ways:	9 <sup>®</sup> 6600 UV-VIS in	
Needle printer		o <sup>®</sup> 6600 UV-VIS in	
Needle printer	one of the following ways:		
Needle printer	one of the following ways: Description	Order no.	

# Appendix

### A.1 Menus

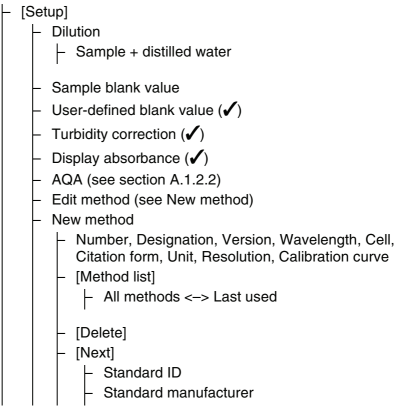
Measuring (see section A.1.1) General settings and functions (see section A.1.2)

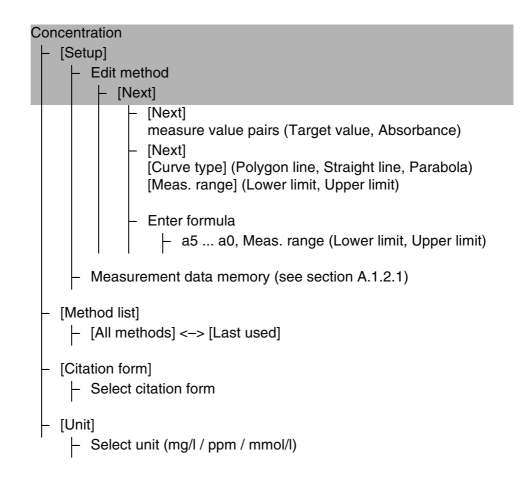
### A.1.1 Measuring

- Concentration (see section A.1.1.1)
- Absorbance / % Transmission (see section A.1.1.2)
- Special / Multi wavelengths
- Spectrum
- Kinetics

### A.1.1.1 Concentration

#### Concentration





#### A.1.1.2 Absorbance / % Transmission

Absorbance / % Transmission

– [Setup]

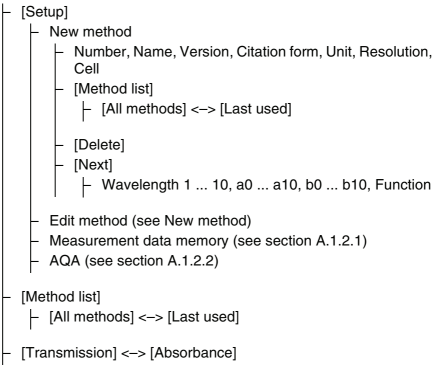
- AQA (see section A.1.2.2)
- Measurement data memory (see section A.1.2.1)

#### [Wavelength]

- Set new wavelength (nm)
- [Transmission] <-> [Absorbance]
- [Reference]
  - Reference absorbance

#### A.1.1.3 Special / Multi wavelengths

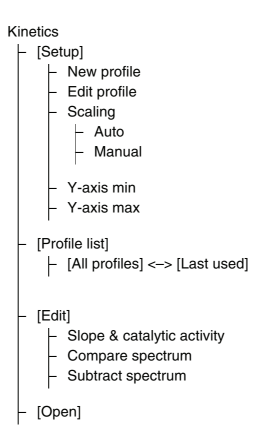
Special / Multi wavelengths



# Spectrum – [Setup] – Wavelength start Wavelength stop - Mode - Absorbance Transmission Smoothing - Yes No Scaling Auto Manual Y-axis min Y-axis max [Edit] Extreme values (zoomed area) Mark points Delete all marks Original values – Integral – Derivative - Compare spectrum Add spectrumSubtract spectrum - Divide spectrum (ratio) Add fixed value Multiply fixed value [Zoom] [Original] – [xy\_max] [Open]

### A.1.1.4 Spectrum

### A.1.1.5 Kinetics



### A.1.2 General settings and functions

- [General setup] (see section A.1.2.1)
- [Logout]
- [AQA] (see section A.1.2.2)
- [Info]

### A.1.2.1 General setup

[General setup] – Language – Date/Time – Date – Time
<ul> <li>Display settings</li> <li>Contrast [%]</li> </ul>
<ul> <li>User management</li> <li>[Setup]</li> <li>Deactivate user management</li> <li>Change password</li> </ul>
– [Add] – [Delete] – [Change]
<ul> <li>Measured value memory</li> <li>[Setup]</li> <li>AutoStore ( <ul> <li>Filter</li> <li>Mode</li> <li>Absorbance / Transmission</li> <li>Concentration</li> <li>Multi wavelength</li> <li>User</li> <li>ID</li> <li>Date</li> <li>from to</li> <li>[Reset entry]</li> <li>[Reset all]</li> </ul> </li> </ul>
<ul> <li>Selected values: invert selection (</li> <li>Delete memory (selected values only)</li> <li>Delete memory (all values)</li> </ul>
<ul> <li>[Single value] &lt;-&gt; [List]</li> <li>[Delete]</li> </ul>

General setup]
<ul> <li>Software/methods update</li> <li>USB memory device</li> <li>PC</li> <li>Cancel</li> </ul>
– Reset
<ul> <li>Reset configuration</li> <li>Delivery condition</li> </ul>
<ul> <li>Delivery condition</li> </ul>
<ul> <li>Data transfer/Printer</li> </ul>
<ul> <li>Decimal separator for csv-Files</li> </ul>
<ul> <li>Point (12.34)</li> <li>Comma (12,34)</li> </ul>
– Comma (12,34)
<ul> <li>Data format (print)</li> </ul>
<ul> <li>Short</li> <li>Extended</li> </ul>
– Extended
<ul> <li>Baudrate for RS232 printer (1200 19200)</li> </ul>
<ul> <li>Exchange methods/profiles</li> <li>Save data to USB memory device</li> <li>Unlock application packages</li> </ul>

A.1.2.2 AQA

```
[AQA]
  - AQA1 setup
      – Mode

    AQA1 inactive

            Weeks

    Interval

    Lock instrument (No/Yes)

      - Configuration of tests ...
          - PhotoCheck
            CertiPUR UV-VIS 1 ...
  - AQA2 setup
      – Mode
           - AQA2 inactive
            Weeks
            Measurements

    Lock methods (No/Yes)

        Method...

    [All methods] <-> [Last used]

    Method

          - AQA2
           Interval

    Target value

            Tolerance
            Standard ID
    AQA3/MatrixCheck setup
     Maximum difference
  - AQA1 check
  - AQA2 check
     [All methods] <-> [Last used]
   AQA3/MatrixCheck
   AQA1 info
    AQA2 info
```

### A.2 Glossary

Absorbance	Logarithmic dimension for the absorption of the sample; negative decadal logarithm of the transmission.
Analysis instructions	The exact proceeding to carry out the detection procedure is described in the analysis instructions.
AQA	Analytical Quality Assurance.
AQA labeling	In the documentation, measured values are given an AQA labeling (AQA1 or AQA2), depending on whether or not the measurement was carried out with AQA and with which AQA level.
AQA1	1st step of the analytical quality assurance: Monitoring of the instrument.
AQA2	2nd step of the analytical quality assurance: Monitoring of the total system.
AQA3	3rd step of the analytical quality assurance: Check of whether the photometric determination is disturbed by other sample ingredients (sample matrix). The MatrixCheck can be carried out by spiking or diluting:
AutoSelector	Plastic cylinder with bar code. It is inserted in the round cell shaft and transmits the code for a reagent test set to the photometer.
Bar code	Optical code (black and white bars) of the method that can be read by light barriers in the photometer.
Baseline	Reference value for the spectrum of reference absorbances or reference transmissions.
Cell	Vessel to take a liquid sample for measurement in a photometer. The cell material (mostly glass) must have certain optical features to be suitable for photometry.
Citation forms	Different forms of representing a measured concentration value that can be derived from each other. The method for the determination of phosphate, e.g. supplies a mea- sured value for phosphorous P. This measured value can alterna- tively be given in the citation forms PO4, PO4-P or P2O5.
CombiCheck	Multiparameter standards used to check the total system for a method.
Concentration	Mass or amount of a dissolved substance per volume, e.g. in g/l or mol/l.
Correlation coefficient	Specifies the extent of the linear relationship of value pairs when determining the zero point and slope for a user-defined method.

Detection procedure	The detection procedure designates the general principle of how a sample is brought into a form suitable for measurement. Different methods can be based on the same detection procedure.
Kinetics	Measurement over a period of time.
MatrixCheck	see AQA3.
Measured value	The measured value is the special value of a measured parameter to be determined. It is given as a combination of the numerical value and unit (e.g. 3 m; 0.5 s; 5.2 A; 373.15 K).
Method	A method comprises a chemical detection procedure and special method data (calibration line) that is required to evaluate the mea- surement results. How to carry out the method up to measuring with the photometer is described in the analysis instructions. The photoLab <sup>®</sup> 6600 UV-VIS contains a database with methods. Fur- thermore, user-defined methods can be entered in the database as well.
PhotoCheck standard	Stable color solution with defined absorbance values for the check of the photometer.
Reagent blank value	The evaluation of the photometric measurement always refers to the comparison value of a test sample without the substance to be determined (reagent blank value). Thus the influence of the basic absorbance of the reagents on photometric measurement is compensated for.
Recovery	The recovery rate is the found measured value divided by the default value (percentage). Example: Default value 20 mg/l; Found 19.7 mg/l => recovery 0.985 or recovery rate 98.5%.
Reference absorbance	With the reference absorbance, the basic absorbance stored in the photometer can be replaced by a measurement of your own.
Reset	Restoring the original condition of all settings of a measuring system.
Sample blank value	Absorbance value of the original test sample, handled according to prescription but without color reagent.
Spectrum	Distribution of the intensity, transmission or absorbance depending on the wavelength.
Standard	Sample with a defined concentration of the analyte to be determined.
Test sample	Designation of the test sample ready to be measured. Normally, a test sample is made by processing the original sample. The test sample and original sample are identical if the test sample was not processed.

Test set (test)	A test set contains all reagents that are required for the photometric determination of the sample according to the analysis instructions.
Transmission	Part of the light that goes through the sample.
Turbidity	Light attenuation caused by diffuse scattering at undissolved sub- stances.
Zero adjustment	Adjusting a photometer with a water-filled cell.

### A.3 List of trademarks

Trademark	Owner
ActiveSync <sup>®</sup>	Microsoft Corporation
CertiPUR <sup>®</sup>	Merck KGaA
Microsoft <sup>®</sup>	Microsoft Corporation
Spectroquant <sup>®</sup>	Merck KGaA
Windows <sup>®</sup>	Microsoft Corporation

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