



# NanoPhotometer® P-Class User Manual P 300 / P 330 / P 360

Version 2.1



**P 360**



**P 330**



**P 300**





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## Declaration of conformity for the NanoPhotometer® P-Class (P300/P330/P360)

This is to certify that the Implen NanoPhotometer® P-Class conforms to the requirements of the following Directives:

2006/95/EC	Low Voltage Equipment Safety Directive.
98/79/EC	In Vitro Diagnostic Medical Devices Directive.
2004/108/EC	EMC Directive.
2002/95/EC	Restrictions on the use of certain Hazardous Substances in Electrical and Electronic Equipment (ROHS).
1995/5/EC	Radio and Telecommunications Terminal Equipment Directive (instruments fitted with Bluetooth accessory only).
2002/96/EC	EC Directive on Waste Electrical and Electronic Equipment (WEEE) 2003/108/EC & 2008/34/EC. By ensuring this product is disposed of correctly, you will help prevent potential negative consequences for the environment and human health, which could otherwise be caused by inappropriate waste handling of this product.

Standards to which conformity is declared, where relevant, are as follows:

EN61010-1:2001	Safety requirements for electrical equipment for measurement, control and laboratory use. General requirements.
EN61010-2-101:2002	Particular requirements for IVD medical equipment.
EN61326-1:2006	Electromagnetic compatibility - generic emission standard electrical equipment for measurement, control & laboratory use.

For further information, including unpacking, positioning and installation of the products please refer to the user manual.

Signed:

Dated: October 1, 2011



Dr. Thomas Sahiri  
Managing Director  
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## 1. ESSENTIAL SAFETY NOTES

There are a number of warning labels and symbols on your instrument. These are there to inform you where potential danger exists or particular caution is required. Before commencing installation, please take time to familiarise yourself with these symbols and their meaning.



Caution (refer to accompanying documents).  
Background colour yellow, symbol and outline black.

### Unpacking, Positioning and Installation

- Check the contents of the package against the delivery note. If any shortages are discovered, inform your supplier immediately.
- Inspect the instrument for any signs of damage caused in transit. If any damage is discovered, inform your supplier immediately.
- Ensure your proposed installation site conforms to the environmental conditions for safe operation:  
Indoor use only.  
Temperature range 5°C to 35°C. Note that if you use the instrument in a room subjected to extremes of temperature change during the day, it may be necessary to recalibrate (by switching off and then on again) once thermal equilibrium has been established (2-3 hours).  
Maximum relative humidity of 80% up to 31°C decreasing linearly to 50% at 40°C.
- The instrument must be placed on a stable, level bench or table that can take its weight (< 4.5 kg) so that air can circulate freely around the instrument.
- The equipment should be positioned such that in the event of an emergency the mains plug can be easily located and removed
- This equipment must be connected to the power supply with the power cord supplied. It can be used on 90 – 240 V, 50-60 Hz supplies.
- If the instrument has just been unpacked or has been stored in a cold environment, it should be allowed to come to thermal equilibrium for 2-3 hours in the laboratory before switching. This will prevent calibration failure as a result of internal condensation.
- Switch on the instrument via the keypad (🔘) after it has been plugged in. The instrument will perform a series of self-diagnostic checks.
- Please read through this user manual prior to use.
- Please contact your original supplier in the first instance if you experience technical or sample handling difficulties.

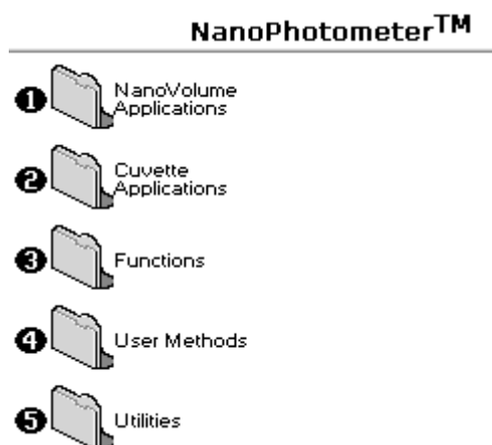
If this equipment is used in a manner not specified or in environmental conditions not appropriate for safe operation, the protection provided by the equipment may be impaired and instrument warranty withdrawn.

## 2. INTRODUCTION

### 2.1 Your spectrophotometer

Your spectrophotometer is a simple-to-use UV/Visible instrument with a CCD array detector (1024 pixels). It has no moving parts, which is the basis of the rapid scanning operating system.

The user interface is built around folders which are displayed on the main screen when the instrument is switched on. Different folders are numbered and opened by using the associated number key on the keypad. After switching on the NanoPhotometer® a self-calibration check is performed and the default main screen "NanoPhotometer®" is offering the choice of:



Keypad number	Description
1	Life Science methods such as nucleic acid assays and protein assays using the NanoPhotometer® P-Class Submicroliter Cell
2	Life Science methods such as nucleic acid assays, protein assays and cell density using cuvettes
3	General spectroscopic methods
4	Contains nine folders that can store user adapted methods (up to 81)
5	Instrument set up (date, time, number format), Output Options and Baseline Compensation set up.

The instrument is equipped with a standard USB port (back) and the P330/P360 also with a USB flash drive port (right side). The NanoPhotometer® P-Class Software Package is necessary to connect the NanoPhotometer® P-Class to a PC or to read the saved data on the USB flash drive. The software enables the user to “print through” the PC directly to the printer that is connected to it. Data may be stored as Excel spreadsheet (report and/or table format), EMF graphics file, a comma delimited (csv) data file, a tab delimited (txt) data file or in native NanoPhotometer® P-Class Software format for later access (see also NanoPhotometer® P-Class PVC Installation and User Manual).

Alternatively, results may be saved on a SD Memory Card (P300 only) or sent to the PC via a Bluetooth (P300 only) accessory; these can either be supplied pre-installed or are available as an optional accessory if the need for the use arises after installation of the product.

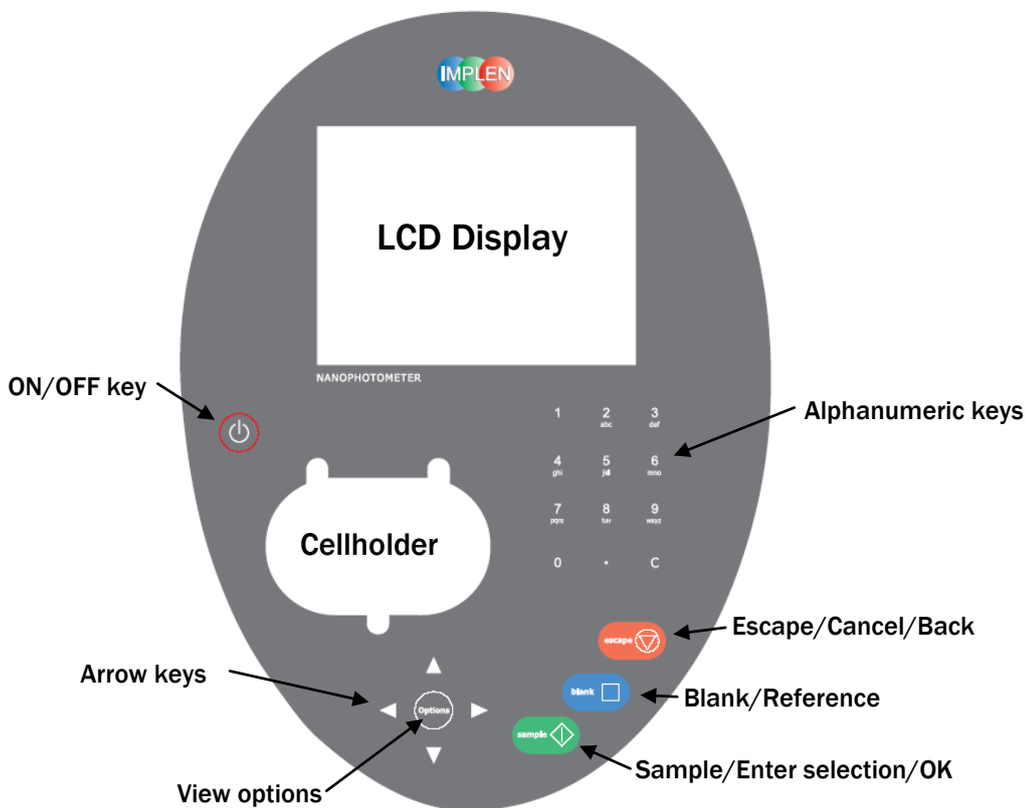
A thermal built-in printer is available for the instrument; this may either be supplied pre-installed or is available as an optional accessory if the need for its use arises after installation of the product.

### 2.2 Sample handling tips

- The NanoPhotometer® P-Class includes an integrated vortexer (P 330 / P 360 only) to assure a good homogeneity of the sample. It is recommended to mix every sample before a measurement.
- Note that the light beam is directed from RIGHT to LEFT through the cell chamber; therefore please ensure the measurement cell is inserted in the correct alignment.
- Insert the measurement cell always in the same direction.
- The cell holder supplied with the instrument accepts the NanoPhotometer® P-Class Submicroliter Cell and standard 10 mm pathlength quartz, glass or plastic cells.
- The optical height of the NanoPhotometer® P-Class is 15 mm.
- The minimum volume that can be used is 0.3 µl with the NanoPhotometer® P-Class Submicroliter Cell.
- 12 mm test tubes may be used (e.g. for cell cultures), however they are not recommended as higher quality data is produced by using disposable cuvettes for the analysis. If used, align the indicator line on 12 mm test tubes in the same direction to ensure reproducible positioning of the tube. Note that test tubes do not last forever, and that the surface becomes scratched and blemished through repetitive use; if this is the case they should be replaced.

### 2.3 Keypad and display for NanoPhotometer® P 300

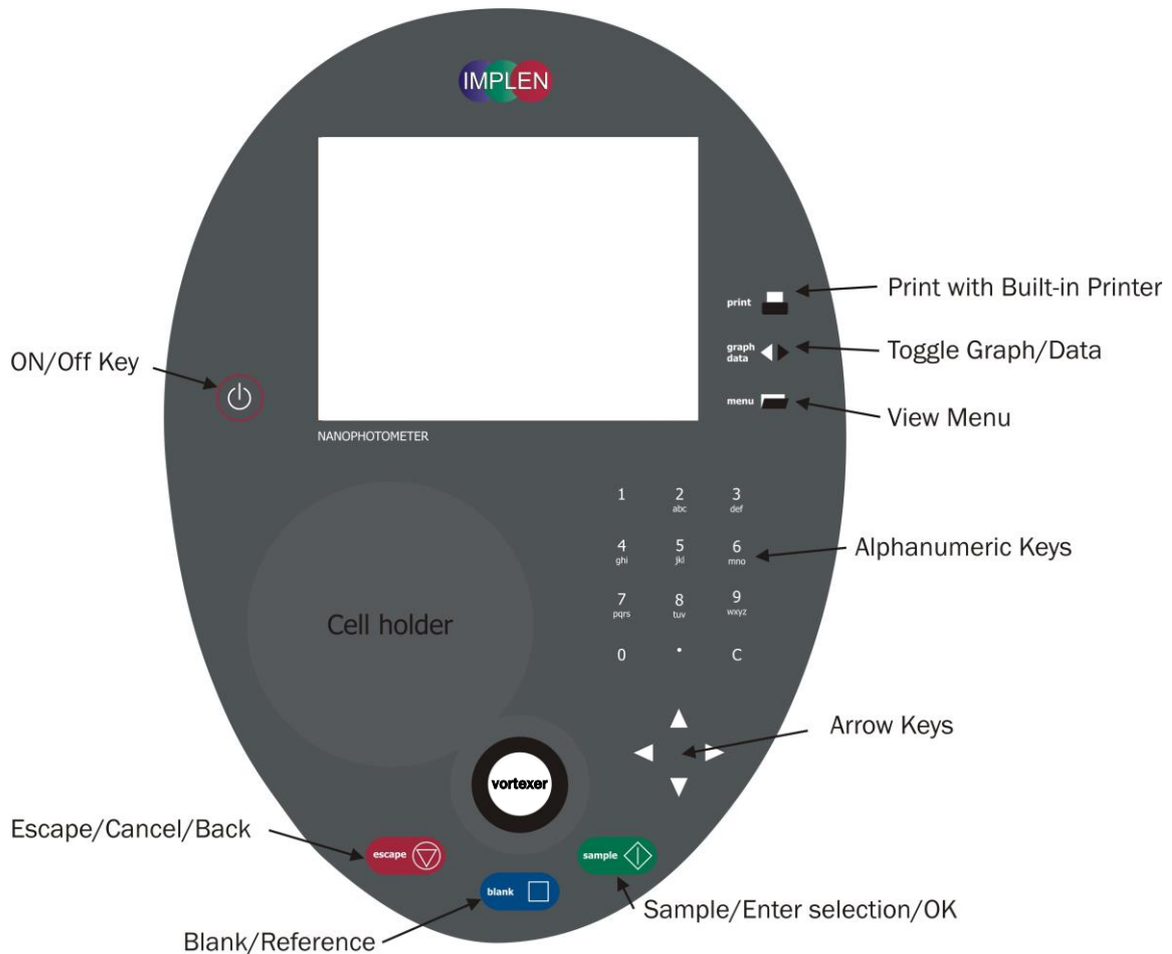
The back-lit liquid crystal display is very easy to navigate around using the alphanumeric entry and navigation arrow keys on the hard wearing, spill proof membrane keypad.



Key	Action
On/off key	Turns the instrument on/off.
Arrow keys	Use the four arrow keys to navigate around the display and select the required setting from the active (highlighted) option.
View Options	View options for that application mode. Some of these are common to all applications and described on page 8. Menu unique to an application are described in the relevant section of the NanoPhotometer® P-Class User Manual.
Alphanumeric keys	Use these to enter parameters and to write text descriptions where appropriate, or required. Use repeated key presses to cycle through lower case, number and upper case. Leave for 1 second before entering next character. Use C button to backspace and 1 to enter a space.
Escape/Cancel/Back:	Escape from a selection and return to the previous folder. Cancel a selection. Stop making measurements.
Blank/Reference	Set reference to 0.000 A or 100%T on a reference solution at the current wavelength in the mode selected. When in scan mode, does a reference scan.
Sample/Enter selection/OK:	Enter, or confirm a selection. Take a measurement.

## 2.4 Keypad and display for NanoPhotometer® P 330 /P 360

The back-lit liquid crystal display is very easy to navigate around using the alphanumeric entry and navigation arrow keys on the hard wearing, spill proof membrane keypad.



Key	Action
On/Off Key	Turns the instrument on/off.
Arrow Keys	Use the four arrow keys to navigate around the display and select the required setting from the active (highlighted) option.
View Menu	View menu for that application mode. Some of these are common to all applications and described on page 8. Menu unique to an application are described in the relevant section of the NanoPhotometer® P-Class User Manual.
Alphanumeric Keys	Use these to enter parameters and to write text descriptions where appropriate, or required. Use repeated key presses to cycle through lower case, number and upper case. Leave for 1 second before entering next character. Use C button to backspace and 1 to enter a space.
Escape/Cancel/Back:	Escape from a selection and return to the previous folder. Cancel a selection. Stop making measurements.
Blank/Reference	Set reference to 0.000 A or 100%T on a reference solution at the current wavelength in the mode selected. When in scan mode, does a reference scan.
Sample/Enter Selection/OK:	Enter, or confirm a selection. Take a measurement.
Print (P 330 and P 360 only)	Prints the results shown on the screen on the built-in printer, if a built-in printer is connected to the NanoPhotometer®.
Graph/Data (P 330 and P 360 only)	Toggle graph on/off. The graph shows a wavescan plot across the range 220 nm to 400 nm (for Dye methods 220 nm to 750 nm) with cursors denoting 230, 260, 280 and 320 nm (Nucleic Acid methods) and 260, 280 and 320 nm (Protein methods).

## 2.5 Menu/Options

(select using key pad numbers)

Options for P 300:

1	Parameters...
2	Print
3	Graph
4	Print Data Only
7	Sample Number...
8	Save Method...
9	Printer Settings...

After each measurement the following Options are possible:

- 1) Return to parameter screen.
- 2) Print the results via selected method.
- 3) Toggle graph on/off. Graph shows a wavescan plot across the range 220 nm to 400 nm (for Dye methods 220 nm to 750 nm) with cursors denoting 230, 260, 280 and 320 nm
- 4) Toggle on/off the graph in the print-out or saved file.
- 7) Define the sample number you wish to start from.
- 8) Save the parameters as a method.
- 9) Open printer settings, possibility to change the printer settings within the method as described in 7.3 Output Options / Printer.

Exit **Options** by pressing **Escape** , OR wait

Menu options for P 330/P 360:

1	Parameters...
2	Data Transfer
4	Output Data Only
7	Sample Number...
8	Save Method...
9	Output Options...

After each measurement the following options are possible in the Menu:

- 1) Return to parameter screen.
- 2) Transfer the results via selected Output Option.
- 4) Toggle on/off the graph in the print-out or saved file.
- 7) Define the sample number you wish to start from.
- 8) Save the parameters as a method.
- 9) Open Output Options settings, possibility to change the Output Options settings within the method as described in 7.3 Output Options / Printer

Exit **Menu** by pressing **Escape** , OR wait

### 3. THE NANOPHOTOMETER® P-CLASS SUBMICROLITER CELL

With its innovative optical pathway the cell is designed for optimum measurement results with submicroliter samples ranging from 0.3 µl up to 5 µl of undiluted sample. Due to a pathlength of 0.04 mm, 0.1 mm, 0.2 mm, 1 mm and 2 mm the cell is offering an automatic dilution of 1/250, 1/100, 1/50, 1/10 and 1/5 in comparison to a standard cuvette measurement. Because the measurements are processed with undiluted samples, the reproducibility of the results is extremely high. If desired, samples can be retrieved after the measurement for further processing. The NanoPhotometer® P-Class Submicroliter Cell can be used for all UV/Vis analysis utilizing the wavelength range of 190 nm to 1,100 nm.

The NanoPhotometer® P-Class Submicroliter Cell is delivered for version P 300 with one lid with a pathlength of 0.2 mm (Lid 50), for version P 330 with two lids pathlength 0.2 mm (Lid 50) and 1 mm (Lid 10) and for version P 360 with three lids pathlength 0.04 mm (Lid 250), 0.2 mm (Lid 50) and 0.1 mm (Lid 10). Lid 5 (2 mm pathlength), Lid 100 (0.1 mm pathlength) and Lid 250 (0.04 mm) can be ordered optionally. The dilution factor (lid factor) is printed on the lid. Please make sure that you use the appropriate lid for your sample.



#### 3.1 Technical instructions



**Step 1** Insert the NanoPhotometer® P-Class Submicroliter Cell into the cell holder with the cell windows facing the light beam. We recommend facing the Implen logo to the front. The light beam is directed from RIGHT to LEFT as indicated with small arrows. Insert the NanoPhotometer® P-Class Submicroliter Cell always in the same direction.



**Step 2** Use the integrated vortexer (P 330 / P 360 only) to mix your sample well to achieve an accurate homogeneity of the sample.



**Step 3** Pipette the appropriate sample volume onto the centre of the measuring window. **Warning!!** Do not overfill the well.

Lid	Sample volume	Pathlength	Dilution
5 (optional)	3.5 - 5 µl	2 mm	1:5
10 (optional for P300)	1 - 3 µl	1 mm	1:10
50	0.3 - 2 µl	0.2 mm	1:50
100 (optional)	0.3 - 2 µl	0.1 mm	1:100
250 (optional)	0.3 - 2 µl	0.04 mm	1:250



**Step 4** Make sure that for the measurements the lid fits exactly onto the positioning supports mounted to the body of the cell. Take measurement. **Remember** to consider the lid factor in your instrument software. Please refer to the NanoPhotometer® P-Class User Manual for detailed information.



**Step 4** Take the lid off and retrieve the sample with a pipette for further applications if desired. Remove sample residues from the measurement window and the mirror in the lid. Clean the measurement window and mirror in the lid well with a slightly wet fluff-free tissue. Use water, 70% ethanol or isopropanol. Do not use aggressive solvents like strong acids or bases or organic solvents at any time.

**Important Note:** Residual fluffs must be removed for optimum performance

Your cell is ready for the next sample.

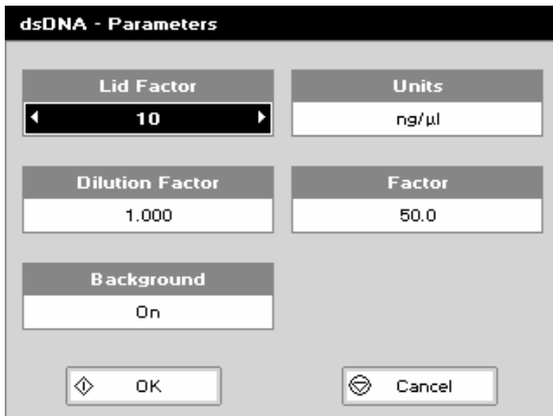
**Operation Limitations:** Do not autoclave the unit! Do not use an ultrasound bath to clean! Do not drop in water or solvent bath. The unit is water resistant, but not water proof!

### 3.2 Software instructions

The **NanoVolume Applications** and **Cuvette Applications** are very similar concerning the analysis of dsDNA, ssDNA, RNA, Oligonucleotides, protein UV and protein dye analysis. This section describes the specific features which have to be considered using the NanoPhotometer® P-Class Submicroliter Cell. For general information please follow the detailed instructions under Nanovolume Applications and Cuvette Applications.

The procedure is as follows:

#### Exemplary Parameter Screen



#### Parameter Screen

- Step 1** Press 1 to select **NanoVolume Applications** folder
- Step 2** Press 1 to select **Nucleic Acids** folder OR 2 to select **Protein** folder.
- Step 3** Select the method you want to use by pressing the corresponding number.
- Step 4** Select the **Lid Factor** using the left and right arrows.

Lid	Sample volume	Pathlength	Dilution
5 (optional)	3.5 – 5 µl	2 mm	1:5
10 (optional for P 300)	1 – 3 µl	1 mm	1:10
50	0.3 – 2 µl	0.2 mm	1:50
100 (optional)	0.3 – 2 µl	0.1 mm	1:100
250 (optional)	0.3 – 2 µl	0.04 mm	1:250

- Step 5** Select subsequent parameters and specifications as described under **4. Nanovolume Applications and Cuvette Applications**.

After the selections are confirmed the results screen displays in top left corner the chosen Lid and the required sample volume.

**Important Information:**

If the absorbance value of the sample is not in the linear range the following **“Warning messages”** will appear and **“Instruction”** will be displayed in the top left corner of the result screen.

	Message:	Answer YES	Answer NO
Lid 5	Concentration too low.		
	Concentration too high. Do you want to change the lid factor?	Please change to lid 10 and press sample. (automatic change of lid factor lid 5 to lid 10 in the software for calculation)	No changes
Lid 10	Concentration too low.		
	Concentration too high. Do you want to change the lid factor?	Please change to lid 50 and press sample. (automatic change of lid factor lid 10 to lid 50 in the software for calculation)	No changes
Lid 50	Concentration too low. Do you want to change the lid factor?	Please change to lid 10, apply a minimum of 1µl of sample and press sample. (automatic change of lid factor lid 50 to lid10 in the software for calculation)	No changes
	Concentration too high. Dilute sample or change to lid 100.		
Lid 100	Concentration too low. Do you want to change the lid factor?	Please change to lid 50 and press sample. (automatic change of lid factor lid 100 to lid 50 in the software for calculation)	No changes
	Concentration too high. Dilute sample or change to lid 250.		
Lid 250	Concentration too low. Do you want to change the lid factor?	Please change to lid 100 and press sample. (automatic change of lid factor lid 250 to lid 100 in the software for calculation)	No changes
	Concentration too high. Dilute sample.		

Lid	Required volume	Warning message	Instruction
		Abs too low	Sample concentration is too low
		Abs is too high	change to lid
		Abs too low	Sample concentration is too low (or change to lid 5 if available)
		Abs is too high	change to lid
		Abs too low	change to lid
		Abs is too high	Physical dilution of the sample is necessary (or change to lid 100 if available)
		Abs too low	change to lid
		Abs is too high	Physical dilution of the sample is necessary (or change to lid 250 if available)
		Abs too low	change to lid
		Abs is too high	Physical dilution of the sample is necessary

\*Some of the lids are only optional available. Lid delivery content for P 300 is Lid 50, for P 330 Lid 10 and Lid 50 and for P360 is Lid 10, Lid 50 und Lid 250.

## 4. NANOVOLUME APPLICATIONS AND CUVETTE APPLICATIONS

The NanoPhotometer® P-Class offers a complete solution for NanoVolume and standard volume applications. With the NanoPhotometer® P-Class Submicroliter Cell the required sample volume ranges from 0.3 µl to a maximal sample volume of 5 µl. Standard volume applications can be performed with 10 mm pathlength quartz, glass or plastic cuvettes.

### Note:

Within the Utilities folder the user has the possibility to select various options that define data output (please see also 7.3 Output Options / Printer).

Folder	Application	Recommended Measurement Cell
<b>Nucleic Acids</b>		
DNA	Concentration, purity check and dye incorporation for DNA samples	Submicroliter Cell / Cuvette
RNA	Concentration, purity check and dye incorporation for RNA samples	Submicroliter Cell / Cuvette
Oligo	Concentration, purity check and dye incorporation for Oligo samples	Submicroliter Cell / Cuvette
<b>Protein</b>		
Protein UV (Christian-Warburg)	Protein determination at 280 nm	Submicroliter Cell / Cuvette
Protein Dye	Protein determination at 280 nm and dye incorporation	Submicroliter Cell / Cuvette
BCA	Protein determination at 562 nm	Cuvette
Bradford	Protein determination at 595 nm	Cuvette
Lowry	Protein determination at 750 nm	Cuvette
Biuret	Protein determination at 546 nm	Cuvette

### Cell Count

OD600	Cell density at 600 nm	Cuvette
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The **NanoVolume Applications** folder and the **Cuvette Applications** folder contain different sub folders:

**Nucleic Acids**, **Protein** and **OD 600** (Cell Density). Contents of these sub folders are detailed below.

### 4.1 Characterization of DNA, RNA and Oligonucleotides

#### 4.1.1 General Information

##### Nucleic Acid Quantification (NAQ)

- Nucleic acids can be quantified at 260 nm because it is well established that a solution of dsDNA in a 10 mm pathlength cell with an optical density of 1.0 has a concentration of 50 µg/ml, ssDNA of 37 µg/ml or 40 µg/ml in the case of RNA. Oligonucleotides have a corresponding factor of 33 µg/ml, although this does vary with base composition; this can be calculated if the base sequence is known. Please refer to 12.1 Nucleic acid quantification for further details.
- The instrument uses factors 50, 37, 40 and 33 as default settings for dsDNA, ssDNA, RNA and Oligonucleotides, respectively, and compensation factors for dilution and use of cells which do not have 10 mm pathlength. Dilution factor and cell pathlength can be entered.

##### Nucleic Acid Purity Checks

- Nucleic acids extracted from cells are accompanied by protein, and extensive purification is required to separate the protein impurity. The 260/280 ratio gives an indication of purity; it is only an indication, however, and not a definitive assessment. Pure DNA and RNA preparations have expected ratios of  $\geq 1.8$  and  $\geq 2.0$ , respectively; deviations from this indicate the presence of impurity in the sample, but care must be taken in interpretation of results.
- The 260 nm reading is taken near the top of a broad peak in the absorbance spectrum for nucleic acids, whereas the 280 nm reading is taken on a steep slope (i.e. small changes in wavelength cause large changes in absorbance). Consequently, small variations in wavelength at 280 nm will have a greater effect on the 260/280 ratio than variations will at 260 nm. Thus different instruments of the same and different types may give slightly

different ratios due to variations in wavelength accuracy. But each instrument will give consistent results within itself.

- Concentration also affects 260/280 readings. If a solution is too dilute, the readings will be at the instrument's detection limit, and results may vary as there is less distinction of the 260 peak and the 280 slope from the background absorbance. This is one reason why the Abs 260 value by using the submicroliter cell (NanoVolume applications) should stay between than 0.01 and 1.50 for accurate measurements.
- An elevated absorbance at 230 nm can indicate the presence of impurities as well; 230 nm is near the absorbance maximum of peptide bonds and also indicates buffer contamination since TRIS, EDTA and other buffer salts absorb at this wavelength. When measuring RNA samples, the 260/230 ratio should be > 2.0; a ratio lower than this is generally indicative of contamination with guanidinium thiocyanate, a reagent commonly used in RNA purification and which absorbs over the 230 - 260 nm range. A wavelength scan of the nucleic acid is particularly useful for RNA samples.
- The instrument can display 260/280 and 260/230 ratios.

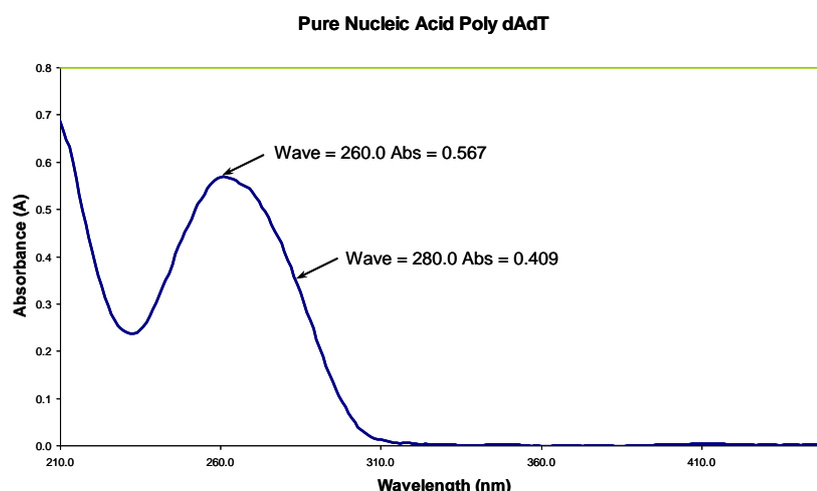
### Fluorescent dye incorporation

- To determine the dye incorporation rate, the absorbance reading at the wavelength reported for maximum absorbance of the fluorescence dye is used. The corresponding extinction coefficient of the dye is used in the Lambert-Beer Law to determine the dye concentration ( $c = A / (e * d)$ ). Comparing these values with the DNA concentration gives a dye incorporation rate. For further details please refer to 12.2 Nucleic acid fluorescent dye incorporation.

### Use of Background Correction

- Background correction at a wavelength totally separate from the nucleic acid and protein peaks at 260 and 280 nm, respectively, is sometimes used to compensate for the effects of background absorbance. The wavelength used is 320 nm and it can allow for the effects of turbidity, high absorbance buffer solution and the use of reduced aperture cells.
- If it is used, there will be different results from those when unused, because Abs 320 is subtracted from Abs 260 and Abs 280 prior to use in equations:  
 Concentration = (Abs 260 - Abs 320) \* Factor  
 Abs ratio = (Abs 260 - Abs 320) / (Abs 280 - Abs 320)  
 Abs ratio = (Abs 260 - Abs 320) / (Abs 230 - Abs 320)
- If your laboratory has not used background correction before, set this option to OFF.
- The use of background correction can remove variability due to handling effects of low volume disposable cells.

### Spectral scan of nucleic acid



**Note:**

- absorbance maximum near 260 nm and absorbance minimum near 230 nm
- flat peak near 260 nm and steep slope at 280 nm
- very little absorbance at 320 nm

Operation of the instrument for Nucleic Acid measurements is described in the following sections.

DNA and RNA are very similar, whilst in Oligo it is possible to calculate the factor from the composite bases by entering the proportions of the 4 bases.

### 4.1.2 Analysis of dsDNA, ssDNA and RNA

The procedure is as follows:

#### Parameter Screen NanoVolume Applications

dsDNA - Parameters

Lid Factor	Units
10	ng/ul
Dilution Factor	Factor
1.000	50.0
Background	
On	

OK Cancel

#### Cuvette Applications

dsDNA - Parameters

Pathlength	Units
10 mm	ng/ul
Dilution Factor	Factor
1.000	50.0
Background	
On	

OK Cancel

#### Results Screen

dsDNA

A230	0.053 A	Sample
A260	0.174 A	
A280	-0.001 A	
A320	-0.001 A	
A260/A280		Concentration
....		
A260/A230		Units
3.241		

87.5

μg/ml

#### Parameter Screen

- Step 1** Press 1 for **NanoVolume** OR 2 for **Cuvette** folder
- Step 2** Press 1 to select **Nucleic Acids** folder
- Step 3** Press 1 to select **dsDNA** mode OR 2 to select **ssDNA** mode OR 3 to select **RNA** mode
- Step 4** Using the NanoVolume Applications select the **Lid Factor** as described in the “Average Detection Range Sheet” and under 3.2.
- Step 5** Enter the **Dilution Factor** using the keypad numbers. Range 1.00 to 9,999. Use the C button to backspace and clear the last digit entered OR press **Menu/Options** to enter the dilution factor screen. Enter the volume of the sample using the keypad numbers. Range 0.01 to 9,999. Enter the volume of the diluent using the keypad numbers. Range 0.01 to 9,999. Press **OK** to calculate the dilution factor and return to the Parameters screen OR press **Cancel** to cancel the selections and return to the Parameters screen.
- Step 6** **Background** correction at 320 nm is recommended to be switched On.
- Step 7** Select the **Units** of measurement using the left and right arrows. Options: μg/ml, ng/μl, μg/μl.
- Step 8** Enter the **Factor** using the keypad numbers. Default value is 50 for dsDNA, 37 for ssDNA and 40 for RNA, range is 0.01 to 9,999.
- Step 9** Press **OK** to enter the Results screen OR **Cancel** to return to the **Nucleic Acids** folder.

#### Results Screen

- Step 10** Apply/insert the reference sample. Press the **Blank** Key. This will be used for all subsequent samples until changed.
- Step 11** Apply/insert sample and press **Sample**. This measures at the selected wavelengths and displays the results. The sample concentration, the ratio of A260/A280 and A260/A230 are calculated (corrected by the background wavelength value if selected).
- Step 12** If the absorbance value of the sample is not in the linear range a “Warning message” will pop up and “Instruction” will be displayed in the top left corner of the result screen. Please refer to 3.2 Software instructions/important information on page 11 for further information.
- Step 13** Repeat for all samples.
- Step 14** Press **Menu/Options** to display available options which are described on page 8.
- Step 15** Press **Escape** and confirm with **Yes** to return to the **Nucleic Acids** folder.

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on 6 (P 300) and 7 (P330 / P 360).

### 4.1.3 Analysis of Oligonucleotides

The procedure is as follows:

#### Parameter Screen

##### NanoVolume Applications

Oligo - Parameters		
Lid Factor	Units	
10	pmol/µl	
Dilution Factor	A	10
1.000	C	10
Background	G	10
On	T	10
<input type="button" value="OK"/> <input type="button" value="Cancel"/>		

#### Cuvette Applications

Oligo - Parameters		
Pathlength	Units	
10 mm	pmol/µl	
Dilution Factor	A	10
1.000	C	10
Background	G	10
On	T	10
<input type="button" value="OK"/> <input type="button" value="Cancel"/>		

#### Results Screen

Oligo	
A230	0.056 A
A260	0.176 A
A280	0.003 A
A320	0.003 A
<input type="button" value="A260/A280"/>	
<input type="button" value="A260/A230"/>	
Sample: 4	
Concentration: <b>57.1</b>	
Units: µg/ml	

#### Parameter Screen

- Step 1** Press 1 for **NanoVolume** OR 2 for **Cuvette** folder.
- Step 2** Press 1 to select **Nucleic Acids** folder.
- Step 3** Press 4 to select **Oligo** mode.
- Step 4** Using the NanoVolume Applications select the **Lid Factor** as described in the "Average Detection Range Sheet" and under 3.2.
- Step 5** Enter the **Dilution Factor** using the keypad numbers. Range 1.00 to 9,999. Use the C button to backspace and clear the last digit entered **OR** press **Menu/Options** to enter the dilution factor screen. Enter the volume of the sample using the keypad numbers. Range 0.01 to 9,999. Enter the volume of the diluent using the keypad numbers. Range 0.01 to 9,999. Press **OK** to calculate the dilution factor and return to the Parameters screen **OR** press **Cancel** to cancel the selections and return to the Parameters screen.
- Step 6** **Background** correction at 320 nm is recommended to be switched on.
- Step 7** Select the **Units** of measurement using the left and right arrows. Options: µg/ml, ng/µl, µg/µl and pmol/µl.
- Step 8** Enter the **Factor** using the keypad numbers. Default value is 33, range is 0.01 to 9,999.
- Step 9** If pmol/µl is selected there are two options to set the factor
  1. A selection table denoting the ratios of the 4 bases according to the oligo sequence. Enter the proportions of bases present using the keypad numbers and up and down arrows to move between boxes. Default is 10 for each, range is 0 to 9,999.
  2. Enter the known extinction factor of the oligo used: factor range 0.01 to 9,999 for ratio = [1 / extinction coefficient \* 10<sup>-6</sup>].
- Step 10** Press **OK** to enter the Results screen **OR** **Cancel**  to return to the **Nucleic Acids** folder.

#### Results Screen

- Step 11** Apply/insert the reference sample. Press **Blank** Key. This will be used for all subsequent samples until changed.
- Step 12** Apply/insert sample and press **Sample** . This measures at the selected wavelengths and displays the results. The sample concentration and the ratio of A260/A280 and A260/A230 are calculated (corrected by the background wavelength value if selected).
- Step 13** If the absorbance value of the sample is not in the linear range a **"Warning message"** will pop up and **"Instruction"** will be displayed in the top left corner of the result screen. Please refer to 3.2 Software instructions/important information on page 11 for further information.
- Step 14** Repeat for all samples.
- Step 15** Press **Menu/Options** to display available Options which are described on page 8.
- Step 16** Press **Escape**  and confirm with **Yes** to return to the **Nucleic Acids** folder.

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on page 6 (P 300) and 7 (P330 / P 360).

#### 4.1.4 Dye Incorporation for dsDNA, ssDNA, RNA and Oligonucleotides

The dye incorporation methods are similar to the dsDNA, ssDNA, RNA and Oligonucleotide methods. This section describes the specific features concerning the dye incorporation. For general information please follow the detailed instructions under **Analysis of dsDNA, ssDNA and RNA and Oligonucleotides**.

To determine the dye incorporation rate, the absorbance reading at the wavelength reported for maximum absorbance of the fluorescence dye is used. For further details please refer to 12.2 Nucleic acid fluorescent dye incorporation.

The procedure is as follows:

##### Parameter Screen NanoVolume Applications

**dsDNA-Dye - Parameters**

Lid Factor	Units
10	µg/ml
Dilution Factor	Factor
1.000	50.0
Dye Correction	Dye-Type
On	Alexa Fluor 350

OK Cancel

##### Cuvette Applications

**dsDNA-Dye - Parameters**

Pathlength	Units
10 mm	µg/ml
Dilution Factor	Factor
1.000	50.0
Dye Correction	Dye-Type
On	Alexa Fluor 350

OK Cancel

##### dsDNA-Dye - Dye Parameters

**dsDNA-Dye - Dye Parameters**

Dye-Type	Dye Abs Max
Custom	300 nm
Dye Ext. Coefficient	Dye Correction
10000	0.000

OK Cancel

##### Parameter Screen

- Step 1** Press 1 for *NanoVolume* OR 2 for *Cuvette* folder.
- Step 2** Press 1 to select *Nucleic Acids* folder.
- Step 3** Press 5, 6, 7 or 8 to select one of the dye incorporation methods.
- Step 4** Using the NanoVolume Applications select the *Lid Factor* as described in the “Average Detection Range Sheet” and under 3.2.
- Step 5** Select *Dilution Factor*, *Units* and *Factor* as described under 4.1.2.
- Step 6** Select whether the *Dye correction* (calculation of the dye-dependent correction factor) is used or not with the left and right arrows. The *Background* correction is always calculated in the Dye methods.
- Step 7** Select the appropriate *Dye Type*. 10 different Alexa Fluors, 4 Cy-Dyes, 6 Oyster-Dyes and Texas Red are programmed with their corresponding maximum absorbance wavelength, dye-dependent correction factor at 260 nm and dye-dependent extinction coefficient. For further details please refer to 12.2 Nucleic acid fluorescent dye incorporation.
- Step 8** If using *Custom Dye* maximum absorbance wavelength of the custom dye, dye-dependent extinction coefficient and dye-dependent correction factor at 260 nm have to be entered.  
 Ranges are:  
*Dye Abs Max*: 300 nm to 950 nm  
*Dye Ext. Coefficient*: 10,000 to 9,999,999  
*Dye Correction*: 0.000 to 0.999

**Results Screen**

dsDNA-Dye	
A230	0.254 A
A260	0.494 A
A280	0.035 A
A320	0.004 A
A550	0.707 A
A650	0.420 A
ADye(345)	0.004 A
A260/A280	15.8
A260/A230	1.960
<b>Sample</b>	
7	
<b>Concentration</b>	
245.00 µg/ml	
<b>FOI (Alexa Fluor 350)</b>	
0.00	
<b>Dye Concentration</b>	
0.00 pmol/µl	

**Results Screen**

- Step 9** Apply/insert the reference sample. Press **Blank** Key. This will be used for all subsequent samples until changed.
- Step 10** Apply/insert sample and press **Sample** . This measures at the selected wavelengths and displays the results. The sample and dye concentration, the FOI and the ratio of A260/A280 and A260/A230 are calculated (corrected by the background if selected).
- Step 11** If the absorbance value of the sample is not in the linear range a **“Warning message”** will pop up and **“Instruction”** will be displayed in the top left corner of the result screen. Please refer to 3.2 Software instructions/important information on page 11 for further information.
- Step 12** Repeat for all samples.
- Step 13** Press **Menu/Options** to display available Options which are described on page 8.
- Step 14** Press **Escape** and confirm with **Yes** to return to the **Nucleic Acids** folder.

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on page 6 (P 300) and 7 (P330 / P360).

## 4.2 Protein Determination

### 4.2.1 General Information

#### Protein determination at 280 nm (NanoVolume Applications and Cuvette Applications)

- Protein can be determined in the near UV at 280 nm due to absorption by tyrosine, tryptophan and phenylalanine amino acids; Abs 280 varies greatly for different proteins due to their amino acid content, and consequently the specific A280 factor for a particular protein must be determined.
- The protein concentration can be calculated the following way:  

$$C_{\text{prot.}} = \text{Abs. 280} * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$
 With background correction:  

$$C_{\text{prot.}} = (\text{Abs. 280} - \text{Abs. 320}) * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$
 This equation can be applied to other proteins if the corresponding factors are known (please note that the factor used by the NanoPhotometer® P-Class is the reciprocal value of the extinction coefficient (l/g\*cm) from a protein). The instrument can determine protein concentration at 280 nm and uses the above equation as default; the factors can be changed, and the use of background correction at 320 nm is optional.
- The A280 Factor is based on the extinction coefficient of the protein [molecular weight/molar extinction coefficient (M<sup>-1</sup>\*cm<sup>-1</sup>) or 1/extinction coefficient (l/g\*cm)].  
 In the software are the following protein A280 factors pre-programmed:  
 BSA (bovine serum albumin), serum albumin (mouse and human), lysozyme, IgG and OD 1 for more information about the factors see 11.3 Protein quantification.  
 There is also the possibility to enter custom factors. For correct calculation the following settings are needed, either the extinction coefficient (l/g\*cm) or the molar extinction coefficient (M<sup>-1</sup>\*cm<sup>-1</sup>) and the molecular weight (g/mol) of the protein.
- Rapid measurements such as this at 280 nm are particularly useful after isolation of proteins and peptides from mixtures using spin and HiTrap columns by centrifuge and gravity, respectively.

#### Protein determination at 280 nm and degree of labelling (NanoVolume Applications and Cuvette Applications)

- To determine the degree of labelling, the absorbance reading at the wavelength reported for maximum absorbance of the fluorescence dye is used. The corresponding extinction coefficient of the dye is used in the Lambert-Beer Law to determine the dye concentration ( $c = A / (e * d)$ ). Absorbance values and extinction coefficients are used to calculate the dye per protein ratio. For further details please refer to 12.4 Protein fluorescent dye incorporation.

#### Colorimetric Bradford, Biuret, BCA and Lowry protein determination (Cuvette Applications)

- The **Bradford** method depends on quantifying the binding of a dye, Coomassie Brilliant Blue, to an unknown protein and comparing this binding to that of different, known concentrations of a standard protein at 595 nm; this is usually BSA (bovine serum albumin).
- The **Biuret** method depends on reaction between cupric ions and peptide bonds in an alkali solution, resulting in the formation of a complex absorbing at 546 nm.
- The **BCA** method also depends on reaction between cupric ions and peptide bonds, but in addition combines this reaction with the detection of cuprous ions using bicinchoninic acid (BCA), giving an absorbance maximum at 562 nm. The BCA process is less sensitive to the presence of detergents used to break down cell walls.
- The **Lowry** method is based on the Biuret reaction. Under alkaline conditions the divalent copper ion forms a complex with peptide bonds in which it is reduced to a monovalent ion. Monovalent copper ion and the radical groups of tyrosine, tryptophan, and cysteine react with Folin reagent to produce an unstable product that becomes reduced to molybdenum/tungsten blue. Bound reagent changes colour from yellow to blue. This binding is compared with those derived from a standard protein at 750 nm; this is usually BSA (bovine serum albumin).
- Detailed protocols are supplied with these assay kits, and must be closely followed to ensure accurate results are obtained.
- A linear regression analysis of the calibration standard data points is calculated; the result, together with the correlation coefficient, can be printed out. A correlation coefficient of between 0.95 and 1.00 indicates a good straight line.

## 4.2.2 Protein UV Method

The procedure is as follows:

### Parameter Screen

#### NanoVolume Applications

Protein-UV - Parameters	
Lid Factor ◀ 10 ▶	Protein BSA
Dilution Factor 1.000	Units µg/ml
Background On	
◊ OK	⊗ Cancel

#### Cuvette Applications

Protein UV - Parameters	
Pathlength ◀ 10 mm ▶	Protein BSA
Dilution Factor 1.000	Units µg/ml
Background On	
◊ OK	⊗ Cancel

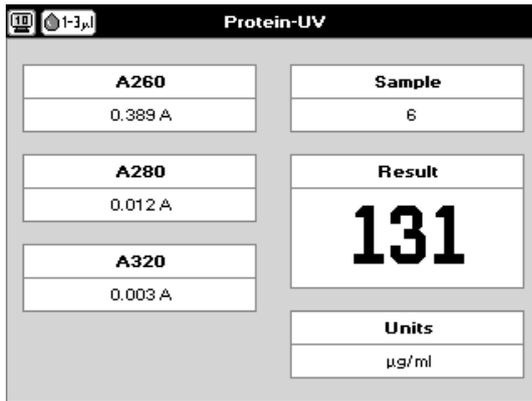
Protein UV - Parameters	
Protein ◀ Molar Ext. Coefficient ▶	Wavelength 200 nm
	Molar Ext. coefficient 47790
	Molecular Weight 69323.398
◊ OK	⊗ Cancel

Protein UV - Parameters	
Protein ◀ Ext Coefficient [l/g*cm] ▶	Wavelength 200 nm
	Ext. coefficient [l/g*cm] 0.690
◊ OK	⊗ Cancel

### Parameter Screen

- Step 1** Press 1 for *NanoVolume* OR 2 for *Cuvette* folder.
- Step 2** Press 2 to select *Protein* folder.
- Step 3** Press 1 to select *Protein UV* mode.
- Step 4** Using NanoVolume Applications select the *Lid Factor* as described in the “Average Detection Range Sheet” or under 3.2. A minimum of **1.5 µl sample volume** (for lid 10) is recommended.
- Step 17** Enter the *Dilution Factor* using the keypad numbers. Range 1.00 to 9,999. Use the C button to backspace and clear the last digit entered. **OR** press *Menu/Options* to enter the dilution factor screen. Enter the volume of the sample using the keypad numbers. Range 0.01 to 9,999. Enter the volume of the diluent using the keypad numbers. Range 0.01 to 9,999. Press **OK** ◊ to calculate the dilution factor and return to the Parameters screen **OR** press **Cancel** ⊗ to cancel the selections and return to the Parameters screen.
- Step 5** Select whether the *Background* correction at 320 nm is used or not with the left and right arrows. It is recommended to switch on the *Background* correction.
- Step 6** Select the *Protein* (BSA (default), Serum Albumin (mouse), Serum Albumin (human), IgG, Lysozyme, Custom or OD 1).
- Step 7** If using *Custom Protein* there are two possibilities to enter the correct factors:  
Molar extinction coefficient ( $M^{-1} * cm^{-1}$ ):  
 Ranges are:  
*Wavelength:* 200 nm to 340 nm  
*Molar extinction coefficient ( $M^{-1} * cm^{-1}$ ):* 10,000 to 9,999,999  
*Molecular weight:* 0.001 to 9,999,999
- Extinction coefficient (l/g \* cm):  
 Ranges are:  
*Wavelength:* 200 nm to 340 nm  
*Extinction coefficient (l/g \* cm):* 0.001 to 9,999
- Step 8** Select the *Units* of measurement using the left and right arrows. Options: mg/ml, µg/ml, ng/µl and µg/µl.
- Step 9** Press **OK** ◊ to enter the Results screen **OR** **Cancel** ⊗ to return to the *Protein* folder.

**Results Screen**



**Results Screen**

- Step 10** Apply/insert the reference sample. Press **Blank** Key. This will be used for all subsequent samples until changed.
- Step 11** Apply/insert sample and press **Sample** . This measures at both 260 and 280 nm wavelengths and displays the result. Protein concentration is calculated (corrected by background wavelength value if selected).
- Step 12** If the absorbance value of the sample is not in the linear range a **“Warning message”** will pop up and **“Instruction”** will be displayed in the top left corner of the result screen. Please refer to 3.2 Software instructions/important information on page 11 for further information.
- Step 13** Repeat for all samples.
- Step 14** Press **Menu/Options** to display available Options which are described on page 8.
- Step 15** Press **Escape** and confirm with **Yes** to return to the **Protein** folder

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on page 6 (P 300) and 7 (P330 / P 360).

### 4.2.3 Protein UV Dye Method

The procedure is as follows:

#### Parameter Screen NanoVolume Applications

#### Cuvette Applications

#### Parameter Screen

- Step 1** Press 1 for *NanoVolume* OR 2 for *Cuvette* folder.
- Step 2** Press 2 to select *Protein* folder.
- Step 3** Press 2 to select *Protein dye* mode.
- Step 4** Using NanoVolume Applications select the **Lid Factor** as described in the “Average Detection Range Sheet” and under 3.2. A minimum of **1.5 µl sample** volume (for lid 10) is recommended.
- Step 5** Enter the **Dilution Factor** using the keypad numbers. Range 1.00 to 9,999. Use the C button to backspace and clear the last digit entered. **OR** press **Menu/Options** to enter the dilution factor screen. Enter the volume of the sample using the keypad numbers. Range 0.01 to 9,999. Enter the volume of the diluent using the keypad numbers. Range 0.01 to 9,999. Press **OK** to calculate the dilution factor and return to the Parameters screen **OR** press **Cancel** to cancel the selections and return to the Parameters screen.
- Step 6** Select whether the **Dye correction** (calculation of the dye-dependent correction factor) is used or not with the left and right arrows. The **Background** correction is always calculated in the Dye methods.
- Step 7** Select the **Protein** (BSA (default), Serum Albumin (mouse), Serum Albumin (human), IgG, Lysozyme, Custom or OD 1).
- Step 8** If using **Custom Protein** there are two possibilities to enter the correct factors (see also page 18 protein UV method):
  - Molar extinction coefficient ( $M^{-1} * cm^{-1}$ ):**
    - Ranges are:
    - Wavelength:** 200 nm to 340 nm
    - Molar extinction coefficient ( $M^{-1} * cm^{-1}$ ):** 10,000 to 9,999,999
    - Molecular weight:** 0.001 to 9,999,999
  - Extinction coefficient ( $l/g * cm$ ):**
    - Ranges are:
    - Wavelength:** 200 nm to 340 nm
    - Extinction coefficient ( $l/g * cm$ ):** 0.001 to 9,999
- Step 9** Select the **Units** of measurement using the left and right arrows. Options: mg/ml, µg/ml, ng/µl and µg/µl.
- Step 10** Enter the protein-dependent extinction coefficient. Range is 10,000 to 9,999,999.
- Step 11** Press **OK** to store the chosen parameters and to enter the next screen **OR** **Cancel** to return to the **Protein** folder.
- Step 12** Select the appropriate Dye Type. 4 different AlexaFluors, 2 Cy-Dyes, 2 DyLight Dyes, FITC, Pacific Blue, r-PE and Texas Red are programmed with their corresponding maximum absorbance wavelength, dye-dependent extinction co-efficient and dye-dependent correction factor at 280 nm. For further details please refer to 12.4 Protein fluorescent dye incorporation.
- Step 13** If using **Custom Dye** maximum absorbance wavelength of the custom dye, dye-dependent extinction coefficient and dye-dependent correction factor at 280 nm have to be entered. Ranges are:
  - Dye Abs Max:** 300 nm to 950 nm
  - Dye Ext. Coefficient:** 10,000 to 9,999,999
  - Dye Correction:** 0 to 0.999

**Results Screen**

Protein Dye	
A260	0.494 A
A280	0.035 A
A320	0.003 A
ADye(346)	0.003 A
<b>Sample</b>	
7	
<b>Protein Concentration</b>	
464 µg/ml	
<b>Dye Concentration</b>	
0.00 pmol/µl	
<b>Degree of Labelling</b>	
0.00	

**Results Screen**

- Step 14** Apply/insert the reference sample. Press **Blank** Key. This will be used for all subsequent samples until changed.
- Step 15** Apply/insert sample and press **Sample** . This measures at 260nm, 280nm, 320nm and the dye specific wavelength and displays the result. Protein concentration (corrected by background wavelength value if selected), dye concentration and degree of labelling is calculated.
- Step 16** If the absorbance value of the sample is not in the linear range a **“Warning message”** will pop up and **“Instruction”** will be displayed in the top left corner of the result screen. Please refer 3.2 Software instructions/important information on page 11 for further information.
- Step 17** Repeat for all samples.
- Step 18** Press **Menu/Options** to display available Options which are described on page 8.
- Step 19** Press **Escape** and confirm with **Yes** to return to the **Protein** folder.

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on 6 (P 300) and 7 (P330 / P 360).

### 7.3.4 BCA Assay

The colorimetric BCA assay is not recommended with the Submicroliter Cell. Please use Cuvette Applications. The procedure is as follows:

#### Parameter Screen

#### Standards Screen

#### Parameter Screen

- Step 1** Press 2 to select **Cuvette** folder.
- Step 2** Press 2 to select **Protein** folder.
- Step 3** Press 2 to select **BCA** mode.
- Step 4** The default **Wavelength** setting is 562 nm.
- Step 5** Enter the number of **Standard** concentration points (1-9) to be used in the curve using the keypad numbers or left and right arrows.

- Step 6** **Units:** The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows (µg/ml, µg/µl, pmol/µl, mg/ml, mmol/l, µmol/l, g/l, mg/l, µg/l, U/l, %, ppm, ppb, conc or none). These units can also be edited once **OK** is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters **OR Cancel** .

- Step 7** Press **Next** to enter the next screen.

- Step 8** Select the **Calibration** mode, either **Standards** (measure prepared standards), **Manual** (keypad data entry) or **New Standards** (previous values are blanked, new standard can be measured).

- Step 9** (if **Standards** selected) Select the number of **Replicates** using the left and right arrows. This determines the number of standards to be measured and averaged at each standard concentration point. Can be Off (1), 2 or 3.

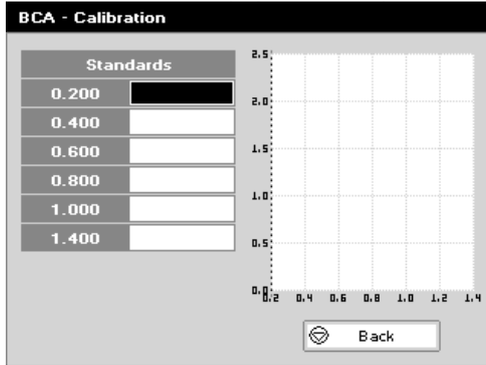
- Step 10** Press **Next** to enter the Standards screen **OR** press **Cancel** to cancel selections and return to the **Protein** folder.

#### Standards Screen

- Step 11** Enter the concentration values by using the keypad numbers and the up and down arrows to move between the different standard boxes. Range 0.001 to 9,999. C button backspaces and clears the last digit entered.

- Step 12** Press **Next** to enter the Calibration screen. If there are duplicate or non-monotonic (increasing) entries the unit will beep and highlight the incorrect entry. **OR** press **Back** to return to the Parameter screen.

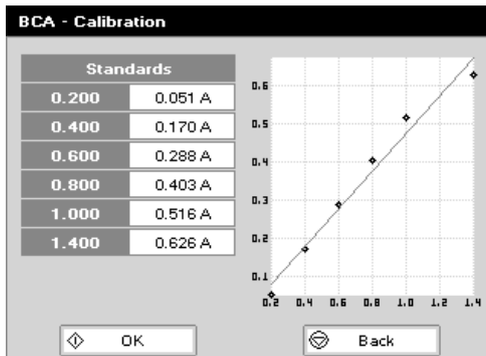
**Calibration Screen (replicates off)**



**Calibration Screen (replicates off)**

**Step 13** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

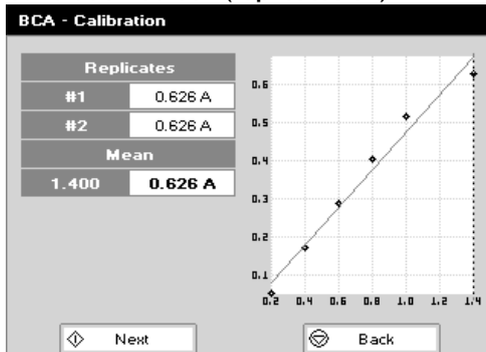
**Step 14** Insert the standard (use C to clear previously stored results before measuring). Press **Sample**  $\blacklozenge$  to measure the standard and store the result.



**Step 15** Repeat for all standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 16** When all standards are measured press **OK**  $\blacklozenge$  to accept the calibration and go to the Results screen (see below) **OR** press **Back**  $\square$  to cancel selections and return to the Standards screen.

**Calibration Screen (replicates on)**



**Calibration Screen (replicates on)**

**Step 17** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

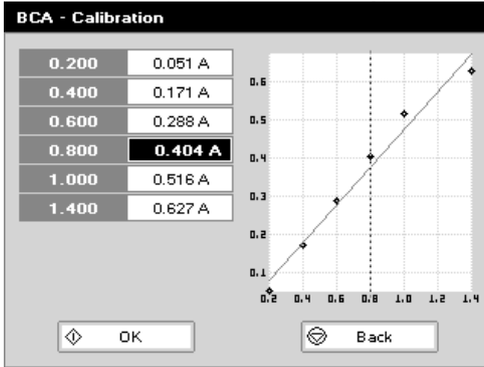
**Step 18** Press **Next**  $\blacklozenge$  to display the replicate entry boxes. Use C to clear previously stored results before measuring.

**Step 19** Insert the standard and press **Sample**  $\blacklozenge$  to measure the standard and store the result.

**Step 20** Repeat for all replicates and standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 21** Press **OK**  $\blacklozenge$  to accept the calibration and go to the Results screen (see below) **OR** press **Back**  $\blacklozenge$  to return to the Standards screen.

**Calibration Screen (manual entry)**



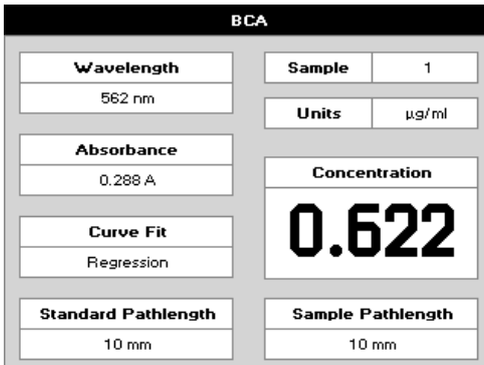
**Calibration Screen (manual entry)**

Shows previously entered calibration values and allows values to be entered via the keypad.

**Step 22** The highlighted box can be edited in order to enter an absorbance value corresponding to a given concentration value using the keypad numbers. Range 0.001 to 9,999. Use C to backspace and clear the last digit entered and the up and down arrows to move between boxes.

**Step 23** Press **OK** to accept the calibration and go to the Results screen (see below) OR press **Back** to return to the Standards screen.

**Results screen**



**Results screen**

**Step 24** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.

**Step 25** Insert the sample and press **Sample** . The concentration of the sample is taken and displayed.

**Step 26** Repeat for all samples.

**Step 27** Press **Menu/Options** to display available options which are described below.

**Step 28** Press **Escape** and confirm with **Yes** to return to the **Protein** folder. Query needs confirmation to avoid unintended escaping the application.

**NanoPhotometer® P 300**

**Options (select using key pad numbers)**

- 1) Parameters...
- 2) Print
- 3) Graph
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Printer Settings...

**Options (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle graph on/off. Displays the calibration graph, cursors give values for last measured sample.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in 7.3 Output Options / Printer

Exit **Options** by pressing **Escape** , or wait.

**NanoPhotometer® P 330 / P360**

**Menu (select using key pad numbers)**

- 1) Parameters...
- 2) Data Transfer
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Output Options...

**Menu (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in 7.3 Output Options / Printer

Exit **Menu** by pressing **Escape** , or wait.

### 7.3.5 Bradford Assay

The colorimetric Bradford assay is not recommended with the Submicroliter Cell. Please use Cuvette Applications. The procedure is as follows:

#### Parameter Screen

#### Standards Screen

#### Parameter Screen

- Step 1** Press 2 to select **Cuvette** folder.  
**Step 2** Press 2 to select **Protein** folder.  
**Step 3** Press 3 to select **Bradford** mode.  
**Step 4** The default **Wavelength** setting is 595 nm.  
**Step 5** Enter the number of **Standard** concentration points (1-9) to be used in the curve using the keypad numbers or left and right arrows.

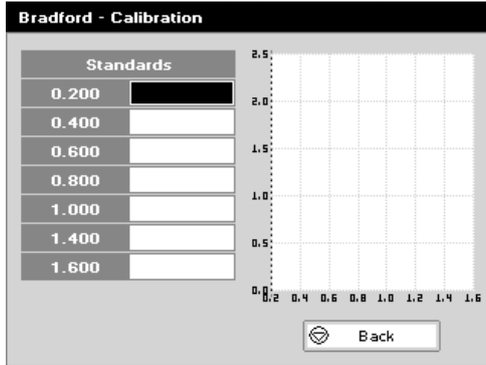
- Step 6** **Units:** The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows ( $\mu\text{g/ml}$ ,  $\mu\text{g}/\mu\text{l}$ ,  $\text{pmol}/\mu\text{l}$ ,  $\text{mg/ml}$ ,  $\text{mmol/l}$ ,  $\mu\text{mol/l}$ ,  $\text{g/l}$ ,  $\text{mg/l}$ ,  $\mu\text{g/l}$ ,  $\text{U/l}$ ,  $\%$ ,  $\text{ppm}$ ,  $\text{ppb}$ ,  $\text{conc}$  or  $\text{none}$ ). These units can also be edited once **OK** is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters **OR Cancel** .

- Step 7** Press **Next** to enter the next screen.  
**Step 8** Select the **Calibration** mode, either **Standards** (measure prepared standards), **Manual** (keypad data entry) or **New Standards** (previous values are blanked, new standard can be measured).  
**Step 9** (if **Standards** selected) Select the number of **Replicates** using the left and right arrows. This determines the number of standards to be measured and averaged at each standard concentration point. Can be Off (1), 2 or 3.  
**Step 10** Press **Next** to enter the Standards screen **OR** press **Cancel** to cancel selections and return to the **Protein** folder.

#### Standards Screen

- Step 11** Enter the concentration values by using the keypad numbers and the up and down arrows to move between the different standard boxes. Range is 0.001 to 9,999. C button backspaces and clears the last digit entered.  
**Step 12** Press **Next** to enter the Calibration screen. If there are duplicate or non-monotonic (increasing) entries the unit will beep and highlight the incorrect entry. **OR** press **Back** to return to the Parameter screen.

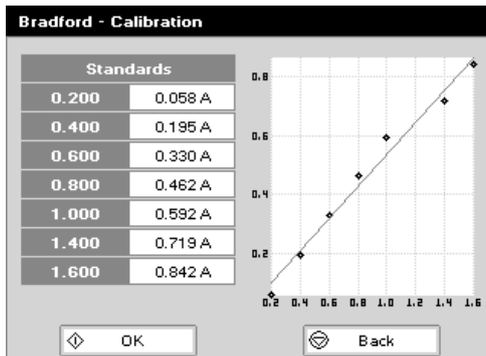
**Calibration Screen (replicates off)**



**Calibration Screen (replicates off)**

**Step 13** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

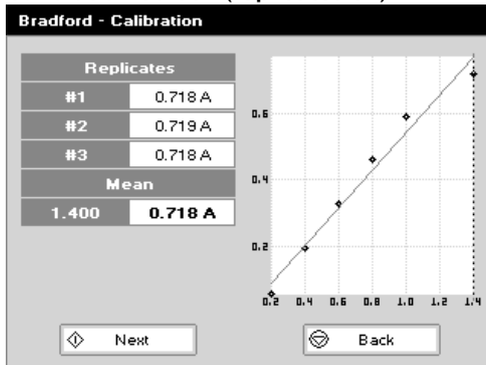
**Step 14** Insert the standard (use C to clear previously stored results before measuring). Press **Next** to measure the standard and store the result.



**Step 15** Repeat for all standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 16** When all standards are measured press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to cancel selections and return to the Standards screen.

**Calibration Screen (replicates on)**



**Calibration Screen (replicates on)**

**Step 17** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

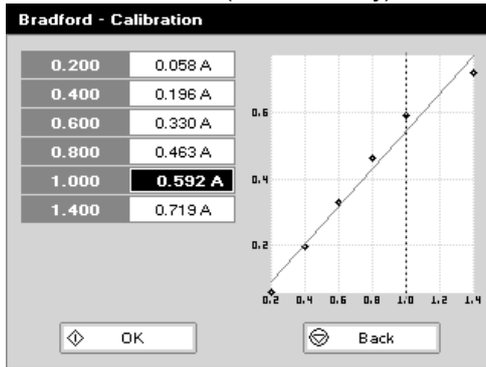
**Step 18** Press **Replicates** to display the replicate entry boxes. Use C to clear previously stored results before measuring.

**Step 19** Insert the standard and press **Sample** to measure the standard and store the result.

**Step 20** Repeat for all replicates and standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 21** Press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

**Calibration Screen (manual entry)**



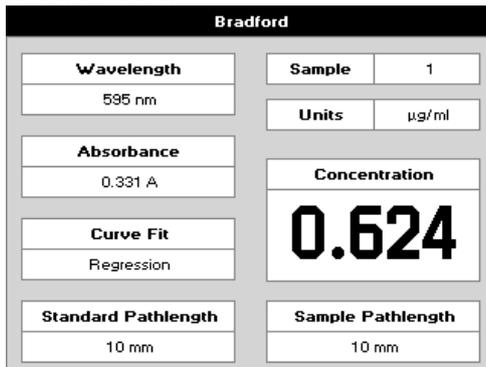
**Calibration Screen (manual entry)**

Shows previously entered calibration values and allows values to be entered via the keypad.

**Step 22** The highlighted box can be edited in order to enter an absorbance value corresponding to a given concentration value using the keypad numbers. Range is 0.001 to 9,999. Use C to backspace and clear the last digit entered and the up and down arrows to move between boxes.

**Step 23** Press **OK** to accept the calibration and go to the Results screen (see below) OR press **Back** to return to the Standards screen.

**Results screen**



**Results screen**

**Step 24** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.

**Step 25** Insert the sample and press **Sample** . The concentration of the sample is taken and displayed.

**Step 26** Repeat for all samples.

**Step 27** Press **Menu/Options** to display available Options which are described below.

**Step 28** Press **Escape** and confirm with **Yes** to return to the **Protein** folder. Query needs confirmation to avoid unintended escaping the application.

**NanoPhotometer® P 300**

**Options (select using key pad numbers)**

- 1) Parameters...
- 2) Print
- 3) Graph
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Printer Settings...

**Options (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle graph on/off. Displays the calibration graph, cursors give values for last measured sample.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Options** by pressing **Escape** , or wait.

**NanoPhotometer® P 330 / P360**

**Menu (select using key pad numbers)**

- 1) Parameters...
- 2) Data Transfer
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Output Options...

**Menu (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in **7.3 Output Options / Printer**

Exit **Menu** by pressing **Escape** , or wait.

### 7.3.6 Lowry Assay

The colorimetric Lowry assay is not recommended with the Submicroliter Cell. Please use Cuvette Applications. The procedure is as follows:

#### Parameter Screen

#### Standards Screen

#### Parameter Screen

- Step 1** Press 2 to select **Cuvette** folder.
- Step 2** Press 2 to select **Protein** folder.
- Step 3** Press 4 to select **Lowry** mode.
- Step 4** The default **Wavelength** setting is 750 nm.
- Step 5** Enter the number of **Standard** concentration points (1-9) to be used in the curve using the keypad numbers or left and right arrows.

- Step 6** **Units:** The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows (µg/ml, µg/µl, pmol/µl, mg/ml, mmol/l, µmol/l, g/l, mg/l, µg/l, U/l, %, ppm, ppb, conc or none). These units can also be edited once **OK** is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters **OR Cancel** .

- Step 7** Press **Next** to enter the next screen.

- Step 8** Select the **Calibration** mode, either **Standards** (measure prepared standards), **Manual** (keypad data entry) or **New Standards** (previous values are blanked, new standard can be measured).

- Step 9** (if **Standards** selected) Select the number of **Replicates** using the left and right arrows. This determines the number of standards to be measured and averaged at each standard concentration point. Can be Off (1), 2 or 3.

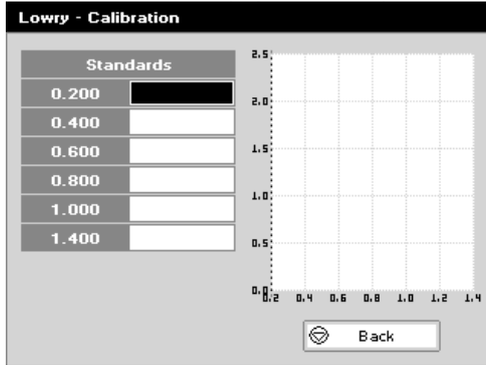
- Step 10** Press **Next** to enter the Standards screen **OR** press **Cancel** to cancel selections and return to the **Protein** folder.

#### Standards Screen

- Step 11** Enter the concentration values by using the keypad numbers and the up and down arrows to move between the different standard boxes. Range 0.001 to 9,999. C button backspaces and clears the last digit entered.

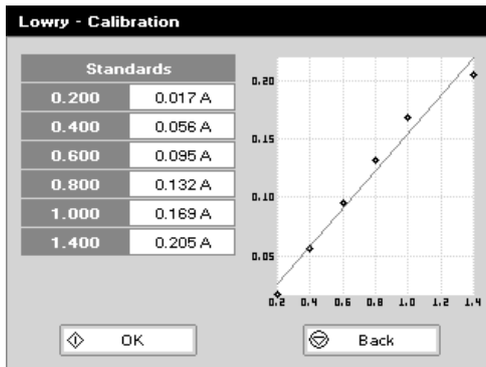
- Step 12** Press **Next** to enter the Calibration screen. If there are duplicate or non-monotonic (increasing) entries the unit will beep and highlight the incorrect entry. **OR** press **Back** to return to the Parameter screen.

**Calibration Screen (replicates off)**



**Step 13** **Calibration Screen (replicates off)** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

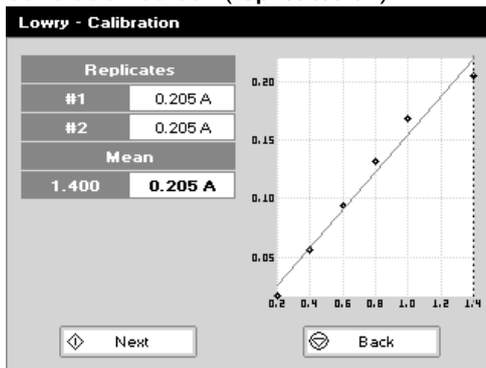
**Step 14** Insert the standard (use C to clear previously stored results before measuring). Press **Sample** to measure the standard and store the result.



**Step 15** Repeat for all standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 16** When all standards are measured press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to cancel selections and return to the Standards screen.

**Calibration Screen (replicates on)**



**Calibration Screen (replicates on)**

**Step 17** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

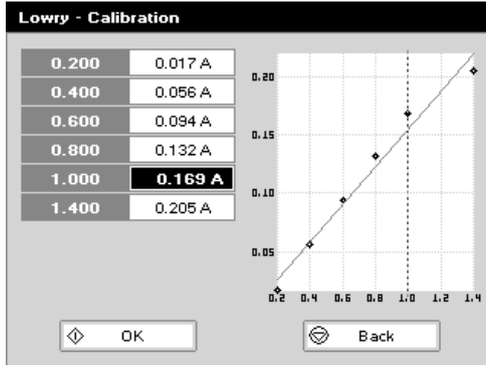
**Step 18** Press **Replicates** to display the replicate entry boxes. Use C to clear previously stored results before measuring.

**Step 19** Insert the standard and press **Sample** to measure the standard and store the result.

**Step 20** Repeat for all replicates and standards. A graph will display the results and the fitted curve as the measurements are input. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 21** Press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

### Calibration Screen (manual entry)



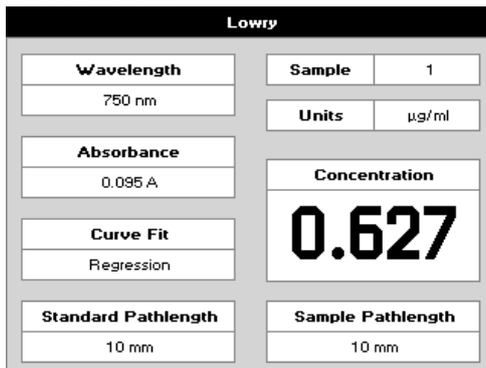
### Calibration Screen (manual entry)

Shows previously entered calibration values and allows values to be entered via the keypad.

**Step 22** The highlighted box can be edited in order to enter an absorbance value corresponding to a given concentration value using the keypad numbers. Range is 0.001 to 9,999. Use C to backspace and clear the last digit entered and the up and down arrows to move between boxes.

**Step 23** Press **OK** to accept the calibration and go to the Results screen (see below) OR press **Back** to return to the Standards screen.

### Results screen



### Results screen

**Step 24** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.

**Step 25** Insert the sample and press **Sample** . The concentration of the sample is taken and displayed.

**Step 26** Repeat for all samples.

**Step 27** Press **Menu/Options** to display available Options which are described below press **Escape** and confirm with **OK** to return to the **Protein** folder. Query needs confirmation to avoid unintended escaping the application.

### NanoPhotometer® P 300

#### Options (select using key pad numbers)

- 1) Parameters...
- 2) Print
- 3) Graph
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Printer Settings...

#### Options (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle graph on/off. Displays the calibration graph, cursors give values for last measured sample.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in 7.3 Output Options / Printer

Exit **Options** by pressing **Escape** , or wait.

### NanoPhotometer® P 330 / P360

#### Menu (select using key pad numbers)

- 1) Parameters...
- 2) Data Transfer
- 4) Edit Sample Pathlength
- 7) Sample Number...
- 8) Save Method...
- 9) Output Options...

#### Menu (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in 7.3 Output Options / Printer

Exit **Menu** by pressing **Escape** , or wait.

### 7.3.7 Biuret Assay

The colorimetric Biuret assay is not recommended with the Submicroliter Cell. Please use Cuvette Applications. The procedure is as follows:

#### Parameter Screen

#### Standards Screen

#### Parameter Screen

- Step 1** Press 2 to select **Cuvette** folder.
- Step 2** Press 2 to select **Protein** folder.
- Step 3** Press 5 to select **Biuret** mode.
- Step 4** The default **Wavelength** setting is 546 nm.
- Step 5** Enter the number of **Standard** concentration points (1-9) to be used in the curve using the keypad numbers or left and right arrows.

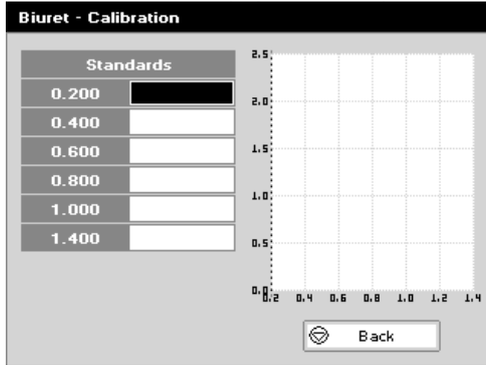
- Step 6** **Units:** The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows ( $\mu\text{g/ml}$ ,  $\mu\text{g}/\mu\text{l}$ ,  $\text{pmol}/\mu\text{l}$ ,  $\text{mg/ml}$ ,  $\text{mmol/l}$ ,  $\mu\text{mol/l}$ ,  $\text{g/l}$ ,  $\text{mg/l}$ ,  $\mu\text{g/l}$ ,  $\text{U/l}$ ,  $\%$ ,  $\text{ppm}$ ,  $\text{ppb}$ ,  $\text{conc}$  or  $\text{none}$ ). These units can also be edited once **OK** is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters **OR Cancel** .

- Step 7** Press **Next** to enter the next screen.
- Step 8** Select the **Calibration** mode, either **Standards** (measure prepared standards), **Manual** (keypad data entry) or **New Standards** (previous values are blanked, new standard can be measured).
- Step 9** (if **Standards** selected) Select the number of **Replicates** using the left and right arrows. This determines the number of standards to be measured and averaged at each standard concentration point. Can be Off (1), 2 or 3.
- Step 10** Press **Next** to enter the Standards screen **OR** press **Cancel** to cancel selections and return to the **Protein** folder.

#### Standards Screen

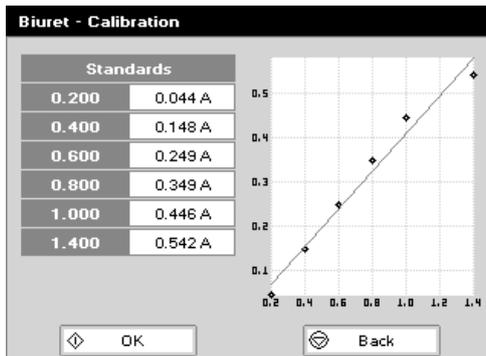
- Step 11** Enter the concentration values by using the keypad numbers and the up and down arrows to move between the different standard boxes. Range is 0.001 to 9,999. C button backspaces and clears the last digit entered.
- Step 12** Press **Next** to enter the Calibration screen. If there are duplicate or non-monotonic (increasing) entries the unit will beep and highlight the incorrect entry **OR** press **Back** to return to the Parameter screen.

**Calibration Screen (replicates off)**



**Step 13** **Calibration Screen (replicates off)** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

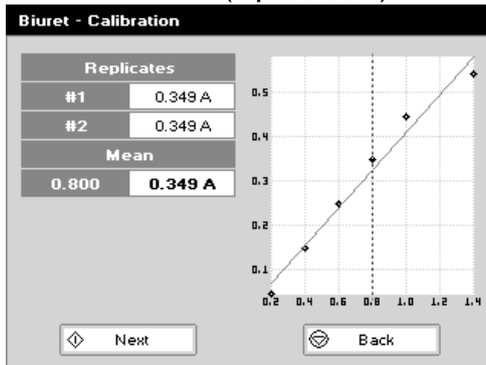
**Step 14** Insert the standard (use C to clear previously stored results before measuring). Press **Sample** to measure the standard and store the result.



**Step 15** Repeat for all standards. A graph will display the results and the fitted curve as the measurements are made. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 16** When all standards are measured press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to cancel selections and return to the Standards screen.

**Calibration Screen (replicates on)**



**Calibration Screen (replicates on)**

**Step 17** This shows the calibration values and allows standards to be measured. Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

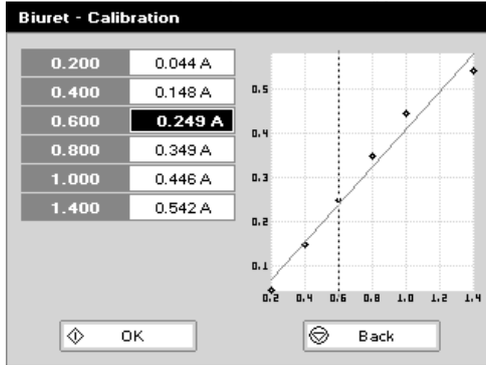
**Step 18** Press **Replicates** to display the replicate entry boxes. Use C to clear previously stored results before measuring.

**Step 19** Insert the standard and press **Sample** to measure the standard and store the result.

**Step 20** Repeat for all replicates and standards. A graph will display the results and the fitted curve as the measurements are input. Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

**Step 21** Press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

**Calibration Screen (manual entry)**

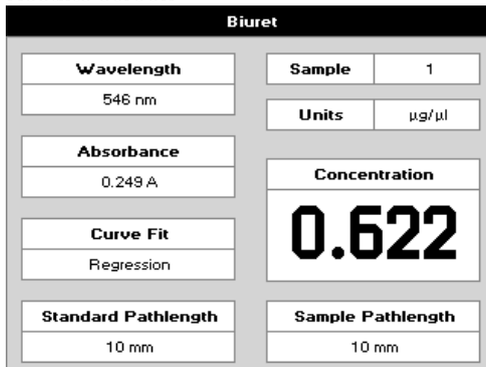


**Calibration Screen (manual entry)**

Shows previously entered calibration values and allows values to be entered via the keypad.

- Step 22** The highlighted box can be edited in order to enter an absorbance value corresponding to a given concentration value using the keypad numbers. Range is 0.001 to 9,999. Use C to backspace and clear the last digit entered and the up and down arrows to move between boxes.
- Step 23** Press **OK**  to accept the calibration and go to the Results screen (see below) **OR** press **Back**  to return to the Standards screen.

**Results screen**

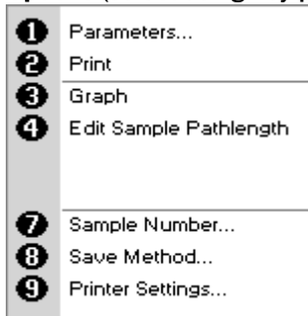


**Results screen**

- Step 24** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.
- Step 25** Insert the sample and press **Sample** . The concentration of the sample is taken and displayed.
- Step 26** Repeat for all samples.
- Step 27** Press **Menu/Options** to display available Options which are described below.
- Step 28** Press **Escape**  and confirm with **Yes**  to return to the **Protein** folder. Query needs confirmation to avoid unintended escaping the application.

**NanoPhotometer® P 300**

**Options (select using key pad numbers)**



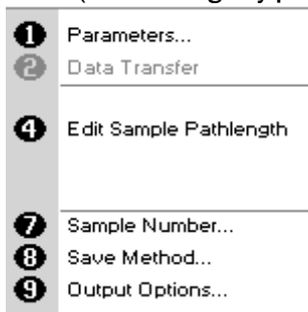
**Options (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle graph on/off. Displays the calibration graph, cursors give values for last measured sample.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Options** by pressing **Escape** , or wait.

**NanoPhotometer® P 330 / P360**

**Menu (select using key pad numbers)**



**Menu (select using key pad numbers)**

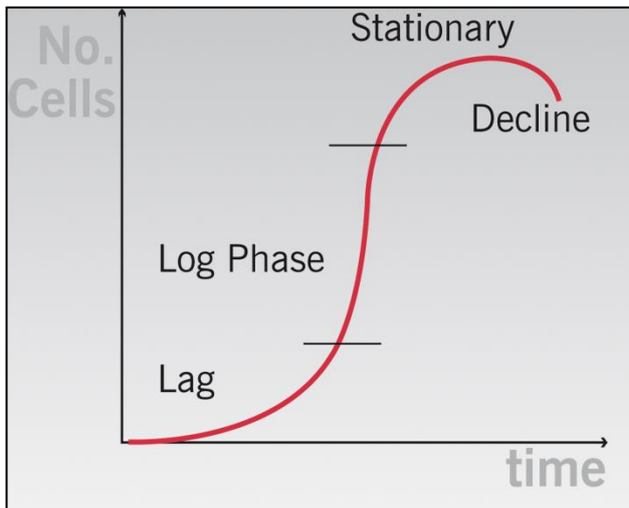
- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in **7.3 Output Options / Printer**

Exit **Menu** by pressing **Escape** , or wait.

## 4.3 Bacterial Cell Culture Measurement (OD600)

### 4.3.1 General Information

- The stage of growth of a bacterial culture needs to be monitored to ensure that the cells are harvested at the optimum point for the greatest density of live cells. An exemplary growth curve is given below. Cells should be harvested towards the end of the log phase. The optical density of the sample indicates when this point has been reached. This value varies dependent on the cells being grown. Routinely the cells are grown until the absorbance at 600 nm (known as OD 600) reaches approximately 0.4 prior to induction or harvesting. A linear relationship exists between cell number (density) and OD 600 up to approx. 0.6



- It is important to note that for turbid samples such as cell cultures, the absorbance measured is due to light scattering, and **not** the result of molecular absorption. The amount of scatter is affected by the optics of the system (distance between the cell holder and instrument exit slit, geometry of this slit and the monochromator optics). Different spectrophotometer types therefore give different responses for the same turbid sample; to compare results, they must be normalized using calibration curves.
- A calibration curve can be determined by comparing measured OD 600 to expected OD 600. Expected OD 600 is determined by counting cell number using an alternative technique (for example microscope slide method) and converting to OD 600 using the rule of thumb that  $1 \text{ OD } 600 = 5 \times 10^8 \text{ cells/ml}$  for E. Coli.
- Additionally your NanoPhotometer® P-Class is coming with a correction factor of 1 as default. To compare OD values between different spectrophotometer, you have to determine the constant deviation between the Absorbance values for the same sample within those instruments and use this factor within the setting "correction factor" of your NanoPhotometer® P-Class Software.
- The use of 10 mm pathlength disposable cells is recommended for optical density measurements of cell culture solutions; to prevent the suspension settling too quickly and giving an OD that changes with time, glycerol should be added to the sample.
- The Submicroliter Cell is not recommended for optical density measurements of cell culture solutions.

### 4.3.2 Analysis of Bacterial Growth

The procedure is as follows:

#### Parameter Screen

OD 600 - Parameters

Wavelength	600 nm
Correction	1.000
Units	OD

OK Cancel

OD 600 - Parameters

Wavelength	600 nm	Factor	500
Correction	1.000	Multiplier	x 1,000,000
Units	cells/ml		

OK Cancel

#### Results Screen

OD 600

Wavelength	950 nm	Sample	10
Absorbance	0.010 A	cells/ml	5.000
		Multiplier	x 1,000,000

#### Parameter Screen

- Step 1** Press 2 to select **Cuvette Applications**.
- Step 2** Press 3 to select **OD 600**.
- Step 3** Select the **Wavelength**. Default value is 600 nm. Range is 200 nm to 950 nm.
- Step 4** Enter the **Correction** factor to compensate for different optical configurations between this and other instruments. Default value is 1.
- Step 5** Select the **Units**. Options are OD or cells/ml. If cells/ml is selected two further parameters are displayed.

- Step 6** (if cells/ml selected) Enter the **Factor** using the keypad numbers. Range 0.00 to 9,999. C button backspaces and clears the last digit entered.
- Step 7** (if cells/ml selected) Select the **Multiplier** using the left and right arrows. Options are 1,000 or 1,000,000. **Factor** and **Multiplier** define the conversion of the measured OD to the number of cells per millilitre (e.g.: 1 OD 600 = 5 x 10<sup>8</sup> cells/ml)
- Step 8** Press **OK** to enter the Results screen **OR** press **Cancel** to cancel selections and return to the **Cuvette Applications** folder.

#### Results Screen

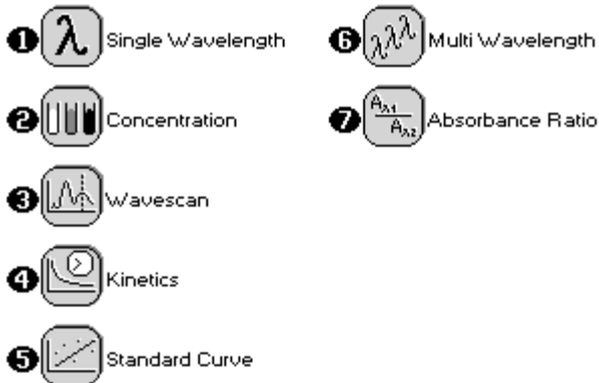
- Step 9** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.
- Step 10** Insert the sample and press **Sample** . The wavelength, absorbance and OD600 value is displayed.
- Step 11** Repeat for all samples.
- Step 12** Press **Menu/Options** to display available Options which are described below.
- Step 13** Press **Escape** and confirm with **Yes** to return to the **Cuvette Applications** folder. Query needs confirmation to avoid unintended escaping the application.

To change parameters, print or save methods press the Menu/Options button. The options menu will be opened. For further explanation please see 2.3 Keypad and display on page 6 (P 300) and 7 (P 330 / P 360).

## 5. FUNCTIONS

Survey of the available Functions:

### Functions



Key pad number	Description
1	Absorbance or %T (transmission) at a single user defined wavelength.
2	Colorimetric assay at a single wavelength based on a simple Factor entered or calculated from a single standard.
3	Spectral plot between two user defined wavelengths. Range 200-950 nm, with user configurable peak finding function.
4	Kinetic colorimetric assay either rate or end value based.
5	Colorimetric assay at a single wavelength based on a user programmed curve.
6	Absorbance or %T (transmission) at up to 5 user defined wavelengths.
7	Ratio of absorbance values at two user specified wavelengths.

### Menu/Options

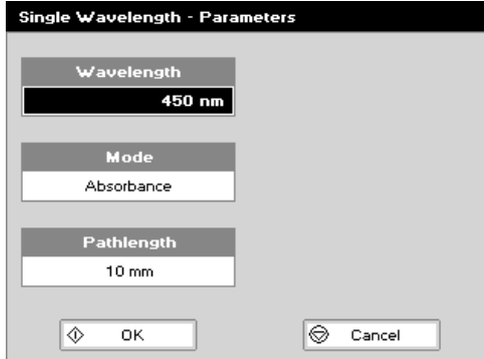
Within each function the user has the possibility to select various options that define the way results are treated. If not using a stored method, it is advisable to check that these options have been appropriately set for your experiment when coming to the instrument. Note that setting the “History” parameter to On (see Preferences later) will cause the instrument to store its last settings. If the “History” parameter is turned Off, all parameters and selections will return to their default settings when leaving that application. (Unless it has been saved as a method).

### 5.1 Single Wavelength – Abs and %T

This makes simple absorbance (A) and % transmission (%T) measurements on samples, measuring the amount of light that has passed through a sample relative to a reference (this can be air).

The procedure is as follows:

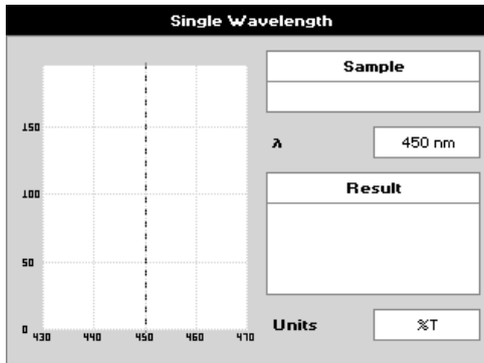
#### Parameter Screen



#### Parameter Screen

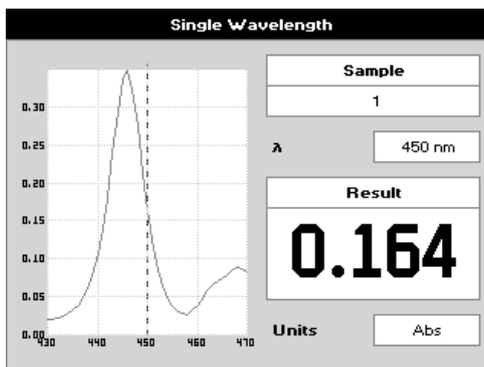
- Step 1** Press 3 to select **Functions**.
- Step 2** Press 1 to select **Single Wavelength**
- Step 3** Set **Wavelength** by using keypad numbers or left and right arrows.
- Step 4** Select the **Mode, Absorbance** or **%Transmission**, using the left and right arrows.
- Step 5** To enter the results screen with the selected parameters press **OK** **OR** cancel the selections and return to the **Functions** folder by pressing **Cancel** .

#### Results Screen



#### Results Screen

- Step 6** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 7** Insert sample and press **Sample** .
- Step 8** Repeat for all samples.



- Step 9** The result at the selected wavelength is displayed on the screen.
- Step 10** Use the left and right arrows to move the cursor and display the value at the cursor position (+/- 15nm from set wavelength).
- Step 11** Press **Menu/Options** to display available options which are described below.
- Step 12** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application folder.

**NanoPhotometer® P 300**

**Options (select using key pad numbers)**

1	Parameters...
2	Print
3	Abs/%T
4	Print Graph
7	Sample Number...
8	Save Method...
9	Printer Settings...

**Options (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle between Absorbance and %T mode.
- 4) Print graph – greyed out if no data are available.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Options** by pressing **Escape** , or wait.

**NanoPhotometer® P 330 /P 360**

**Menu (select using key pad numbers)**

1	Parameters...
2	Data Transfer
3	Abs/%T
4	Print Graph
7	Sample Number...
8	Save Method...
9	Output Options...

**Menu (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 3) Toggle between Absorbance and %T mode.
- 4) Print graph – greyed out if no data are available.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in **7.3 Output Options / Printer**.

Exit **Menu** by pressing **Escape** , or wait.

## 5.2 Concentration

This makes simple concentration measurements on samples, by measuring the amount of light that has passed through a sample relative to a reference (this can be air). Concentration is obtained by multiplying the measured absorbance at a specific wavelength by a factor. The factor may be known in advance, or may be calculated by the instrument by measuring a standard of known concentration.

The procedure is as follows:

### Parameter Screen

### Results Screen (if using a Factor)

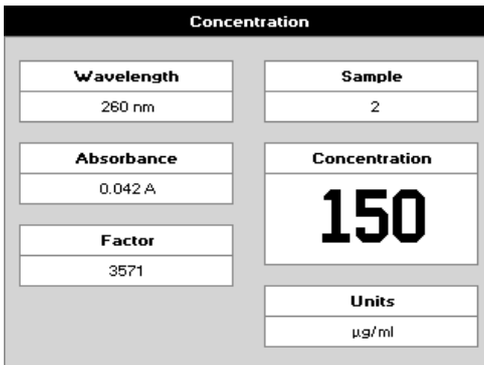
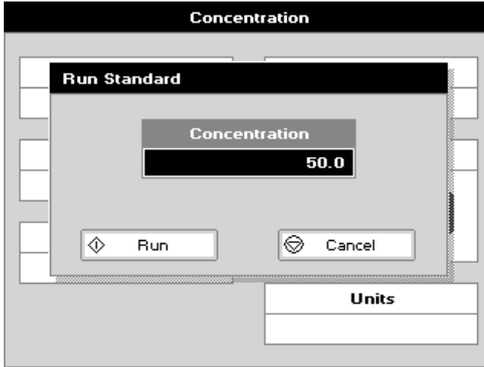
### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 2 to select **Concentration**
- Step 3** Set **Wavelength** by using keypad numbers or left and right arrows.
- Step 4** Select the **Mode**: Factor (user entered) or Standard (factor is calculated from a calibration sample), using the left and right arrows.
- Step 5** (if Factor is selected) Enter the **Factor** using the keypad numbers. Range 0.001 to 9,999. Use the C button to delete the last digit entered.
- Step 6** (if Standard is selected) Enter the concentration using keypad numbers. Range 0.01 to 9,999. Use the C button to delete the last digit entered.
- Step 7** **Units**: The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows (µg/ml, µg/µl, pmol/µl, mg/ml, mmol/l, µmol/l, g/l, mg/l, µg/l, U/l, %, ppm, ppb, conc or none). These units can also be edited once **OK** is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters or **Cancel** .
- Step 8** To enter the results screen with the selected parameters press **OR** cancel the selections and return to the **Functions** folder by pressing **Cancel** .

### Results Screen (if using a Factor)

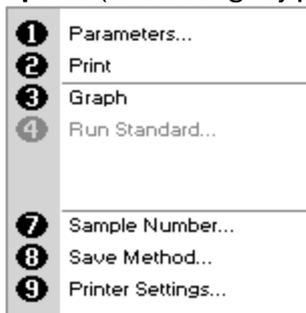
- Step 9** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 10** Insert sample and press **Sample** .

**Results Screen (if using standard mode)**



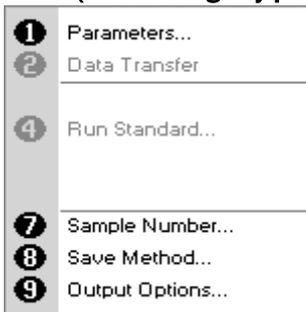
**NanoPhotometer® P 300**

**Options (select using key pad numbers)**



**NanoPhotometer® P 330 / P 360**

**Menu (select using key pad numbers)**



**Results Screen (if using standard mode)**

**Step 11** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

**Step 12** Press **Sample** to display the Run Standard screen.

**Step 13** Run the standard by pressing **Run** OR Press **Cancel** to return to the measure screen.

**Step 14** Insert the sample and press **Sample** . The concentration of the sample is displayed. Results shown as — indicate the concentration is out of range.

**Step 15** Repeat for all samples.

**Step 16** Press **Menu/Options** to display available options which are described below.

**Step 17** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application.

**Options (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggles on/off, displaying a graph of wavescan +/- 20 nm from selected wavelength.
- 4) Return to Run Standard screen.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Options** by pressing **Escape** , or wait.

**Menu (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 4) Return to Run Standard screen.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in **7.3 Output Options / Printer**

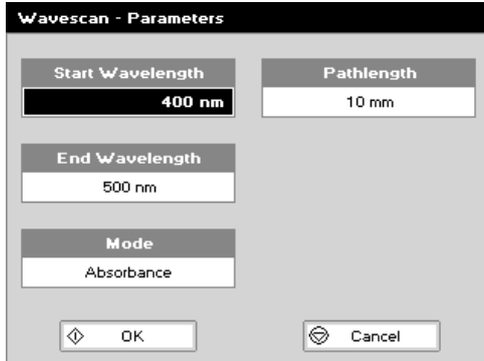
Exit **Menu** by pressing **Escape** , or wait.

### 5.3 Wavescan

An absorption spectrum can be obtained from your instrument, enabling simple identification of peak height and position.

The procedure is as follows:

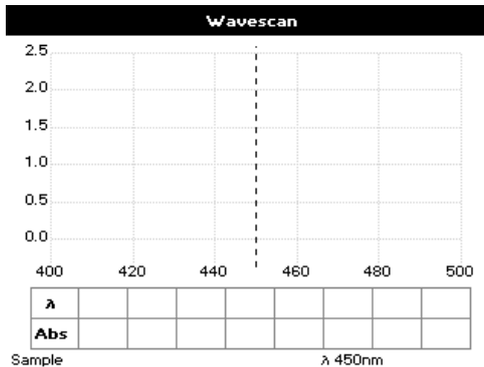
#### Parameter Screen



#### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 3 to select **Wavescan**.
- Step 3** Set **Start Wavelength** by using keypad numbers or left and right arrows.
- Step 4** Set **End Wavelength** by using keypad numbers or left and right arrows.
- Step 5** Select the **Mode**, **Absorbance** or **%Transmission**, using the left and right arrows.
- Step 6** To enter the measurements screen with the selected parameters press **OK** **OR** cancel the selections and return to the **Functions** folder by pressing **Cancel** .

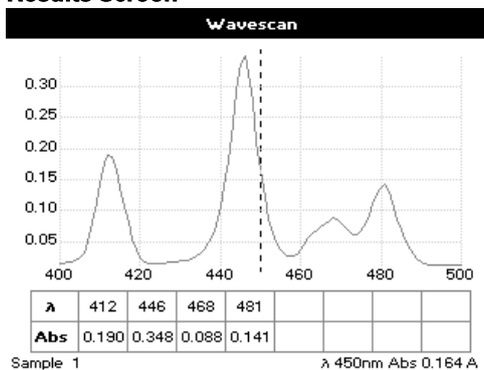
#### Measurement Screen



#### Measurement Screen

- Step 7** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 8** Insert sample and press **Sample** .
- Step 9** Repeat for all samples.

#### Results Screen



#### Results Screen

A graph of the wavescan is displayed, along with a table of Absorbance/%T at each peak. Up to eight peaks can be shown. Use the left and right arrows to move the cursor along the graph. When it reaches a peak the peak height and width of the peak is displayed at the top of the screen. To zoom in on the wavelength scale, use the up arrow. This auto-scales on the Absorbance/%T scale (dependent on the Graph Scale option) and this is retained for subsequent measurements. To zoom out again, use the down arrow.

- Step 10** Press **Menu/Options** to display available options which are described next.

- Step 11** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application.

**Menu (select using key pad numbers)**

- 1 Parameters...
- 2 Data Transfer
- 3 Abs/%T
- 4 Peak Detection...
- 6 Graph Scale...
- 7 Sample Number...
- 8 Save Method...
- 9 Output Options...

**Menu (select using key pad numbers)**

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options / Printer settings.
- 3) Toggle between Absorbance and %T mode.
- 4) Displays Peak Detection Parameter Screen. See description below.
- 5) Manually adds a peak position to the peak table in the results screen at the position set by the cursor. If the cursor is returned to this position the legend "User Defined Peak" is displayed at the top of the scan and this option changes to Delete Peak...
- 6) Displays Graph Scale Parameter Screen. See description below.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (Favourites/Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options/Printer settings, possibility to change the Output settings within the method as described in 7.3 Output Options / Printer

Exit *Menu/Options* by pressing **Escape** , or wait.

**Peak Detection (Shortcut button 4)**

**Peak Detection**

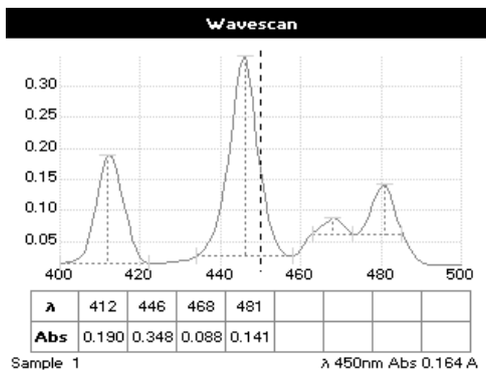
<p>Auto-detect Peaks</p> <p><input type="checkbox"/> Yes</p>	<p>Peak Detect on Zoom</p> <p><input type="checkbox"/> Yes</p>
<p>Min Pk Height</p> <p>0.01</p>	<p>Sort Peaks By...</p> <p>Wavelength</p>
<p>Min Pk Width</p> <p>5 nm</p>	<p>Draw Peaks</p> <p>No</p>
<p><input type="button" value="OK"/></p>	<p><input type="button" value="Cancel"/></p>

**Peak Detection (Shortcut button 4)**

**Auto Detect Peaks:** Turns on and off the automatic peak detection. The following options determine how peaks are detected:

**Minimum Peak Height:** Minimum height the peak has to be above the higher of the two adjacent minima for the peak to be detected.

**Minimum Peak Width:** Minimum width of the peak as determined by the difference in wavelength between the highest of the two adjacent minima and the opposing intersection of that higher minimum level and the peak profile. (See the screen displayed below).



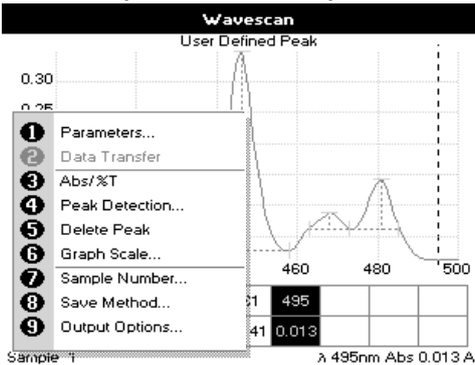
**Peak Detect on Zoom:** Determines whether peaks are re-assessed and tabulated when the user zooms into a region of the wavescan. If **Off**, leaves the peak detection as determined on the un-zoomed display.

**Sort Peaks by...:** Determines the sequence that peaks are reported by. Can be wavelength, peak height or peak width.

**Draw Peaks:** Switches display of peak cursors on and off. These show vertical dashed lines displaying the measured peak height and horizontal dashed lines showing the peak width.

Pressing **Cancel**  ignores the selection, pressing **OK**  accepts them.

### Add Peak (Shortcut button 5)



### Add Peak (Shortcut button 5)

Adds a user defined peak at the current cursor position. The entry is then displayed in inverse colouring to discriminate between user defined peaks and auto-detect peaks. When the cursor is positioned over the user defined peak a legend “User Defined Peak” appears at the top of the graph. The option then changes to Delete Peak to enable the user to remove the peak.

### Graph Scale (Shortcut button 6)

### Graph Scale (Shortcut button 6)

This enables the user to set up a defined graph by defining the limits in either or both of the x and y axes.

#### Zoom mode:

This sets up the operation of the Zoom keys (up and down arrows). The “x & y axis” expands the display around the cursor measurement point, whilst the other options select the absorbance or wavelength axes respectively. With x or y axis limits set to on, zooming out will only be permitted to the set limits.

#### x/y axis limits:

Setting “x (or y) axis limits” to “On” activates the start and finish points of the desired graph to user defined specific wavelengths and/or absorbance values.

Pressing **Cancel** ignores the selection; pressing **OK** accepts it and displays the required graph.

## 5.4 Kinetics

Simple kinetics studies, where the change in absorbance needs to be followed as a function of time at a fixed wavelength, can be readily performed.

Reagent test kits are routinely used for the enzymatic determination of compounds in food, beverage and clinical laboratories by measuring NAD<sup>+</sup> / NADH conversion at 340 nm. The change in absorbance over a specified time period can be used to provide useful information when an appropriate factor, defined in the reagent kit protocol, is applied. Reaction rate and enzyme activity can be calculated if the factor used takes account of the absorbance difference per unit time, as opposed to the absorbance difference *per se*. For this reason, the change in absorbance per minute ( $\Delta A/\text{min}$ ), concentration ( $\Delta A/\text{min} \times \text{factor}$ ) and **correlation coefficient** (calculated from a best fit of the data points) are displayed. They may not be relevant for simple kinetics experiments.

The procedure to define a new method is as follows:

### Parameter Screen

**Kinetics - Parameters 1**

Wavelength	Delay Time
340 nm	0 Seconds
	Duration
	1 Minute
	Interval
	10 Seconds

Next Cancel

**Kinetics - Parameters 2**

Mode	Pathlength
Delta A	10 mm
Units	
Factor	
1.000	

OK Back

**Kinetics - Parameters 2**

**Units**

Units
U/l
DP
Auto

OK Cancel

### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 4 to select **Kinetics**
- Step 3** **Wavelength**: Enter all numerical values using the keypad numbers or the left and right arrows.
- Step 4** **Delay time**: Enter the delay time in seconds before the first measurement is taken. This can be a maximum of 600 seconds (10 minutes).
- Step 5** **Duration**: Enter the time in minutes over which measurements are taken. This can be a maximum of 60 minutes.
- Step 6** **Interval**: Enter the interval time in seconds between measurements using the left and right arrows. Options are: 5, 10, 15, 20, 30 or 60 seconds.
- Step 7** Press **Next** to go to the next parameters screen **OR** Press **Cancel** to return to the **Functions** folder.
- Step 8** Select the measurement **Mode** using the left and right arrows.  
**Delta A**: change in absorbance over the measurement duration (or selected period).  
**Final A**: absorbance at the end of the measurement duration (or selected time).  
**Slope**: rate of change of absorbance over the measurement duration or selected period.
- Step 9** **Units**: The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows ( $\mu\text{g}/\text{ml}$ ,  $\mu\text{g}/\mu\text{l}$ ,  $\text{pmol}/\mu\text{l}$ ,  $\text{mg}/\text{ml}$ ,  $\text{mmol}/\text{l}$ ,  $\mu\text{mol}/\text{l}$ ,  $\text{g}/\text{l}$ ,  $\text{mg}/\text{l}$ ,  $\mu\text{g}/\text{l}$ ,  $\text{U}/\text{l}$ , %, ppm, ppb, conc or none). These units can also be edited once OK is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2 Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters **OR** **Cancel** .
- Step 10** Set the **Factor** by which the result is multiplied to give the amount in the chosen range using the left and right arrows. Range of 0.01 to 9,999.
- Step 11** Press **Next** to enter the Results screen **OR** press **Cancel** to return to the Parameters screen.



## 5.5 Standard Curve

The construction of a multi-point calibration curve from standards of known concentration to quantify unknown samples is a fundamental use of a spectrophotometer; this instrument has the advantage of being able to store this curve as a method, using up to 9 standards.

To include a zero concentration standard, include this in the number of standards to be entered and enter 0.00 for concentration; use a reagent blank when required to enter the zero standard.

The procedure is as follows:

### Parameter Screen

### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 5 to select **Standard Curve**
- Step 3** Select the **Wavelength** using the keypad numbers or left and right arrows.
- Step 4** Enter the number of **Standard** concentration points to be used in the curve (1-9).

- Step 5** **Units:** The user can enter a text string up to 8 characters long. To access a list of pre-defined units press the **Menu/Options** key and then use the left/right arrows (µg/ml, µg/µl, pmol/µl, mg/ml, mmol/l, µmol/l, g/l, mg/l, µg/l, U/l, %, ppm, ppb, conc or none). These units can also be edited once OK is pressed. This screen also allows the number of displayed **Decimal Points (DP)** to be selected, from 0 to 2. Note that the result will always be fixed to 5 significant figures regardless of how many decimal points are selected (so 98768.2 will display as 98768 even with 1 decimal point selected). Press **OK** to store the chosen parameters or **Cancel** .

- Step 6** Select the type of **Curve Fit** using the left and right arrows. Options: straight line regression, a zero regression (this forces the straight line through the origin), interpolated or cubic spline.
- Step 7** Select the **Calibration** mode: either **Standards** (measure prepared standards) or **Manual** (keypad data entry) or **New Standards** (previous values are blanked, new standard can be measured).
- Step 8** (if standards selected) Select the number of standards to be measured and averaged at each standard concentration point. Can be Off (1), 2 or 3.
- Step 9** Press **Next** to enter the Standards screen **OR** Press **Cancel** to cancel selections and return to the **Functions** folder.

## Standard Screen

Standard Curve - Standards

Std. 1 10.0	Std. 4 40.0
Std. 2 20.0	Std. 5 50.0
Std. 3 30.0	Std. 6 60.0

Next Back

## Calibration Screen (replicates off)

Standard Curve - Calibration

Standards	
10.0	
20.0	
30.0	
40.0	
50.0	
60.0	

Back

Standard Curve - Calibration

Standards	
10.0	0.015 A
20.0	0.049 A
30.0	0.082 A
40.0	0.115 A
50.0	0.147 A
60.0	0.179 A

OK Back

## Calibration Screen (replicates on)

Standard Curve - Calibration

Replicates	
#1	0.082 A
#2	0.082 A
Mean	
30.0	0.082 A

Next Back

## Standards screen

- Step 10** Enter the concentration values by using the keypad numbers and the up and down arrows to move between the different standard boxes. Range 0.001 to 9,999.
- Step 11** Press **Next** to enter the Calibration screen. If any duplicate or non-monotonic (increasing entries) are present the unit will beep and highlight the incorrect entry. **OR** press **Back** to return to the Parameter screen.

## Calibration Screen (replicates off)

- Step 12** This shows the calibration values and allows standards to be measured.
- Step 13** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 14** Insert the standard (use C to clear previously stored results before measuring) and press **Sample** to measure the standard and store the result.

- Step 15** Repeat for all standards. A graph will display the results and the fitted curve as the measurements are input.
- Step 16** Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.

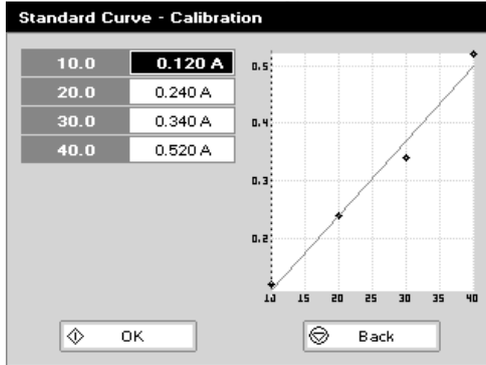
- Step 17** Press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

## Calibration Screen (replicates on)

This shows the calibration values and allows standards to be measured.

- Step 18** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 19** Press **Replicates** to display the replicate entry boxes. Use C to clear previously stored results before measuring.
- Step 20** Insert the standard and press **Sample** to measure the standard and store the result.
- Step 21** Repeat for all replicates and standards. A graph will display the results and the fitted curve as the measurements are input.
- Step 22** Use the up and down arrows to select a standard to be repeated if a poor reading has been obtained. Use C to clear the previous reading.
- Step 23** Press **Next** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

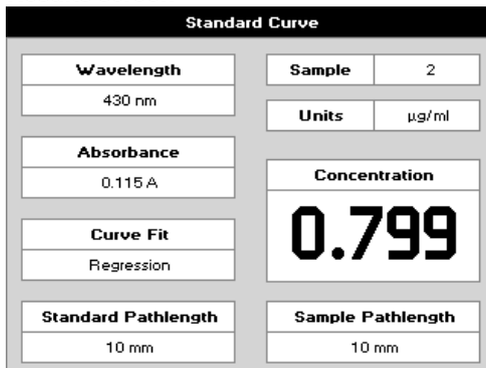
### Calibration (Manual entry)



### Calibration (Manual entry)

- Step 24** Shows previously entered calibration values and allows values to be entered via the keypad.
- Step 25** The highlighted box can be edited in order to enter an absorbance value corresponding to a given concentration value using the keypad numbers. Range 0.001 to 9,999. Use C to backspace and clear the last digit entered and the up and down arrows to move between boxes.
- Step 26** Press **OK** to accept the calibration and go to the Results screen (see below) **OR** press **Back** to return to the Standards screen.

### Results screen



### Results screen

- Step 27** Insert the reference sample and press **Blank** key. This will be used for all subsequent samples until changed.
- Step 28** Insert the sample and press **Sample** . The concentration of the sample is taken and displayed.
- Step 29** Repeat for all samples.
- Step 30** Press **Menu/Options** to display available options which are described below.
- Step 31** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application.

### Menu (select using key pad numbers)

- 1 Parameters...
- 2 Data Transfer
- 4 Edit Sample Pathlength
- 7 Sample Number...
- 8 Save Method...
- 9 Output Options...

### Menu (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options / Printer settings.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options/Printer settings, possibility to change the Output / Printer settings within the method as described in 7.3 **Output Options / Printer**

Exit **Menu/Options** by pressing **Escape** , or wait.

## 5.6 Multiple Wavelength

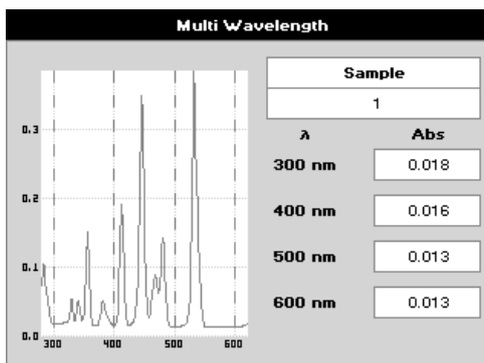
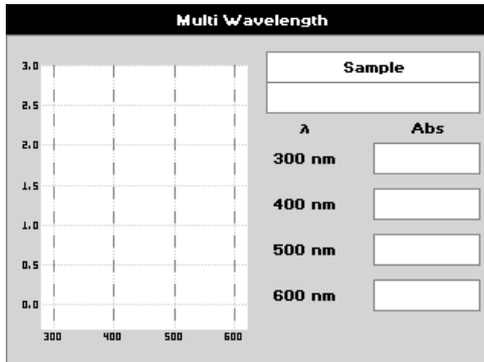
This makes up to 5 absorbance measurements on the same sample.  
The procedure is as follows:

### Parameter Screen

### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 6 to select **Multi Wavelength**
- Step 3** Select the number of **Wavelengths**.
- Step 4** Press **OK** to enter the next screen
- Step 5** Enter the **first Wavelength** using either the number keys or the left and right arrows.
- Step 6** Enter the **second Wavelength** as above and repeat for the number of wavelengths selected (up to 5).
- Step 7** Press **OK** to enter the results screen **OR** press **Cancel** to return to the **Functions** folder.

### Results Screen



### Results Screen

- Step 8** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.
- Step 9** Insert sample and press **Sample** .
- Step 10** Repeat for all samples. A scan plot covering the range of wavelengths selected (with cursors at the relevant wavelengths) and a table of values is displayed.
- Step 11** Press **Menu/Options** to display available options which are described below.
- Step 12** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application.

### NanoPhotometer® P 300

#### Options (select using key pad numbers)

1	Parameters...
2	Print
3	Graph
4	Edit Sample Pathlength
7	Sample Number...
8	Save Method...
9	Printer Settings...

#### Options (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Print result via selected method.
- 3) Toggle graph on/off. Displays the calibration graph, cursors give values for last measured sample.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open printer settings, possibility to change the printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Options** by pressing **Escape** , or wait.

### NanoPhotometer® P 330 / P360

#### Menu (select using key pad numbers)

1	Parameters...
2	Data Transfer
4	Edit Sample Pathlength
7	Sample Number...
8	Save Method...
9	Output Options...

#### Menu (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options settings, possibility to change the Output settings within the method as described in **7.3 Output Options / Printer**

Exit **Menu** by pressing **Escape** , or wait.

## 5.7 Absorbance Ratio

This makes simple absorbance ratio measurements on samples, measuring the amount of light that has passed through a sample relative to a blank (this can be air) at two wavelengths.

The procedure is as follows:

### Parameter Screen

### Parameter Screen

- Step 1** Press 3 to select **Functions**.
- Step 2** Press 7 to select **Absorbance Ratio**
- Step 3** Enter the **first Wavelength** by using the keypad numbers or the left and right arrows.
- Step 4** Enter the **second Wavelength** as above.
- Step 5** Select whether a **Background** correction is applied to both wavelengths 1 and 2 using the left and right arrows.
- Step 6** (If background correction is On) Enter the **third Wavelength**, from which the background correction will be obtained.
- Step 7** Press **Next** to enter the next screen **OR** press **Cancel** to return to the **Functions** folder.
- Step 8** (Dilution Factor known) Enter a **Dilution factor** by using the keypad numbers within the range 1.00 – 9,999.
- Step 9** (Calculate Dilution Factor) Press the **Menu/Options** key. Enter the **Volume** of the sample (range 0.01 – 9,999), using the keypad numbers. Enter the volume of **Diluent** (range 0.01-9,999) by using the keypad numbers.
- Step 10** Press **OK** to calculate the dilution factor and return to the Parameters screen **OR** press **Cancel** to cancel selections.
- Step 11** Select units of measurement, using left and right arrows. Options are: µg/ml, ng/µl, µg/µl.
- Step 12** Enter the factor using the keypad numbers (Range 0.001 to 9,999).
- Step 13** Press **OK** to enter the results screen **OR** **Cancel** to return to the **Functions** folder.

### Results Screen

Absorbance Ratio	
260 nm 0.258 A	Sample 1
280 nm 0.166 A	Ratio <b>1.554</b>
	Concentration 12.9 µg/ml

### Results Screen

**Step 14** Insert the reference sample. Press **Blank** key. This will be used for all subsequent samples until changed.

**Step 15** Insert sample and press **Sample**

**Step 16** Repeat for all samples. The absorbance at the selected wavelengths is measured and the ratio between wavelengths 1 and 2 is calculated (both corrected by the background wavelength value if this was selected).

**Step 17** Press **Menu/Options** to display available options which are described below

**Step 18** Press **Escape** and confirm with **Yes** to return to the **Functions** folder. Query needs confirmation to avoid unintended escaping the application.

### Menu (select using key pad numbers)

<b>1</b>	Parameters...
<b>2</b>	Data Transfer
<hr/>	
<b>7</b>	Sample Number...
<b>8</b>	Save Method...
<b>9</b>	Output Options...

### Menu (select using key pad numbers)

- 1) Return to parameters screen.
- 2) Transfer the results via selected Output Options/Printer settings.
- 7) Sample number – add a prefix to the sample number and reset the incrementing number to the desired value.
- 8) Save method – use the left and right arrows to select a folder to store in (User Methods 1-9), press the down arrow and enter name.
- 9) Open Output Options / Printer settings, possibility to change the Output Options / Printer settings within the method as described in **7.3 Output Options / Printer**

Exit **Menu** by pressing **Escape** , or wait.

## 6. USER METHODS

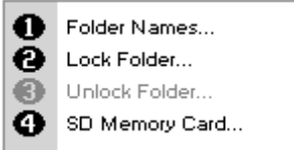
These folders are the storage locations for any user modified Applications (Methods) that are saved in the **Options Menu**. They are accessible from the home folders page. The folder enables the user to quickly select any frequently used Methods. Up to 9 Methods may be stored in the folder.

### User Methods



Folder names can be renamed, locked, unlocked and saved to the SD memory card using the **Options Menu**.

#### Menu / Options (select using key pad numbers)



#### Rename Folder Names

1. Press 1 to select Folder Names.
2. Select the method to be renamed using the left and right arrows.
3. Enter the new name.
4. Press **OK** to save the new name **OR Cancel** to return to the User Methods folder.

#### Lock Method

1. Press 2 to select Lock Folder.
2. Select the method to be locked using the left and right arrows.
3. Select a pass code using the keypad numbers or left and right arrows.
4. Press **OK** to lock the method **OR Cancel** to return to the **User Methods** folder.

#### Unlock Method

1. Press 3 to select Unlock Folder.
2. Select the method to be unlocked using the left and right arrows.
3. Enter the pass code using the keypad numbers or left and right arrows.
4. Press **OK** to unlock the method **OR Cancel** to return to the **User Methods** folder

#### SD Memory Card

Individual or all methods can be copied on the SD Memory Card and can be restored back into the same instrument at a later date. For further details, please refer to the NanoPhotometer® P-Class Accessory manual.

1. Press 4 to select SD Memory Card.
2. Four options are available:
 

Backup folder	generates a copy of an individual folder on the SD Memory Card
Restore folder	restores an individual folder from the SD Memory Card to the instrument
Backup all folders	generates a copy of all folders on the SD Memory Card
Restore all folders	restores all folders from the SD Memory Card to the instrument
3. Select the method to be saved using the left and right arrows.
4. Press **OK** to save the method **OR Cancel** to return to the **User Methods** folder.

**Methods - Methods 1**



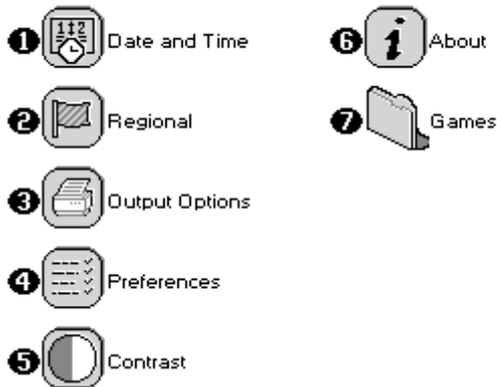
**Delete Method**

1. Select the method to be deleted using the key pad numbers.
2. Select **Menu/Options** and press 1 Delete Method.
3. Press **OK** to delete the method **OR Cancel** to return to **User Methods** folder.

**7. UTILITIES**

Survey of the available Utilities:

**Utilities**



Key pad number	Description
1	Set correct time and date
2	Select preferred number format
3	Printer/output options
4	Select screen layout (themes), history, and Baseline Compensation
5	Adjust screen contrast & brightness
6	Serial number and software version

## 7.1 Date and Time

The procedure is as follows:

- Step 1** Enter the **Day** using the keypad numbers or left and right arrows.
- Step 2** Enter the **Month** as above.
- Step 3** Enter the **Year**.
- Step 4** Enter the **Hour**.
- Step 5** Enter the **Minute**. Seconds are zeroed when OK is pressed.
- Step 6** Press **OK** to store the settings and return to the **Utilities** folder **OR** press **Cancel** to return to the **Utilities** folder without storing the time.

## 7.2 Regional

Sets Number Format.

The procedure is as follows:

- Step 1** Set the **Number Format** (decimal point style). Options are “,” or “.”.
- Step 2** Press **OK** to store the settings and return to the **Utilities** folder **OR** press **Cancel** to return to the **Utilities** folder without storing the settings.

## 7.3 Output Options / Printer

### 7.3.1 NanoPhotometer® P 300

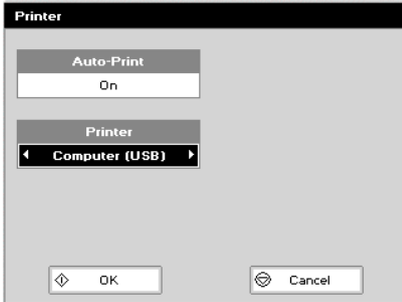
The NanoPhotometer® P 300 offers a selection of data-output accessories. Please proceed to Folder 5 **Utilities**, press 3 **Printer** to set the requested reporting settings:

**Data-output to Built-in printer:**

- Step 1** Select whether **Auto-print** is on or off using the left and right arrows. When auto-print is set as “on” the results are automatically printed after a measurement is taken. When the setting is “off” printing has to be initiated manually. This can also be set using the **Options** key in each application or method. The default is OFF.
- Step 2** Select printer module: **Built-In** Press **OK** to store the settings and return to the **Utilities** folder **OR** Press **Cancel** to return to the **Utilities** folder without storing the settings.

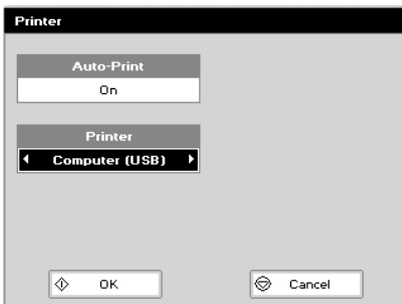
### Direct Data-output to PC:

Connection between NanoPhotometer® and PC is possible with a USB or Bluetooth connection:



- Step 1** Select whether **Auto-print** is on or off using the left and right arrows. When auto-print is set as “on” the results are automatically printed after a measurement is taken. When setting is “off” printing has to be initiated manually. This can also be set using the **Options** key in each application or method. The default is OFF.
- Step 2** Select how the data are sent. Options are **Computer USB**  or **Bluetooth** . Press **OK**  to store the settings and return to the **Utilities** folder **OR** Press **Cancel**  to return to the **Utilities** folder without storing the settings.

### Data-output to SD Memory Card:



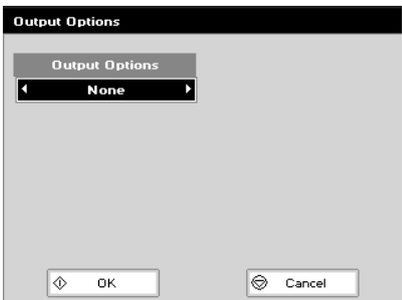
- Step 1** Select whether **Auto-print** is on or off using the left and right arrows. When auto-print is “on” the results are automatically printed after a measurement is taken. When it is set to “off”, printing has to be initiated manually. This can also be set using the **Options** key in each application or method. The default is OFF.
- Step 2** Select printer module: **SD Memory Card** . Press **OK**  to store the settings and return to the **Utilities** folder **OR** Press **Cancel**  to return to the **Utilities** folder without storing the settings.

## 7.3.2 NanoPhotometer® P 330 /P 360

The NanoPhotometer® P-Class offers a selection of data-output accessories. Please proceed to Folder 5 **Utilities**, press 3 **Output Options** to set the favoured reporting settings:

### Data-output to Built-in printer:

Printing on the built-in printer is possible independent from the **Output Options** setting with the **Print** key on the keypad. Pressing the **Print** key prints the results shown on the result screen on the built-in printer. If no built-in printer is connected to the NanoPhotometer® P-Class the **Print** key results in a warning message.



If **None** in the **Output Options** is selected it is only possible to print with the built-in printer or save the data on a connected USB flash drive. No data transfer to a PC is possible. To transfer the data on the result screen to a PC press 9 for **Output Options**, change the **Output Options** to the device of choice. Press **OK**  to store the settings and return to the result screen. Press **Menu** or directly 2 for **Data Transfer** to transfer the data to the PC. All further measurements are transferred and saved automatically.

### Direct Data-output to USB flash drive:

If a USB flash drive is connected to the USB flash drive port (right side of the NanoPhotometer®) all measured data are automatically saved on the USB flash drive, independent from the Output Options settings. Please exit the method first before disconnecting the USB flash drive. Otherwise the data can be lost.

On the right corner of the display is the status of the USB flash drive shown:



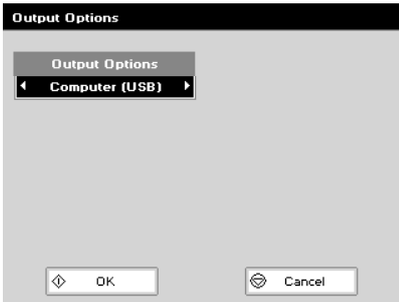
USB flash drive is connected

Data are written to the USB flash drive

Open file on the USB flash drive. Please exiting the method first, before disconnecting the USB flash drive.

**Direct Data-output to PC:**

Connection between NanoPhotometer® P-Class and PC is possible with a USB cable connection:



Select how the data are send. Options are **Computer (USB)** or **None**. Press **OK** to store the settings and return to the **Utilities** folder **OR** press **Cancel** to return to the **Utilities** folder without storing the settings.

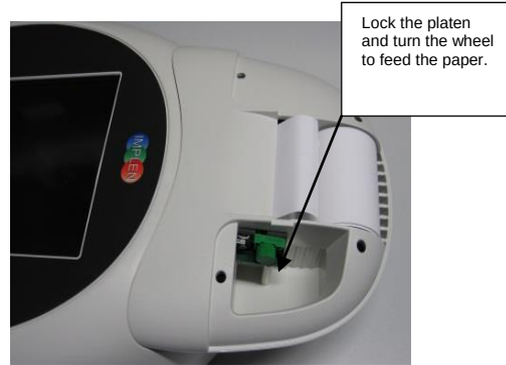
If **Computer (USB)** is selected all measured data are sent automatically to the PC. The file is closed and saved once the method is left by pressing **Cancel** or parameters are changed in the parameter settings.

**7.3.3 Loading / changing the printer paper**

**1. Lift off the cover**

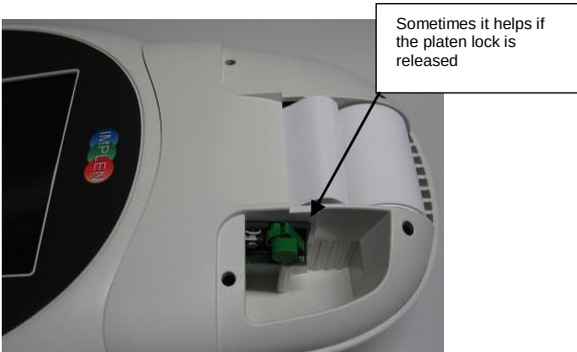


**3. Paper gripped**



**2. Feed in the paper**

Normally the paper is automatically gripped.

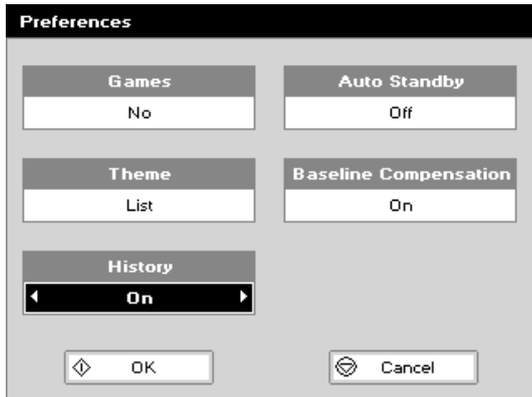


**4. Cover replaced**



## 7.4 Preferences

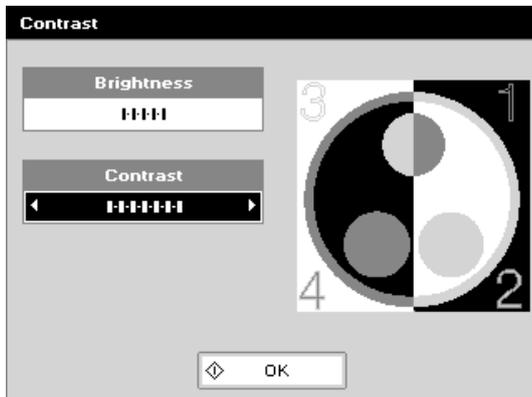
Sets user preferences.  
The procedure is as follows:



- Step 1** Select **Games** function. This determines whether the games folder is displayed or not; Options are yes or no.
- Step 2** Define the **Screen layout** (Theme) of folders. Options are either a grid format or a list.
- Step 3** Select **History** whether to use previously entered parameters (memory function) set On or to return to default settings set Off.
- Step 4** Select whether to use a **Standby** mode after defined periods. Options are 1 hour, 2 hours, at night or off.
- Step 5** Select **Baseline Compensation** to improve value stability and to overcome background effects.
- Step 6** Press **OK** to store the settings and return to the **Utilities** folder **OR** press **Cancel** to return to the **Utilities** folder without storing the settings.

## 7.5 Contrast

Ambient temperature can affect the display. This function can optimize the display for local conditions.  
The procedure is as follows:



- Step 1** Adjust the **Brightness** using the left and right arrows.
- Step 2** Adjust the **Contrast** using the left and right arrows.
- Step 3** Press **OK** to store the settings and return to the **Utilities** folder.

## 7.6 About



Displays the instrument serial number and software version.

Press **OK** to close the window and return to the **Utilities** folder or wait.

## 8. MAINTENANCE

### 8.1 Maintenance-free Technology

The NanoPhotometer® P-Class technology is maintenance-free. Regular maintenance or calibration is not necessary.

For facilities that are working according to national as well as international guidelines and standards – such as Good Laboratory Practice (GLP), Good Manufacturing Practice (GMP) or ISO9000-9004 - the proper performance of a spectrophotometer has to be tested and proved on regular individually set intervals. Implen provides certified NanoPhotometer® P-Class secondary standards as an optional accessory. These NanoPhotometer® P-Class Didymiumglassfilters are suitable for the control and documentation of the wavelength accuracy and the photometric accuracy of your system. IQ/OQ documentation is also available (only for P 330 and P 360). Please contact your local Implen office or an authorized Implen partner for further information.

Support agreements that help to fulfil the demands of regulatory guidelines concerning GLP/GMP:

- calibration, certification using filters traceable to international standards (during production and quality control)
- certified engineers and calibrated test equipment
- approved to ISO 9001 standard
- automatic self-diagnostic calibration test during start of the NanoPhotometer® P-Class, result is documented in each data output file
- possibility to save a PVC file (no data manipulation possible, hardcopy printout)

### 8.2 Lamp Replacement

The xenon lamp should not need replacement until after several years of use. In the unlikely event that it does need replacing, this should be undertaken by a service engineer from your supplier.

### 8.3 Mixer replacement

The mixer should not need replacement until after several years of use. In the unlikely event that it does need replacing, this should be undertaken by a service engineer from your supplier.



**Caution**

Do not open the base plate of the vortexer.  
Do not break the seal.

### 8.4 Exchange of the gaiter

If the gaiter is damaged please exchange it immediately. The gaiter can be ordered separately, item number is S-330360-G.



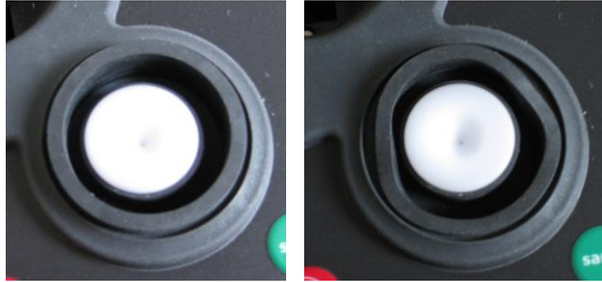
**Caution**

Switch the instrument off before removing the gaiter  
Do not look into the gap between the mixer and the housing, UV exposure

1. Remove the defect gaiter:



2. Place the new gaiter in the gap and in a first step place inner flute and in the second step the outer flute.



## 8.5 Cleaning and general care of the instrument

### External cleaning

Switch off the instrument and disconnect the power cord.

Use a soft damp cloth.

Clean all external surfaces.

A mild liquid detergent may be used to remove stubborn marks.

**NanoPhotometer® P-Class *design edition*** (glossy anthracite):

All painted surfaces of the NanoPhotometer® P-Class can be cleaned with a soft damp cloth and approved cleaning or disinfectant solutions.



### Caution

Product damage by wrong cleaning or disinfection  
 Desinfection or cleaning only by wiping, no spraying.  
 Use no solvents or aggressive chemicals.

Approved disinfectant solutions:

Apesin disinfection spray (Tana Chemie GmbH)

Incidin Liquid and Incidin Foam (Ecolab)

Lysoformin Spezial (Lysoform Dr. Hans Rosemann GmbH)

Observe all necessary precautions if dealing with hazardous samples or solvents.

### Changing cell holder or removal for cleaning

This can be removed by undoing the appropriate screws on the bottom of the instrument.

The symbol  on the product, or on the documents accompanying the product, indicates that this appliance may not be treated as household waste. Instead it shall be handed over to the applicable collection point for the recycling of electrical and electronic equipment. Disposal must be carried out in accordance with local environmental regulations for waste disposal.

## Error messages and Trouble shooting

## 8.6 Error messages

Error text in display	Explanation	Solution
If this icon is  displayed...exit the application before removing the USB memory stick.	<ul style="list-style-type: none"> <li>• USB flash drive was removed before leaving the method</li> <li>• The saved file may be damaged, because it was not closed correctly</li> </ul>	<ul style="list-style-type: none"> <li>• To avoid data loss please exit the method first before disconnecting the USB flash drive</li> </ul>
Calibration failure (UV on Reference channel)	<ul style="list-style-type: none"> <li>• Submicroliter cell or cuvette in the cell holder by switching on the instrument.</li> <li>• Instrument was too cold.</li> </ul>	<ul style="list-style-type: none"> <li>• Remove cuvette or submicroliter cell from the cell holder, turn off the instrument and start again.</li> <li>• Did you start the NanoPhotometer® P-Class directly after delivery? If yes, turn it of and wait 30 min before switching the unit on.</li> <li>• If both suggestions doesn't help disconnect the power completely for at least 10 seconds and then reconnect and restart the system again.</li> </ul>
Cell holder obstructed	<ul style="list-style-type: none"> <li>• Submicroliter cell or cuvette in the cell holder by switching on the instrument.</li> </ul>	<ul style="list-style-type: none"> <li>• Remove cuvette or submicroliter cell from the cell holder, turn off the instrument and start again.</li> </ul>
No light on Reference channel	<ul style="list-style-type: none"> <li>• Submicroliter cell or cuvette in the cell holder by switching on the instrument.</li> <li>• Cell holder has been removed from the instrument and placed back in the wrong position.</li> </ul>	<ul style="list-style-type: none"> <li>• Remove cuvette or submicroliter cell from the cell holder, turn off the instrument and start again.</li> <li>• Remove the cell holder and place back in the right position.</li> <li>• If both suggestions doesn't help disconnect the power completely for at least 10 seconds and then reconnect and restart the system again.</li> </ul>
Calibration problem (UV/IR) on Reference channel	<ul style="list-style-type: none"> <li>• Low light levels.</li> </ul>	<ul style="list-style-type: none"> <li>• Disconnect the power for quite some time to make sure that the problem is not occurring due to low energy sent to the Xenon flash lamp.</li> </ul>
Waiting for software update (Keyboard)	<ul style="list-style-type: none"> <li>• The instrument has been turned on and off too fast.</li> </ul>	<ul style="list-style-type: none"> <li>• Disconnect the power completely for at least 10 seconds and then reconnect and restart the system again.</li> </ul>
Calibration problem: possible lamp failure	<ul style="list-style-type: none"> <li>• Insufficient provision of electricity.</li> <li>• Power supply does not deliver 18V.</li> </ul>	<ul style="list-style-type: none"> <li>• Check all connections if the cables etc. are sticking right.</li> <li>• Check correct power supply 18 V is attached.</li> <li>• If possible try another power supply.</li> </ul>

Please contact the Implen Support Team (support@implen.de, Phone +49-89-7263718 20) if none of the mentioned solution helps to solve the problem or if another error message should appear on the NanoPhotometer® P-Class display.

**8.7 Trouble shooting**

Symptom	Solution
Instrument failing start up calibration.	Check that the correct power supply 18 V, 1.2 A is being used and ensure that the connector is pushed in fully.
Instrument switching off after calibration.	User may be keeping their finger on the ON/OFF button too long, so that the instrument receives both signals and switches off after the calibration
Instrument intermittently switches off during measurement.	Faulty or loose power input connection. Check voltage output of power supply.
Thermal paper printer displays random or endless numbers after paper exchange.	Paper roll was maybe changed when the instrument is still in one of the programs. User needs to move to the main menu before exchanging the paper roll. Switch the device Off and disconnect the power supply, reconnect it after about 5 minutes, Firmware update (version 2.4.2) recommended.
Thermal paper printer prints a plenty of paper, normal sound, no error message.	Thermal Printer paper was probably put the wrong way, check correct position of the printer paper.
Thermal printer automatic roller stops.	Please contact the Implen support team!

Please contact the Implen Support Team (support@implen.de, Phone +49-89-7263718 20) if none of the mentioned solution helps to solve the problem or if another symptom should occur.

**9. ACCESSORIES**

Certified Didymiumglassfilter for the control of wavelength and photometric accuracy for the NanoPhotometer®	P-3034-F
Thermal Paper Printer module for the NanoPhotometer®	P-31-P
Replacement Paper Rolls for Thermal Paper Printer (pack of 5)	P-31-Pa
Bluetooth connection module (for P300 only)	P-33-B
SD Memory Card module (for P300 only)	P-34-SD
Dust cover for the NanoPhotometer®	P-3034-D
Lid 5, 2 mm lid for NanoPhotometer® Submicroliter Cell	P-3034-L2
Lid 100, 1 mm lid for NanoPhotometer® Submicroliter Cell	P-3034-L1
Lid 50, 0.2 mm lid for NanoPhotometer® Submicroliter Cell	P-3034-L02
Lid 100, 0.1 mm lid for NanoPhotometer® Submicroliter Cell	P-3034-L01
Lid 250, 0.04 mm lid for NanoPhotometer® Submicroliter Cell	P-3034-L004
Gaiter for the NanoPhotometer® P330 / P360	S-330360-G

## 10. SPECIFICATION AND WARRANTY

### Technical Specifications:

#### Spectrophotometer

Wavelength range	190 – 1,100 nm
Wavelength scan range	200 – 950 nm
System start up time	Less than 5 seconds, no warm up necessary
Measure time for full scan range	3.5 seconds
Wavelength reproducibility	< ± 0.2 nm
Wavelength accuracy	± 2 nm
Bandwidth	Better than 5 nm
Stray light	< 0.5% at 220 nm using NaI and 340 nm using NaNO <sub>2</sub>
Photometric range	-0.3 – 2.499 A 0-199% T
Detection Range	dsDNA: 0.5 ng/μl to 125 ng/μl, BSA: 0.02 mg/ml to 3.6 mg/ml
Absorbance reproducibility	±0.003 A (0 to 0.5 A), ±0.007 A (0.5-1.0 A) @ 260 nm
Absorbance accuracy	±0.005 A or ±1% of the reading, whichever is the greater
Zero stability	±0.003 A/hour after 20 min warm up @ 340 nm
Noise	0.002 A rms at 0 A @ 260 nm, 0.005 A (pk to pk) at 0 A @ 260 nm
Optical arrangement	Dual channel Czerny Turner with flat grating, 1024 pixel CCD array, concave mirrors
Lamp	Xenon flash lamp
Lifetime	10 <sup>9</sup> flashes, up to 10 years
Cell types	15 mm centre height (z-height), outside dimension 12.5 mm x 12.5 mm

#### NanoVolume application

Detection range	dsDNA: 2 ng/μl to 18,750 ng/μl, BSA: 0.08 mg/ml to 543 mg/ml
Absorbance range	0.01 – 1.5 (10 mm equivalent: 0.05 – 375)
Minimum sample size	0.3 μl
Path lengths	0.04 mm, 0.1 mm, 0.2 mm, 1 mm and 2 mm
Virtual dilution factors	5, 10, 50, 100 and 250

#### Other technical data

Vortexer	2,800 rpm; tube size up to 2.0 ml
Cuvette storage	capacity for eight 10 mm cells
Photometric mode	Abs, %T, concentration, scan, ratio, multi wavelength, kinetics in Abs x factor / min
Method storage	Up to 81 methods in user methods
Built-in methods	Nucleic acid, microarray (labelling efficiency), protein and cell density
Display formats	320 x 240 pixels
Size	140 mm x 275 mm x 380 mm
Weight	< 4.5 kg
Operating voltage	90-250 V, 50/60 Hz, Max 30 VA
Output ports	Availability of USB flash drive port, USB cable, SD Memory Card or Bluetooth for easy PC connection and data export on selected models; built-in printer
Input Port	USB cable for firmware updates
Performance verification	Auto diagnostics when switched on

Specifications are measured after the instrument has warmed up at a constant ambient temperature and are typical of a production unit. As part of our policy of continuous development, we reserve the right to alter specifications without notice.

#### Warranty

- IMPLEN guarantees that the product supplied has been thoroughly tested to ensure that it meets its published specification. The warranty included in the conditions of supply is valid for 12 month for the NanoPhotometer® P 300 / P 330 and 24 months for the NanoPhotometer® P 360 only if the product has been used according to the instructions supplied. Implen or your supplier can accept no liability for loss or damage, however caused, arising from the faulty or incorrect use of this product.

## 11. SOLVENT COMPATIBILITY

Most solvents typically used in life science laboratories are compatible with the NanoPhotometer® P-Class submicroliter cell surface and the lid surfaces.

The following solvents are compatible for use on the submicroliter cell at room temperature:

- acetone (up to 5%)
- acetonitrile
- benzene
- butanol
- carbon tetrachloride
- chloroform
- ethanol
- ether
- HEPES
- hexane
- isopropanol
- MES
- methanol
- methylenchlorid
- MOPS
- phenol (up to 1%)
- n-propanol
- toluene
- buffers containing phosphate or chloride salts  
e.g. PBS-buffer pH 4-10 or citrate or borate
- low concentrated acids and bases



Please note that all kind of high concentrated acids and bases are not recommended. It is recommended to wipe off every sample immediately upon completion of a measurement. For more specific solvents, please contact the Implen support team ([support@implen.de](mailto:support@implen.de)) to check the compatibility.

## 12. APPENDIX

### 12.1 Nucleic acid quantification

For determination nucleic acid concentration in solution the absorbance at wavelength 260 nm is used. The function describing the concentration to absorbance relation is a modification of the Lambert-Beer equation.

$$c_{\text{nuc}} = \text{Abs. 260} * \text{factor}_{\text{nuc}} * \text{lid factor} * \text{dilution factor}$$

With background correction:

$$c_{\text{nuc}} = (\text{Abs. 260} - \text{Abs. 320}) * \text{factor}_{\text{nuc}} * \text{lid factor} * \text{dilution factor}$$

$c_{\text{nuc}}$	nucleic acid concentration (ng/μl)
Abs. 260	absorbance (AU) of nucleic acids
$\text{factor}_{\text{nuc}}$	substance specific factor for nucleic acids (ng * cm/μl) (ds DNA 50, ssDNA 37, RNA 40, Oligo 33)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.

### 12.2 Nucleic acid fluorescent dye incorporation

To determine the nucleic acid concentration and the dye concentration after probe labelling a modification of the Lambert-Beer equation is used. Background correction is always calculated; possibility to switch the dye correction on and off.

- **Calculation of the fluorescence nucleic acid concentration**

$$c_{\text{nuc}} = (\text{Abs. 260} - \text{Abs. 320}) * \text{factor}_{\text{nuc}} * \text{lid factor} * \text{dilution factor}$$

With dye correction:

$$c_{\text{nuc}} = [(\text{Abs. 260} - \text{Abs. 320}) - (\text{CF}_{\text{dye}} * (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}))] * \text{factor}_{\text{nuc}} * \text{lid factor} * \text{dilution factor}$$

$c_{\text{nuc}}$	nucleic acid concentration (ng/μl)
Abs. 260	absorbance (AU) of nucleic acids
$\text{CF}_{\text{dye}}$	dye-dependent correction factor at 260 nm
$\text{Abs.}_{\text{max, dye}}$	absorbance at absorption maximum of the dye (AU)
$\text{factor}_{\text{nuc}}$	substance specific factor for nucleic acids (ng * cm/μl) (ds DNA 50, ssDNA 37, RNA 40, Oligo 33)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.

- **Calculation of the dye concentration**

$$c_{\text{dye}} = ((\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}) * \text{lid factor} * \text{dilution factor}) / (\epsilon_{\text{dye}} * 10^{-6})$$

$c_{\text{dye}}$	dye concentration (pmol/μl)
$\text{Abs.}_{\text{max, dye}}$	absorbance at absorption maximum of the dye (AU)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.
$\epsilon_{\text{dye}}$	dye-dependent molar extinction coefficient ( $\text{M}^{-1} * \text{cm}^{-1}$ )

- **Calculation of the frequency of incorporation (FOI) of dye per 1,000 bases:**

**Formula for dsDNA:**

$$\text{FOI} = (6.49 * (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}) / [\epsilon_{\text{dye}} * 10^{-6} * (\text{Abs. 260} - \text{Abs. 320})]$$

With dye correction:

$$\text{FOI} = (6.49 * (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}) / [\epsilon_{\text{dye}} * 10^{-6} * (\text{Abs. 260} - \text{Abs. 320} - \text{CF}_{\text{dye}} * (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}))]$$

**Formula for ssDNA:**

$$FOI = (8.77 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320)]$$

With dye correction:

$$FOI = (8.77 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320 - CF_{dye} * (Abs._{max, dye} - Abs. 320))]$$

**Formula for RNA:**

$$FOI = (8.11 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320)]$$

With dye correction:

$$FOI = (8.11 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320 - CF_{dye} * (Abs._{max, dye} - Abs. 320))]$$

**Formula for Oligonucleotides:**

$$FOI = (9.83 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320)]$$

With dye correction:

$$FOI = (9.83 * (Abs._{max, dye} - Abs. 320)) / [\epsilon_{dye} * 10^{-6} * (Abs. 260 - Abs. 320 - CF_{dye} * (Abs._{max, dye} - Abs. 320))]$$

Abs. <sub>max, dye</sub>  
 $\epsilon_{dye}$

absorbance at absorption maximum of the dye (AU)  
 dye-dependent extinction coefficient ( $M^{-1} * cm^{-1}$ )

The following dye types and parameters are pre-programmed in the NanoPhotometer® P-Class.

Dye Type	Absorption maximum Dye (nm)	Dye-dependent extinction coefficient $\epsilon_{dye}$	Dye-dependent correction factor (260 nm) $CF_{Dye}$
Alexa Fluor 350	345	18,400	0.25
Alexa Fluor 488	492	62,000	0.30
Alexa Fluor 532	525	82,300	0.24
Alexa Fluor 546	555	104,000	0.21
Alexa Fluor 555	555	150,000	0.04
Alexa Fluor 568	576	93,000	0.45
Alexa Fluor 594	588	80,400	0.43
Alexa Fluor 647	650	239,000	0.00
Alexa Fluor 660	660	107,000	0.00
Alexa Fluor 680	680	164,000	0.00
Cy3	550	150,000	0.08
Cy3.5	581	150,000	0.08
Cy5	649	250,000	0.05
Cy5.5	675	250,000	0.05
Oyster-500	503	78,000	0.29
Oyster-550	553	150,000	0.05
Oyster-556	560	155,000	0.03
Oyster-645	649	220,000	0.05
Oyster-650	653	200,000	0.04
Oyster-656	660	200,000	0.04
Texas Red	593	85,000	0.23

\*In all formulas the molar dye-dependent extinction coefficient is used.

### 12.3 Protein quantification

For determination of protein concentration in solution the absorbance at wavelength 280 nm is used. The function describing the concentration to absorbance relation is a modification of the Lambert-Beer equation.

$$c_{\text{prot.}} = \text{Abs. 280} * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$

With background correction:

$$c_{\text{prot.}} = (\text{Abs. 280} - \text{Abs. 320}) * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$

$c_{\text{prot}}$	protein concentration (mg/ml)
Abs. 280	absorbance (AU) of proteins
A280 factor	Default setting is BSA; molecular weight <sub>prot.</sub> / molar extinction coefficient ( $M^{-1} * cm^{-1}$ ) <sub>prot.</sub> or 1 / extinction coefficient (l/g*cm)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.

A280 factors pre-programmed:

	BSA	Serum albumin (mouse)	Serum albumin (human)	IgG	Lysozyme
<b>A280 factor</b>	<b>1.45</b>	<b>1.57</b>	<b>1.76</b>	<b>0.71</b>	<b>0.38</b>
Molecular Weight [g/mol]	69,323.4	68,692.5	69,365.7	150,000	14,300
Molar extinction coefficient [ $M^{-1} * cm^{-1}$ ]	47,790	43,780	39,310	210,000	37,500

### 12.4 Protein fluorescent dye incorporation

To determine the protein concentration and the dye concentration after labelling a modification of the Lambert-Beer equation is used. Background correction is always calculated; possibility to switch the dye correction on and off.

- **Calculation of labelled protein concentration**

$$c_{\text{prot}} = (\text{Abs. 280} - \text{Abs. 320}) * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$

With dye correction:

$$c_{\text{prot}} = [\text{Abs. 280} - \text{Abs. 320} - \text{CF}_{\text{dye}} * (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320})] * \text{A280 factor} * \text{lid factor} * \text{dilution factor}$$

$c_{\text{prot}}$	protein concentration (mg/ml)
Abs. 280	absorbance (AU) of proteins
$\text{CF}_{\text{dye}}$	dye-dependent correction factor at 280 nm (to be delivered from dye-supplier)
$\text{Abs.}_{\text{max, dye}}$	absorbance at absorption maximum of the dye (AU)
A280 factor	molecular weight <sub>prot.</sub> / molar extinction coefficient ( $M^{-1} * cm^{-1}$ ) <sub>prot.</sub> or 1 / extinction coefficient (l/g*cm)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.

- **Calculation of fluorescence dye concentration [pmol/μl]**

$$c_{\text{dye}} = (\text{Abs.}_{\text{max, dye}} - \text{Abs. 320}) * \text{lid factor} * \text{dilution factor} / (\epsilon_{\text{dye}} * 10^{-6})$$

$c_{\text{dye}}$	dye concentration (pmol/μl)
$\text{Abs.}_{\text{max, dye}}$	absorbance at absorption maximum of the dye (AU)
lid factor	virtual dilution factor 5, 10, 50, 100 and 250; dependent on the used dilution lid.
$\epsilon_{\text{dye}}$	dye-dependent molar extinction coefficient ( $M^{-1} * cm^{-1}$ )

- **Calculation of degree of labelling (D/P)**

$$\text{degree of labelling} = (\text{Abs.}_{\text{max, dye}} - \text{Abs. } 320) * \epsilon_{\text{prot}} / [(\text{Abs. } 280 - \text{Abs. } 320) * \epsilon_{\text{dye}}]$$

With dye correction:

$$\text{degree of labelling} = (\text{Abs.}_{\text{max, dye}} - \text{Abs. } 320) * \epsilon_{\text{prot}} / [(\text{Abs. } 280 - \text{Abs. } 320 - ((\text{Abs.}_{\text{max, dye}} - \text{Abs. } 320) * \text{CF}_{\text{dye}})) * \epsilon_{\text{dye}}]$$

<b>Abs. max, dye</b>	absorbance at absorption maximum of the dye (AU)
<b><math>\epsilon_{\text{prot}}</math></b>	protein-dependent molar extinction coefficient ( $M^{-1} * cm^{-1}$ )
<b><math>CF_{\text{dye}}</math></b>	dye-dependent correction factor at 280 nm (to be delivered from dye-supplier)
<b><math>\epsilon_{\text{dye}}</math></b>	dye-dependent molar extinction coefficient ( $M^{-1} * cm^{-1}$ )

The following dye types and parameters are pre-programmed in the NanoPhotometer® P-Class.

Dye Type	Absorption maximum Dyes (nm)	Dye-dependent extinction coefficient $\epsilon_{\text{dye}}$	Dye-dependent correction factor (280 nm) $CF_{\text{Dye}}$
Alexa Fluor 350	346	19,000	0.19
Alexa Fluor 405	401	34,500	0.70
Alexa Fluor 488	495	71,000	0.11
Alexa Fluor 647	650	239,000	0.03
Cy3	550	150,000	0.05
Cy5	649	250,000	0.05
DyLight 649	654	250,000	0.04
DyLight 488	493	70,000	0.15
FITC	495	68,000	0.30
Pacific Blue	416	46,000	0.20
r-PE	566	200,000	0.18
Texas Red	595	80,000	0.18

\*In all formulas the molar dye-dependent and the molar protein-dependent extinction coefficient is used.