# Fluorimeters Models 6270, 6280 & 6285 Operating Manual

#### **Safety**

This is important information; please read carefully before installing or using this instrument.

- 1. The 62 Series Fluorimeters are designed for operation by **trained personnel** that are aware of the principles and applications involved. For further help and advice please contact your local distributor
- The fluorimeter is a sensitive electronic and optical instrument designed for use in a laboratory environment. Careful adherence to the installation instructions must be observed. If in doubt contact a relevant and competent authority for advice before proceeding.
- 3. In addition to observing the instructions detailed in the Operating Manual and Service Manual for this instrument all installation, operating and service personnel must be aware of, and employ, a safe system of work.
- 4. Voltage levels hazardous to life are present in this instrument, for personal safety only trained engineers aware of the risk and avoidance of electric shock should remove protective covers from the instrument.
- 5. This instrument is designed for minimal maintenance, which must be carried out carefully following the **procedures detailed in this manual**. All safety instructions in these procedures, as well as those defined locally for the **area or environment** where the work is being carried out must be observed.
- 6. Other than for those items defined in the maintenance procedures herein there are no user serviceable items in this instrument. Removal of covers and attempted adjustment or service by unqualified personnel will invalidate any warranty and incur additional charges for repair.
- 7. Reference should always be made to the **Health and Safety Data** for any chemicals or reagents used. All available information, advice and warnings on the handling, storage, use and disposal of such must be carefully observed. When not available this data must be requested from the supplier before proceeding in any way.
- 8. It is important that **good laboratory practice** is observed when handling samples, chemicals, reagents and ancillary equipment in order to carry out measurement and analysis with this instrument. Suitable **safety and personal protective equipment** must be used at all times.
- 9. If it is suspected that safety protection has been impaired in any way, the fluorimeter must be made **inoperative and secured** against any intended operation. The fault condition must be reported to the **appropriate servicing authority**. In all such reports the **model number and serial number** of the fluorimeter must be quoted.

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**Glossary of Terms** 

**EC Declaration of Conformity** 

#### Introduction

#### 1.1 Instrument Description

Three models are available to cover a wide range of applications. The model 6280 covers the most sensitive determinations with emission wavelengths up to 650nm. Where higher emission wavelengths will be utilised the model 6285 with its redenhanced detector is applicable. For less sensitive applications with a broader wavelength range the model 6270 will meet the necessary requirements.

All three models offer intuitive operation with a user interface based on logical menus that can be navigated from the simple keypad. Up to 20 methods can be created without restriction and be saved for future use, they can also be locked against accidental change by password entry, ensuring data integrity.

The permanent time and date tag on every stored reading further enhances Good Laboratory Practice, while calibration reminders and operator identity can also be entered to support conformance and traceability of operation.

With press-to-read operation and Total energy transfer (TET) technology the output of the high-energy xenon lamp is maximised and its expected life extended so that it should never need replacing in normal use.

The high quality optics is complemented by the Intelligent Filter Modules (IFM) that can be identified by the system, this enables error messages to be generated and displayed should the wrong filters be fitted or their positions be incorrect.

All models are powered from an external, universal mains adapter suitable for use from 90Vac to 264Vac, while the 12V dc input enables use of the fluorimeters in vehicles or from suitable battery packs.

Where temperature controlled fluorescence studies are necessary an optional electrically heated sample holder is easily fitted while for continuous flow analysis an external sipper pump and a wide range of flow through cuvettes are also available.

#### 1.2 Good Practice Guidelines

- 1. The selection of the optimum excitation and emission wavelengths (filters) is critical to achieving the best performance from the analysis.
- 2. All fluorimeters should be sited in a clean, dry, dust free environment. When in use ambient temperature and light levels should remain as constant as possible.
- 3. Adherance to Standard Operating Procedures (SOP) and Good Laboratory Practice (GLP) should be maintained, with regular calibration checks and a suitable Quality Control (QC) programme.
- 4. The correct selection of cuvettes is imperative for accurate and reproducible results:
  - a) Ensure all cuvettes used are compatible with fluorimetric measurements where the emission beam is at 90° to the excitation source. Typical examples have 4 clear sides.
  - b) Ensure any native fluorescence from the cuvette material is minimal at the analysis wavelengths.
  - c) Plastic cuvettes should be used once only.
  - d) Glass and quartz cuvettes should be thoroughly cleaned after use. Discard when scratches or marks are evident in their optical surfaces.
  - e) Ensure any cleaning agents used do not fluoresce at the analytical wavelengths and are thoroughly rinsed away before drying.
  - f) Ensure the cuvettes used are compatible with the constituents of both the samples and standards they are to hold. Plastic cuvettes are not compatible with some organic solvents.
  - g) Cuvettes must be handled with care; by the top and non-optical surfaces only. Any finger marks must be removed by using a suitable cleaning process.
  - h) Flow through cuvettes must be selected with additional consideration for the sample type, aspirated volume, pumping system and rinse cycle, as well as the waste handling to be used.
- 5. The high sensitivity of fluorimetric analysis means that all glassware used in the preparation of samples and standards must be totally free from contamination.
- 6. Chemicals and reagents used in sample and standard preparation should be of the highest grade of purity (AR Grade) and all should be checked for excessive background fluorescence at the analytical wavelengths.
- 7. Samples and working standards should not be stored due to the effects of evaporation, as well as chemical and photo-degradation. Only prepare samples and working standards when they are required for analysis.
- 8. Fluorescence is inversely proportional to temperature. Ensure that all samples and standards have equilibrated to ambient temperature before analysis. If in doubt, use a temperature controlled cuvette holder.
- 9. Refrigerated or cold samples will form micro-bubbles on the cuvette wall as they warm up. These are a common cause of drift in readings. Ensure all samples and standards have equilibrated to ambient temperature before analysis.

- 10. Check the linear range for each method and, where necessary, use a multi-point calibration or calibration curve.
- 11. Be aware of the effects of quenching and, where necessary, use sample dilution or extraction methods to eliminate this.
- 12. Monitor the blank during and between batches to identify any increase due to contamination.
- 13. Sources of contamination to be considered include cleaning agents, microorganisms, particles in suspension, stop-cock grease, filter paper residues and plasticisers leached from containers, caps or sealing materials.
- 14. Many fluorescent assays are pH dependent. Ensure the pH of all samples and standards is within specified limits before carrying out the analysis.

#### **Getting Started**

#### 2.1 Unpacking

Remove the universal 12V power supply (with UK, US and EU leads) and the pack of 100 disposable cuvettes from the packaging.

Remove the fluorimeter from the carton by lifting it in the centre between the two support cheeks; do not lift it by the support cheeks.

Place all items on a clean workbench then remove the support cheeks and the polythene bag from the fluorimeter.

Any shortages or damage must be reported to your local distributor or the manufacturer as soon as possible.

Keep all packing materials in case the unit has to be re-shipped at a later date. It is important that when re-packing the instrument it is first sealed in a strong, clean polythene bag to protect it from the dust and particles that are present in all packing materials.

#### 2.2 Installation

#### 2.21 Location

In ideal circumstances the installation environment will be clean, dry and dust free with the instrument protected from extreme variations in ambient lighting and temperature change. For field use it is recommended that the instrument is used in the optional storage case for additional protection.

Where conditions are less than ideal, maintenance and cleaning must be carried out regularly and additional protection offered where possible. The optional dust cover should always be fitted when the unit is not being used or is stored for short periods.

#### 2.22 Supply Voltage

The fluorimeter is powered by a low voltage dc power supply that operates from a 90-264Vac mains supply. The universal power supply is supplied with 3 mains leads for UK, EU and US sockets. The correct lead for your supply should be selected.

#### 2.23 Mains Connections

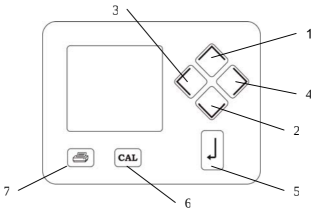
The leads supplied have a moulded on plug. However, if this is removed for any reason the wires in the mains lead are colour coded to conform to the internationally recognised standard such that:

UK CONNECTIONS		<b>US CONNECTIONS</b>	
BROWN	LIVE	BLACK	LIVE
BLUE	NEUTRAL	WHITE	NEUTRAL
GREEN/YELLOW	EARTH	GREEN	EARTH

#### Safety

When disposing of any removed plug the connectors must be removed or made incapable of insertion into a mains socket.

#### 2.24 Keypad Functions



**1. UP ARROW key** - used to navigate through menus, to increase values and for paging up in stored results.

2. DOWN ARROW key - used to navigate through menus, to decrease values

and for paging down in stored results.

3. LEFT ARROW key - used to navigate through menus and to highlight the

selected digit when setting values.

**4. RIGHT ARROW key** - used to navigate through menus and to highlight the

selected digit when setting values.

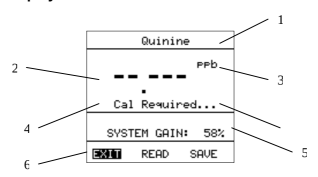
5. ENTER key - used to accept the highlighted menu option.6. CAL key - initiates a calibration sequence from within the

measurement mode.

**7. PRINT key** - initiates a print from the measurement display or stored

results. Sends data to RS232 serial port.

#### 2.25 Instrument Display



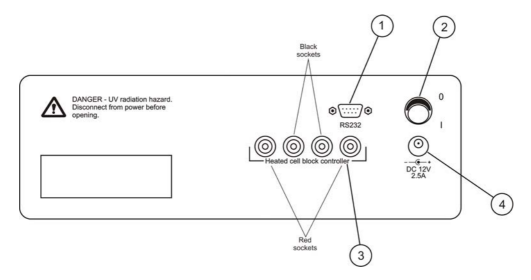
- **1. Method Name** this will appear on all measurement screens, with the exception of Raw Fluorescence.
- 2. Results display provides direct readout of standards and sample results.
- **3. Units of measure** shows selected measurement units: ppm, RFU, U/ml, mU/l, U/l, µM/l, mM/l, M/l, µg/ml, mg/ml, µg/dl, mg/dl, g/dl, ng/l, mg/l, g/l, none, %, ppb
- **4. Status message** shows the current instrument status, such as reading ... or printing ... and provides a reminder when calibration is due.
- **5. Gain** shows the current photomultiplier tube gain setting (0-100%). Gain should be optimised either manually or by using auto set gain for each method.

Note: This option is not available on the Model 6270.

**6. Menu Options** - **EXIT** - allows the user to return to the main menu, **READ** - press read to read the sample or standard, **SAVE** - saves the currently displayed result to instrument memory

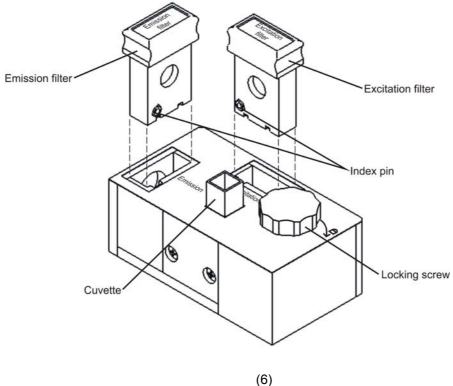
(5)

#### 2.26 **Rear Panel Layout**



- 1. Output Socket – 9 way output socket for RS232
- 2. **Switch** – Power On/Off switch
- 3. **Connection Sockets** – 4 x pin sockets for heated cell block controller.
- 4. Power In Socket – Connection socket for 12V DC universal mains adapter

#### 2.27 **Sample Chamber Filter and Cuvette Positioning**



#### 2.28 Power on and Self-Tests

Connect the mains supply cable to the rear panel mains input socket and plug the other end into a suitable mains supply socket.

Lift the sample chamber lid on the instrument and ensure that there is no sample or other item present in the sample holder, close the lid.

Switch on the supply socket, then the instrument, using the Power switch located on the rear panel.

The instrument will then perform a power on self-test protocol. The following screen will be shown for approx. 3 seconds:

Please ensure sample chamber door remains closed during power on tests

Followed by:

Barloworld
6270
Fluorimeter
Jenway

Barloworld
6280
Fluorimeter
Jenway

Barloworld
6285
Fluorimeter
Jenway

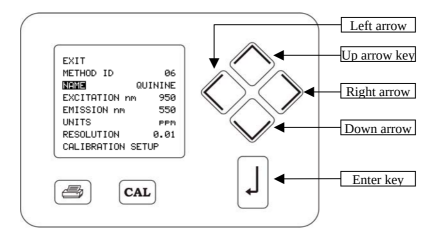
On successful completion of these tests the Main Menu screen will be displayed. For optimum performance a 10-minute warm-up period is required if the ambient temperature is below 10°C. The unit must be re-calibrated and sample measurement repeated if this calibration check shows excessive drift.

Should a problem occur during the self-tests an information box or error support message will be displayed. For assistance please refer to the troubleshooting section in this manual.

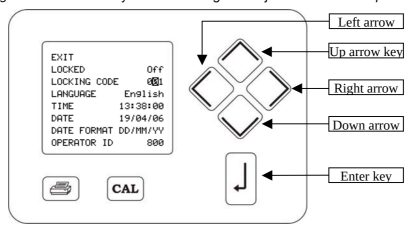
#### **Example Menu Operation**

A common operating system is used throughout this and similar Jenway products, a brief overview of navigating through the menu system with the cursor keys follows;

Use the *up* and *down* arrow keys to highlight a menu option, press the *enter* key to confirm.

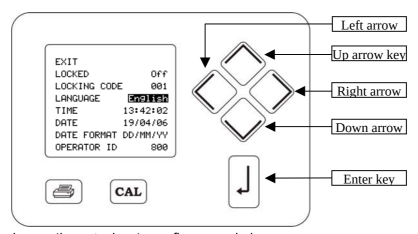


Use the right and left arrow keys to select a digit for adjustment with the up and down arrow



keys, press the enter key to confirm the setting when ALL digits have been correctly set.

Use the up or down arrow keys to browse through pre-set options, when the correct selection



is displayed press the *enter* key to confirm your choice.

#### Fluorescence Theory

The interaction between electromagnetic radiation and matter provides a useful, qualitative and quantitative analytical tool, known as spectroscopy. The region of the electromagnetic spectrum, to which matter under investigation is subjected to, defines the type of transitions that occur within the molecules.

Fluorimetry uses radiation from the UV-Vis region of the electromagnetic spectrum to study transitions between electronic levels in a molecule or atom. The absorption of energy from light radiation (photons) by a molecule or atom, promotes electrons from a low energy ground state to a higher energy excited level. This is known as excitation and the amount of energy transferred to the molecule or atom will depend on two main factors. The composition of the matter under investigation and the energy and wavelength of the radiation, have a significant effect on the transitions of electrons.

The molecule or atom converts the excitation energy to vibrational or light energy and the electron returns to its ground state. Vibrational energy is transferred through movement and collision with other molecules, but energy not lost in this way is released as light radiation. The light emission is known as fluorescence and if some energy has been removed through vibration, it will be of a lower energy and longer wavelength than the excitation energy. The wavelength and intensity of the emitted radiation is dependent on the structure and composition of the molecule and the excitation wavelength used.

Relationship between concentration and fluorescence

The fluorescence signal F and concentration C of the matter under investigation, are related by:

$$F = KQP_0 (1-10^{-6} E_{bC})$$

Where

= A constant characteristic of the instrument (Including instrument electronics, pH and Temperature)

Q = Quantum efficiency (= Photons emitted/Photons absorbed)

 $P_0$  = Power of incident radiation

 $\varepsilon$  = Molar absorptivity of the species (matter)

b = Absorption path length

If the concentration of the matter in question is low (dilute),  $\mathcal{E}bC$  is small. The relationship is then linear and the equation can be written as

 $F = 2.3KQP_0 \&bC$ 

The accuracy of fluorescent measurements is very high because the radiant energy being formed is measured directly. There are also only a few, easily controlled limits on the sensitivity of fluorescence measurements. From the equation above it can be seen that adjustments made to instrument electrical noise and competing radiations, along with physical limitations such as radiation energy, sample volume and cell size affect the measurement sensitivity.

#### **Instrument Menu Options**

In this section a brief explanation is given for the main menu options available. These will be covered in greater depth in the sections appropriate to their usage.

#### **Measure Screen**



METHOD NAME - user set current method name or DEFAULT METHOD

LAST READING - result from last read or --.---

UNITS OF MEASURE - as set in MEASURE SETUP menu

**SYSTEM GAIN%** - 0 to 100% as set up in MEASURE SETUP menu. **Note**: This option is not available on the Model 6270.

**CALIBRATION REMINDER** (flashing) - user alert indicating that a calibration is due **TIMED READING COUNTDOWN** - number of seconds until the next auto read is performed

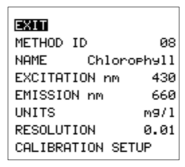
**READ** – allows the user to take a reading of sample/standard currently in the sample chamber

**SAVE** – allows the user to save the last reading to instrument memory

To exit this menu highlight **EXIT** using the right and left arrow keys and press enter to confirm. This will return you to the main menu options.

#### Measure Setup menu

Enter **MEASURE SETUP** menu by selecting **SETUP** from the **MAIN** menu. The following screen will be shown:



**EXIT** – allows user to return to the previous menu **METHOD ID** – used to index individual methods. (0 to 19)

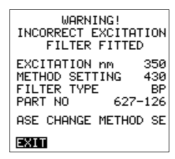
**NAME** – user entered method name (up to a maximum of 19 characters)



**EXCITATION NM** – user set excitation wavelengths. Actual filter data is checked against this setting during measurement. If an incorrect filter is detected a warning will be displayed.

**EMISSION NM** – user set emission wavelengths. Actual filter data is checked against this setting during measurement. If an incorrect filter is detected a warning will be displayed.

**Note:** If the excitation or emission filter modules do not match the current method settings on entry to the measure mode (or prior to taking a reading or calibration) then a warning is displayed:

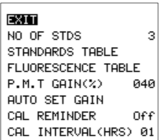


A scrolling message will appear 'please change method setting or fit correct filter module'. No measurements can be performed until this has been corrected. If a filter is missing the instrument will not allow measurement to take place. Pressing the EXIT key will return the screen to the setup menu options.

**UNITS** – allows user to select the units appropriate to the test. ppm, RFU, U/ml, mU/l, U/l,  $\mu$ M/l, mM/l,  $\mu$ g/ml, mg/ml,  $\mu$ g/dl, mg/dl, g/dl, ng/l,  $\mu$ g/l, mg/l, g/l, none, %, ppb

**RESOLUTION** – 0.01, 0.1, 1, 0.001

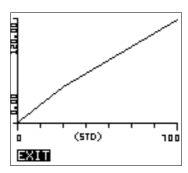
**CALIBRATION SETUP** – this menu is not displayed when RFU (Raw Fluorescence) units are selected.



**EXIT** – allows the user to return to the previous menu **NO OF STDS** – allows selection of 1-6 standards (blank is always included) **STANDARDS TABLE** – user entered calibration standards

**FLUORESCENCE TABLE** – shows results of last calibration RFU (raw fluorescence) values. Tabular and graphical displays available

FLUORE:	SCENCE TABLE
F1 F2 F3	2.385 41.229 118.916
EXIT	VIEW CURVE



**P.M.T GAIN(%)** - 0 to 100% gain setting for system photomultiplier tube. **Note**: This option is not available on the Model 6270.

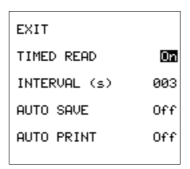
**AUTO SET GAIN** - uses the highest user entered calibration standard to automatically set the PMT gain to give as close as possible to 95% full scale emission signal. **Note**: This option is not available on the Model 6270.

**CAL REMINDER** - enables or disables the measure mode prompt to calibrate at the set calibration interval

**CAL INTERVAL(HRS)** - the number of hours between calibrations can be user set from 1-999 hours.

#### **Results Storage Menu**

From the **MAIN MENU** press the *down* arrow key to move the highlight on the display to cover **RESULTS STORAGE**, then press the *enter* key. The following display will be shown:



**EXIT** - allows the user to return to the previous menu

**TIMED READ** - enables automatic measurements to be taken at a set interval when in measure mod

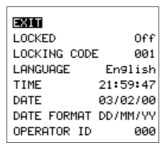
**INTERVAL** - time interval between the end of one reading and the automatic start of the next reading in seconds

**AUTO SAVE** - when enabled, automatically saves to instrument memory after each reading is performed (either manually or timed)

**AUTO PRINT** - when enabled, automatically prints to RS232 connected device after each reading is performed (either manually or timed)

#### **Instrument Setup Menu**

From the **MAIN MENU** press the *down* arrow key to move the highlight on the display to cover **INSTRUMENT SETUP**, then press the *enter* key. The following display will be shown:



**EXIT** - allows the user to return to the previous menu

**LOCKED** - when enabled, prevents access to set up menus without the PIN code (settings of current method to be viewed but not changed). Calibrations can still be performed when the instrument is locked.

**LOCKING CODE** - 3 digit code that must be entered to unlock the instrument **LANGUAGE** - choice of English, French, German, Italian and Spanish

**TIME** - set to time stamp readings and calibrations

**DATE** – set to date stamp readings and calibrations

**DATE FORMAT** - DD/MM/YY or MM/DD/YY (If the date format is changed to MM/DD/YY the changes are reflected straight away). Used to identify saved/printed results

**OPERATOR ID** – 3 digit code can be entered from 000 to 999

#### **Filter Status Menu**

This menu provides a summary of the current method wavelength setting and excitation and emission filters.

From the **MAIN MENU** press the *down* arrow key to move the highlight on the display to cover **FILTER STATUS**, then press the *enter* key. The following display will be shown:

EXCITATION	425
METHOD SETTIN	NG 425
FILTER TYPE	BP
PART NO	627-124
EMISSION	610
METHOD SETTIN	NG 610
FILTER TYPE	COL
PART NO	627-127
EXIT	

**EXCITATION** - wavelength in nm of current excitation filter

METHOD SETTING - wavelength in nm set in MEASURE SETUP menu

**FILTER TYPE** - 3 types of filter are available: **BP** = Bandpass,

**COL** = Cut-Off Longpass and **NPB** = Narrow Bandpass

PART NO - Jenway part number of filter module

**EMISSION** - wavelength in nm of current emission filter

METHOD SETTING - wavelength in nm set in MEASURE SETUP menu

**FILTER TYPE** - 3 types of filter are available: **BP** = Bandpass,

**COL** = Cut-Off Longpass and **NPB** = Narrow Bandpass

PART NO - Jenway part number of filter module

**EXIT** - allows the user to return to the previous menu

#### **Instrument Settings**

**Note:** Using the left arrow key to move from the settings on the right hand side of the menu will abort any adjustments you have made and return the value or selection to its previous settings. The *enter* key must be used to confirm any changes.

All settings listed below are accessed from the **INSTRUMENT SETUP** menu as shown below:

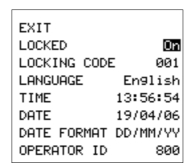
EXIT	
LOCKED	Off
LOCKING CODE	001
LANGUAGE	En9lish
TIME	21:59:47
DATE	03/02/00
DATE FORMAT	DD/MM/YY
OPERATOR ID	999

The 62 Series Fluorimeters offer a number of important functions to control or restrict use of the instrument. This can be useful in developing GLP procedures or for controlling usage in multi-user installations.

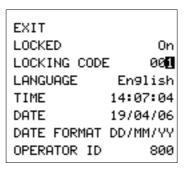
The instrument **LOCKING CODE** restricts access to the set up menus without the entry of a 3 digit security code. When the **LOCKED** function is enabled, settings can only be viewed, not changed. Calibrations can still be performed when the instrument is locked.

#### Instrument Lock

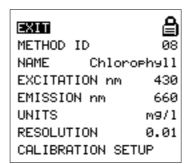
- 1. Use the down arrow key to select **LOCKED** from the menu and press the enter key.
- 2. The instrument lock can be turned on or off using the *up* or *down* keys. Press enter to confirm your selection.
- 3. With the instrument lock set to **Off** all functions of the instrument are available to all users.
- 4. With the instrument lock set to **On** all methods will be locked without the entry of a locking code.



5. To set or change the locking code, press the *down* arrow key to select **LOCKING CODE**, then press the *enter* key.



- 6. Adjust the highlighted digit using the *up* or *down* arrow keys, then select the other digits using the *right* or *left* arrow keys. Adjust each in turn until the code of your choice is displayed. Press the *enter* key to confirm this, ensuring that you can remember the number selected.
- 7. All methods are now locked against adjustment to the set parameters and a padlock icon will be displayed at the top of the method settings to indicate this. It is not possible to move between methods.
- 8. Access to the locking code from the instrument setup menu will now require the entry of the locking code first.
- 9. To exit the **INSTRUMENT SETUP** menu use the *up* or *down* arrow keys to highlight **EXIT**, then press the *enter* key. To set additional parameters continue as detailed in the following text.





#### Language

1. Press the *down* arrow key to move the highlight to **LANGUAGE** and then press the *enter* key.

EXIT	
LOCKED	Off
LOCKING CODE	991
LANGUAGE	En9lish
TIME	14:30:55
DATE	19/04/06
DATE FORMAT	DD/MM/YY
OPERATOR ID	800

Off
001
En9lish
14:27:59
19/04/06
DD/MM/YY
800

2. Use the *up* or *down* arrow keys to view the alternative languages available (English, Spanish, Italian, German or French).

- 3. When the language of your choice is highlighted press the *enter* key to confirm. The language used on the display will now be changed to match your choice.
- 4. To exit the **INSTRUMENT SETUP** menu use the *up* or *down* arrow keys to highlight **EXIT**, then press the *enter* key. To set additional parameters continue as detailed in the following text.

#### **Time**

1. Press the *down* arrow key to move the highlight to **TIME** and press the *enter* key.

EXIT	
LOCKED	Off
LOCKING CODE	001
LANGUAGE	En9lish
TIME	14:41:05
DATE	19/04/06
DATE FORMAT	DD/MM/YY
OPERATOR ID	800

EXIT	
LOCKED	Off
LOCKING CODE	E 001
LANGUAGE	En9lish
TIME	14:4 <b>8</b> :05
DATE	19/04/06
DATE FORMAT	DD/MM/YY
OPERATOR ID	800

- 2. Use the *up* and *down* arrow keys to set the highlighted digit of the time to the correct value. Use the *right* and *left* arrow keys to highlight each digit in turn for adjustment with the *up* and *down* keys until the correct time is set.
- 3. Press the enter key to confirm the settings and start the clock running from the set value.

**Note:** The clock can only be set in the 24-hour system, i.e; 1.00 p.m. = 13.00.00

4. To exit the **INSTRUMENT SETUP** use the *up* or *down* arrow keys to highlight **EXIT**, the press the *enter* key. To set additional parameters continue as detailed in the following text.

#### **Date**

EXIT	
LOCKED	Off
LOCKING CODE	E 001
LANGUAGE	English
TIME	14:47:08
DATE	19/04/06
DATE FORMAT	DD/MM/YY
OPERATOR ID	800

EXIT	
LOCKED	Off
LOCKING CODE	901
LANGUAGE	En9lish
TIME	14:49:01
DATE	19/04/06
DATE FORMAT	DD/MM/YY
OPERATOR ID	800

EXIT	
LOCKED	Off
LOCKING CODE	001
LANGUAGE	En9lish
TIME	14:49:01
DATE	19/04/06
DATE FORMAT	MM/DD/YY
OPERATOR ID	800

- 1. Press the *down* arrow key to highlight **DATE**, then press the *enter* key.
- 2. Use the *up* and *down* arrow keys to set the highlighted digit of the date to the correct value. Use the *right* and *left* arrow keys to highlight each digit in turn for adjustment with the *up* and *down* arrow keys until the correct date is set.
- 3. Press the *enter* key to confirm the setting and start the date running from the set value.

- 4. Use the *down* arrow key to select **DATE FORMAT**. Press the *up* or *down* arrow keys to view the two alternative date formats. Press the *enter* key to confirm your choice. The previously set date will automatically change with your selection in this menu option.
- 5. To exit the **INSTRUMENT SETUP** use the *up* or *down* arrow keys to highlight **EXIT**, then press the *enter* key. To set additional parameters continue as detailed in the following text.

#### **Operator ID**

**OPERATOR ID** is a 3-digit code that can be quickly and easily entered from the **INSTRUMENT SETUP** menu. This ID will appear in the header of all results printed or down loaded from the fluorimeter, identifying the operator that carried out the analysis. When using this facility all potential users should be allocated individual 3 digit codes.

Operators should enter their code using the following procedure:

1. Use the *up* or *down* arrow keys to select **OPERATOR ID** from the **INSTRUMENT SETUP** menu, then press the *enter* key.

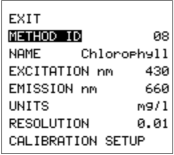
EXIT	
LOCKED	Off
LOCKING CODE	001
LANGUAGE	En9lish
TIME	14:56:57
DATE	04/19/06
DATE FORMAT	MM/DD/YY
OPERATOR ID	800

EXIT	
LOCKED	Off
LOCKING CODE	001
LANGUAGE	En9lish
TIME	14:58:42
DATE	04/19/06
DATE FORMAT	MM/DD/YY
OPERATOR ID	82 <b>7</b>

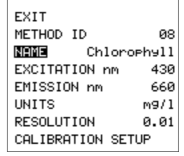
- 2. Use the *right* or *left* arrow keys to highlight each digit in turn. Use the *up* or *down* arrow keys for adjustment until the new operator ID is set.
- 3. Confirm the operator ID by pressing the *enter* key. This current ID will be used in the header of all result printouts and results data transferred to a PC or other serial device.
- 4. To exit the **INSTRUMENT SETUP** menu use the *up* or *down* arrow keys to highlight **EXIT**, then press the *enter* key. This will return you to the main menu.

#### Setting up a method

- 1. Enter **MEASURE SETUP** menu by selecting **SETUP** from the **MAIN** menu.
- 2. Select a suitable **METHOD ID** (0-19).
- Give the method a NAME (up to 19 characters) using the alphanumeric screen accessed by selecting the NAME option of the MEASURE SETUP menu.

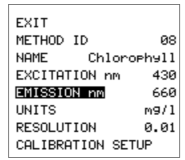






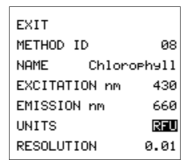
- 4. Return to the **MEASURE SETUP** menu and set the **EXCITATION** wavelength, in nm for the test you wish to perform. The excitation filter module fitted for this test will need to be the same wavelength.
- 5. In the same way set the **EMISSION** filter wavelength, ensuring this is the same wavelength as the emission filter module.
- By scrolling through the available units, select the required UNITS OF MEASURE.

EXIT
METHOD ID 08
NAME Chlorophyll
EXCITATION nm 430
EMISSION nm 660
UNITS m9/1
RESOLUTION 0.01
CALIBRATION SETUP

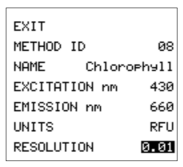


EXIT
METHOD ID 08
NAME Chlorophyll
EXCITATION nm 430
EMISSION nm 660
UNITS **M9/1**RESOLUTION 0.01
CALIBRATION SETUP

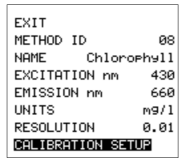
7. If you wish to make raw fluorescence measurements, then select **RFU** from the units of measure. If **RFU** is used, then there is no need to calibrate before taking measurements so the **CALIBRATION SETUP** menu (at the bottom of the **MEASURE SETUP** menu) is removed.

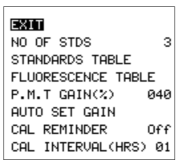


8. The number of decimal places (**RESOLUTION**) to which the measurements will be displayed can then be set at the next option in the **MEASURE SETUP** menu. The fluorimeter will auto range resolution for measurements too large to be displayed in the set resolution.

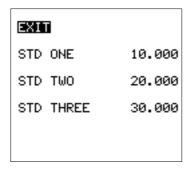


9. To define the calibration parameters, select the last option of the **MEASURE SETUP** menu, **CALIBRATION SETUP**.



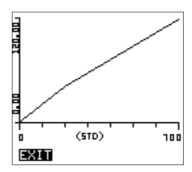


- 10. Once in the **CALIBRATION SETUP** menu, set the **NUMBER OF STANDARDS** that will be used at calibration. The number can be set between 1 and 6, but this does not include the blank (no fluorescence) standard, which is required for all calibrations.
- 11. The concentration values of the standards (not including the blank) can then be set in the **STANDARDS TABLE**.

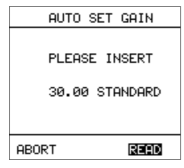


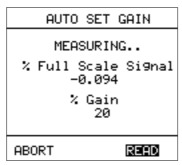
12. The menu option below this is the FLUORESCENCE TABLE. The default, raw fluorescence values will be shown here until a calibration is successfully completed. Once a calibration has been performed, the raw fluorescence data will be displayed here. From the FLUORESCENCE TABLE it is also possible to view the calibration curve, by choosing the VIEW CURVE option. As with the data table, this will show default values until a valid calibration has been completed.

FLUORE	SCENCE TABLE
F1 F2 F3	2.385 41.229 118.916
EXIT	VIEW CURVE



- 13. Set the photomultiplier tube gain (**P.M.T. GAIN** %) to the required percentage, for this test, if known. If not known, use the **AUTO SET GAIN** option as below. **Note**: This option is not available on the Model 6270.
- 14. To find the optimum gain for a given test, select the **AUTO SET GAIN** menu option and follow the instructions on the instrument screen. If you accept the gain suggested by the instrument, this will then automatically update the gain setting in the **CALIBRATION SETUP** menu. **Note**: This option is not available on the Model 6270.





AUTO SET GAIN			
% Full Scale Signal 89.722			
% Gain <b>66</b>			
ABORT REPEAT			

15. To ensure that calibrations are regularly performed the final two **CALIBRATION SETUP** menu options can be employed. If you wish for a calibration reminder to appear in the measure screen, at regular intervals set the **CAL REMINDER** to **On**. The interval between calibration and the reminder being shown is set using **CAL INTERVAL** to the required number of hours using the up and down arrow keys.

EXIT
NO OF STDS 3
STANDARDS TABLE
FLUORESCENCE TABLE
P.M.T GAIN(%) 040
AUTO SET GAIN
CAL REMINDER 07
CAL INTERVAL(HRS) 01

EXIT
NO OF STDS 3
STANDARDS TABLE
FLUORESCENCE TABLE
P.M.T GAIN(%) 040
AUTO SET GAIN
CAL REMINDER OFF
CAL INTERVAL(HRS) 06

16. The method is now fully set and will be the available in the **MEASURE** screen.

NO OF STDS 3
STANDARDS TABLE
FLUORESCENCE TABLE
P.M.T GAIN(%) 066
AUTO SET GAIN
CAL REMINDER On
CAL INTERVAL(HRS) 02

#### Section 7

#### Using a stored method

- 1. From the MAIN menu, select the SETUP menu and then MEASURE SETUP.
- 2. By setting the **METHOD ID** to the appropriate number (0-19) the saved parameters such as the method name and wavelengths will appear and be ready for use.
- 3. **EXIT** the setup menus and return to the **MAIN** menu.
- 4. Ensure that the excitation and emission filters, appropriate to selected method, are fitted in the sample chamber.
- 5. Enter the **MEASURE** mode.
- 6. The name of the selected method will be shown at the top of the screen, with the PMT gain % as set for that method also displayed. **Note**: This option is not available on the Model 6270.
- 7. No concentration will be shown as no measurement has been taken, but the selected units of measure will be displayed.

#### Calibration

- 1. Before beginning the calibration, ensure that close to hand, are the appropriate cuvette/s, blank solution and standard solutions (as set in the **CALIBRATION SETUP** of the **MEASURE SETUP** menu).
- 2. Using the steps as detailed in the **'Using a stored method**' section, select the required method.
- 3. Once in measure mode, press the **CAL** key once to initiate the calibration sequence.
- 4. The instrument display will show the message "Please insert blank..."



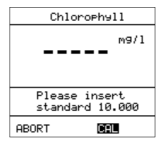
- 5. Taking care not to mark the optical surfaces, place the cuvette containing the blank solution in the cuvette holder of the sample chamber and close the lid.
- Measure the fluorescence of the blank by pressing either the CAL key or with the CAL option highlighted on the instrument screen. Press the enter key to confirm.
- 7. The instrument screen will show the status message "**Reading...**" for the duration of the measurement.



8. Once complete, the concentration (zero, as this is the blank) will be shown as the large number on screen. Below this are the raw fluorescence and the percentages of full-scale emission for the signal and reference.



- 9. The options at the bottom of the screen will have also changed. The ACCEPT and REPEAT options relate to the calibration step just completed. If the concentration and other data from the calibration are correct and satisfactory, select ACCEPT to continue through the calibration sequence to the first standard. If the previous step has given unacceptable figures, then select REPEAT and perform the previous step again. ABORT allows the whole calibration sequence to be cancelled and the instrument to return to the measure screen.
- 10. Once the blank calibration has been accepted, the instrument will display the message "Please insert standard x.xxx" and the concentration of the first standard, as set in the STANDARDS TABLE of the MEASURE/CALIBRATION SETUP.



- 11. Remove the cuvette of Blank solution from the sample chamber, replace with a cuvette of the requested standard and close the lid. If using reusable cuvettes, ensure that proper cleaning procedures are employed to make certain that there is no carry over of solutions or contamination.
- 12. As with the blank, calibrate the instrument with this first standard by using the **CAL** option on screen or **CAL** key.
- 13. As with the blank, the concentration of the standard (as set in the **STANDARDS TABLE**) will be shown on screen with the raw fluorescence and the percentages of full-scale emission for the signal and reference.
- 14. Once again, review this data and if acceptable use the **ACCEPT** option to move on to any further calibration standards in the same way. If not the **REPEAT** function can be used.



- 15. The sequence of calibrating with each standard will continue in this fashion (points 10-13) until each standard that has been set, has been used. At this point the instrument will give a summary of the calibration by displaying the raw fluorescence for each standard in a table. An option to view the calibration curve, a plot of the concentration of the standards against their raw fluorescence, is given at the bottom of the screen (**VIEW CURVE**).
- 16. On exiting the calibration summary the instrument is returned to the measure screen and the status message "Cal required..." will have been removed.
- 17. Measurements of unknown concentration samples can now be made.

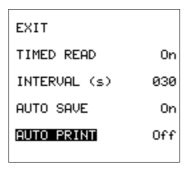
#### **Performing measurements**

- 1. Using the steps as detailed in the 'Using a stored method' to select the required method.
- 2. Prior to taking any measurements make certain that the correct excitation or emission filter modules are fitted and that a valid calibration (as detailed in Section 8) has been completed.
- 3. Before taking any measurements it is worth considering the utilisation of functions such as **AUTO SAVE** and **AUTO PRINT** to provide easy access and analysis of results at a later time. These are fully explained in the following section 'Saving results' and 'Printing results'.
- 4. If readings are required at regular timed intervals, the **TIMED READ** facility can be applied. The use and setting of this is detailed separately in section '**Timed readings**' below.
- 5. Once in measure mode, place the sample (solution of unknown concentration) to be measured into the cuvette holder of the sample chamber and close the lid.
- 6. Move the on screen highlight, using the left and right arrow keys, to the **READ** option and press the enter key.
- 7. A status message "**Reading...**" will be shown throughout the duration of the measurement.
- 8. Once the measurement is complete, the calculated concentration of the sample will be shown, at the resolution and with the units of measure as set in **MEASURE SETUP**.
- 9. This measurement can then be saved or printed as detailed in the sections below entitled 'Saving results' and 'Printing results'.
- 10. Further measurements can be taken by use of the **READ** option.

NOTE: AUTO SAVE, AUTO PRINT and TIMED READ are all related to the selected method.

#### **Timed readings**

- To ensure that all measurements are taken at the required interval, select the required method as described in section 'Using a stored method' above, then return to the main menu.
- 2. Enter the **SETUP** menu and select the **RESULTS STORAGE** section.
- 3. Moving down to the first parameter, **TIMED READ**, set this to **On**.
- 4. The interval between the end of one read and the beginning of the next read can then be set, by selection of **INTERVAL**. Using the *up* and *down* arrow keys, set the numerical figure to the required period, in seconds, between readings.



- 5. The instrument can now be returned to the **MEASURE** mode.
- 6. Place the cuvette of sample in the sample chamber and close the lid.
- 7. To begin taking timed measurements, highlight **READ** and then press the enter key. At this point the first measurement will be taken, indicated by the status message "**Reading...**".
- 8. Once this first read is complete, a further message will be shown which displays a countdown in seconds, to the next read. When this timer reaches zero, the next reading will be taken.



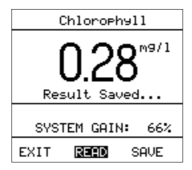
- 9. It is especially important when interval times used are short; to use either the AUTO SAVE or AUTO PRINT in conjunction with timed readings. This enables full automation of result collection without the user having to manually record readings for the entire duration of the testing.
- 10. To cease the timer readings, move the highlight to **EXIT** and press the enter key. This will return the instrument to the main menu.

NOTE: AUTO SAVE, AUTO PRINT and TIMED READ are all related to the selected method.

#### Saving results

#### Saving results

1. When a reading has been taken in the MEASURE mode, the concentration result is displayed on screen and can be saved to the instrument internal memory by selecting the SAVE option on the right hand side of the MEASURE screen. To confirm the save is successful the message "Result Saved..." is shown briefly on screen.



 A maximum of 100 results can be saved in this way, for each method. If more than 100 results are required, the results area must be cleared before the 101<sup>st</sup> result is taken. Please see section "Stored results/Results mode" for details of how to achieve this.

#### Auto save

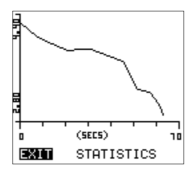
- To remove the need to manually select SAVE after each result, AUTO SAVE can be activated. After every read operation, either timed or manual, the concentration will be automatically saved.
- 2. To ensure every result is saved, select the required method as detailed in **'Using a stored method'** and then return to the **MAIN** *menu*.
- 3. Enter the **SETUP** menu and select the **RESULTS STORAGE** section.
- 4. Move to AUTO SAVE and set to On.
- 5. Using the **EXIT** options, return to the **MAIN** menu and then move to the **MEASURE** screen.
- 6. Each read operation will now have the result automatically saved and this is indicated on screen by the status message "Result saved..."
- 7. A maximum of 100 results can be saved in this way, for each method. If more than 100 results are required, the results area must be cleared before the 101<sup>st</sup> result is taken. Please see section "Stored results/Results mode" for details of how to achieve this.

#### Stored results/Results mode

- 1. To access the saved results for the method currently selected in **MEASURE SETUP**, return to the **MAIN** menu. Using the *down* arrow key, move the highlight to **RESULTS** and press the enter key.
- 2. A table of all saved results, for the method named at the top of the screen will be shown along with the time at which the **READ** operation occurred. The results are presented in chronological order, beginning with the most recent.
- 3. If the number of stored results exceeds seven, a small down arrow will be shown on the right hand side of the screen to indicate further screens of results. These other results are accessed by use of the *down* arrow key. The full range of results can be viewed by use of the *up* and *down* arrow keys to move to the screen of interest.

Chlorophyll			
2.89	15:21:01		
3.06	15:20:59		
3.24	15:20:55		
3.28	15:20:50		
3.69	15:20:43		
3.87	15:20:29		
3.88	15:20:25 ↓		
EXIT	GRAPH CLEAR		

- 4. The *Clear* function on this screen will remove ALL stored results for the *Method* named at the top of the screen. Once *Clear* has been utilised, the results cannot be recovered.
- 5. A graphical representation of the stored results plotted against time is available. This is accessed by the **GRAPH** option at the bottom of the results table screen.



6. The first (earliest) result is set as time zero and the times of subsequent results are calculated from this, but purely for the graph.

7. Further analysis of the saved results can be made by accessing the **STATISTICS** screen. To move to **STATISTICS**, use the option at the bottom of the **GRAPH** screen.

STATISTICS			
SLOPE :	-0.019		
R SQUARED	: 0.922		
EXIT			

8. The **STATISTICS** screen shows two numbers calculated from the saved results. The **SLOPE** reveals the extent of change of fluorescence over time. The **R-squared** value gives an indication of the linearity of the saved results.

#### Section 12

#### **Printing results**

#### **Printing**

1. When a reading has been taken in the **MEASURE** mode, the concentration/raw fluorescence result is displayed on screen. This can be printed to a serial printer or PC if attached via the RS232. The fluorimeter is designed for use with the optional accessories of the Porti printer (P/N 037-701) or DataWay PC software (P/N 050-501).

#### Porti printer

- 1. Connect the Porti printer to the RS232 socket on the back panel of the instrument using the supplied cable.
- 2. The printer can be used without connection to a mains power supply for periods of time by means of the integral battery. If the battery is low or mains power is preferred, plug in the power supply pack using the appropriate cables, as provided with the printer.
- 3. Ensure the printer is fitted with a roll of thermal paper. Paper can be advanced by use of the *Feed* button on the printer.
- 4. Switch on the printer with the black button and the left hand side LED will light. The LED will be green when mains power is used or the battery is well charged. If the LED shows flashes of red and no mains power supply is used, this indicates the battery level is low. Further instructions on the Porti printer can be found in the operator manual included with the printer.
- 5. Now to print the result displayed on the instrument screen just press the *Print* button on the fluorimeter.

#### DataWay

- 1. Connect the supplied cable between the PC and the instrument RS232 socket.
- 2. If using for the first time, insert the DataWay CD into the CD drive of the PC. The CD will automatically run and installation instructions will be shown on the PC screen.
- 3. Once installed the software can be opened either from the programs menu or with the DataWay icon on the PC desktop.
- 4. Open the DataWay software. A pop-up box will appear as the software starts up. Tick the box to the left hand side of the instrument model number and then press the search button.
- 5. When the software has found the attached instrument, the right hand side 'Connect' button will be available. Click on this to connect DataWay.
- 6. Once connected, click the 'Ok' and DataWay will fully open.
- 7. For further help or greater detail on using DataWay, please consult the DataWay help file.
- 8. Click on the Data Manager tab of the DataWay screen.
- 9. Now to print the result displayed on the instrument screen just press the *Print* button on the fluorimeter or the 'take reading' button in DataWay.
- 10. For instruction on saving data, plotting graphs and other features of DataWay, please consult the DataWay help file.

#### Auto print

- 1. To remove the need to manually select **PRINT** after each result, **AUTO PRINT** can be activated. After every read operation, either timed or manual, the concentration will be automatically printed.
- 2. Select the required accessory to which to print results to and setup as detailed above in either the 'Porti printer' or 'Dataway' section.
- 3. To ensure every result is printed, select the required method as detailed in 'Using a stored method' and then return to the main menu.
- 4. Enter the setup menu and select the **results storage** section.
- 5. Move to **AUTO PRINT** and set to **On**.
- 6. Using the **EXIT** options, return to the main menu and then move to the **MEASURE** screen.

Each read operation will now instigate the result being automatically printed, to the connected device.

#### **Maintenance & Troubleshooting**

#### 8.1 General

The 62 Series Fluorimeters are designed to give optimum performance with minimum maintenance. It is only necessary to keep the external surfaces clean and free from dust. The sample area should always be kept clean and any accidental spillage should be wiped away immediately.

To give added protection when not in use, the unit should be disconnected from the mains supply and covered with the optional dust cover (630 028).

For longer-term storage or re-shipment it is recommended that the unit be returned to the original packing case.

#### 8.2 Xenon Lamp Replacement

In the highly unlikely event of lamp failure it will be necessary to contact your local distributor or service agent for assistance. There are no serviceable parts within this product. Under no circumstances should any attempt be made to effect repair as this will invalidate the product Warranty.

627 147

627 148

627 149

627 150

627 151

### **Optional Accessories**

#### 14.1 Accessories

Bandpass Fi	Iters	
627 126	UG1	320-380nm
627 124	BG28	380-500nm
627 125	VG9	480-580nm
Cut-off Filter	s (low cut, tra	nsmission blocked below stated wavelength)
627 127	Kodak 29	610nm
627 128	Ilford 201	545nm
627 129	Kodak 8	475nm
627 130	Kodak 2B	395nm
627 131	Glass	305nm
Interference	Filters (10nm	bandpass)
627 141	Wavelength 2	250nm
627 132	Wavelength 2	254nm
627 142	Wavelength 2	260nm
627 143	Wavelength 2	265nm
627 144	Wavelength 2	270nm
627 145	Wavelength 2	280nm
627 146	Wavelength 2	290nm

Wavelength 295nm

Wavelength 300nm

Wavelength 305nm

Wavelength 310nm

Wavelength 313nm

**627 133** Wavelength 340nm Wavelength 350nm Wavelength 360nm

627 138 Wavelength 365nm
627 158 Wavelength 370nm
627 134 Wavelength 380nm
627 159 Wavelength 390nm

**627 135** Wavelength 400nm Wavelength 405nm Wavelength 410nm

**627 162** Wavelength 415nm **627 163** Wavelength 420nm

**627 164** Wavelength 430nm Wavelength 436nm

**627 166** Wavelength 440nm Wavelength 450nm

Interference F	Filters (continued)
627 167	Wavelength 460nm
627 136	Wavelength 470nm
627 168	Wavelength 480nm
627 169	Wavelength 490nm
627 170	Wavelength 500nm
627 171	Wavelength 510nm
627 137	Wavelength 515nm
627 172	Wavelength 520nm
627 173	Wavelength 530nm
627 174	Wavelength 532nm
627 175	Wavelength 540nm
627 176	Wavelength 546nm
627 177	Wavelength 550nm
627 178	Wavelength 560nm
627 179	Wavelength 570nm
627 180	Wavelength 577nm
627 181	Wavelength 580nm
627 182	Wavelength 590nm
627 183	Wavelength 600nm
627 184	Wavelength 610nm
627 185	Wavelength 620nm
627 186	Wavelength 630nm
627 187	Wavelength 633nm
627 188	Wavelength 640nm
627 189	Wavelength 650nm
627 190	Wavelength 656nm
627 191	Wavelength 660nm
627 192	Wavelength 670nm
627 193	Wavelength 680nm
627 194	Wavelength 690nm
627 195	Wavelength 694nm
627 196	Wavelength 700nm

## Alternative filters available on request

### Sample Cells

A wide range of cells, in addition to those listed, is available to special order. Details supplied on request.

035 120	100µl quartz cell
035 121	500µl glass cell
035 122	500µl quartz cell
060 247	Pack 100 (10mm) plastic (3ml) cuvettes
060 253	10x10mm glass cell (UV)
060 254	10x10mm glass cell (Visible)
060 255	Flow cell

#### **Additional Items**

628 200	Heated cell system comprising heated cell holder, controller, 2 leads
021 057	12V connection lead
630 028	Dust Cover
050 501	DataWay PC Software CD-ROM with interface cables and instructions
033 290	Carry Case
021 056	Universal 12V power supply with UK, EU and US mains leads.

# **Specification & Data**

# 10.1 Technical Specification

Specification		6270	6280	6285
•				
Light source		Pulsed Xenor	n Lamp with press-to-r	ead operation
Expected Lifetime			ment required in norm	
Sensitivity (Quinine	Sulphate)	1μg/ml	<1ng/ml	<1ng/ml
Dynamic Range (Quinine Sulphate)		5.2x 10 <sup>4</sup>	5.2 x 10 <sup>6</sup>	5.2 x 10 <sup>6</sup>
Detector		Photodiode	Photomultiplier	Photomultiplier
Wavelength Range		190 to 1100nm	190 to 650nm	190 to 850nm
Wavelength Selection	on	Intelligent Filter Modules (IFM)		
Raw Fluorescence		✓	<b>✓</b>	<b>✓</b>
	Calibration	Blank + %	Blank + %	Blank + %
Concentration		✓	<b>✓</b>	~
	Calibration	Blank + up to 6 standards	Blank + Up to 6 standards	Blank + up to 6 standards
Kinetics (Timed Rea	adings)	In both above modes	In both above modes	In both above modes
	Time Limits	1 reading every 3 to 999 seconds		
Methods Stored		20	20	20
Results Stored		100/method	100/method	100/method
GLP		Method lock with password access Time and date stamped results Last calibration date and time stored Calibration Interval set Calibration due reminder User ID		
Outputs			RS232 Serial Port	
PC Software		Compatible with DataWay (option)		
Sample Handling		10mm Square cuvette holder (standard) Heated Sample Holder (Option) Sipper Pump (Option)		
Power Supply		90 to 230Vac universal mains adapter		
		12Vdc input (cigarette lighter cable and battery pack options)		
Size		365(w) x 272 (d) x 160(h) mm		
Weight		6Kgs		

#### **Glossary of Terms**

**Filters** - There are three types of filter commonly used in fluorimeters, Band-pass filters, Cut-Off filters and Interference filters. The first two offer a low-cost general purpose option while the more expensive interference filters offer precise selection of excitation or emission wavelengths. Often one type will be used for excitation and another for emission depending on the proximity of interfering wavelengths.

An excitation filter is fitted between the light source and the sample to limit the wavelengths reaching it to those absorbed by the analyte but not by any other substance in the sample matrix.

An emission filter is fitted between the sample and detector to restrict the wavelengths reaching the detector to those due to the fluorescence of the analyte, while blocking those due to scatter and stray light.

**Band-pass filters** - can be used to isolate excitation or emission wavelengths, their broad transmittance spectra enable them to be used in many applications. However when used for both excitation and emission great care must be taken that their transmittance ranges do not overlap. Most useful of those available are the UG1 filter with transmittance from 320 to 380nm, BG28 from 380 to 500nm and VG9 from 480 to 580nm.

**Cut-off filters** - (low cut) will pass all wavelengths above their designated cut-off point, this makes them ideal as an emission filter, blocking any stray radiation from the shorter wavelength excitation source; but beware of scatter which will be at longer wavelengths and so will be transmitted with the fluorescence.

**Interference filters** - are available across the full wavelength range and are specified by their peak wavelength and spectral bandwidth (SBW). With typical SBWs of 10nm their performance is close to that of a monochromator and all unwanted wavelengths can be excluded.

### **TET** - Total Energy Transfer

**IFM** - Intelligent Filter Modules. Filters are pre-programmed with their peak wavelength and type. These are identified by the system, enabling error messages to be generated and displayed should the wrong filters be fitted or their positions be incorrect.

## **EC Declaration of Conformity**

Jenway Models 6270, 6280 and 6285 Fluorimeters comply with the following European Standards:

EN 50081-1: 1992 Electromagnetic compatibility - Generic emission standard

EN 50082-1: 1992 Electromagnetic compatibility – Generic immunity standard

EN 61010-1: 2001 Safety requirements for electrical equipment for measurement, control and laboratory use

Following the provision of:

EMC Directive – 89/336/EEC and Low Voltage Directive – 73/23/EEC



Thank you for reading this data sheet.

For pricing or for further information, please contact us at our UK Office, using the details below

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Please note - Product designs and specifications are subject to change without notice. The user is responsible for determining the suitability of this product.